

Package ‘forestmangr’

January 24, 2021

Type Package

Title Forest Mensuration and Management

Version 0.9.3

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Description

Processing forest inventory data with methods such as simple random sampling, stratified random sampling and systematic sampling. There are also functions for yield and growth predictions and model fitting, linear and nonlinear grouped data fitting, and statistical tests. References: Kershaw Jr., Ducey, Beers and Husch (2016). <doi:10.1002/9781118902028>.

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URL <https://github.com/sollano/forestmangr#readme>

BugReports <https://github.com/sollano/forestmangr/issues>

Depends R(>= 3.3)

Imports dplyr(>= 0.7.0), ggplot2(>= 2.0), ggthemes, tidyr, broom,
purrr, forcats, plyr, tibble(>= 3.0.0), systemfit, ggpmisc,
rlang, utils, car, stats, methods, magrittr, minpack.lm,
FinCal, formattable, scales, ggdendro,
gridExtra,tidysselect,shiny,miniUI

Suggests covr, knitr, rmarkdown

VignetteBuilder knitr

Date 2021-01-24

Encoding UTF-8

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RoxygenNote 7.1.1

NeedsCompilation no

Repository CRAN

Date/Publication 2021-01-24 22:00:02 UTC

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<code>average_tree_curve</code>	<i>Generate the curve of a forest's average tree using the Kozak taper model</i>
---------------------------------	--

Description

Generate a ggplot curve of a forest's average tree using the Kozak taper model (Kozak, Munro and Smith, 1969).

Usage

```
average_tree_curve(  
  df,  
  d,  
  dbh,  
  h,  
  th,  
  facet = NA,  
  color = NA,  
  mirror = TRUE,  
  eq = TRUE  
)
```

Arguments

<code>df</code>	A data frame.
<code>d</code>	Quoted name of the section diameter variable, in cm.
<code>dbh</code>	Quoted name of the diameter at breast height variable, in cm.

h	Quoted name of the section height variable, in meters.
th	Quoted name of the total height variable, in meters.
facet	Optional argument. If supplied with the Quoted name of a factor variable(s), this variable is used to divide the plot into facets. Default: NA.
color	Quoted name of a variable. If supplied, this variable will be used to classify the data by color. Default: NA.
mirror	if TRUE, the plot will be mirrored, to resemble the shape of a tree. Default: TRUE
eq	if TRUE, Kozak's taper model is adjusted and the equation is shown on the plot. Default TRUE

Value

A ggplot object.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

References

Kozak, A., Munro, D. D. and Smith, J. H. G. (1969) Taper Functions and their Application in Forest Inventory, *The Forestry Chronicle*, 45, pp. 278–283.

Examples

```
library(forestmangr)
data("exfm7")
exfm7

# Get the data's average tree curve inserting the section diameter and height, total height
# and dbh variables:
average_tree_curve(df = exfm7, d = "di_wb", dbh = "DBH", h = "hi", th = "TH")

# It's possible to get the average tree curve of each strata with the facet argument,
# and divide the data by color with the color argument:
average_tree_curve(df=exfm7,d="di_wb",dbh="DBH",h="hi",th="TH",facet="STRATA",color="GENCODE")
```

 bdq_meyer

Classify a forest for selective cutting using the Meyer BDq method

Description

This function can be used to plan and execute selective cuttings of a native forest, without damaging the forest's natural structure.

Usage

```
bdq_meyer(  
  df,  
  plot,  
  dbh,  
  plot_area,  
  class_interval = 5,  
  dbh_min = 5,  
  licourt_index = 2,  
  output = "table"  
)
```

Arguments

df	A data frame.
plot	Quoted name of the plot variable. used to differentiate the plot's trees, and calculate the number of sampled plots.
dbh	Quoted name of the diameter at breast height variable, in cm.
plot_area	Quoted name of the plot area variable, or a numeric vector with the plot area value. The plot area value must be in square meters.
class_interval	Numeric value for the class interval used to classify the data. Default: 5.
dbh_min	Numeric value for minimum diameter value to be considered in the classification. dbh values smaller than this will be dismissed from the classification. Default: 5.
licourt_index	Numeric value for the start licourt index used. Default: 2.
output	Character value for the desired output. Can be either "table" for the classified data table, "model" to get a lm object with the linear model fitted, "coefs" to get a vector with the Meyer coefficients, or "full", to get a list with all results. Default: "table".

Value

a data frame, a lm object, a vector or a list, according to the output argument.

Author(s)

Eric Bastos Gorgens <e.gorgens@gmail.com>

References

Souza, A. L. and Soares, C. P. B. (2013) Florestas Nativas: estrutura, dinamica e manejo. Vicosa: UFV.

Examples

```

library(forestmangr)
data("exfm20")
exfm20

# To get the table with the regulated forest:
bdq_meyer(exfm20, "transect", "dbh", 1000)

# Use different class interval values to get different results:
bdq_meyer(exfm20, "transect", "dbh", 1000, class_interval = 10)

# It's possible to get different outputs:
bdq_meyer(exfm20, "transect", "dbh", 1000, output="model")
bdq_meyer(exfm20, "transect", "dbh", 1000, output="coefs")
bdq_meyer(exfm20, "transect", "dbh", 1000, output="full")

```

bias_per

Bias of an estimator in percentage

Description

Function for calculating the bias of an estimator.

Usage

```
bias_per(df, y, yhat, na.rm = TRUE)
```

Arguments

df	a data frame.
y	Quoted name of the variable representing the observed values in the data frame. If a data frame is not provided, y can also be a numeric vector.
yhat	Quoted name of the variable representing the estimated values in the data frame. If a data frame is not provided, yhat can also be a numeric vector.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds. default: TRUE

Details

Function for calculating the bias of an estimator, given the observed values, and the estimated values.

Value

Numeric vector with the bias value, in percentage.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

See Also

other statistics to evaluate estimators: [rmse_per](#) for the Root mean square error of an estimator

Examples

```
library(forestmangr)
data("exfm11")
exfm11

# Bias of an estimator, given the data frame and quoted variable names:
bias_per(exfm11, "TH", "TH_EST3")

# Bias of an estimator, given the vectors for observed and estimated values:
bias_per(y = exfm11$TH, yhat = exfm11$TH_EST3)
```

check_names

Check if character vector contains variable names

Description

Function used to check if a string, or a character vector contains variable names of a given data frame.

Usage

```
check_names(df, var_names, boolean = TRUE)
```

Arguments

df	a data frame.
var_names	Character vector to be compared with the data frame names.
boolean	Boolean object used to define if the output is going to be a boolean object TRUE, or a string FALSE. Default: TRUE.

Details

Function used to check if a string, or a character vector contains variable names of a given data frame. This functions is mainly used to error-proof other functions of this package,

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```
library(forestmangr)

check_names(iris, "Species")
check_names(iris, "Species", boolean = FALSE )

check_names(iris, c("Especies", "Setal.Width") )
check_names(iris, c("Especies", "Setal.Width"), boolean = FALSE)
```

classify_site	<i>Classify inventory data based on site index</i>
---------------	--

Description

Use the site variable to classify a forest management data.

Usage

```
classify_site(df, site, nc = 3, plot, .groups = NA)
```

Arguments

df	A data frame.
site	Quoted name for the site variable.
nc	number of categories used to classify the data. If 3, a additional column will be created with levels Lower, Middle and Upper, referencing the 3 categories. If not, only numbers will be used to differentiate the categories. Default: 3.
plot	Quoted name for the plot variable.
.groups	Optional argument. Quoted name(s) of grouping variables used to fit multiple regressions, one for each level of the provided variable(s). Default NA.

Value

A data frame classified based on the site index.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

See Also

other sampling functions: [fit_clutter](#) for fitting Clutter's Growth and Yield, and [est_clutter](#) for estimating Clutter's Growth and Yield model variables.

Examples

```
library(forestmangr)
data("exfm17")
exfm17

# Classify data into 3 classes:
ex_class <- classify_site(exfm17, "S", 3, "plot")
head(ex_class ,15)
```

class_center	<i>Classify a given variable and get center of class</i>
--------------	--

Description

This function can be used to divide the data into classes, based on minimum value and class interval of a given variable, and create a column with the center of each class.

Usage

```
class_center(df, y, ci = 3, ymin = 5)
```

Arguments

df	A data frame.
y	Quoted name of a variable, or a vector to be classified.
ci	Numeric value for the class interval used to classify the data. Default: 3.
ymin	Numeric value for minimum value value to be considered in the classifications. dbh values smaller than this will be dismissed from the classification. Default: 5.

Value

if df is supplied, a data frame containing the supplied data with a new column for the center of classes; if df is missing, a vector with the center of class.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```

library(forestmangr)
library(dplyr)
data("exfm20")
exfm20

# n
# Number of individuals per ha per diameter class
class_center(df = exfm20, y = "dbh", ci = 10, ymin = 10)

exfm20 %>%
mutate(CC = class_center(y = dbh, ci = 10, ymin = 10))

```

diameter_class	<i>Divide data into diameter classes, and get number of observations</i>
----------------	--

Description

This function can be used to divide data into diameter classes, get the number of observations, number of observations per ha, and check number of species individuals, volume and G in each diameter class. It's also possible to spread the diameter classes as columns.

Usage

```

diameter_class(
  df,
  dbh,
  plot = NA,
  plot_area,
  ci = 5,
  dbhmin = 5,
  species = NA,
  volume = NA,
  NI_label = "NI",
  cc_to_column = FALSE,
  G_to_cc = FALSE,
  cctc_ha = TRUE,
  keep_unused_classes = FALSE
)

```

Arguments

df	A data frame.
dbh	Quoted name of the diameter at breast height variable, in cm.
plot	Optional parameter. Quoted name of the plot variable. used to differentiate the plots trees, and calculate the number of sampled plots. Default NA.

plot_area	Optional parameter. Quoted name of the plot area variable, or a numeric vector with the plot area value. The plot area value must be in square meters. Default NA.
ci	Numeric value for the class interval used to classify the data. Default: 5.
dbhmin	Numeric value for minimum diameter value to be considered in the classifications. dbh values smaller than this will be dismissed from the classification. Default: 5.
species	Optional parameter. Quoted name of the scientific names variable, or any variable used to differentiate the different species found in data. If supplied, will be used to classify the species in the diameter data. Default NA.
volume	Optional parameter. Quoted name of the volume variable. If supplied, will be used classify the volume variable in the different diameter classes. Also, if cc_to_column is TRUE, the center of class columns will be filled with volume values, instead of number of individuals. Default NA.
NI_label	Label used for Species not identified. This parameter works along with species. The level supplied here will not be considered in the classification. Default "NI".
cc_to_column	If TRUE, will spread the center class column as multiple columns, one for each class. The value that fills these columns, by default is the number of individuals found in each class, but this can be changed by using other arguments. Default FALSE.
G_to_cc	If TRUE, and cc_to_column is also TRUE, will fill the center of class columns with basal area values, instead of number of individuals. Default FALSE.
cctc_ha	If TRUE, will calculate values per hectare for number of individuals per class, basal area per class and volume per class (if supplied). These values will also be used to fill the center of class columns, if cc_to_column is TRUE. Default TRUE.
keep_unused_classes	Some diameter classes may end up empty, depending on the maximum value of diameter and the class interval used. If this is TRUE, those classes will not be removed from the final data frame. Default FALSE.

Value

A data frame containing the supplied data divided into diameter classes.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```
library(forestmangr)
data("exfm20")
exfm20

# n
# Number of individuals per ha per diameter class
```

```
diameter_class(df = exfm20, dbh = "dbh", ci = 10, dbhmin = 10, volume = "vol")

# Number of individuals per ha per diameter class per species
diameter_class(exfm20,"dbh", "transect", 10000, 10, 10, "scientific.name")

# Number of individuals per ha per diameter class, with each diameter class as a column
diameter_class(exfm20,"dbh", "transect", 10000, 10, 10, "scientific.name", cc_to_column=TRUE)

# G
# Basal area per ha per diameter class, with each diameter class as a column
diameter_class(exfm20,"dbh", "transect",10000,10,10,"scientific.name",
cc_to_column=TRUE,G_to_cc=FALSE)

# Volume
# Volume per ha per diameter class
diameter_class(exfm20,"dbh", "transect", 10000, 10, 10, "scientific.name",volume = "vol")

# Volume per ha per diameter class, with each diameter class as a column
diameter_class(exfm20,"dbh","transect",10000,10,10,"scientific.name","vol",cc_to_column=TRUE)
```

dom_height

Calculate the dominant height of forest inventory data plots

Description

This function is used to get a data frame with Dominant height values for each plot in an forest inventory data.

Usage

```
dom_height(
  df,
  th,
  dbh,
  plot,
  obs,
  dom,
  .groups,
  merge_data = FALSE,
  dh_name = "DH"
)
```

Arguments

df	A data frame.
th	Quoted name of the total height variable.

dbh	Quoted name of the diameter at breast height variable. Used to filter out trees with no diameter measurement.
plot	Quoted name of the plot variable. used to differentiate the data's plots. If this argument is missing, the defined groups in the data frame will be used, If there are no groups in the data, the function will fail.
obs	Quoted name of the observations variable. This will be used to tell which trees are dominant, i.e. it's the variable that tells the type of tree; if it is normal, dominant, suppressed, etc. If this argument is not supplied, the function will calculate the average value of 2 trees with bigger height values in each plot, and use that as the dominant value.
dom	Character value for the dominant tree code used in the observations variable supplied in the obs argument. This is used alongside the obs argument to differentiate dominant trees from the others.
.groups	Optional argument. Quoted name(s) of grouping variables that can be added to differentiate subdivisions of the data. Default: NA.
merge_data	If TRUE, will merge the original data frame with the dominant height table. Default: FALSE.
dh_name	Character value for the name of the dominant height variable created. Default: "DH"

Value

A data frame with the the dominant height values by plot.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```
library(forestmangr)
data("exfm9")
exfm9

# Let's say we need to get the dominant height (DH) values for a forest inventory data.
# If we don't have a variable that tells which trees are dominant, it's ok. We can
# still estimate DH using this function. It will average the top 2 trees of each plot:
dom_height(df=exfm9, th="TH", dbh="DBH", plot="PLOT")

# Of course, if we do have a variable that differentiate the dominant trees, it's
# best we use it. For that we use the obs argument, and the dom argument.
# In this data, the OBS variable is used to tell the type of tree.
# Let's check the levels in our OBS variable, to see which one is associated
# with dominant trees.

levels(as.factor(exfm9$OBS))

# So, the "D" level must be the one that tells which trees are dominant. Let's use it:
dom_height(df=exfm9, th="TH", dbh="DBH", plot="PLOT", obs="OBS", dom="D")
```

```

# If there are subdivisions of the data, like different strata, they can be included in the
# .groups argument:
dom_height(df=exfm9, th="TH", dbh="DBH", plot="PLOT", obs="OBS", dom="D", .groups="STRATA")

# It's possible to automatically bind the dominant heights table to the original data,
# using the merge_data argument:

dom_height(df=exfm9, th="TH", dbh="DBH", plot="PLOT", obs="OBS",
dom="D", .groups="STRATA", merge_data=TRUE)

```

est_clutter	<i>Estimate future and present basal area, volume, TCA, CMI and MMI values of the Clutter Growth and Yield Model</i>
-------------	--

Description

This function estimates the present the present value of basal area for each class using either the class mean, or a linear quadratic model, and then uses it's value to calculate the basal area from Clutter's growth and yield model.

Usage

```

est_clutter(
  df,
  age,
  basal_area,
  site,
  category,
  coeffs,
  method = "average",
  annual_increment = FALSE,
  gray_scale = TRUE,
  output = "table"
)

```

Arguments

df	A data frame.
age	A numeric vector with the desired age range to be used in the estimation, or a Quoted name for the age variable.
basal_area	Quoted name for the basal area variable.
site	Quoted name for the average site variable.
category	Quoted name for the category variable.

coeffs	Numeric vector or a data frame with the fitted values of Clutter's growth and yield model. It must be a named vector, with b0,b1,b2,b3,a0 and a1 as names. a1 is not obligatory.
method	Method used for estimating the present basal area of each class. It can either be the class' average basal area "average", or an estimated value from a linear quadratic model of site as a function of basal area "model". Default: "average".
annual_increment	If TRUE, changes the labels from Mean Monthly Increment (MMI) and Current Monthly Increment (CMI) to Mean Annual Increment (MAI) and Current Annual Increment (CAI). Default FALSE.
gray_scale	If TRUE, the plot will be rendered in a gray scale. Default: "TRUE".
output	Type of output the function should return. This can either be "plot", for the estimation plots, "table", for a data frame with the estimated values, and "full" for a list with the plot and 2 more data frames. "table".

Value

A data frame, a ggplot object or a list, according to output.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

See Also

other sampling functions: [fit_clutter](#) for fitting the clutter growth and Yield model, and [classify_site](#) for classifying data according to site.

Examples

```
library(forestmangr)
data("exfm17")
exfm17

clutter <- fit_clutter(exfm17, "age", "DH", "B", "V", "S", "plot")
clutter

# Classify data into 3 classes:
ex_class <- classify_site(exfm17, "S", 3, "plot")
head(ex_class ,15)

# Estimate basal area using the average basal area as the initial basal area,
# volume, Mean Monthly Increment (MMI) and Current Monthly Increment (CMI)
# values using Clutter's model:
est_clutter(ex_class,20:125, "B", "S", "category_", clutter, "average")

# For a more detailed output, including a plot, use output="full":
est_clutter(ex_class,20:125, "B", "S", "category_", clutter, output="full")
```

```
# Estimate basal area using an estimated basal area as the initial basal area:  
est_clutter(ex_class,20:125,"B","S","category_",clutter,"model")  
  
# age can be a variable:  
est_clutter(ex_class,"age","B","S","category_", clutter,"model")
```

exfm1

Stratified random inventory pilot data

Description

In this data, each observation is a plot.

Usage

```
data(exfm1)
```

Format

A data frame with 22 observations and 4 variables:

STRATA stratum number

STRATA_AREA area of each strata, in hectares

PLOT_AREA area of plots, in square meters

VWB volume with bark, in cubic meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Source

Soares, C. P. B., Paula Neto, F. and Souza, A. L. (2012) Dendrometria e Inventario Florestal. 2nd ed. Vicosa: UFV.

exfm10	<i>Inventory data of an eucalyptus forest Brazil with dominant height variable</i>
--------	--

Description

In this data, each observation is a tree.

Usage

```
data(exfm10)
```

Format

A data frame with 900 observations and 14 variables:

MAP map numbers

PROJECT project number

STRATA stratum number

GENCODE genetic code of plots

STRATA_AREA area of each strata, in hectares

PLANTING_DATE date of planting

SPACING Spacing used in the plots, in meters

PLOT plot number

MEASUREMENT_DATE date of measurement

PLOT_AREA area of plots, in square meters

PIT pit number

DBH diameter at breast height, in meters

TH total height, in meters

OBS quality of trees, N = normal tree, D = dominant tree, F = a failure, or dead tree

DH dominant height, in meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm11	<i>Observed height values of trees, and estimated values using 3 different models</i>
--------	---

Description

In this data, each observation is a tree.

Usage

```
data(exfm11)
```

Format

A data frame with 199 observations and 6 variables:

STRATA stratum number

PLOT plot number

TH total height, in meters

TH_EST1 total estimated height with model 1, in meters

TH_EST2 total estimated height with model 2, in meters

TH_EST3 total estimated height with model 3, in meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm12	<i>Inventory data of an eucalyptus forest in Brazil, with age and site variables</i>
--------	--

Description

In this data, each observation is a plot. The site was estimated using the Chapman & Richards model.

Usage

```
data(exfm12)
```

Format

A data frame with 139 observations and 8 variables:

strata stratum number
plot plot number
age average age of plots, in months
DH dominant height, in meters
N number of individuals
V volume of trees, in cubic meters
B basal area, in square meters
S site variable, in meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm13

Experimental data on nitrogen effect in species fertilizing

Description

In this data, each observation is a tree.

Usage

data(exfm13)

Format

A data frame with 36 observations and 7 variables:

species species common name
trat treatment number
esp species number
N quantity of nitrogen used, in grams
N2 squared quantity of nitrogen used, in squared grams
block block number
dbh diameter at breast height, in meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm14	<i>Inventory data of an eucalyptus forest in Brazil, with age and dominant height variables</i>
--------	---

Description

In this data, each observation is a plot.

Usage

```
data(exfm14)
```

Format

A data frame with 859 observations and 4 variables:

strata stratum number

plot plot number

age average age of plots, in months

dh dominant height, in meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm15	<i>Simplified Inventory data of an eucalyptus forest in Brazil</i>
--------	--

Description

In this data, each observation is a tree.

Usage

```
data(exfm15)
```

Format

A data frame with 900 observations and 7 variables:

STRATA stratum number

STRATA_AREA area of each strata, in hectares

PLOT plot number

PLOT_AREA area of plots, in square meters

DBH diameter at breast height, in meters

TH total height, in meters

OBS quality of trees, N = normal tree, D = dominant tree, F = a failure, or dead tree

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm16

Inventory data of an eucalyptus forest in Brazil, with age variable

Description

In this data, each observation is a plot.

Usage

```
data(exfm16)
```

Format

A data frame with 139 observations and 8 variables:

strata stratum number

plot plot number

age average age of plots, in months

DH dominant height, in meters

N number of individuals

V volume of trees, in cubic meters

B basal area, in square meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm17

Inventory data of an eucalyptus forest in Brazil, with age and site variables

Description

In this data, each observation is a plot. The site was estimated using the Schumacher model.

Usage

```
data(exfm17)
```

Format

A data frame with 139 observations and 8 variables:

strata stratum number

plot plot number

age average age of plots, in months

DH dominant height, in meters

N number of individuals

V volume of trees, in cubic meters

B basal area, in square meters

S site variable, in meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm18

Experimental data on nitrogen effect in species fertilizing

Description

In this data, each observation is a tree.

Usage

`data(exfm18)`

Format

A data frame with 877 observations and 6 variables:

Plot plot number

Species species scientific name

Tree tree number

Trunk trunk number

CBH circumference at breast height, in meters

DBH diameter at breast height, in meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm19

Volume of felled trees, measured in the Smalian method

Description

In this data, each observation is a section of a tree.

Usage

```
data(exfm19)
```

Format

A data frame with 16 observations and 3393 variables:

STRATA stratum number

TREE number of trees

DBH diameter at breast height, in meters

TH total height, in meters

CSA cross section area with bark, in square meters

VWB volume with bark, in cubic meters

FFWB form factor for each section

FFWB_mean average form factor value

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm2

Stratified random inventory definite data

Description

In this data, each observation is a plot.

Usage

```
data(exfm2)
```

Format

A data frame with 57 observations and 4 variables:

STRATA stratum number

STRATA_AREA area of each strata, in hectares

PLOT_AREA area of plots, in square meters

VWB volume with bark, in cubic meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Source

Soares, C. P. B., Paula Neto, F. and Souza, A. L. (2012) Dendrometria e Inventario Florestal. 2nd ed. Vicosa: UFV.

exfm20

Inventory data of a natural forest in Brazil

Description

In this data, each observation is a tree.

Usage

```
data(exfm20)
```

Format

A data frame with 12295 observations and 18 variables:

cod area code

transect plot number

tree tree number

common.name species common name

scientific.name species scientific name

family species family name

dbh diameter at breast height, in meters

canopy.pos canopy position

light level of light received by the tree

dead tells if the tree is dead or not

Hcom commercial height, in meters

Htot total height, in meters
date date of measurement
utm.east utm east position value
utm.north utm north position value
vol volume of trees, in cubic meters
plot.area plot area, in square meters
total.area total area, in hectares

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm21

Inventory data of an eucalyptus forest in Brazil

Description

In this data, each observation is a tree.

Usage

data(exfm21)

Format

A data frame with 900 observations and 13 variables:

STRATA stratum number
STRATA_AREA area of each strata, in hectares
PLOT plot number
PLOT_AREA area of plots, in square meters
DBH diameter at breast height, in meters
TH total height, in meters
OBS quality of trees, N = normal tree, D = dominant tree, F = a failure, or dead tree
DH dominant height, in meters
TH_EST estimated total height, in meters
CSA cross sectional area, in square meters
AGE average age of plots, in months
VWB volume with bark, in cubic meters
VWOB volume without bark, in cubic meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm22

Revenue data if an eucalyptus forest

Description

In this data, each observation is the year's revenue.

Usage

```
data(exfm22)
```

Format

A data frame with 8 observations and 3 variables:

year Revenue year

cost cost values for that year, in dollars

revenue revenue values for that year, in dollars

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm3

Simple random inventory pilot data

Description

In this data, each observation is a plot.

Usage

```
data(exfm3)
```

Format

A data frame with 10 observations and 3 variables:

TOTAL_AREA total area, in hectares

PLOT_AREA area of plots, in square meters

VWB volume with bark, in cubic meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Source

Soares, C. P. B., Paula Neto, F. and Souza, A. L. (2012) Dendrometria e Inventario Florestal. 2nd ed. Vicosa: UFV.

exfm4

Simple random inventory definite data

Description

In this data, each observation is a plot.

Usage

```
data(exfm4)
```

Format

A data frame with 25 observations and 3 variables:

TOTAL_AREA total area, in hectares

PLOT_AREA area of plots, in square meters

VWB volume with bark, in cubic meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Source

Soares, C. P. B., Paula Neto, F. and Souza, A. L. (2012) Dendrometria e Inventario Florestal. 2nd ed. Vicosa: UFV.

exfm5

Systematic inventory data

Description

In this data, each observation is a plot.

Usage

```
data(exfm5)
```

Format

A data frame with 18 observations and 3 variables:

TOTAL_AREA total area, in hectares

PLOT_AREA area of plots, in square meters

VWB volume with bark, in cubic meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Source

Soares, C. P. B., Paula Neto, F. and Souza, A. L. (2012) Dendrometria e Inventario Florestal. 2nd ed. Vicosa: UFV.

exfm6

Stratified random inventory definite data 2

Description

In this data, each observation is a plot.

Usage

`data(exfm6)`

Format

A data frame with 10 observations and 14 variables:

GENCODE genetic code of plots

MAP map numbers

STRATA stratum number

PLOT plot number

AGE average age of plots, in months

STRATA_AREA area of each strata, in hectares

PLOT_AREA area of plots, in square meters

DBH diameter at breast height, in meters

q quadratic diameter, in meters

TH total height, in meters

DH dominant height, in meters

G basal area, in square meters

VWB volume with bark, in cubic meters

VWOB volume without bark, in cubic meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm7

Data of felled trees sections, measured in the Smalian method

Description

In this data, each observation is a section of a tree.

Usage

```
data(exfm7)
```

Format

A data frame with 11 observations and 3393 variables:

MAP map numbers

PROJECT Project's name

SPACING Spacing used in the plots, in meters

STRATA stratum number

GENCODE genetic code of plots

TREE number of trees

DBH diameter at breast height, in meters

TH total height, in meters

hi height of sections, in meters

di_wb diameter of sections with bark, in centimeters

bark_t bark of thickness, in millimeters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm8

Data of felled trees sections, measured in the Huber method

Description

In this data, each observation is a section of a tree.

Usage

```
data(exfm8)
```

Format

A data frame with 10 observations and 596 variables:

CLONE Clone number

STRATA stratum number

TREE number of trees

LOG log number

DBH diameter at breast height, in meters

TH total height, in meters

hi height of sections, in meters

di_wb diameter of sections with bark, in centimeters

bark_t bark of thickness, in millimeters

sec_length section length, in meters

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

exfm9

Inventory data of an eucalyptus forest in minas gerais, Brazil

Description

In this data, each observation is a tree.

Usage

```
data(exfm9)
```

Format

A data frame with 900 observations and 14 variables:

MAP map numbers

PROJECT project number

STRATA stratum number

GENCODE genetic code of plots

STRATA_AREA area of each strata, in hectares

PLANTING_DATE date of planting

SPACING Spacing used in the plots, in meters

PLOT plot number

MEASUREMENT_DATE date of measurement

PLOT_AREA area of plots, in square meters

PIT pit number

DBH diameter at breast height, in meters

TH total height, in meters

OBS quality of trees, N = normal tree, D = dominant tree, F = a failure, or dead tree

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

fit_clutter

Fit the Clutter model for growth and yield

Description

Fit the Clutter model for growth and yield using the two stage least squares method (2SLS).

Usage

```
fit_clutter(  
  df,  
  age,  
  dh,  
  basal_area,  
  volume,  
  site,  
  plot,  
  .groups = NA,  
  model = "full",  
  keep_model = FALSE  
)
```

Arguments

df	A data frame.
age	Quoted name for the age variable.
dh	Quoted name for the dominant height variable.
basal_area	Quoted name for the basal area variable.
volume	Quoted name for the volume area variable.
site	Quoted name for the site variable.
plot	Quoted name for the plot variable.
.groups	Optional argument. Quoted name(s) of grouping variables used to fit multiple regressions, one for each level of the provided variable(s). Default NA.
model	Character variable for the type of the model fitted. If "full", the full model will be used. if "mod", a modified model will be fitted, where the X3 variable is excluded from the regression. Default: full.
keep_model	If TRUE a variable with the regression model will be kept in the data frame. Default: FALSE.

Value

A data frame with the regression coefficients.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

References

Clutter, J. L. (1963) Compatible Growth For Loblolly by the Southeastern, Forest Science, 9(3), pp. 354–371. Sullivan, A. D. and Clutter, J. L. (1972) A Simultaneous Growth and Yield for Loblolly Pine, Forest Science, 18(1), pp. 76–86. Campos, J. C. C. and Leite, H. G. (2017) Mensuracao Florestal: Perguntas e Respostas. 5a. Vicosa: UFV.

See Also

other sampling functions: [est_clutter](#) for estimating the Clutter growth and yield model variables, and [classify_site](#) for classifying data according to site.

Examples

```
library(forestmangr)
data("exfm17")
exfm17

# To fit the Clutter model we just need to define the data, and age, dominant height,
# basal area, volume, site and plot variables:
fit_clutter(exfm17, "age", "DH", "B", "V", "S", "plot")
```



```
# To fit the alternate model (without a1) just use model="mod":
fit_clutter(exfm17, "age", "DH", "B", "V", "S", "plot", model="mod")

# To keep the regression model, use keep_model=TRUE:
fit_clutter(exfm17, "age", "DH", "B", "V", "S", "plot", keep_model=TRUE)
```

forest_structure	<i>Get the forest horizontal, vertical and internal structure</i>
------------------	---

Description

This function calculates the horizontal structure of a given forest inventory data, with information like absolute frequency, relative frequency, absolute density, relative density, absolute dominance, relative dominance, importance value index, and coverage value index. If additional variables are supplied, the vertical and internal structures are also provided.

Usage

```
forest_structure(
  df,
  species,
  dbh,
  plot,
  plot_area,
  vertical_est = NA,
  internal_est = NA,
  NI_label = ""
)
```

Arguments

df	A data frame.
species	Quoted name of the scientific names variable, or any variable used to differentiate the different species found in data.
dbh	Quoted name of the diameter at breast height variable, in cm.
plot	Quoted name of the plot variable. used to differentiate the plot's trees, and calculate the number of sampled plots.
plot_area	Quoted name of the plot area variable, or a numeric vector with the plot area value. The plot area value must be in square meters.
vertical_est	Optional argument. Quoted name of the vertical strata variable, or the height variable. If this is a factor variable, it's levels will be used to classify the forest vertically. If it's a height variable, the vertical strata will be created based on it's mean and standard deviation values. Default: NA.
internal_est	Optional argument. Quoted name of the internal strata variable. Default: NA.
NI_label	Label used for Species not identified. This parameter works along with species. The level supplied here will not be considered in the classification. Default "".

Value

a data frame with the forest's structure.

Author(s)

Eric Bastos Gorgens <e.gorgens@gmail.com>

References

Souza, A. L. and Soares, C. P. B. (2013) Florestas Nativas: estrutura, dinamica e manejo. Vicosa: UFV.

Examples

```
library(forestmangr)
data("exfm20")
exfm20

# Get the forest's horizontal structure:
forest_structure(exfm20, "scientific.name", "dbh", "transect", 10000)

# area plot as a variable name:
forest_structure(exfm20, "scientific.name", "dbh", "transect", "plot.area")

# Get the forest's horizontal and vertical structure.
# The vertical structure variable can either be the height variable,
# or a factor variable with the horizontal strata:
forest_structure(exfm20, "scientific.name", "dbh", "transect", 10000, "canopy.pos")

# Get the forest's horizontal, vertical and internal structure:
forest_structure(exfm20, "scientific.name", "dbh", "transect", 10000, "canopy.pos", "light")
```

graybill_f

Graybill F Test

Description

Hypothesis test as described by Graybill (1976).

Usage

```
graybill_f(df, Y1, Yj, signif = 0.05, output = "simple")
```

Arguments

df	A data frame.
Y1	Quoted name of the standard variable.
Yj	Quoted name of the proposed variable.
signif	Numeric value for the significance level used in the test. Default: 0.05.
output	Defines the type of output. If "simple", a simple data frame is created, with only essential information about the test. If "table", more information is provided, and if "full", a data frame with informations about the test and both variables is created. Default: "simple".

Details

This test is used to compare two variables, usually a proposed method, and a standard variable. This test is popular among forestry engineers, specially because, since it considers all data in it's analysis, it's usually more precise than a standard mean t-test. If the data has outliers, the mean may not represent the data correctly, so Graybill F test is specially useful for heterogeneous data.

A simple model regression is applied, and it's significance is evaluated by applying Graybill F test for the parameters estimate, according to the methodology described by Graybill (1976).

Value

A data frame. Its dimensions will vary, according to the output argument.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

References

- Campos, J. C. C. and Leite, H. G. (2017) Mensuracao Florestal: Perguntas e Respostas. 5a. Vicosa: UFV.
- Graybill, F. A. (1976) Theory and application of the linear model. Massachusetts: Ouxburg 239 Press.
- Leite, H. G. and Oliveira, F. H. T. (2006) Statistical procedure to test identity between analytical methods, Communications in Soil Science and Plant Analysis, 33(7-8), pp. 1105-1118.

Examples

```
library(forestmangr)
data("exfm11")
exfm11

# The data frame exfm11 contains a height variable called "TH". This will be our
# standard value. We'll compare it to height estimated using different hypsometric equations.
# These are variables "TH_EST1" and "TH_EST2":
graybill_f( exfm11,"TH", "TH_EST1")

# TH_EST1 is statistically different from "TH".
```

```
# It's possible to alter the test's significance level using the signif argument:
graybill_f( exfm11,"TH", "TH_EST1", signif = 0.01)

# Different output options are available through the output argument:
graybill_f( exfm11,"TH", "TH_EST2", output="table")
graybill_f( exfm11,"TH", "TH_EST2", output="full")
```

guide_curve	<i>Get the guide curve plot for growth and yield analysis of inventory data</i>
-------------	---

Description

Get the guide curve for growth and yield analysis of inventory data using the factor method, and different statistical models.

Usage

```
guide_curve(
  df,
  dh,
  age,
  age_index,
  n_class = 4,
  model = "Schumacher",
  start_chap = c(b0 = 23, b1 = 0.03, b2 = 1.3),
  start_bailey = c(b0 = 3, b1 = -130, b2 = 1.5),
  round_classes = FALSE,
  font = "serif",
  gray_scale = TRUE,
  output = "plot"
)
```

Arguments

df	A data frame.
dh	Quoted name for the dominant height variable.
age	Quoted name for the age variable.
age_index	Numeric value for the age index.
n_class	Numeric value for the number of classes used to divide the data. Default 4.
model	model used to fit dh as a function of age. The models available are "Schumacher", "Curtis", "Chapman-Richards" and "Bailey-Clutter". Default: "Schumacher".
start_chap	Numeric vector with the start values for the Chapman-Richards model. This must be a named vector, with b0, b1 and b2 as parameter names. Default: c(b0=23,b1=0.03,b2 = 1.3).

start_bailey	Numeric vector with the start values for the Bailey-Clutter model. This must be a named vector, with b0, b1 and b2 as parameter names. Default: c(b0=3, b1=-130, b2 = 1.5).
round_classes	If TRUE, class values will be rounded to the nearest 5. Default TRUE.
font	Type of font used in the plot. Default: "serif".
gray_scale	If TRUE, the plot will be rendered in a gray scale. Default: "TRUE".
output	Type of output the function should return. This can either be "plot", for the guide curve plot, "table", for a data frame with the data used on the guide curve plot, and "full" for a list with 2 ggplot2 objects, one for residual plot and other for plot curves, a lm object for the regression, a data frame with quality of fit variables, the dominant height index, the class table used, and the table used for the guide curve plot. Default "plot".

Value

A data frame, a ggplot object, or a list, varying according to the "output" argument.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```
data("exfm14")
exfm14

# To get a guide curve plot for this data, we simply need to input
# dominant height and age variables, age index, and number of classes to be used:
guide_curve(exfm14, "dh", "age", 72, 5)

# if we want to get the table used to get the plot, we can choose the output "table":
guide_curve(exfm14, "dh", "age", 72, 5, output = "table")

# Other models are available for use, such as Curtis, Chapman Richards, and Bailey:
# CR and BC models are non linear, and thus need start values. There are default values,
# but they may fail, depending on the data used, so it's recommended to try start values that
# are ideal for the data used:
guide_curve(exfm14, "dh", "age", 72, 5,
  model = "Chapman-Richards", start_chap = c(b0=23, b1=0.03, b2 = 1.3))

# Or, to get more information on the analysis, such as details on the regression,
# bias, rmse, plot for residuals and more (cpu taxing):
## Not run:
guide_curve(exfm14, "dh", "age", 72, 5, output = "full")

## End(Not run)
```

`huberwb`*Calculate the volume with bark of trees using the Huber method*

Description

Function used to calculate the volume with bark of trees using the Huber method. This function has integration with `dplyr`, so it can be used inside a pipe, along with the `group_by` function.

Usage

```
huberwb(df, di, section_length, tree, .groups = NA, di_mm_to_cm = FALSE)
```

Arguments

<code>df</code>	A data frame.
<code>di</code>	Quoted name of the section diameter variable, in centimeters.
<code>section_length</code>	Quoted name of the section length variable, in meters
<code>tree</code>	Quoted name of the tree variable. used to differentiate the trees' sections. If this argument is <code>NA</code> , the defined groups in the data frame will be used. Default: <code>NA</code> .
<code>.groups</code>	Optional argument. Quoted name(s) of additional grouping variables that can be added to differentiate subdivisions of the data. If this argument is <code>NA</code> , the defined groups in the data frame will be used. Default: <code>NA</code> .
<code>di_mm_to_cm</code>	Boolean argument that, if <code>TRUE</code> , converts the <code>di</code> argument from milliliters to centimeters. Default: <code>FALSE</code> .

Value

Data frame with volume values by section.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

References

Campos, J. C. C. and Leite, H. G. (2017) Mensuracao Florestal: Perguntas e Respostas. 5a. Vicosa: UFV.

See Also

Complementary functions: [huberwob](#), For calculation of volume without bark using the Huber method, [smalianwb](#), for calculation of volume with bark using the Smalian method, [smalianwob](#), for calculation of volume without bark the Smalian method.

Examples

```

library(forestmangr)
data("exfm8")
exfm8

# Calculate the volume with bark using the Huber method:
huberwb(exfm8,"di_wb", "sec_length", "TREE")

# Using pipes:
library(dplyr)

exfm8 %>%
  group_by(TREE) %>%
  huberwb("di_wb", "sec_length")

```

huberwob

Calculate the volume without bark of trees using the Huber method

Description

Function used to calculate the volume without bark of trees using the Huber method. This function has integration without dplyr, so it can be used inside a pipe, along with the `group_by` function.

Usage

```

huberwob(
  df,
  di,
  section_length,
  bt,
  tree,
  .groups = NA,
  di_mm_to_cm = FALSE,
  bt_mm_to_cm = FALSE
)

```

Arguments

<code>df</code>	A data frame.
<code>di</code>	Quoted name of the section diameter variable, in centimeters.
<code>section_length</code>	Quoted name of the section length variable, in meters
<code>bt</code>	Quoted name of the bark thickness variable, in centimeters.
<code>tree</code>	Quoted name of the tree variable. used to differentiate the trees' sections. If this argument is <code>NA</code> , the defined groups in the data frame will be used. Default: <code>NA</code> .

<code>.groups</code>	Optional argument. Quoted name(s) of additional grouping variables that can be added to differentiate subdivisions of the data. If this argument is NA, the defined groups in the data frame will be used. Default: NA.
<code>di_mm_to_cm</code>	Boolean argument that, if TRUE, converts the di argument from milliliters to centimeters. Default: FALSE.
<code>bt_mm_to_cm</code>	Boolean argument that, if TRUE, converts the bt argument from milliliters to centimeters. Default: FALSE.

Value

Data frame with volume values by section.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

References

Campos, J. C. C. and Leite, H. G. (2017) Mensuracao Florestal: Perguntas e Respostas. 5a. Vicosa: UFV.

See Also

Complementary functions: [huberwb](#), For calculation of volume with bark using the Huber method, [smalianwb](#), for calculation of volume with bark using the Smalian method, [smalianwob](#), for calculation of volume without bark the Smalian method.

Examples

```
library(forestmangr)
data("exfm8")
exfm8

# Calculate the volume without bark using the Huber method:
huberwob(exfm8,"di_wb", "sec_length", "bark_t", "TREE")

# Using pipes:
library(dplyr)

exfm8 %>%
  group_by(TREE) %>%
  huberwob("di_wb", "sec_length", "bark_t")
```

ident_model	<i>Identity of a Model Test</i>
-------------	---------------------------------

Description

Function for using the Identity of a Model test, as described by Regazzi (1999).

Usage

```
ident_model(
  df,
  factor,
  reduced_model,
  filter = NA,
  gray_scale = TRUE,
  signif = 0.05,
  font = "serif",
  output = "table"
)
```

Arguments

<code>df</code>	a data frame.
<code>factor</code>	Quoted name of the factor variable used to differentiate the data projects in the test.
<code>reduced_model</code>	Quoted or unquoted reduced model used in the test. The variables mentioned in the model must exist in the provided data frame. X and Y sides of the model must be separated by "~".
<code>filter</code>	Optional argument. If supplied with levels present in <code>factor</code> , only these levels will be used in the test. NA.
<code>gray_scale</code>	If TRUE a gray scale will be used in the plots. Default: FALSE.
<code>signif</code>	Numeric value for the significance level used in the test. Default: 0.05.
<code>font</code>	font family used in the plots. Can be either "serif" for Times New Roman or "sans" for arial unicode MS. Default: "serif".
<code>output</code>	Defines the type of output. If "table" an anova table with the identity of model test is provided, if "plot" a ggplot plot/object representing the test is created, if "table_plot", both anova table and plot are provided, and if "full", a list is provided, with details on the dummies variables created, the reduced and complete models, the anova table and the plot. Default: "table"

Value

A data frame, a ggplot object, or a list, varying according to the output argument.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Marcio leles Romarco de Oliveira <marcioromarco@gmail.com>

References

Regazzi, A. J. (1999) Teste para verificar a identidade de modelos de regressao e a igualdade de parametros no caso de dados de delineamentos experimentais, Ceres, 46(266), pp. 383–409.

Examples

```
library(forestmangr)
data("exfm13")
exfm13

# The objective is to know if the diameter's behavior is similar among 3 species.
# For this we'll use a quadratic model. We'll use nitrogen (N) as our X variable.

ident_model(exfm13, "species", dbh ~ N + N2)

# This test shows that there are differences between the species.
# We can get more details on this using a different output, that will also
# give us a plot:

ident_model(exfm13, "species", dbh ~ N + N2, output = "table_plot")

# This gives us only the plot:
ident_model(exfm13, "species", dbh ~ N + N2, output = "table_plot")

# And this gives us additional information on the test:
ident_model(exfm13, "species", dbh ~ N + N2, output = "full")

# Looking at the plot, it seems that 2 species are behaving very similar, while
# the Pequi species is different from the other 2. We can confirm this by running
# the test in a paired fashion, using the filter argument:

ident_model(exfm13, "species", dbh ~ N + N2,
  filter = c("PEQUI", "SUCUPIRA-PRETA"), output = "table_plot")

ident_model(exfm13, "species", dbh ~ N + N2,
  filter = c("PEQUI", "VINHATICO"), output = "table_plot")

ident_model(exfm13, "species", dbh ~ N + N2,
  filter = c("SUCUPIRA-PRETA", "VINHATICO"), output = "table_plot")

# As we imagined, a single model can be used to describe the behavior of
# the "Sucupira-preta" and "Vinhatico" species,
# and a second model is needed to explain the Pequi Variable.

# It's possible to apply a color scale to the plots, and also change it's font to arial:
```

```
ident_model(exfm13, "species", dbh ~ N + N2,output="plot",gray_scale=FALSE,font="sans")
```

inv

Calculate the inverse of a number

Description

This function returns the inverse of a numeric vector.

Usage

```
inv(x)
```

Arguments

x A numeric vector

Details

This function is mainly used when fitting statistical models. If one of the variables in a model is an inverse of a vector, the `lm` function does not properly compute the variable, if `1/vector` is inserted directly in the model, leading to the need of creating a separate variable. This function allows the user to get the inverse of a given numeric vector inside the model, without the need to create a new variable.

Value

a numeric vector containing the inverse of `x`.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```
library(forestmangr)
data("exfm15")
exfm15

# Get the inverse of a vector
inv(iris$Petal.Length)

# Fit a model that contains the inverse of a variable, without the need to
# create a new variable for the inverse:
lm(log(TH) ~ inv(DBH), exfm15 )
# or
lm_table(exfm15, log(TH) ~ inv(DBH) )
```

lm_resid	<i>Fit linear regressions by group, with the option of removing outliers using a interactive plot of residuals.</i>
----------	---

Description

With this function it's possible to fit linear regressions by a grouping variable, and evaluate each equation via a interactive plot of residuals, and get a data frame. with each column as a coefficient and quality of fit variables, and other output options. Works with dplyr grouping functions.

Usage

```
lm_resid(df, model, output_mode = "table", est.name = "est", group_print = NA)
```

Arguments

df	A data frame.
model	A linear regression model, with or without quotes. The variables mentioned in the model must exist in the provided data frame. X and Y sides of the model must be separated by "~".
output_mode	Selects different output options. Can be either "table", "merge", "merge_est" and "nest". See details for explanations for each option. Default: "table".
est.name	Name of the estimated y value. Used only if est.name = TRUE. Default: "est".
group_print	This argument is only used internally by another function. Please ignore.

Details

this function uses lm_table as a basis, but calls a plot of residuals for each fitted model, for the user to evaluate. If one decides to remove any of the points, one can click and drag, and then click on the 'remove points' button. After that, one must simply click 'done' and the coefficients will be printed out.

It's possible to use the output argument to get a merged table if output="merge", that binds the original data frame and the fitted coefficients. If output="merge_est" we get a merged table as well, but with y estimated using the coefficients. If the fit is made using groups, this is taken into account, i.e. the estimation is made by group.

If output="nest", a data frame with nested columns is provided. This can be used if the user desires to get a customized output.

Value

A data frame. Different data frame options are available using the output argument.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```

if (interactive() ){
  library(forestmangr)
  library(dplyr)

  data("exfm19")

  # Fit SH model:
  lm_resid_group(exfm19, log(VWB) ~ log(DBH) + log(TH))
}

```

lm_resid_group	<i>Fit linear regressions by group, with the option of removing outliers using a interactive plot of residuals.</i>
----------------	---

Description

With this function it's possible to fit linear regressions by a grouping variable, and evaluate each equation via a interactive plot of residuals, and get a data frame. with each column as a coefficient and quality of fit variables, and other output options. Works with dplyr grouping functions.

Usage

```
lm_resid_group(df, model, .groups, output_mode = "table", est.name = "est")
```

Arguments

df	A data frame.
model	A linear regression model, with or without quotes. The variables mentioned in the model must exist in the provided data frame. X and Y sides of the model must be separated by "~".
.groups	Quoted name(s) of grouping variables used to fit multiple regressions, one for each level of the provided variable(s). Default NA.
output_mode	Selects different output options. Can be either "table", "merge", "merge_est" and "nest". See details for explanations for each option. Default: "table".
est.name	Name of the estimated y value. Used only if est.name = TRUE. Default: "est".

Details

this function uses lm_table as a basis, but calls a plot of residuals for each fitted model, for the user to evaluate. If one decides to remove any of the points, one can click and drag, and then click on the 'remove points' button. After that, one must simply click 'done' and the coefficients will be printed out.

It's possible to use the output argument to get a merged table if output="merge", that binds the original data frame and the fitted coefficients. If output="merge_est" we get a merged table as

well, but with y estimated using the coefficients. If the fit is made using groups, this is taken into account, i.e. the estimation is made by group.

If `output="nest"`, a data frame with nested columns is provided. This can be used if the user desires to get a customized output.

Value

A data frame. Different data frame options are available using the `output` argument.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```
if (interactive() ){
  library(forestmangr)
  library(dplyr)

  data("exfm19")

  # Fit SH model by group:
  lm_resid_group(exfm19, log(VWB) ~ log(DBH) + log(TH), "STRATA")
}
```

lm_table

Fit linear regressions by group, and get different output options.

Description

With this function it's possible to fit linear regressions by a grouping variable, and get a data frame with each column as a coefficient and quality of fit variables, and other output options. Works with `dplyr` grouping functions.

Usage

```
lm_table(
  df,
  model,
  .groups = NA,
  output = "table",
  est.name = "est",
  keep_model = FALSE
)
```

Arguments

df	A data frame.
model	A linear regression model, with or without quotes. The variables mentioned in the model must exist in the provided data frame. X and Y sides of the model must be separated by "~".
.groups	Optional argument. Quoted name(s) of grouping variables used to fit multiple regressions, one for each level of the provided variable(s). Default NA.
output	Selects different output options. Can be either "table", "merge", "merge_est" and "nest". See details for explanations for each option. Default: "table".
est.name	Name of the estimated y value. Used only if est.name = TRUE. Default: "est".
keep_model	If TRUE, a column containing lm object(s) is kept in the output. Useful if the user desires to get more information on the regression. Default: FALSE.

Details

With this function there's no more need to use the do function when fitting a linear regression in a pipe line. It's also possible to easily make fit multiple regressions, specifying a grouping variable. In addition to that, the default output sets each coefficient as a column, making it easy to call coefficients by name or position when estimating values.

It's possible to use the output argument to get a merged table if output="merge", that binds the original data frame and the fitted coefficients. If output="merge_est" we get a merged table as well, but with y estimated using the coefficients. If the fit is made using groups, this is taken into account, i.e. the estimation is made by group.

If output="nest", a data frame with nested columns is provided. This can be used if the user desires to get a customized output.

Value

A data frame. Different data frame options are available using the output argument.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```
library(forestmangr)
library(dplyr)

data("exfm19")
exfm19

# Fit Schumacher and Hall model for volume estimation, and get
# coefficient, R2 and error values:

lm_table(exfm19, log(VWB) ~ log(DBH) + log(TH))

# Fit SH model by group:
```

```

lm_table(exfm19, log(VWB) ~ log(DBH) + log(TH), "STRATA")

# This can also be done using dplyr::group_by:
exfm19 %>%
  group_by(STRATA) %>%
  lm_table(log(VWB) ~ log(DBH) + log(TH) )

# It's possible to merge the original data with the table containing the coefficients
# using the output parameter:
fit <- lm_table(exfm19, log(VWB) ~ log(DBH) + log(TH), "STRATA", output = "merge")
head(fit)

# It's possible to merge the original data with the table,
# and get the estimated values for this model:
fit <- lm_table(exfm19, log(VWB) ~ log(DBH) + log(TH), "STRATA",
  output = "merge_est", est.name = "VWB_EST")
head(fit)

# It's possible to further customize the output,
# unnesting the nested variables provided when output is defined as "nest":
lm_table(exfm19, log(VWB) ~ log(DBH) + log(TH), "STRATA", output = "nest")

```

nls_table

Fit non-linear regressions by group, using LM algorithm and get different output options.

Description

With this function it's possible to fit non-linear regressions using Levenberg-Marquardt or Gauss-Newton algorithms by a grouping variable, and get a data frame with each column as a coefficient and quality of fit variables, and other output options. Works with dplyr grouping functions.

Usage

```

nls_table(
  df,
  model,
  mod_start,
  .groups = NA,
  output = "table",
  est.name = "est",
  replace = FALSE,
  keep_model = FALSE,
  global_start,
  algorithm = "LM"
)

```


Arguments

df	A data frame.
model	A linear regression model, with or without quotes. The variables mentioned in the model must exist in the provided data frame. X and Y sides of the model must be separated by "~".
mod_start	A vector or data frame, with start values for coefficients used in the model. This can be a data frame containing the same group variables used in the .groups argument, and the start values.
.groups	Optional argument. Quoted name(s) of grouping variables used to fit multiple regressions, one for each level of the provided variable(s). Default NA.
output	Selects different output options. Can be either "table", "merge", "merge_est" and "nest". See details for explanations for each option. Default: "table".
est.name	Name of the estimated y value. Used only if est.name = TRUE. Default: "est".
replace	If TRUE, models that don't converge on a grouped regression fit will be replaced by coefficients fitted using all data. Default: FALSE.
keep_model	If TRUE, a column containing lm object(s) is kept in the output. Useful if the user desires to get more information on the regression. Default: FALSE.
global_start	Optional argument. A vector or data frame, with start values for the global fit regression used when "replace" is TRUE.
algorithm	Algorithm to be used in the non-linear regression. It can be "LM" (Levenberg-Marquardt, more robust) or "GN" (Gauss-Newton, less robust, uses nls default algorithm). Default: "LM".

Details

This function Levenberg-Marquardt algorithm as default for fitting non-linear regression models. Also, with this function there no more need to use the do function when fitting a linear regression in a pipe line. It's also possible to easily make fit multiple regressions, specifying a grouping variable. In addition to that, the default output sets each coefficient as a column, making it easy to call coefficients by name or position when estimating values. The Levenberg-Marquardt fit uses [nlsLM](#).

Value

A data frame. Different data frame options are available using the output argument.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```
library(forestmangr)
library(dplyr)
data("exfm14")
exfm14
```

```

# Fit Chapman & Richards non-linear model for dominant Height:
nls_table(exfm14, dh ~ b0 * (1 - exp( -b1 * age ) )^b2,
          mod_start = c( b0=23, b1=0.03, b2 = 1.3 ) )

# Fit CR model by strata:
nls_table(exfm14,dh ~ b0 * (1 - exp( -b1 * age ) )^b2,
          mod_start = c( b0=23, b1=0.03, b2 = 1.3 ) ,
          .groups = "strata") %>%
  as.data.frame

# or, using group_by

exfm14 %>%
  group_by(strata) %>%
  nls_table(dh ~ b0 * (1 - exp( -b1 * age ) )^b2,
            mod_start = c( b0=23, b1=0.03, b2 = 1.3 ) )

# If there are multiple start values, for each strata, they can be supplied like so:
tab_coef <- data.frame(strata = c(1:20, 24,25,27,28,30,31,33,35,36,37),
                       rbind(
                         data.frame(b0 = rep(23, 20), b1 = rep(0.03, 20), b2 = rep(1.3, 20) ),
                         data.frame(b0 = rep(23, 10), b1 = rep(0.03, 10), b2 = rep(.5, 10) )))

tab_coef

nls_table(exfm14, dh ~ b0 * (1 - exp( -b1 * age ) )^b2,
          mod_start = tab_coef,
          .groups = "strata" )
# mod_start needs to be a data frame in this case.

# It's possible to bind the coefficients to the original data,
# to estimate y. We'll also estimate bias and rmse for this estimation.

# This can also be done directly using "merge_est" as output:
nls_table(exfm14,dh ~ b0 * (1 - exp( -b1 * age ) )^b2,
          mod_start = tab_coef ,
          .groups = "strata",
          output = "merge_est",
          est.name = "dh_est" ) %>%
  mutate(
    bias = bias_per(y = dh, yhat = dh_est),
    rmse = rmse_per(y = dh, yhat = dh_est) ) %>%
  head(15)

# It's possible to further customize the output, using nested columns:
nls_table(exfm14,dh ~ b0 * (1 - exp( -b1 * age ) )^b2,
          mod_start = tab_coef ,
          .groups = "strata",
          output = "nest" )

# It's possible to use Gauss-Newton's algorithm. In this case,
# some regressions will not converge.
exfm14 %>%

```

```

group_by(strata) %>%
nls_table(dh ~ b0 * (1 - exp( -b1 * age ) )^b2,
          mod_start = c( b0=23, b1=0.03, b2 = 1.3 ),algorithm="GN" )

# If some regressions don't converge, it's possible to fill those NAs with
# regression coefficients from a general fit, using the entire data:
nls_table(exfm14,dh ~ b0 * (1 - exp( -b1 * age ) )^b2,
          mod_start = c( b0=23, b1=0.03, b2 = 1.3 ),
          .groups = "strata",
          replace = TRUE,
          algorithm="GN" )

```

npv_irr

Calculate Net Present Value and other economic variables

Description

Get the net present value, internal rate of return, and other economic variables, given cost and revenue values.

Usage

```

npv_irr(
  df,
  year,
  cost,
  revenue,
  rate,
  output = "full",
  sens_limits = c(1, 30),
  big_mark = ",",
  dec_mark = ".",
  prefix = "$"
)

```

Arguments

df	A data frame.
year	Quoted name of the year variable.
cost	Quoted name of the costs variable.
revenue	Quoted name of the revenue variable.
rate	Numeric value of the yearly rate to be used.
output	Selects different output options. It can be either "full" for a list containing a sensibility plot and a data frame with a single observation and one column for each variable, or "simple" for a two column data frame with one observation for each calculated variable. Default: "simple".

sens_limits	Selects the rate range used in the sensibility plot. This is a numeric vector with two elements, the initial and final rate to be used as range. These can vary between 0 and 100. Default: c(1, 30).
big_mark	Selects thousands separator. Can be either ".", " " or ", ". Default: " , ".
dec_mark	Selects decimal separator. Can be either " , " or " . " . Default: " . " .
prefix	selects the prefix for the y axis in the sensibility plot. Can be either "\$" or "R\$". Default: "\$".

Value

A data frame, or a list, according to output.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```
## Not run:
library(forestmangr)
data(exfm22)

npv_irr(exfm22, "year", "cost", "revenue", rate=8.75)

# To also get a sensibility plot, use

npv_irr(exfm22, "year", "cost", "revenue", rate=8.75, output="full")

## End(Not run)
```

plot_summarise	<i>Summarize forest inventory data</i>
----------------	--

Description

Get informations about forest inventory plots, like number of individuals, mean DBH, q, height, basal area, volume, etc.

Usage

```
plot_summarise(
  df,
  plot,
  plot_area,
  dbh,
  th,
  .groups,
  total_area,
```

```

    vwb,
    vwob,
    dh,
    age,
    dec_places = 4
  )

```

Arguments

df	A data frame.
plot	Quoted name of the plot variable. used to differentiate the data's plots. If this argument is missing, the defined groups in the data frame will be used, If there are no groups in the data, the function will fail.
plot_area	Quoted name of the plot area variable, or a numeric vector with the plot area value. The plot area value must be in square meters.
dbh	Optional parameter. Quoted name of the diameter at breast height variable. If supplied, will be used to calculate the mean diameter per plot, quadratic diameter (q), basal area and basal area per hectare. Default NA.
th	Optional parameter. Quoted name of the total height variable. If supplied, will be used to calculate the mean total height, and the dominant height variable, if the dh is NA. Default NA.
.groups	Optional argument. Quoted name(s) of grouping variables that can be added to differentiate subdivisions of the data. Default: NA.
total_area	Optional argument. Quoted name of the total area variable, or a numeric vector with the total area value. The total area value must be in hectares. Default: NA.
vwb	Optional parameter. Quoted name of the volume with bark variable. If supplied, will be used to calculate the total vwb per plot, and vwb per hectare per plot. Default NA.
vwob	Optional parameter. Quoted name of the volume without bark variable. If supplied, will be used to calculate the total vwob per plot, and vwob per hectare per plot. Default NA.
dh	Optional parameter. Quoted name of the dominant height variable. If supplied, will be used to calculate the mean dominant height per plot. If not, the ht variable supplied will be used to calculate the average of the top two trees of each plot, and use that as dh. Default: NA.
age	Optional parameter. Quoted name of the age variable. If supplied, will be used to calculate the average age per plot. Default: NA.
dec_places	Numeric value for the number of decimal places to be used in the output tables. Default: 4.

Value

A data frame with informations per plot.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```

library(forestmangr)
data("exfm21")
exfm21

# Obligatory arguments. Basic informations about the plot.
plot_summarise(exfm21, "PLOT", 810)

# Area values can be numeric, or a variable name
plot_summarise(exfm21, "PLOT", "PLOT_AREA")

# With DBH supplied, we get the mean diameter, quadratic diameter,
# basal area and basal area per hectare:
plot_summarise(exfm21, "PLOT", "PLOT_AREA", "DBH")

# With TH supplied, we get the mean total height and dominant height
plot_summarise(exfm21, "PLOT", "PLOT_AREA", "DBH", "TH_EST")

# With strata supplied, we divide the data into 2 strata
plot_summarise(exfm21, "PLOT", "PLOT_AREA", "DBH", "TH_EST", "STRATA")

# The strata area can also be supplied
plot_summarise(exfm21, "PLOT", "PLOT_AREA", "DBH", "TH_EST", "STRATA", "STRATA_AREA")

# With VWB supplied, we get the total vwb, and vwb per hectare
plot_summarise(exfm21, "PLOT", "PLOT_AREA", "DBH", "TH_EST", "STRATA", "STRATA_AREA",
  "VWB")

# With VWOB supplied, we get the total vwob, and vwob per hectare
plot_summarise(exfm21, "PLOT", "PLOT_AREA", "DBH", "TH_EST", "STRATA", "STRATA_AREA",
  "VWB", "VWOB")

# If the data already has a dominant height variable, it can also be supplied here
plot_summarise(exfm21, "PLOT", "PLOT_AREA", "DBH", "TH_EST", "STRATA", "STRATA_AREA",
  "VWB", "VWOB", "DH")

# With the AGE variable supplied, we get the average age of each plot
plot_summarise(exfm21, "PLOT", "PLOT_AREA", "DBH", "TH_EST", "STRATA", "STRATA_AREA",
  "VWB", "VWOB", "DH", "AGE")

```

pow

Raise a numeric vector to a given power

Description

This function returns a numeric vector raised to a given power.

Usage

```
pow(x, y)
```

Arguments

- x A numeric vector.
- y A numeric value for the power x should be raised to.

Details

This function is mainly used when fitting statistical models. If one of the variables in a model is a variable raised to a given power, the `lm` function does not properly compute the variable, if `vector^power` is inserted directly in the model, leading to the need of creating a separate variable. This function allows the user to get the power of a given numeric vector to `y` inside the model, without the need to create a new variable.

Value

a numeric vector containing `x` to the power of `y`.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```
library(forestmangr)
data("exfm15")
head(exfm15)

# Raise a numeric vector to the power of 2:
pow(iris$Petal.Length, 2)

# Fit a model that contains the dbh squared, without the need to create a new variable:
lm(log(TH) ~ DBH + pow(DBH,2), exfm15 )
# or
lm_table(exfm15, log(TH) ~ DBH + pow(DBH,2) )
```

resid_plot

Calculate residual values and create plots

Description

Function for creating plots and tables for residual values from observed and estimated values.

Usage

```
resid_plot(
  df,
  obs,
  ...,
  type = "scatterplot",
  point_size = 3,
  color = NA,
  nrow = NA,
  ncol = NA,
  lim_y = NA,
  xlab = "Observed values",
  clab = NA,
  font = "serif",
  legend_pos = "bottom",
  gray_scale = TRUE,
  res_table = FALSE
)
```

Arguments

df	A data frame.
obs	Quoted name of the observed values variable.
...	Quoted name(s) for the estimated values variable(s). Multiple variables must be separated by comma.
type	Character object for the type of plot created. The available plots are: "scatterplot", "histogram", "histogram_curve" and "versus". Default: "scatterplot".
point_size	Numeric value for the point size in scatter plots. Default: 3.
color	Quoted name of a variable. If supplied, this variable will be used to classify the data by color. Default: NA.
nrow	Numeric value for number of rows in the plot matrix. If not supplied, the plots will be automatically sorted. Default: NA.
ncol	Numeric value for number of columns in the plot matrix. If not supplied, the plots will be automatically sorted. Default: NA.
lim_y	Numeric value for the y axis upper and lower limit. If NA, the biggest residual value is used. Default: NA.
xlab	Character value for the x label used in some plots. Default: "Observed values".
clab	Character value for the color label used, if a color variable is supplied. If not supplied, the color variable name will be used. Default: NA.
font	Type of font used in the plot. Default: "serif".
legend_pos	Position of legend, when a color variable is supplied. This can either be "left", "right", "top" or "bottom". Default: "bottom".
gray_scale	If TRUE, the plot will be rendered in a gray scale. Default: "TRUE".
res_table	If TRUE, the function will return a data frame with observed, estimated, and residual values. Default: FALSE.

Value

A ggplot object, or, if `res_table = TRUE`, a data frame.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```
library(forestmangr)
data("exfm11")
exfm11

# Specifying the observed and estimated variables, we get a scatter plot
# for the percentage residuals:
resid_plot(exfm11, "TH", "TH_EST1")

# It's possible to get other types of plots, with the type argument:
resid_plot(exfm11, "TH", "TH_EST1", type = "histogram_curve")
resid_plot(exfm11, "TH", "TH_EST1", type = "versus")

# It's possible to add a factor variable as color in the plots:
resid_plot(exfm11, "TH", "TH_EST1", "TH_EST2", color="STRATA",
xlab="Total Height (m)", clab="Strata", gray_scale=FALSE)

# If there are more estimated values variables, they can also be used
# in the comparison:
resid_plot(exfm11, "TH", "TH_EST1", "TH_EST2", "TH_EST3")

# It's possible to rearrange the plots with ncol and nrow:
resid_plot(exfm11, "TH", "TH_EST1", "TH_EST2", "TH_EST3", ncol=1)

# It's possible to get the residuals table used to generate these plots, with res_table=TRUE:
head( resid_plot(exfm11, "TH", "TH_EST1", "TH_EST2", res_table = TRUE) )
```

rmse_per

RMSE of an estimator in percentage

Description

Function for calculating the Root-Mean-Square-Error of an estimator.

Usage

```
rmse_per(df, y, yhat, na.rm = TRUE)
```

Arguments

df	a data frame.
y	Quoted name of the variable representing the observed values in the data frame. If a data frame is not provided, y can also be a numeric vector.
yhat	Quoted name of the variable representing the estimated values in the data frame. If a data frame is not provided, yhat can also be a numeric vector.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds. default: TRUE

Details

Function for calculating the Root-Mean-Square-Error of an estimator, given the observed values, and the estimated values.

Value

Numeric vector with the RMSE value, in percentage.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

See Also

other statistics to evaluate estimators: [bias_per](#) for the bias of an estimator

Examples

```
library(forestmangr)
data("exfm11")
exfm11

# RMSE of an estimator, given the data frame and quoted variable names:
rmse_per(exfm11, "TH", "TH_EST3")

# RMSE of an estimator, given the vectors for observed and estimated values:
rmse_per(y = exfm11$TH, yhat = exfm11$TH_EST3)
```

rm_empty_col

Remove empty columns

Description

This function removes columns filled with NA or 0 from a dataframe.

Usage

```
rm_empty_col(x)
```

Arguments

x A dataframe

Details

This function is mainly used inside other functions, to remove optional variables when they are not opted in.

Value

a dataframe.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```
library(forestmangr)
library(dplyr)
data("exfm15")
head(exfm15)

exfm15 %>%
  mutate(emptycol=NA) %>%
  rm_empty_col
```

round_df

Round all numeric variables of a data frame to a given digit

Description

This function allows the user to round all numeric values of a data frame, directly, even if the data frame contains non-numeric variables (which would throw an error in the [round](#) function).

Usage

```
round_df(df, digits, rf = "round")
```

Arguments

df	A data frame.
digits	Numeric vector for the desired number of digits.
rf	Type of round to be used. It can either be "ceiling", "floor", "trunc", "signif", or "round". Default "round".

Value

A data frame, with all the numeric variables rounded up to the number given to digits.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

Examples

```
library(forestmangr)

# Round all numeric variables
round_df(iris)

# Round all numeric variables using the floor function
round_df(iris, rf="floor")

# Do not run
# trying this with the the base function throws an error:

# round(iris)
```

similarity_matrix *Get the similarity matrix of an area*

Description

Calculates the Jaccard similarity index and Sorensen similarity index.

Usage

```
similarity_matrix(
  df,
  species,
  comparison,
  NI_label = "",
  index = "Sorensen",
  dendrogram = FALSE,
  n_groups = 3
)
```

Arguments

df	A data frame.
species	Quoted name of the scientific names variable, or any variable used to differentiate the different species found in data. If supplied, will be used to classify the species in the diameter data.
comparison	Quoted name of the variable containing levels to be compared with each other.
NI_label	Label used for Species not identified. This parameter works along with species. The level supplied here will not be considered in the classification. Default "".
index	Character value for the desired index to be used. Can be either "Jaccard", for a matrix based on the Jaccard index of similarity, "Sorensen", for a matrix based the Sorensen index of similarity, or "all", for a list with matrices for both indexes. Default: "Sorensen".
dendrogram	If TRUE, a dendrogram will also be created. Default: FALSE.
n_groups	Number of groups in the dendrogram. Default 3.

Value

a matrix object with a similarity matrix, or a list, according to the "index" and "dendrogram" arguments.

Author(s)

Eric Bastos Gorgens <e.gorgens@gmail.com>

References

Souza, A. L. and Soares, C. P. B. (2013) Florestas Nativas: estrutura, dinamica e manejo. Vicosa: UFV.

Examples

```
library(forestmangr)
data("exfm20")
exfm20

# To get the similarity matrix of an area, we simply need to provide
# the species variable name, and a subdivision variable name, like
# transect. By default we get a a matrix based on the Sorensen index:
similarity_matrix(exfm20, "scientific.name", "transect")

# To get the similarity matrix of Jaccard, use the index argument:
similarity_matrix(exfm20, "scientific.name", "transect", index = "Jaccard")

# To get a dendrogram with the matrix, use dendrogram=TRUE:
similarity_matrix(exfm20, "scientific.name", "transect", index = "Jaccard", dendrogram = TRUE)

# To get a list with both matrices, use index="all":
similarity_matrix(exfm20, "scientific.name", "transect", index = "all")
```

```
# If the data supplied only has 2 levels, a paired comparison is made instead:
ex_pair <- exfm20[exfm20$transect %in% c("T01", "T02") , ]
ex_pair

similarity_matrix(ex_pair, "scientific.name", "transect", index = "all")
```

smalianwb

Calculate the volume with bark of trees using the Smalian method

Description

Function used to calculate the volume with bark of trees using the Smalian method. This function has integration with dplyr, so it can be used inside a pipe, along with the group_by function.

Usage

```
smalianwb(
  df,
  di,
  hi,
  tree,
  .groups = NA,
  di_mm_to_cm = FALSE,
  hi_cm_to_m = FALSE
)
```

Arguments

df	A data frame.
di	Quoted name of the section diameter variable, in centimeters.
hi	Quoted name of the section height variable, in meters
tree	Quoted name of the tree variable. used to differentiate the trees' sections. If this argument is NA, the defined groups in the data frame will be used
.groups	Optional argument. Quoted name(s) of additional grouping variables that can be added to differentiate subdivisions of the data. If this argument is not supplied, the defined groups in the data frame will be used. Default: NA.
di_mm_to_cm	Boolean argument that, if TRUE, converts the di argument from milliliters to centimeters. Default: FALSE.
hi_cm_to_m	Boolean argument that, if TRUE, converts the hi argument from centimeters to meters. Default: FALSE.

Value

Data frame with volume values by section.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

References

Campos, J. C. C. and Leite, H. G. (2017) Mensuracao Florestal: Perguntas e Respostas. 5a. Vicosa: UFV.

See Also

Complementary functions: [smalianwob](#), For calculation of volume without bark using the Smalian method, [huberwb](#), for calculation of volume with bark using the Huber method, [huberwob](#), for calculation of volume without bark the Huber method.

Examples

```
library(forestmangr)
data("exfm7")
exfm7

# Calculate the volume with bark using the Smalian method:
smalianwb(exfm7,"di_wb", "hi", "TREE")

# Using pipes:
library(dplyr)

exfm7 %>%
  group_by(TREE) %>%
  smalianwb("di_wb", "hi")
```

smalianwob

Calculate the volume without bark of trees using the Smalian method

Description

Function used to calculate the volume without bark of trees using the Smalian method. This function has integration with dplyr, so it can be used inside a pipe, along with the `group_by` function.

Usage

```
smalianwob(
  df,
  di,
  hi,
  bt,
  tree,
  .groups = NA,
```

```

    di_mm_to_cm = FALSE,
    hi_cm_to_m = FALSE,
    bt_mm_to_cm = FALSE
  )

```

Arguments

df	A data frame.
di	Quoted name of the section diameter variable, in centimeters.
hi	Quoted name of the section height variable, in meters
bt	Quoted name of the bark thickness variable, in centimeters.
tree	Quoted name of the tree variable. used to differentiate the trees' sections. If this argument is NA, the defined groups in the data frame will be used. Default: NA.
.groups	Optional argument. Quoted name(s) of additional grouping variables that can be added to differentiate subdivisions of the data. If this argument is NA, the defined groups in the data frame will be used. Default: NA.
di_mm_to_cm	Boolean argument that, if TRUE, converts the di argument from milliliters to centimeters. Default: FALSE.
hi_cm_to_m	Boolean argument that, if TRUE, converts the hi argument from centimeters to meters. Default: FALSE.
bt_mm_to_cm	Boolean argument that, if TRUE, converts the bt argument from milliliters to centimeters. Default: FALSE.

Value

Data frame with volume values by section.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

References

Campos, J. C. C. and Leite, H. G. (2017) Mensuracao Florestal: Perguntas e Respostas. 5a. Vicosia: UFV.

See Also

Complementary functions: [smalianwb](#), For calculation of volume with bark using the Smalian method, [huberwb](#), for calculation of volume with bark using the Huber method, [huberwob](#), for calculation of volume without bark the Huber method.

Examples

```

library(forestmangr)
data("exfm7")
exfm7

# Calculate the volume without bark using Smalian's method:
smalianwob(exfm7,"di_wb", "hi", "bark_t", "TREE")

# Using pipes:
library(dplyr)

exfm7 %>%
  group_by(TREE) %>%
  smalianwob("di_wb", "hi", "bark_t")

```

species_aggreg

Get the aggregation state of species

Description

Get the aggregation state of species according to the Payandeh, Hazen and Morista methods.

Usage

```
species_aggreg(df, species, plot, NI_label = "")
```

Arguments

df	A data frame.
species	Quoted name of the scientific names variable, or any variable used to differentiate the different species found in data.
plot	Quoted name of the plot variable. used to differentiate the plots trees, and calculate the number of sampled plots.
NI_label	Label used for Species not identified. This parameter works along with species. The level supplied here will not be considered in the classification. Default "".

Value

a data frame with the aggregation classification.

Author(s)

Eric Bastos Gorgens <e.gorgens@gmail.com>

References

Souza, A. L. and Soares, C. P. B. (2013) Florestas Nativas: estrutura, dinamica e manejo. Vicosa: UFV.

Examples

```
library(forestmangr)
data("exfm20")
exfm20

# Get the aggregation indexes of species:
species_aggreg(exfm20, "scientific.name", "transect")
```

species_diversity *Get the species diversity indexes*

Description

Calculate the diversity of species for the following indexes: Shannon, Simpson, Equitability, Pielou and Jentsch.

Usage

```
species_diversity(df, species, plot = NA, NI_label = "", index = "all")
```

Arguments

df	A data frame.
species	Quoted name of the scientific names variable, or any variable used to differentiate the different species found in data. If supplied, will be used to classify the species in the diameter data.
plot	Optional parameter. Quoted name of the plot variable. used to differentiate the plots, and calculate the indexes by plot, or other subdivision variable.
NI_label	Label used for Species not identified. This parameter works along with species. The level supplied here will not be considered in the classification. Default "".
index	Character value for the desired index to be used. Can be either "H" for Shannon's diversity index, "S" for Total number of species in the community, "Hmax" for the maximum equitability, "J" for Pielou evenness, "QM" for the mixture coefficient of Jentsch, or "all", to get all indexes. Default: "all".

Value

a data frame with the indexes, or a numeric value of the desired index specified in the index argument.

Author(s)

Eric Bastos Gorgens <e.gorgens@gmail.com>

References

Souza, A. L. and Soares, C. P. B. (2013) Florestas Nativas: estrutura, dinamica e manejo. Vicosa: UFV.

Examples

```
library(forestmangr)
data("exfm20")
exfm20

# By default, the function returns all indexes:
species_diversity(exfm20, "scientific.name")

# It's possible to use a subdivision variable, like plot, to get
# the indexes for each subdivision:
species_diversity(exfm20, "scientific.name", "transect")

# To only get one specific index, use the index argument:
species_diversity(exfm20, "scientific.name", index = "H")
species_diversity(exfm20, "scientific.name", index = "S")
species_diversity(exfm20, "scientific.name", index = "Hmax")
species_diversity(exfm20, "scientific.name", index = "J")
```

sprs

Simple Random Sampling

Description

Function for processing forest inventory data using simple random sampling.

Usage

```
sprs(
  df,
  Yi,
  plot_area,
  total_area,
  age = NA,
  .groups = NA,
  alpha = 0.05,
  error = 10,
  dec_places = 4,
  pop = "inf",
  tidy = TRUE
)
```

Arguments

<code>df</code>	a data frame.
<code>Yi</code>	Quoted name of the volume variable, or other variable one desires to evaluate, in quotes.
<code>plot_area</code>	Quoted name of the plot area variable, or a numeric vector with the plot area value. The plot area value must be in square meters.
<code>total_area</code>	Quoted name of the total area variable, or a numeric vector with the total area value. The total area value must be in hectares.
<code>age</code>	Optional parameter. Quoted name of the age variable. Calculates the average age supplied. NA.
<code>.groups</code>	Optional argument. Quoted name(s) of additional grouping variable(s) that, if supplied, will be used to run multiple surveys, one for each level. If this argument is NA, the defined groups in the data frame will be used, if they exist. Default: NA.
<code>alpha</code>	Numeric value for the significance value used in the t-student estimation. Default: 0.05.
<code>error</code>	Numeric value for the minimum admitted error value in the survey, in percentage. Default: 10.
<code>dec_places</code>	Numeric value for the number of decimal places to be used in the output tables. Default: 4.
<code>pop</code>	Character value for the type of population considered in the calculations. This can be either infinite ("inf") or finite ("fin"). Default: "inf".
<code>tidy</code>	Boolean value that defines if the output tables should be tidied up or not. Default: TRUE.

Details

This function allows the user to process inventory data using simple random sampling for finite or infinite populations. It's possible to run multiple sampling analysis using a factor variable indicated in the `.groups()` parameter.

Value

A data frame with the sampling results.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

References

- Campos, J. C. C. and Leite, H. G. (2017) *Mensuração Florestal: Perguntas e Respostas*. 5a. Vicosa: UFV.
- Soares, C. P. B., Paula Neto, F. and Souza, A. L. (2012) *Dendrometria e Inventário Florestal*. 2nd ed. Vicosa: UFV.

See Also

other sampling functions: [strs](#) for stratified random sampling, and [ss_diffs](#) for Systematic Sampling.

Examples

```
library(forestmangr)
data("exfm2")
data("exfm3")
data("exfm4")

# The objective is to sample an area, with an error of 20%.
# First we run a pilot inventory, considering a 20% error and a finite population:
exfm3

sprs(exfm3, "VWB", "PLOT_AREA", "TOTAL_AREA", error = 20, pop = "fin")

# With these results, in order to obtain the desired error, we'll need to sample new
# plots, and run the definitive inventory. Again, we aim for a 20% error, and consider
# the population as finite:
exfm4

sprs(exfm4, "VWB", "PLOT_AREA", "TOTAL_AREA", error = 20, pop = "fin")

# The desired error was met

# area values can be numeric
sprs(exfm4, "VWB", 3000, 46.8, error = 20, pop = "fin")

# Here we run a simple random sampling inventory for each forest subdivision,
# using the STRATA variable as a group variable:
exfm2

sprs(exfm2, "VWB", "PLOT_AREA", "STRATA_AREA", .groups = "STRATA", error = 20, pop = "fin")
```

ss_diffs

Systematic Sampling

Description

Function for processing forest inventory data using systematic sampling.

Usage

```
ss_diffs(
  df,
  Yi,
  plot_area,
```

```

total_area,
age = NA,
.groups = NA,
alpha = 0.05,
error = 10,
dec_places = 4,
tidy = TRUE
)

```

Arguments

df	a data frame.
Yi	Quoted name of the volume variable, or other variable one desires to evaluate, in quotes.
plot_area	Quoted name of the plot area variable, or a numeric vector with the plot area value. The plot area value must be in square meters.
total_area	Quoted name of the total area variable, or a numeric vector with the total area value. The total area value must be in hectares.
age	Optional parameter. Quoted name of the age variable. Calculates the average age supplied. NA.
.groups	Optional argument. Quoted name(s) of additional grouping variable(s) that, if supplied, will be used to run multiple surveys, one for each level. If this argument is NA, the defined groups in the data frame will be used, if they exist. Default: NA.
alpha	Numeric value for the significance value used in the t-student estimation. Default: 0.05.
error	Numeric value for the minimum admitted error value in the survey, in percentage. Default: 10.
dec_places	Numeric value for the number of decimal places to be used in the output tables. Default: 4.
tidy	Boolean value that defines if the output tables should be tidied up or not. Default: TRUE.

Details

This function allows the user to process inventory data using simple random sampling for finite or infinite populations. It's possible to run multiple sampling analysis using a factor variable indicated in the `.groups()` parameter.

Value

A data frame with the sampling results.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

References

- Campos, J. C. C. and Leite, H. G. (2017) Mensuracao Florestal: Perguntas e Respostas. 5a. Vicosa: UFV.
- Soares, C. P. B., Paula Neto, F. and Souza, A. L. (2012) Dendrometria e Inventario Florestal. 2nd ed. Vicosa: UFV.

See Also

other sampling functions: [sprs](#) for Simple Random Sampling, and [strs](#) for stratified random sampling, and

Examples

```
library(forestmangr)
data("exfm2")
data("exfm5")

# We're trying to run a inventory for an area This data was collected systematically,
# but we'll try to run the data using simple random sampling,
# to show the difference between the two methods:
exfm5

sprs(exfm5, "VWB", "PLOT_AREA", "TOTAL_AREA")

# We get a 22% error value. Now, we run this same data
# considering the data as a systematic inventory, using the
# successive differences method:
exfm5

ss_diffs(exfm5, "VWB", "PLOT_AREA", "TOTAL_AREA")

# The error was significantly lowered.

# Area Values can be numeric;
ss_diffs(exfm5, "VWB", 200, 18)

# Here we run a systematic sampling inventory for each forest subdivision,
# using the STRATA variable as a group variable:
exfm2

ss_diffs(exfm2, "VWB", "PLOT_AREA", "STRATA_AREA", .groups = "STRATA")
```

strs

Stratified Random Sampling

Description

Function for processing forest inventory data using stratified random sampling.

Usage

```

strs(
  df,
  Yi,
  plot_area,
  strata_area,
  strata,
  .groups = NA,
  age = NA,
  alpha = 0.05,
  error = 10,
  dec_places = 4,
  pop = "inf",
  tidy = TRUE
)

```

Arguments

df	a data frame.
Yi	Quoted name of the volume variable, or other variable one desires to evaluate, in quotes.
plot_area	Quoted name of the plot area variable, or a numeric vector with the plot area value. The plot area value must be in square meters.
strata_area	Quoted name of the strata area variable, or a numeric vector with the plot strata values. If there are more than 1 area values, it's possible to use a vector with all area values, like so:c(14.4, 16.4, 14.2). The strata area values must be in hectares.
strata	Quoted name of the subdivision variable(s), also known as strata. If this argument is not supplied, the defined groups in the data frame will be used, if they exist.
.groups	Optional argument. Quoted name(s) of additional grouping variable(s) that, if supplied, will be used to run multiple surveys, one for each level. If this argument is NA, the defined groups in the data frame will be used, if they exist. Default: NA.
age	Optional parameter. Quoted name of the age variable. Calculates the average age supplied. NA.
alpha	Numeric value for the significance value used in the t-student estimation. Default: 0.05.
error	Numeric value for the minimum admitted error value in the survey, in percentage. Default: 10.
dec_places	Numeric value for the number of decimal places to be used in the output tables. Default: 4.
pop	Character value for the type of population considered in the calculations. This can be either infinite ("inf") or finite ("fin"). Default: "inf".
tidy	Boolean value that defines if the output tables should be tidied up or not. Default: TRUE.

Details

This function allows the user to process inventory data using stratified random sampling for n forest subdivisions (strata), for finite or infinite populations. It's possible to run multiple sampling analysis using a factor variable indicated in the `.groups()` parameter.

Value

A list containing two data frames, one with information for each strata, and one with the stratified sampling results.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

References

Campos, J. C. C. and Leite, H. G. (2017) *Mensuracao Florestal: Perguntas e Respostas*. 5a. Vicosa: UFV.

Soares, C. P. B., Paula Neto, F. and Souza, A. L. (2012) *Dendrometria e Inventario Florestal*. 2nd ed. Vicosa: UFV.

See Also

other sampling functions: [sprs](#) for Simple Random Sampling, and [ss_diffs](#) for Systematic Sampling.

Examples

```
library(forestmangr)
data("exfm1")
data("exfm2")
data("exfm6")

# The objective is to sample an area, with an error of 5%.
# First we run a pilot inventory, considering a 5% error and a finite population:
exfm1

strs(exfm1, "VWB", "PLOT_AREA", "STRATA_AREA", strata = "STRATA", error = 5, pop = "fin")

# With these results, in order to meet the desired error of 5%, we'll need to sample 24 more plots,
# 4 in stratum 1, 8 in stratum 2, and 12 in stratum 3.
# After sampling the necessary plots, we now run a definitive inventory,
# considering an 5% error and a finite population:
exfm2

strs(exfm2, "VWB", "PLOT_AREA", "STRATA_AREA", strata = "STRATA", error = 5, pop = "fin")

# The desired error value was met.

# Area values can be numeric:
strs(exfm2, "VWB", 1000, c(14.4, 16.4,14.2), strata = "STRATA", error = 5, pop = "fin")
```

```
# Optional variable age, and one stratified sampled inventory for each map:
exfm6

strs(exfm6, "VWB", "PLOT_AREA", "STRATA_AREA", strata="STRATA", .groups = "MAP", age = "AGE")
```

tree_summarise	<i>Calculate the equivalent diameter of trees with more than one trunk</i>
----------------	--

Description

This function uses takes the square root of the diameters squared sum, in order to estimate the equivalent diameter of trees. Other supplied variables are summed up, or averaged, depending on the variable.

Usage

```
tree_summarise(df, dbh, tree, .groups = NA, vwob = NA, vwob = NA)
```

Arguments

df	A data frame.
dbh	Quoted name of the diameter at breast height variable.
tree	Quoted name of the tree variable. used to differentiate the trees' sections. If this argument is missing, the defined groups in the data frame will be used. If there are no groups in the data, the function will fail.
.groups	Optional argument. Quoted name(s) of grouping variables that can be added to differentiate subdivisions of the data. Default: NA.
vwob	Optional argument. Quoted name of the volume with bark variable, in cubic meters. Default: NA.
vwob	Optional argument. Quoted name of the volume without bark variable, in cubic meters. Default: NA.

Value

A data frame with the the equivalent diameter calculated.

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

References

Soares, C. P. B., Paula Neto, F. and Souza, A. L. (2012) Dendrometria e Inventario Florestal. 2nd ed. Vicosa: UFV.

Examples

```
library(forestmangr)
data("exfm18")
exfm18

# Calculate the equivalent diameter of trees with more than one trunk:
eq_diam <- tree_summarise(exfm18, "DBH", tree="Tree", .groups=c("Plot", "Species") )
head(eq_diam, 10)
```

vertical_stratum	<i>Divide data into 3 vertical strata</i>
------------------	---

Description

Get the vertical strata of data based on the height variable. The data will be divided into inferior, medium and superior strata.

Usage

```
vertical_stratum(df, th)
```

Arguments

df	A data frame.
th	Quoted name of the total height variable.

Value

a data frame.

Author(s)

Eric Bastos Gorgens <e.gorgens@gmail.com>

References

Souza, A. L. and Soares, C. P. B. (2013) Florestas Nativas: estrutura, dinamica e manejo. Vicosa: UFV.

Examples

```
library(forestmangr)
data("exfm10")
exfm10

# To classify the data, supply the data frame and the height variable name:
vertical_stratum(exfm10, "TH" )
```

vol_summarise	<i>Summarize volume of trees</i>
---------------	----------------------------------

Description

This function can be used to summarize volume with and without bark of trees in a data frame.

Usage

```
vol_summarise(df, dbh, th, vwb, tree, .groups = NA, vwob = NA)
```

Arguments

df	A data frame.
dbh	Quoted name of the diameter at breast height variable, in cm.
th	Quoted name of the total height variable, in meters.
vwb	Quoted name of the volume with bark variable, in cubic meters.
tree	Quoted name of the tree variable. used to differentiate the trees' sections. If this argument is NA, the defined groups in the data frame will be used. Default: NA.
.groups	Optional argument. Quoted name(s) of additional grouping variables that can be added to differentiate subdivisions of the data.
vwob	Optional argument. Quoted name of the volume without bark variable, in cubic meters. Default: NA. If this argument is NA, the defined groups in the data frame will be used. Default: NA.

Value

A data frame summarized by the .groups variable(s).

Author(s)

Sollano Rabelo Braga <sollanorb@gmail.com>

See Also

Complementary functions: [smalianwb](#), For calculation of volume with bark using the Smalian method, [smalianwob](#), For calculation of volume without bark using the Smalian method, [huberwb](#), for calculation of volume with bark using the Huber method, [huberwob](#), for calculation of volume without bark the Huber method.

Examples

```

library(forestmangr)
data("exfm7")
exfm7

# In order to calculate the volume of each tree, first we
# Calculate the volume by tree section using the Smalian method:
sec_data_vol <- exfm7 %>%
  smalianwb("di_wb", "hi", "TREE") %>%
  smalianwob("di_wb", "hi", "bark_t", "TREE", bt_mm_to_cm = TRUE)

sec_data_vol

# Now, we summarize the tree's volume:
vol_summarise(sec_data_vol, dbh = "DBH", th = "TH", vwb = "VWB",
  tree = "TREE", .groups = "STRATA", vwob = "VWOB")

# It's possible to do everything using pipes:
exfm7 %>%
  smalianwb("di_wb", "hi", "TREE") %>%
  smalianwob("di_wb", "hi", "bark_t", "TREE", bt_mm_to_cm = TRUE) %>%
  vol_summarise("DBH", "TH", "VWB", "TREE", "STRATA", "VWOB")

```

%>% *Pipe*

Description

Pipe an object forward into a function or call expression. See [magrittr](#) for more details.
 Search for quasiquotation for more details.

%T>% *Tee*

Description

Pipe an object forward into a function or call expression. See [magrittr](#) for more details.

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