Package ‘GGally’

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BugReports https://github.com/ggobi/ggally/issues

Description The R package 'ggplot2' is a plotting system based on the grammar of graphics. 'GGally' extends 'ggplot2' by adding several functions to reduce the complexity of combining geometric objects with transformed data. Some of these functions include a pairwise plot matrix, a two group pairwise plot matrix, a parallel coordinates plot, a survival plot, and several functions to plot networks.

Depends R (>= 3.1)
Imports ggplot2 (>= 2.2.0), grDevices, grid, gtable (>= 0.2.0), plyr (>= 1.8.3), progress, RCColorBrewer, reshape (>= 0.8.5), utils
Suggests broom (>= 0.4.0), chemometrics, geosphere (>= 1.5-1), igraph (>= 1.0.1), intergraph (>= 2.0-2), maps (>= 3.1.0), mapproj, network (>= 1.12.0), scagnostics, scales (>= 0.4.0), sna (>= 2.3-2), survival, packageRdocs (>= 0.4.0), rmarkdown, roxygen2, testthat

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NeedsCompilation no

Author Barret Schloerke [aut, cre] (author for ggpairs, ggduo, ggnostic, ggt水流, ggface, and ggally_*. Contributor for all functions.), Jason Crowley [aut] (ggparcoord), Di Cook [aut, ths] (ggscatmat, gglyph), Heike Hofmann [ths], Hadley Wickham [ths],
Francois Briatte [aut] (ggcorr, ggnet, ggnet2),
Moritz Marbach [aut] (ggnet, ggnet2),
Edwin Thoen [aut] (ggsurv),
Amos Elberg [aut] (ggnetworkmap),
Joseph Larmarange [aut] (ggcoef)

Maintainer Barret Schloerke <schloerke@gmail.com>
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R topics documented:

+ gg .................................................. 4
add_ref_boxes ...................................... 5
add_ref_lines ...................................... 5
australia_PISA2012 .................................. 6
brew_colors ........................................ 7
broomify ............................................ 8
find_plot_type ...................................... 8
flea ................................................... 9
fn_switch .......................................... 10
getPlot ............................................. 11
ggally_barDiag .................................... 11
ggally_blank ....................................... 12
ggally_box ......................................... 13
ggally_cor ......................................... 13
ggally_density ..................................... 14
ggally_densityDiag ................................ 15
ggally_denstrip .................................... 16
ggally_diagAxis .................................... 17
ggally_dot .......................................... 18
ggally_dot_and_box ................................ 19
ggally_facetbar .................................... 19
ggally_facetdensity ................................ 20
ggally_facetdensitystrip ........................... 21
ggally_facethist .................................... 21
ggally_na ........................................... 22
ggally_nostic_cooksd ................................ 23
ggally_nostic_hat ................................... 24
ggally_nostic_line .................................. 25
ggally_nostic_resid ................................ 26
ggally_nostic_se_fit ................................ 27
ggally_nostic_sigma ................................ 28
ggally_nostic_std_resid ............................. 29
ggally_points ....................................... 29
ggally_ratio ........................................ 30
ggally_smooth ....................................... 31
ggally_text ......................................... 32
R topics documented:

- ggcoef .................................................. 32
- ggcorr .................................................. 33
- ggduo .................................................. 36
- ggfacet .................................................. 40
- gglegend ................................................. 41
- ggmatrix ................................................. 43
- ggmatrix_gtable ....................................... 45
- gnet ..................................................... 46
- ggnet2 .................................................. 49
- ggnetworkmap ........................................... 54
- ggnostic ............................................... 57
- ggpairs .................................................. 60
- ggparcoord ............................................. 64
- ggscatmat .............................................. 67
- ggsurv .................................................. 68
- ggt ....................................................... 70
- glyphplot .............................................. 71
- glyphs ................................................... 72
- grab_legend ............................................ 73
- happy .................................................... 74
- lowertriangle ......................................... 75
- model_response_variables ......................... 75
- nasa ..................................................... 76
- pigs ..................................................... 77
- print.ggmatrix ......................................... 78
- print_if_interactive .................................. 78
- psychademic .......................................... 79
- putPlot ................................................. 80
- rescale01 ............................................... 81
- scag_order ............................................. 81
- scatmat ............................................... 82
- singleClassOrder ..................................... 83
- skewness ............................................... 83
- str.ggmatrix ........................................... 84
- twitter_spambots .................................... 84
- uppertriangle .......................................... 85
- v1_ggmatrix_theme .................................... 86
- wrap_fn_with_param_arg ............................. 86

Index .................................................. 89
+ gg

Modify a ggmatrix object by adding an ggplot2 object to all plots

Description

This operator allows you to add ggplot2 objects to a ggmatrix object.

Usage

```r
## S3 method for class 'gg'
e1 + e2
```

Arguments

- `e1`: An object of class `ggplot` or `theme`
- `e2`: A component to add to `e1`

Details

If the first object is an object of class `ggmatrix`, you can add the following types of objects, and it will return a modified ggplot object.

- `theme`: update plot theme

The `+` operator completely replaces elements with elements from `e2`.

See Also

`+.gg` and `theme`

Examples

```r
data(tips, package = "reshape")
pm <- ggpairs(tips[, 2:3])
## change to black and white theme
pm + ggplot2::theme_bw()
## change to linedraw theme
# pm + ggplot2::theme_linedraw()
## change to custom theme
# pm + ggplot2::theme(panel.background = ggplot2::element_rect(fill = "lightblue"))
## add a list of information
extra <- list(ggplot2::theme_bw(), ggplot2::labs(caption = "My caption!"))
pm + extra
```
**add_ref_boxes**

Add reference boxes around each cell of the glyphmap.

### Description

Add reference boxes around each cell of the glyphmap.

### Usage

```r
add_ref_boxes(data, var_fill = NULL, color = "white", size = 0.5,
               fill = NA, ...)
```

### Arguments

- **data**
  A glyphmap structure.
- **var_fill**
  Variable name to use to set the fill color
- **color**
  Set the color to draw in, default is "white"
- **size**
  Set the line size, default is 0.5
- **fill**
  fill value used if var_fill is NULL
- **...**
  other arguments passed onto `geom_rect`

**add_ref_lines**

Add reference lines for each cell of the glyphmap.

### Description

Add reference lines for each cell of the glyphmap.

### Usage

```r
add_ref_lines(data, color = "white", size = 1.5, ...)
```

### Arguments

- **data**
  A glyphmap structure.
- **color**
  Set the color to draw in, default is "white"
- **size**
  Set the line size, default is 1.5
- **...**
  other arguments passed onto `geom_line`
Description

About PISA

Usage

data(australia_PISA2012)

Format

A data frame with 8247 rows and 32 variables

Details

The Programme for International Student Assessment (PISA) is a triennial international survey which aims to evaluate education systems worldwide by testing the skills and knowledge of 15-year-old students. To date, students representing more than 70 economies have participated in the assessment.

While 65 economies took part in the 2012 study, this data set only contains information from the country of Australia.

- gender : Factor w/ 2 levels "female","male" : 1 1 2 2 2 1 1 1 2 1 ...
- age : Factor w/ 4 levels "4","5","6","7" : 2 2 2 4 3 1 2 2 2 2 ...
- homework : num 5 5 9 3 2 3 4 3 5 1 ...
- desk : num 1 0 1 1 1 1 1 1 1 1 ...
- room : num 1 1 1 1 1 1 1 1 1 1 ...
- study : num 1 1 1 1 1 1 1 1 1 1 ...
- computer : num 1 1 1 1 1 1 1 ...
- software : num 1 1 1 1 1 1 1 1 1 ...
- internet : num 1 1 1 1 1 1 1 1 1 1 ...
- literature : num 0 0 1 0 1 1 1 1 0 ...
- poetry : num 0 0 1 0 1 1 0 1 1 1 ...
- art : num 1 0 1 0 1 1 0 1 1 1 ...
- textbook : num 1 1 1 1 0 1 1 1 1 ...
- dictionary : num 1 1 1 1 1 1 1 1 1 ...
- dishwasher : num 1 1 1 0 1 1 1 1 1 ...
- PV1MATH : num 562 565 602 520 613 ...
- PV2MATH : num 569 557 594 507 567 ...
• PV3MATH : num 555 553 552 501 585 ...
• PV4MATH : num 579 538 526 521 596 ...
• PV5MATH : num 548 573 619 547 603 ...
• PV1READ : num 582 617 650 554 605 ...
• PV2READ : num 571 572 608 560 557 ...
• PV3READ : num 602 560 594 517 627 ...
• PV4READ : num 572 564 575 564 597 ...
• PV5READ : num 585 565 620 572 598 ...
• PV1SCIE : num 583 627 668 574 639 ...
• PV2SCIE : num 579 600 665 612 635 ...
• PV3SCIE : num 593 574 620 571 666 ...
• PV4SCIE : num 567 582 592 598 700 ...
• PV5SCIE : num 587 625 656 662 670 ...
• SENWGT_STU : num 0.133 0.133 0.141 0.141 0.141 ...
• possessions: num 10 8 12 9 11 11 10 12 12 11 ...

Source


brew_colors

RColorBrewer Set1 colors

Description

RColorBrewer Set1 colors

Usage

brew_colors(col)

Arguments

col standard color name used to retrieve hex color value
broomify  

**Broomify a model**

**Description**

broom::augment a model and add broom::glance and broom::tidy output as attributes. X and Y variables are also added.

**Usage**

broomify(model, lmStars = TRUE)

**Arguments**

- `model`  
  model to be sent to broom::augment, broom::glance, and broom::tidy
- `lmStars`  
  boolean that determines if stars are added to labels

**Value**

broom::augmented data frame with the broom::glance data.frame and broom::tidy data.frame as `broom_glance` and `broom_tidy` attributes respectively. var_x and var_y variables are also added as attributes

**Examples**

```r
data(mtcars)
model <- stats::lm(mpg ~ wt + qsec + am, data = mtcars)
broomified_model <- broomify(model)
str(broomified_model)
```

---

**find_plot_type**

**Find Plot Types**

**Description**

Retrieves the type of plot for the specific columns

**Usage**

find_plot_type(col1Name, col2Name, type1, type2, isAllNa, allowDiag)

---
Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>col1Name</td>
<td>x column name</td>
</tr>
<tr>
<td>col2Name</td>
<td>y column name</td>
</tr>
<tr>
<td>type1</td>
<td>x column type</td>
</tr>
<tr>
<td>type2</td>
<td>y column type</td>
</tr>
<tr>
<td>isAllNa</td>
<td>is.na(data)</td>
</tr>
<tr>
<td>allowDiag</td>
<td>allow for diag values to be returned</td>
</tr>
</tbody>
</table>

Author(s)

Barret Schloerke <schloerke@gmail.com>

---

**flea**

*Historical data used for classification examples.*

Description

This data contains physical measurements on three species of flea beetles.

Usage

data(flea)

Format

A data frame with 74 rows and 7 variables

Details

- species Ch. concinna, Ch. heptapotamica, Ch. heikertingeri
- tars1 width of the first joint of the first tarsus in microns
- tars2 width of the second joint of the first tarsus in microns
- head the maximal width of the head between the external edges of the eyes in 0.01 mm
- aede1 the maximal width of the aedeagus in the fore-part in microns
- aede2 the front angle of the aedeagus (1 unit = 7.5 degrees)
- aede3 the aedeagus width from the side in microns

References

Function switch

Description

Function that allows you to call different functions based upon an aesthetic variable value.

Usage

```
fn_switch(types, mapping_val = "y")
```

Arguments

types

list of functions that follow the ggmatrix function standard: function(data, mapping, ...){ #make ggplotR object }

One key should be a 'default' key for a default switch case.

mapping_val

mapping value to switch on. Defaults to the 'y' variable of the aesthetics list.

Examples

```
ggnostic_continuous_fn <- fn_switch(list(  default = ggally_points,  .fitted = ggally_points,  .se.fit = ggally_nostic_se_fit,  .resid = ggally_nostic_resid,  .hat = ggally_nostic_hat,  .sigma = ggally_nostic_sigma,  .cooksd = ggally_nostic_cooksd,  .std.resid = ggally_nostic_std_resid ))


ggnostic_combo_fn <- fn_switch(list(  default = ggally_box_no_facet,  fitted = ggally_box_no_facet,  .se.fit = ggally_nostic_se_fit,  .resid = ggally_nostic_resid,  .hat = ggally_nostic_hat,  .sigma = ggally_nostic_sigma,  .cooksd = ggally_nostic_cooksd,  .std.resid = ggally_nostic_std_resid ))
```
getPlot

Description

Retrieves the ggplot object at the desired location.

Usage

getPlot(pm, i, j)

## S3 method for class 'ggmatrix'
pm[i, j, ...]

Arguments

pm  ggmatrix object to select from
i   row from the top
j   column from the left
... ignored

Author(s)

Barret Schloerke <schloerke@gmail.com>

Examples

data(tips, package = "reshape")
plotMatrix2 <- ggpairs(tips[, 3:2], upper = list(combo = "denstrip"))
plotMatrix2[, 1, 2]

---

ggally_barDiag  Plots the Bar Plots by Using Diagonal

Description

Plots the bar plots by using Diagonal.

Usage

ggally_barDiag(data, mapping, ..., rescale = FALSE)
Arguments

- **data**
  - data set using

- **mapping**
  - aesthetics being used

- **...**
  - other arguments are sent to `geom_bar`

- **rescale**
  - boolean to decide whether or not to rescale the count output. Only applies to numeric data

Author(s)

Barret Schloerke <schloerke@gmail.com>

Examples

```r
data(tips, package = "reshape")
ggally_barDiag(tips, mapping = ggplot2::aes(x = day))
ggally_barDiag(tips, mapping = ggplot2::aes(x = tip), binwidth = 0.25)
```

Description

Draws nothing.

Usage

```r
ggally_blank(...)  
ggally_blankDiag(...)  
```

Arguments

```r
...  
other arguments ignored
```

Details

Makes a "blank" ggplot object that will only draw white space

Author(s)

Barret Schloerke <schloerke@gmail.com>
**ggally_box**

**Plots the Box Plot**

**Description**

Make a box plot with a given data set. `ggally_box_no_facet` will be a single panel plot, while `ggally_box` will be a faceted plot.

**Usage**

```r
ggally_box(data, mapping, ...)

ggally_box_no_facet(data, mapping, ...)
```

**Arguments**

- `data`: data set using
- `mapping`: aesthetics being used
- `...`: other arguments being supplied to `geom_boxplot`

**Author(s)**

Barret Schloerke <schloerke@gmail.com>

**Examples**

```r
data(tips, package = "reshape")
ggally_box(tips, mapping = ggplot2::aes(x = total_bill, y = sex))
ggally_box(tips, mapping = ggplot2::aes_string(x = "total_bill", y = "sex"))
ggally_box(  
tips,  
  mapping = ggplot2::aes_string(y = "total_bill", x = "sex", color = "sex"),  
  outlier.colour = "red",  
  outlier.shape = 13,  
  outlier.size = 8  
)
```

---

**ggally_cor**

**Correlation from the Scatter Plot**

**Description**

Estimate correlation from the given data.
Usage

ggally_cor(data, mapping, alignPercent = 0.6, method = "pearson",
    use = "complete.obs", corAlignPercent = NULL, corMethod = NULL,
    corUse = NULL, ...)

Arguments

data          data set using
mapping        aesthetics being used
alignPercent   right align position of numbers. Default is 60 percent across the horizontal
method         method supplied to cor function
use            use supplied to cor function
corAlignPercent deprecated. Use parameter alignPercent
corMethod      deprecated. Use parameter method
corUse         deprecated. Use parameter use
...            other arguments being supplied to geom_text

Author(s)

Barret Schloerke <schloerke@gmail.com>

Examples

data(tips, package = "reshape")
ggally_cor(tips, mapping = ggplot2::aes_string(x = "total_bill", y = "tip"))
ggally_cor(tips,
    mapping = ggplot2::aes(x = total_bill, y = tip),
    size = 15,
    colour = 1("red")
)
ggally_cor(tips,
    mapping = ggplot2::aes_string(x = "total_bill", y = "tip", color = "sex"),
    size = 5
)

ggally_density                  Plots the Scatter Density Plot

Description

Make a scatter density plot from a given data.
**ggally_densityDiag**

**Usage**

```r
ggally_density(data, mapping, ...)
```

**Arguments**

- `data`: data set using
- `mapping`: aesthetics being used
- `...`: parameters sent to either `stat_density2d` or `geom_density2d`

**Details**

The aesthetic "fill" determines whether or not `stat_density2d` (filled) or `geom_density2d` (lines) is used.

**Author(s)**

Barret Schloerke <schloerke@gmail.com>

**Examples**

```r
data(tips, package = "reshape")
ggally_density(tips, mapping = ggplot2::aes(x = total_bill, y = tip))
ggally_density(tips, mapping = ggplot2::aes_string(x = "total_bill", y = "tip"))
ggally_density(tips, mapping = ggplot2::aes_string(x = "total_bill", y = "tip", fill = ".level.."))
```

**ggally_densityDiag**  
*Plots the Density Plots by Using Diagonal*

**Usage**

```r
ggally_densityDiag(data, mapping, ..., rescale = FALSE)
```

**Arguments**

- `data`: data set using
- `mapping`: aesthetics being used.
- `...`: other arguments sent to `stat_density`
- `rescale`: boolean to decide whether or not to rescale the count output
**ggally_denstrip**

Plots a tile plot with facets

**Description**

Make Tile Plot as densely as possible.

**Usage**

`ggally_denstrip(data, mapping, ...)`

**Arguments**

- `data` : data set using
- `mapping` : aesthetics being used
- `...` : other arguments being sent to `stat_bin`

**Examples**

```r
data(tips, package = "reshape")
ggally_densityDiag(tips, mapping = ggplot2::aes(x = total_bill))
ggally_densityDiag(tips, mapping = ggplot2::aes(x = total_bill, color = day))
```

```r
data(tips, package = "reshape")
ggally_denstrip(tips, mapping = ggplot2::aes(x = total_bill, y = sex))
ggally_denstrip(tips, mapping = ggplot2::aes_string(x = "total_bill", y = "sex"))
ggally_denstrip(tips, mapping = ggplot2::aes_string(x = "sex", y = "tip", binwidth = "0.2")
) + ggplot2::scale_fill_gradient(low = "grey80", high = "black")
```
ggally_diagAxis

Internal Axis Labeling Plot for ggpairs

Description

This function is used when axisLabels == "internal".

Usage

ggally_diagAxis(data, mapping, label = mapping$x, labelSize = 5,
labelXPercent = 0.5, labelYPercent = 0.55, labelHJust = 0.5,
labelVJust = 0.5, gridLabelSize = 4, ...)

Arguments

data dataset being plotted
mapping aesthetics being used (x is the variable the plot will be made for)
label title to be displayed in the middle. Defaults to mapping$x
labelSize size of variable label
labelXPercent percent of horizontal range
labelYPercent percent of vertical range
labelHJust hjust supplied to label
labelVJust vjust supplied to label
gridLabelSize size of grid labels
... other arguments for geom_text

Author(s)

Jason Crowley <crowley.jason.s@gmail.com> and Barret Schloerke

Examples

data(tips, package = "reshape")
ggally_diagAxis(tips, ggplot2::aes(x=tip))
ggally_diagAxis(tips, ggplot2::aes(x=sex))
**ggally_dot**

*Plots the Box Plot with Dot*

**Description**

Add jittering with the box plot. `ggally_dot_no_facet` will be a single panel plot, while `ggally_dot` will be a faceted plot.

**Usage**

```r
ggally_dot(data, mapping, ...)
ggally_dot_no_facet(data, mapping, ...)
```

**Arguments**

- `data` : data set using
- `mapping` : aesthetics being used
- `...` : other arguments being supplied to `geom_jitter`

**Author(s)**

Barret Schloerke <schloerke@gmail.com>

**Examples**

```r
data(tips, package = "reshape")
ggally_dot(tips, mapping = ggplot2::aes(x = total_bill, y = sex))
ggally_dot(tips, mapping = ggplot2::aes_string(x = "total_bill", y = "sex"))
ggally_dot(
  tips,
  mapping = ggplot2::aes_string(y = "total_bill", x = "sex", color = "sex")
)
ggally_dot(
  tips,
  mapping = ggplot2::aes_string(y = "total_bill", x = "sex", color = "sex", shape = "sex")
) + ggplot2::scale_shape(solid=FALSE)
```
**ggally_dot_and_box**

Plots either Box Plot or Dot Plots

Description

Place box plots or dot plots on the graph

Usage

```r
ggally_dot_and_box(data, mapping, ..., boxPlot = TRUE)
```

Arguments

- `data`: data set using
- `mapping`: aesthetics being used
- `...`: parameters passed to either geom_jitter or geom_boxplot
- `boxPlot`: boolean to decide to plot either box plots (TRUE) or dot plots (FALSE)

Author(s)

Barret Schloerke <schloerke@gmail.com>

Examples

```r
data(tips, package = "reshape")
ggally_dot_and_box(tips, 
  mapping = ggplot2::aes(x = total_bill, y = sex, color = sex),
  boxPlot = TRUE)
ggally_dot_and_box(tips, 
  mapping = ggplot2::aes(x = total_bill, y = sex, color = sex),
  boxPlot = FALSE)
```

**ggally_facetbar**

Plots the Bar Plots Faceted by Conditional Variable

Description

X variables are plotted using geom_bar and faceted by the Y variable.

Usage

```r
ggally_facetbar(data, mapping, ...)
```
Arguments

- **data**: data set using
- **mapping**: aesthetics being used
- **...**: other arguments are sent to geom_bar

Author(s)

Barret Schloerke <schloerke@gmail.com>

Examples

```r
data(tips, package = "reshape")
ggally_facetbar(tips, ggplot2::aes(x = sex, y = smoker, fill = time))
ggally_facetbar(tips, ggplot2::aes(x = smoker, y = sex, fill = time))
```

---

**ggally_facetdensity**  
Plots the density plots by faceting

Description

Make density plots by displaying subsets of the data in different panels.

Usage

```r
ggally_facetdensity(data, mapping, ...)
```

Arguments

- **data**: data set using
- **mapping**: aesthetics being used
- **...**: other arguments being sent to stat_density

Author(s)

Barret Schloerke <schloerke@gmail.com>

Examples

```r
data(tips, package = "reshape")
ggally_facetdensity(tips, mapping = ggplot2::aes(x = total_bill, y = sex))
ggally_facetdensity(
  tips,
  mapping = ggplot2::aes_string(y = "total_bill", x = "sex", color = "sex")
)
```
ggally_facetdensitystrip

Plots a density plot with facets or a tile plot with facets

Description

Make Tile Plot as densely as possible.

Usage

```
ggally_facetdensitystrip(data, mapping, ..., den_strip = FALSE)
```

Arguments

- `data` data set using
- `mapping` aesthetics being used
- `...` other arguments being sent to either `geom_histogram` or `stat_density`
- `den_strip` boolean to decide whether or not to plot a density strip(TRUE) or a facet density(FALSE) plot.

Author(s)

Barret Schloerke <schloerke@gmail.com>

Examples

```
example(ggally_facetdensity)
example(ggally_denstrip)
```

---

ggally_facethist

Plots the Histograms by Faceting

Description

Make histograms by displaying subsets of the data in different panels.

Usage

```
ggally_facethist(data, mapping, ...)
```

Arguments

- `data` data set using
- `mapping` aesthetics being used
- `...` parameters sent to `stat_bin()`
Author(s)

Barret Schloerke <schloerke@gmail.com>

Examples

data(tips, package = "reshape")
ggally_facethist(tips, mapping = ggplot2::aes(x = tip, y = sex))
ggally_facethist(tips, mapping = ggplot2::aes_string(x = "tip", y = "sex"), binwidth = 0.1)

-------------

ggally_na  NA plot

Description

Draws a large NA in the middle of the plotting area. This plot is useful when all X or Y data is NA

Usage

ggally_na(data = NULL, mapping = NULL, size = 10, color = "grey20", ...)
ggally_nadiag(...)  

Arguments

- `data`: ignored
- `mapping`: ignored
- `size`: size of the geom_text 'NA'
- `color`: color of the geom_text 'NA'
- `...`: other arguments sent to geom_text

Author(s)

Barret Schloerke <schloerke@gmail.com>
Description

A function to display stats::cooks.distance.

Usage

```r
ggally_nostic_cooksd(data, mapping, ..., linePosition = 4/nrow(data),
                      lineColor = brew_colors("grey"), lineType = 2)
```

Arguments

data, mapping, ..., lineColor, lineType
parameters supplied to `ggally_nostic_line`

linePosition 4 / n is the general cutoff point for Cook's Distance

Details

A line is added at 4 / n to display the general cutoff point for Cook's Distance.


Value

`ggplot2` plot object

See Also

stats::cooks.distance

Examples

```r
dt <- broomify(stats::lm(mpg ~ wt + qsec + am, data = mtcars))
ggally_nostic_cooksd(dt, ggplot2::aes(wt, .cooksd))
```
Description

A function to display stats::influence’s hat information against a given explanatory variable.

Usage

```
ggally_nostic_hat(data, mapping, ..., linePosition = 2 * sum(data[[deparse(mapping$y)]]))/nrow(data), lineColor = brew_colors("grey"), lineSize = 0.5, lineAlpha = 1, lineType = 2, avgLinePosition = sum(data[[deparse(mapping$y)]]))/nrow(data), avgLineColor = brew_colors("grey"), avgLineSize = lineSize, avgLineAlpha = lineAlpha, avgLineType = 1)
```

Arguments

- `data`, `mapping`, ...
  supplied directly to `ggally_nostic_line`
- `linePosition`, `lineColor`, `lineSize`, `lineAlpha`, `lineType`
  parameters supplied to ggplot2::geom_line for the cutoff line
- `avgLinePosition`, `avgLineColor`, `avgLineSize`, `avgLineAlpha`, `avgLineType`
  parameters supplied to ggplot2::geom_line for the average line

Details

As stated in stats::influence documentation:

hat: a vector containing the diagonal of the 'hat' matrix.

The diagonal elements of the 'hat' matrix describe the influence each response value has on the fitted value for that same observation.

A suggested "cutoff" line is added to the plot at a height of 2 * p / n and an expected line at a height of p / n. If either `linePosition` or `avgLinePosition` is NULL, the respective line will not be drawn.

Value

ggplot2 plot object

See Also

stats::influence

Examples

```
dt <- broomify(stats::lm(mpg ~ wt + qsec + am, data = mtcars))
ggally_nostic_hat(dt, ggplot2::aes(wt, .hat))
```
Description

If a non-null `linePosition` value is given, a line will be drawn before the given `continuous_geom` or `combo_geom` is added to the plot.

Usage

```r
ggally_nostic_line(data, mapping, ..., linePosition = NULL, 
lineColor = "red", lineSize = 0.5, lineAlpha = 1, lineType = 1, 
continuous_geom = ggplot2::geom_point, combo_geom = ggplot2::geom_boxplot, 
mapColorToFill = TRUE)
```

Arguments

- `data`, `mapping` supplied directly to `ggplot2::ggplot(data, mapping)`
- `...` parameters supplied to `continuous_geom` or `combo_geom`
- `linePosition`, `lineColor`, `lineSize`, `lineAlpha`, `lineType` parameters supplied to `ggplot2::geom_line`
- `continuous_geom` `ggplot2` geom that is executed after the line is (possibly) added and if the x data is continuous
- `combo_geom` `ggplot2` geom that is executed after the line is (possibly) added and if the x data is discrete
- `mapColorToFill` boolean to determine if combo plots should cut the color mapping to the fill mapping

Details

Functions with a color in their name have different default color behavior.

Value

`ggplot2` plot object
ggally_nostic_resid  ggnostic - residuals

Description

If non-null pVal and sigma values are given, confidence interval lines will be added to the plot at the specified pVal percentiles of a N(0, sigma) distribution.

Usage

```r
ggally_nostic_resid(data, mapping, ..., linePosition = 0,
  lineColor = brew_colors("grey"), lineSize = 0.5, lineAlpha = 1,
  lineType = 1, lineConfColor = brew_colors("grey"),
  lineConfSize = lineSize, lineConfAlpha = lineAlpha, lineConfType = 2,
  pVal = c(0.025, 0.975), sigma = attr(data, "broom_glance")$sigma,
  se = TRUE, method = "auto")
```

Arguments

data, mapping, ...

parameters supplied to `ggally_nostic_line`

linePosition, lineColor, lineSize, lineAlpha, lineType

parameters supplied to `ggplot2::geom_line`

lineConfColor, lineConfSize, lineConfAlpha, lineConfType

parameters supplied to the confidence interval lines

pVal

percentiles of a N(0, sigma) distribution to be drawn

sigma

sigma value for the pVal percentiles

se

boolean to determine if the confidence intervals should be displayed

method

parameter supplied to `ggplot2::geom_smooth`. Defaults to "auto"

Value

ggplot2 plot object

See Also

`stats::residuals`

Examples

dt <- broomify(stats::lm(mpg ~ wt + qsec + am, data = mtcars))
ggally_nostic_resid(dt, ggplot2::aes(wt, .resid))
ggally_nostic_se_fit  

### Description

A function to display stats::predict's standard errors

### Usage

```r
ggally_nostic_se_fit(data, mapping, ..., lineColor = brew_colors("grey"),
linePosition = NULL)
```

### Arguments

- `data`, `mapping`, `...`, `lineColor` parameters supplied to `ggally_nostic_line`
- `linePosition` base comparison for a perfect fit

### Details

As stated in stats::predict documentation:

If the logical 'se.fit' is 'TRUE', standard errors of the predictions are calculated. If the numeric argument 'scale' is set (with optional "df"), it is used as the residual standard deviation in the computation of the standard errors, otherwise this is extracted from the model fit.

Since the se.fit is TRUE and scale is unset by default, the standard errors are extracted from the model fit.

A base line of 0 is added to give reference to a perfect fit.

### Value

`ggplot2` plot object

### See Also

stats::influence

### Examples

```r
dt <- broomify(stats::lm(mpg ~ wt + qsec + am, data = mtcars))
ggally_nostic_se_fit(dt, ggplot2::aes(wt, .se.fit))
```
ggally_nostic_sigma  ggnostic - leave one out model sigma

Description

A function to display stats::influence’s sigma value.

Usage

```
ggally_nostic_sigma(data, mapping, ..., lineColor = brew_colors("grey"),
linePosition = attr(data, "broom_glance")$sigma)
```

Arguments

data, mapping, ..., lineColor
parameters supplied to `ggally_nostic_line`

linePosition  line that is drawn in the background of the plot. Defaults to the overall model’s sigma value.

Details

As stated in stats::influence documentation:

sigma: a vector whose i-th element contains the estimate of the residual standard deviation obtained when the i-th case is dropped from the regression. (The approximations needed for GLMs can result in this being 'NaN'.)

A line is added to display the overall model’s sigma value. This gives a baseline for comparison

Value

ggplot2 plot object

See Also

stats::influence

Examples

```
dt <- broomify(stats::lm(mpg ~ wt + qsec + am, data = mtcars))
ggally_nostic_sigma(dt, ggplot2::aes(wt, .sigma))
```
**ggally_nostic_std_resid**

**Description**

If non-null pVal and sigma values are given, confidence interval lines will be added to the plot at the specified pVal locations of a N(0, 1) distribution.

**Usage**

```r
ggally_nostic_std_resid(data, mapping, ..., sigma = 1)
```

**Arguments**

- `data`: data
- `mapping`: mapping
- `sigma`: sigma value for the pVal percentiles. Set to 1 for standardized residuals

**Value**

ggplot2 plot object

**See Also**

`stats::rstandard`

**Examples**

```r
dt <- broomify(stats::lm(mpg ~ wt + qsec + am, data = mtcars))
ggally_nostic_std_resid(dt, ggplot2::aes(x, .std.resid))
```

---

**ggally_points**

**Plots the Scatter Plot**

**Description**

Make a scatter plot with a given data set.

**Usage**

```r
ggally_points(data, mapping, ...)
```
Arguments

data | data set using
description | aesthetics being used
other arguments | other arguments are sent to geom_point

Author(s)

Barret Schloerke <schloerke@gmail.com>

Examples

```r
data(mtcars)
ggally_points(mtcars, mapping = ggplot2::aes(x = disp, y = hp))
ggally_points(mtcars, mapping = ggplot2::aes_string(x = "disp", y = "hp"))
ggally_points(mtcars,
  mapping = ggplot2::aes_string(
    x = "disp",
    y = "hp",
    color = "as.factor(cyl)",
    size = "gear"
  )
)
```

---

**ggally_ratio**

*Plots a mosaic plot*

Description

Plots the mosaic plot by using fluctuation.

Usage

```r
ggally_ratio(data, mapping = do.call(ggplot2::aes_string,
  as.list(colnames(data)[1:2])), ..., floor = 0, ceiling = NULL)
```

Arguments

data | data set using
description | aesthetics being used. Only x and y will used and both are required
other arguments | other arguments are passed to geom_tile(...)
floor | don’t display cells smaller than this value
ceiling | max value to scale frequencies. If any frequency is larger than the ceiling, the fill color is displayed darker than other rectangles
ggally_smooth

Author(s)
Barret Schloerke <schloerke@gmail.com>

Examples

```r
data(tips, package = "reshape")
ggally_ratio(tips, ggplot2::aes(sex, day))
ggally_ratio(tips, ggplot2::aes(sex, day)) + ggplot2::coord_equal()
# only plot tiles greater or equal to 20 and scale to a max of 50
  ggally_ratio(
    tips, ggplot2::aes(sex, day),
    floor = 20, ceiling = 50
  ) + ggplot2::theme(aspect.ratio = 4/2)
```

---

**ggally_smooth**

Plots the Scatter Plot with Smoothing

Description

Add a smoothed condition mean with a given scatter plot.

Usage

```r
ggally_smooth(data, mapping, ..., method = "lm")
ggally_smooth_loess(data, mapping, ...)
ggally_smooth_lm(data, mapping, ...)
```

Arguments

data  data set using
mapping aesthetics being used
... other arguments to add to geom_point
method method parameter supplied to `geom_smooth`

Author(s)
Barret Schloerke <schloerke@gmail.com>

Examples

```r
data(tips, package = "reshape")
ggally_smooth(tips, mapping = ggplot2::aes(x = total_bill, y = tip))
ggally_smooth(tips, mapping = ggplot2::aes_string(x = "total_bill", y = "tip"))
ggally_smooth(tips, mapping = ggplot2::aes_string(x = "total_bill", y = "tip", color = "sex"))
```
ggally_text

Description

Plot text for a plot.

Usage

```r
ggally_text(label, mapping = ggplot2::aes(color = "black"), xP = 0.5, yP = 0.5, xrange = c(0, 1), yrange = c(0, 1), ...)
```

Arguments

- `label`: text that you want to appear
- `mapping`: aesthetics that don’t relate to position (such as color)
- `xP`: horizontal position percentage
- `yP`: vertical position percentage
- `xrange`: range of the data around it. Only nice to have if plotting in a matrix
- `yrange`: range of the data around it. Only nice to have if plotting in a matrix
- `...`: other arguments for geom_text

Author(s)

Barret Schloerke <schloerke@gmail.com>

Examples

```r
ggally_text("Example 1")
```
```r
ggally_text("Example\nTwo", mapping = ggplot2::aes(size = 15), color = I("red"))
```

---

ggcoef

---

Description

Plot the coefficients of a model with broom and ggplot2.

Usage

```r
ggcoef(x, mapping = aes_string(y = "term", x = "estimate"), conf.int = TRUE, conf.level = 0.95, exponentiate = FALSE, exclude_intercept = FALSE, vline = TRUE, vline_intercept = "auto", vline_color = "gray50", vline_linetype = "dotted", vline_size = 1, errorbar_color = "gray25", errorbar_height = 0, errorbar_linetype = "solid", errorbar_size = 0.5, ...
```

---
Arguments

x a model object to be tidied with tidy or a data frame (see Details)
mapping default aesthetic mapping
conf.int display confidence intervals as error bars?
conf.level level of confidence intervals (passed to tidy if x is not a data frame)
extponentiate if TRUE, x-axis will be logarithmic (also passed to tidy if x is not a data frame)
exclude_intercept should the intercept be excluded from the plot?
vline print a vertical line?
vline_intercept xintercept for the vertical line. "auto" for x = 0 (or x = 1 if exponentiate is TRUE)
vline_color color of the vertical line
vline_linetype line type of the vertical line
vline_size size of the vertical line
errorbar_color color of the error bars
errorbar_height height of the error bars
errorbar_linetype line type of the error bars
errorbar_size size of the error bars
... additional arguments sent to geom_point

Examples

library(broom)
reg <- lm(Sepal.Length ~ Sepal.Width + Petal.Length + Petal.Width, data = iris)
ggcoef(reg)

d <- as.data.frame(Titanic)
reg2 <- glm(Survived ~ Sex + Age + Class, family = binomial, data = d, weights = d$freq)

# exponentiate is TRUE
ggcoef(reg2, exponentiate = TRUE, exclude_intercept = TRUE, errorbar_height = .2, color = "blue")

Description

Function for making a correlation matrix plot, using ggplot2. The function is directly inspired by Tian Zheng and Yu-Sung Su's corrplot function in the 'arm' package. Please visit http://github.com/briatte/ggcorr for the latest version of ggcorr, and see the vignette at https://briatte.github.io/ggcorr/ for many examples of how to use it.
Usage

ggcorr(data, method = c("pairwise", "pearson"), cor_matrix = NULL, nbreaks = NULL, digits = 2, name = "", low = "#3B9AB2", mid = "#EEEEEE", high = "#F21A00", midpoint = 0, palette = NULL, geom = "tile", min_size = 2, max_size = 6, label = FALSE, label_alpha = FALSE, label_color = "black", label_round = 1, label_size = 4, limits = c(-1, 1), drop = is.null(limits) || identical(limits, FALSE), layout.exp = 0, legend.position = "right", legend.size = 9, ...)

Arguments

data a data frame or matrix containing numeric (continuous) data. If any of the columns contain non-numeric data, they will be dropped with a warning.

method a vector of two character strings. The first value gives the method for computing covariances in the presence of missing values, and must be (an abbreviation of) one of "everything", "all.obs", "complete.obs", "na.or.complete" or "pairwise.complete.obs". The second value gives the type of correlation coefficient to compute, and must be one of "pearson", "kendall" or "spearman". See cor for details. Defaults to c("pairwise", "pearson").

cor_matrix the named correlation matrix to use for calculations. Defaults to the correlation matrix of data when data is supplied.

nbreaks the number of breaks to apply to the correlation coefficients, which results in a categorical color scale. See 'Note'. Defaults to NULL (no breaks, continuous scaling).

digits the number of digits to show in the breaks of the correlation coefficients: see cut for details. Defaults to 2.

name a character string for the legend that shows the colors of the correlation coefficients. Defaults to "" (no legend name).

low the lower color of the gradient for continuous scaling of the correlation coefficients. Defaults to "#3B9AB2" (blue).

mid the midpoint color of the gradient for continuous scaling of the correlation coefficients. Defaults to "#EEEEEE" (very light grey).

high the upper color of the gradient for continuous scaling of the correlation coefficients. Defaults to "#F21A00" (red).

midpoint the midpoint value for continuous scaling of the correlation coefficients. Defaults to 0.

palette if nbreaks is used, a ColorBrewer palette to use instead of the colors specified by low, mid and high. Defaults to NULL.

geom the geom object to use. Accepts either "tile", "circle", "text" or "blank".

min_size when geom has been set to "circle", the minimum size of the circles. Defaults to 2.

max_size when geom has been set to "circle", the maximum size of the circles. Defaults to 6.
ggcorr

- **label**: whether to add correlation coefficients to the plot. Defaults to FALSE.
- **label_alpha**: whether to make the correlation coefficients increasingly transparent as they come close to 0. Also accepts any numeric value between 0 and 1, in which case the level of transparency is set to that fixed value. Defaults to FALSE (no transparency).
- **label_color**: the color of the correlation coefficients. Defaults to "grey75".
- **label_round**: the decimal rounding of the correlation coefficients. Defaults to 1.
- **label_size**: the size of the correlation coefficients. Defaults to 4.
- **limits**: bounding of color scaling for correlations, set limits = NULL or FALSE to remove.
- **drop**: if using nbreaks, whether to drop unused breaks from the color scale. Defaults to FALSE (recommended).
- **layout.exp**: a multiplier to expand the horizontal axis to the left if variable names get clipped. Defaults to 0 (no expansion).
- **legend.position**: where to put the legend of the correlation coefficients: see theme for details. Defaults to "bottom".
- **legend.size**: the size of the legend title and labels, in points: see theme for details. Defaults to 9.
- ... other arguments supplied to geom_text for the diagonal labels.

**Note**

Recommended values for the nbreaks argument are 3 to 11, as values above 11 are visually difficult to separate and are not supported by diverging ColorBrewer palettes.

**Author(s)**

Francois Briatte, with contributions from Amos B. Elberg and Barret Schloerke

**See Also**

cor and corrplot in the arm package.

**Examples**

```r
# Basketball statistics provided by Nathan Yau at Flowing Data.

# Default output.
ggcrr(dt[, -1])

# Labelled output, with coefficient transparency.
ggcrr(dt[, -1],
    label = TRUE,
    label_alpha = TRUE)
```
# Custom options.
ggcorr(
dt[, -1],
  name = expression(rho),
  geom = "circle",
  max_size = 10,
  min_size = 2,
  size = 3,
  hjust = 0.75,
  nbreaks = 6,
  angle = -45,
  palette = "PuOr" # colorblind safe, photocopy-able
)

# Supply your own correlation matrix
ggcorr(
data = NULL,
cor_matrix = cor(dt[, -1], use = "pairwise")
)

---

**ggduo**

*ggduo - A ggplot2 generalized pairs plot for two columns sets of a data.frame*

---

**Description**

Make a matrix of plots with a given data set with two different column sets

**Usage**

```r
ggduo(data, mapping = NULL, columnsX = 1:ncol(data),
columnsY = 1:ncol(data), title = NULL, types = list(continuous =
  "smooth_loess", comboVertical = "box_no_facet", comboHorizontal = "facethist",
  discrete = "ratio"), axisLabels = c("show", "none"),
columnLabelsX = colnames(data[columnsX]),
columnLabelsY = colnames(data[columnsY]), labeller = "label_value",
switch = NULL, xlab = NULL, ylab = NULL, showStrips = NULL,
legend = NULL, cardinality_threshold = 15, legends = stop("deprecated"))
```

**Arguments**

- **data**
  - data set using. Can have both numerical and categorical data.
- **mapping**
  - aesthetic mapping (besides x and y). See `aes()`. If mapping is numeric, columns will be set to the mapping value and mapping will be set to NULL.
- **columnsX, columnsY**
  - which columns are used to make plots. Defaults to all columns.
- **title, xlab, ylab**
  - title, x label, and y label for the graph
types see Details
axisLabels either "show" to display axisLabels or "none" for no axis labels
columnLabelsX, columnLabelsY label names to be displayed. Defaults to names of columns being used.
labeller labeller for facets. See labellers. Common values are "label_value" (default) and "label_parsed".
switch switch parameter for facet_grid. See ggplot2::facet_grid. By default, the labels are displayed on the top and right of the plot. If "x", the top labels will be displayed to the bottom. If "y", the right-hand side labels will be displayed to the left. Can also be set to "both"
showStrips boolean to determine if each plot's strips should be displayed. NULL will default to the top and right side plots only. TRUE or FALSE will turn all strips on or off respectively.
legend May be the two objects described below or the default NULL value. The legend position can be moved by using ggplot2's theme element pm + theme(legend.position = "bottom")

- a numeric vector of length 2 provides the location of the plot to use the legend for the plot matrix’s legend. Such as legend = c(3,5) which will use the legend from the plot in the third row and fifth column
- a single numeric value provides the location of a plot according to the display order. Such as legend = 3 in a plot matrix with 2 rows and 5 columns displayed by column will return the plot in position c(1,2)
- a object from grab_legend() a predetermined plot legend that will be displayed directly

cardinality_threshold maximum number of levels allowed in a character / factor column. Set this value to NULL to not check factor columns. Defaults to 15

legends deprecated

Details

types is a list that may contain the variables ‘continuous’, ‘combo’, ‘discrete’, and ‘na’. Each element of the list may be a function or a string. If a string is supplied, it must implement one of the following options:

- **continuous** exactly one of (‘points’, ‘smooth’, ‘smooth_loess’, ‘density’, ‘cor’, ‘blank’). This option is used for continuous X and Y data.
- **comboHorizontal** exactly one of (‘box’, ‘box_no_facet’, ‘dot’, ‘dot_no_facet’, ‘facethist’, ‘facetdensity’, ‘denstrip’, ‘blank’). This option is used for either continuous X and categorical Y data or categorical X and continuous Y data.
- **comboVertical** exactly one of (‘box’, ‘box_no_facet’, ‘dot’, ‘dot_no_facet’, ‘facethist’, ‘facetdensity’, ‘denstrip’, ‘blank’). This option is used for either continuous X and categorical Y data or categorical X and continuous Y data.
- **discrete** exactly one of (‘facetbar’, ‘ratio’, ‘blank’). This option is used for categorical X and Y data.
- **na** exactly one of (‘na’, ‘blank’). This option is used when all X data is NA, all Y data is NA, or either all X or Y data is NA.
If 'blank' is ever chosen as an option, then ggduo will produce an empty plot.

If a function is supplied as an option, it should implement the function api of function(data, mapping, ...){#make ggplotR plot}.
If a specific function needs its parameters set, wrap(fn, param1 = val1, param2 = val2) the function with its parameters.

Examples

```r
# small function to display plots only if it's interactive
p_ <- GGally::print_if_interactive

data(baseball, package = "plyr")

# Keep players from 1990-1995 with at least one at bat
# Add how many singles a player hit
# (must do in two steps as X1b is used in calculations)
dt <- transform(  
  subset(baseball, year >= 1990 & year <= 1995 & ab > 0),
  X1b = h - X2b - X3b - hr
  )

# Add
# the player's batting average,
# the player's slugging percentage,
# and the player's on base percentage
# Make factor a year, as each season is discrete
dt <- transform(  
  dt,
  batting_avg = h / ab,
  slug = (X1b + 2*X2b + 3*X3b + 4*hr) / ab,
  on_base = (h + bb + hbp) / (ab + bb + hbp),
  year = as.factor(year)
  )

pm <- ggduo(  
  dt,
  c("year", "g", "ab", "lg"),
  c("batting_avg", "slug", "on_base"),
  mapping = ggplot2::aes(color = lg)
  )

# Prints, but
# there is severe overplotting in the continuous plots
# the labels could be better
# want to add more hitting information
p_(pm)

# address overplotting issues and add a title
pm <- ggduo(  
  dt,
  c("year", "g", "ab", "lg"),
  c("batting_avg", "slug", "on_base"),
  columnLabelsX = c("year", "player game count", "player at bat count", "league"),
  columnLabelsY = c("batting avg", "slug %", "on base %")
  )
```

title = "Baseball Hitting Stats from 1990-1995",
mapping = ggplot2::aes(color = lg)

types = list(
  # change the shape and add some transparency to the points
  continuous = wrap("smooth_loess", alpha = 0.50, shape = "+")
),
showStrips = FALSE
);
p_(pm)

# Example derived from:
## R Data Analysis Examples | Canonical Correlation Analysis. UCLA: Institute for Digital
## Research and Education.
## from http://www.stats.idre.ucla.edu/r/dae/canonical-correlation-analysis
# "Example 1. A researcher has collected data on three psychological variables, four
# academic variables (standardized test scores) and gender for 500 college freshman.
# She is interested in how the set of psychological variables relates to the academic
# variables and gender. In particular, the researcher is interested in how many
# dimensions (canonical variables) are necessary to understand the association between
# the two sets of variables."
data(pychademic)
summary(pychademic)

(psych_variables <- attr(pychademic, "psychology"))
(academic_variables <- attr(pychademic, "academic"))

## Within correlation
p_(ggpairs(pychademic, columns = psych_variables))
p_(ggpairs(pychademic, columns = academic_variables))

## Between correlation
loess_with_cor <- function(data, mapping, ..., method = "pearson") {
  x <- eval(mapping$x, data)
  y <- eval(mapping$y, data)
  cor <- cor(x, y, method = method)
  ggally_smooth_loess(data, mapping, ...) +
  ggplot2::geom_label(
    data = data.frame(
      x = min(x, na.rm = TRUE),
      y = max(y, na.rm = TRUE),
      lab = round(cor, digits = 3)
    ),
    mapping = ggplot2::aes(x = x, y = y, label = lab),
    hjust = 0, vjust = 1,
    size = 5, fontface = "bold",
    inherit.aes = FALSE # do not inherit anything from the ...
  )
} )

pm <- ggduo
ggfacet

### Description

**ggfacet** - single ggplot2 plot matrix with facet_grid

### Usage

```r
ggfacet(data, mapping = NULL, columnsX = 1:ncol(data),
         columnsY = 1:ncol(data), fn = ggally_points, ...
         columnLabelsX = names(data[columnsX]),
         columnLabelsY = names(data[columnsY]), xlab = NULL, ylab = NULL,
         title = NULL, scales = "free")
```

### Arguments

- **data**
  data.frame that contains all columns to be displayed. This data will be melted before being passed into the function `fn`
mapping aesthetic mapping (besides x and y). See aes()
columnsX columns to be displayed in the plot matrix
columnsY rows to be displayed in the plot matrix
fn function to be executed. Similar to ggpairs and ggduo, the function may either be a string identifier or a real function that wrap understands.
... extra arguments passed directly to fn
columnLabelsX, columnLabelsY column and row labels to display in the plot matrix
xlab, ylab, title plot matrix labels
scales parameter supplied to ggplot2::facet_grid. Default behavior is "free"

Examples

# Small function to display plots only if it's interactive
p_ <- Ggally::print_if_interactive
library(chemometrics)
data(NIR)
NIR_sub <- data.frame(NIR$sGlcEtOH, NIR$sNIR[,1:3])
str(NIR_sub)
x_cols <- c("X1115.0", "X1120.0", "X1125.0")
y_cols <- c("Glucose", "Ethanol")

# using ggduo directly
p <- ggduo(NIR_sub, x_cols, y_cols, types = list(continuous = "points"))
p_(p)

# using ggfacet
p <- ggfacet(NIR_sub, x_cols, y_cols)
p_(p)

# add a smoother
p <- ggfacet(NIR_sub, x_cols, y_cols, fn = 'smooth_loess')
p_(p)
# same output
p <- ggfacet(NIR_sub, x_cols, y_cols, fn = ggally_smooth_loess)
p_(p)

# Change scales to be the same in for every row and for every column
p <- ggfacet(NIR_sub, x_cols, y_cols, scales = "fixed")
p_(p)

---

**gglegend**

Plot only legend of plot function

**Description**

Plot only legend of plot function
Usage

gglegend(fn)

Arguments

fn this value is passed directly to an empty wrap call. Please see ?wrap for more details.

Value

a function that when called with arguments will produce the legend of the plotting function supplied.

Examples

# display regular plot
ggally_points(iris, ggplot2::aes(Sepal.Length, Sepal.Width, color = Species))

# Make a function that will only print the legend
points_legend <- gglegend(ggally_points)
points_legend(iris, ggplot2::aes(Sepal.Length, Sepal.Width, color = Species))

# produce the sample legend plot, but supply a string that 'wrap' understands
same_points_legend <- gglegend("points")
identical(
  attr(attr(points_legend, "fn"), "original_fn"),
  attr(attr(same_points_legend, "fn"), "original_fn")
)

# Complicated examples
custom_legend <- wrap(gglegend("points"), size = 6)
custom_legend(iris, ggplot2::aes(Sepal.Length, Sepal.Width, color = Species))

# Use within ggpairs
pm <- ggpairs(
  iris, 1:2,
  mapping = ggplot2::aes(color = Species),
  upper = list(continuous = gglegend("points"))
)
# pm

# Place a legend in a specific location
pm <- ggpairs(iris, 1:2, mapping = ggplot2::aes(color = Species))
# Make the legend
pm[1,2] <- points_legend(iris, ggplot2::aes(Sepal.Width, Sepal.Length, color = Species))
pm
**ggmatrix**

- **ggmatrix - A ggplot2 Matrix**

### Description

Make a generic matrix of ggplot2 plots.

### Usage

```r
ggmatrix(plots, nrow, ncol, xAxisLabels = NULL, yAxisLabels = NULL,
         title = NULL, xlab = NULL, ylab = NULL, byrow = TRUE,
         showStrips = NULL, showAxisPlotLabels = TRUE,
         showXAxisPlotLabels = TRUE, showYAxisPlotLabels = TRUE, labeller = NULL,
         switch = NULL, xProportions = NULL, yProportions = NULL, data = NULL,
         gg = NULL, legend = NULL)
```

### Arguments

- **plots**
  - list of plots to be put into matrix
- **nrow, ncol**
  - number of rows and columns
- **xAxisLabels, yAxisLabels**
  - strip titles for the x and y axis respectively. Set to NULL to not be displayed
- **title, xlab, ylab**
  - title, x label, and y label for the graph. Set to NULL to not be displayed
- **byrow**
  - boolean that determines whether the plots should be ordered by row or by column
- **showStrips**
  - boolean to determine if each plot’s strips should be displayed. NULL will default to the top and right side plots only. TRUE or FALSE will turn all strips on or off respectively.
- **showAxisPlotLabels, showXAxisPlotLabels, showYAxisPlotLabels**
  - booleans that determine if the plots axis labels are printed on the X (bottom) or Y (left) part of the plot matrix. If showAxisPlotLabels is set, both showXAxisPlotLabels and showYAxisPlotLabels will be set to the given value.
- **labeller**
  - labeller for facets. See `labellers`. Common values are "label_value" (default) and "label_parsed".
- **switch**
  - switch parameter for `facet_grid`. See `ggplot2::facet_grid`. By default, the labels are displayed on the top and right of the plot. If "x", the top labels will be displayed to the bottom. If "y", the right-hand side labels will be displayed to the left. Can also be set to "both"
- **xProportions, yProportions**
  - Value to change how much area is given for each plot. Either NULL (default), numeric value matching respective length, or `grid::unit` object with matching respective length
- **data**
  - data set using. This is the data to be used in place of ‘ggally_data’ if the plot is a string to be evaluated at print time
ggplot2 theme objects to be applied to every plot

**legend**

May be the two objects described below or the default NULL value. The legend position can be moved by using `ggplot2`'s theme element `pm + theme(legend.position = "bottom")`

- **A numeric vector of length 2** provides the location of the plot to use the legend for the plot matrix’s legend. Such as `legend = c(3,5)` which will use the legend from the plot in the third row and fifth column.
- **A single numeric value** provides the location of a plot according to the display order. Such as `legend = 3` in a plot matrix with 2 rows and 5 columns displayed by column will return the plot in position `c(1,3)`.
- **A object from `grab_legend()`** a predetermined plot legend that will be displayed directly.

**Memory usage**

Now that the `print.ggmatrix` method uses a large `gtable` object, rather than print each plot independently, memory usage may be of concern. From small tests, memory usage flutters around `object.size(data) * 0.3 * length(plots)`. So, for a 80Mb random noise dataset with 100 plots, about 2.4 Gb of memory needed to print. For the 3.46 Mb diamonds dataset with 100 plots, about 100 Mb of memory was needed to print. The benefits of using the `ggplot2` format greatly outweigh the price of about 20.

**Author(s)**

Barret Schloerke <schloerke@gmail.com>

**Examples**

```r
# small function to display plots only if it's interactive
p_ <- GGally::print_if_interactive

plotList <- list()
for (i in 1:6) {
  plotList[[i]] <- ggally_text(paste("Plot ", i, sep = ""))
}

pm <- ggmatrix(
  plotList,
  2, 3,
  c("A", "B", "C"),
  c("D", "E"),
  byrow = TRUE
)
p_(pm)

pm <- ggmatrix(
  plotList,
  2, 3,
  xAxisLabels = c("A", "B", "C"),
  yAxisLabels = NULL,
  byrow = FALSE,
  showXAxisPlotLabels = FALSE
)```
Print ggmatrix object

Description

Specialized method to print the ggmatrix object.

Usage

```r
ggmatrix_gtable(pm, ..., progress = interactive() & & (pm$ncol * pm$nrow) > 15,
    progress_format = "plot: [:plot_i,:plot_j] [:bar]:percent est::eta ")
```

Arguments

- `pm`: ggmatrix object to be plotted
- `...`: ignored
- `progress`: boolean to determine if a progress bar should be displayed. This defaults to interactive sessions only
- `progress_format`: string supplied directly to `progress::progress_bar(format = progress_format)`. Defaults to display the plot number, progress bar, percent complete, and estimated time to finish.

Author(s)

Barret Schloerke <schloerke@gmail.com>

Examples

```r
data(tips, package = "reshape")
pm <- ggpairs(tips, c(1,3,2), mapping = ggplot2::aes_string(color = "sex"))
ggmatrix_gtable(pm)
```
ggnet - Plot a network with ggplot2

Description

Function for plotting network objects using ggplot2, now replaced by the `ggnet2` function, which provides additional control over plotting parameters. Please visit [http://github.com/briatte/ggnet](http://github.com/briatte/ggnet) for the latest version of ggnet2, and [https://briatte.github.io/ggnet](https://briatte.github.io/ggnet) for a vignette that contains many examples and explanations.

Usage

```r
ggnet(net, mode = "fruchtermanreingold", layout.par = NULL, layout.exp = 0, size = 9, alpha = 1, weight = "none", weight.legend = NA, weight.method = weight, weight.min = NA, weight.max = NA, weight.cut = FALSE, group = NULL, group.legend = NA, node.group = group, node.color = NULL, node.alpha = alpha, segment.alpha = alpha, segment.color = "grey50", segment.label = NULL, segment.size = 0.25, arrow.size = 0, arrow.gap = 0, arrow.type = "closed", label = FALSE, label.nodes = label, label.size = size/2, label.trim = FALSE, legend.size = 9, legend.position = "right", names = c("", ""), quantize.weights = FALSE, subset.threshold = 0, top8.nodes = FALSE, trim.labels = FALSE, ...)```

Arguments

- **net**: an object of class `network`, or any object that can be coerced to this class, such as an adjacency or incidence matrix, or an edge list: see `edgeset.constructors` and `network` for details. If the object is of class `igraph` and the `intergraph` package is installed, it will be used to convert the object: see `asNetwork` for details.
- **mode**: a placement method from those provided in the `sna` package: see `gplot.layout` for details. Also accepts the names of two numeric vertex attributes of `net`, or a matrix of numeric coordinates, in which case the first two columns of the matrix are used. Defaults to the Fruchterman-Reingold force-directed algorithm.
- **layout.par**: options to be passed to the placement method, as listed in `gplot.layout`. Defaults to `NULL`.
- **layout.exp**: a multiplier to expand the horizontal axis if node labels get clipped: see `expand_range` for details. Defaults to `0` (no expansion).
- **size**: size of the network nodes. If the nodes are weighted, their area is proportionally scaled up to the size set by `size`. Defaults to `9`.
- **alpha**: a level of transparency for nodes, vertices and arrows. Defaults to `1`.
- **weight**: the weighting method for the nodes, which might be a vertex attribute or a vector of size values. Also accepts "indegree", "outdegree", "degree" or "freeman" to size the nodes by their unweighted degree centrality ("degree"
and "freeman" are equivalent): see degree for details. All node weights must be positive. Defaults to "none" (no weighting).

weight.legend the name to assign to the legend created by weight. Defaults to NA (no name).

weight.method see weight

weight.min whether to subset the network to nodes with a minimum size, based on the values of weight. Defaults to NA (preserves all nodes).

weight.max whether to subset the network to nodes with a maximum size, based on the values of weight. Defaults to NA (preserves all nodes).

weight.cut whether to cut the size of the nodes into a certain number of quantiles. Accepts TRUE, which tries to cut the sizes into quartiles, or any positive numeric value, which tries to cut the sizes into that many quantiles. If the size of the nodes do not contain the specified number of distinct quantiles, the largest possible number is used. See quantile and cut for details. Defaults to FALSE (does nothing).

group the groups of the nodes, either as a vector of values or as a vertex attribute. If set to mode on a bipartite network, the nodes will be grouped as "actor" if they belong to the primary mode and "event" if they belong to the secondary mode.

group.legend the name to assign to the legend created by group.

node.group see group

node.color a vector of character strings to color the nodes with, holding as many colors as there are levels in node.group. Defaults to NULL, which will assign grayscale colors to each group.

node.alpha transparency of the nodes. Inherits from alpha.

segment.alpha the level of transparency of the edges. Defaults to alpha, which defaults to 1.

segment.color the color of the edges, as a color value, a vector of color values, or as an edge attribute containing color values. Defaults to "grey50".

segment.label the labels to plot at the middle of the edges, as a single value, a vector of values, or as an edge attribute. Defaults to NULL (no edge labels).

segment.size the size of the edges, in points, as a single numeric value, a vector of values, or as an edge attribute. Defaults to 0.25.

arrow.size the size of the arrows for directed network edges, in points. See arrow for details. Defaults to 0 (no arrows).

arrow.gap a setting aimed at improving the display of edge arrows by plotting slightly shorter edges. Accepts any value between 0 and 1, where a value of 0.05 will generally achieve good results when the size of the nodes is reasonably small. Defaults to 0 (no shortening).

arrow.type the type of the arrows for directed network edges. See arrow for details. Defaults to "closed".

label whether to label the nodes. If set to TRUE, nodes are labeled with their vertex names. If set to a vector that contains as many elements as there are nodes in net, nodes are labeled with these. If set to any other vector of values, the nodes are labeled only when their vertex name matches one of these values. Defaults to FALSE (no labels).
The degree centrality measures that can be produced through the weight argument will take the directedness of the network into account, but will be unweighted. To compute weighted network measures, see the tnet package by Tore Opsahl (help("tnet", package = "tnet")).

Author(s)

Moritz Marbach and Francois Briatte, with help from Heike Hoffmann, Pedro Jordano and Ming-Yu Liu

See Also

ggnet2 in this package, gplot in the sna package, and plot.network in the network package

Examples

library(network)

# random adjacency matrix
x <- 10
ndyads <- x * (x - 1)
density <- x / ndyads
m <- matrix(0, nrow = x, ncol = x)
dimnames(m) <- list(letters[1:x], letters[1:x])
m[ row(m) != col(m) ] <- runif(n * dyads) < density
m

# random undirected network
n <- network::network(m, directed = FALSE)

ngnet2(n, label = TRUE, alpha = 1, color = "white", segment.color = "black")

# random groups
g <- sample(letters[1:3], 10, replace = TRUE)

# color palette
p <- c("a" = "steelblue", "b" = "forestgreen", "c" = "tomato")

ngnet2(n, node.group = g, node.color = p, label = TRUE, color = "white")

# edge arrows on a directed network
ngnet2(network(m, directed = TRUE), arrow.gap = 0.05, arrow.size = 10)

---

**ggnet2**  
**ggnet2 - Plot a network with ggplot2**

**Description**

Function for plotting network objects using ggplot2, with additional control over graphical parameters that are not supported by the `ggnet` function. Please visit [http://github.com/briatte/ggnet](http://github.com/briatte/ggnet) for the latest version of ggnet2, and [https://briatte.github.io/ggnet](https://briatte.github.io/ggnet) for a vignette that contains many examples and explanations.

**Usage**

```r
ngnet2(net, mode = "fruchtermanreingold", layout.par = NULL,
layout.exp = 0, alpha = 1, color = "grey75", shape = 19, size = 9,
max.size = 9, na.rm = NA, palette = NULL, alpha.palette = NULL,
alpha.legend = NA, color.palette = palette, color.legend = NA,
shape.palette = NULL, shape.legend = NA, size.palette = NULL,
size.legend = NA, size.zero = FALSE, size.cut = FALSE, size.min = NA,
size.max = NA, label = FALSE, label.alpha = 1, label.color = "black",
label.size = max.size/2, label.trim = FALSE, node.alpha = alpha,
node.color = color, node.label = label, node.shape = shape,
node.size = size, edge.alpha = 1, edge.color = "grey50",
edge.lty = "solid", edge.size = 0.25, edge.label = NULL,
edge.label.alpha = 1, edge.label.color = label.color,
edge.label.fill = "white", edge.label.size = max.size/2, arrow.size = 0,
arrow.gap = 0, arrow.type = "closed", legend.size = 9,
legend.position = "right", ...)```
Arguments

net an object of class network, or any object that can be coerced to this class, such as an adjacency or incidence matrix, or an edge list: see edgeset.constructors and network for details. If the object is of class igraph and the intergraph package is installed, it will be used to convert the object: see asNetwork for details.

mode a placement method from those provided in the sna package: see gplot.layout for details. Also accepts the names of two numeric vertex attributes of net, or a matrix of numeric coordinates, in which case the first two columns of the matrix are used. Defaults to the Fruchterman-Reingold force-directed algorithm.

layout.par options to be passed to the placement method, as listed in gplot.layout. Defaults to NULL.

layout.exp a multiplier to expand the horizontal axis if node labels get clipped: see expand_range for details. Defaults to 0 (no expansion).

alpha the level of transparency of the edges and nodes, which might be a single value, a vertex attribute, or a vector of values. Also accepts "mode" on bipartite networks (see 'Details'). Defaults to 1 (no transparency).

color the color of the nodes, which might be a single value, a vertex attribute, or a vector of values. Also accepts "mode" on bipartite networks (see 'Details'). Defaults to grey75.

shape the shape of the nodes, which might be a single value, a vertex attribute, or a vector of values. Also accepts "mode" on bipartite networks (see 'Details'). Defaults to Y (solid circle).

size the size of the nodes, in points, which might be a single value, a vertex attribute, or a vector of values. Also accepts "indegree", "outdegree", "degree" or "freeman" to size the nodes by their unweighted degree centrality ("degree" and "freeman" are equivalent): see degree for details. All node sizes must be strictly positive. Also accepts "mode" on bipartite networks (see 'Details'). Defaults to 9.

max_size the maximum size of the node when size produces nodes of different sizes, in points. Defaults to 9.

na.rm whether to subset the network to nodes that are not missing a given vertex attribute. If set to any vertex attribute of net, the nodes for which this attribute is NA will be removed. Defaults to NA (does nothing).

palette the palette to color the nodes, when color is not a color value or a vector of color values. Accepts named vectors of color values, or if RColorBrewer is installed, any ColorBrewer palette name: see brewer_pal and http://colorbrewer2.org/ for details. Defaults to NULL, which will create an array of grayscale color values if color is not a color value or a vector of color values.

alpha.palette the palette to control the transparency levels of the nodes set by alpha when the levels are not numeric values. Defaults to NULL, which will create an array of alpha transparency values if alpha is not a numeric value or a vector of numeric values.

alpha.legend the name to assign to the legend created by alpha when its levels are not numeric values. Defaults to NA (no name).
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>color.palette</td>
<td>the name to assign to the legend created by palette. Defaults to NA (no name).</td>
</tr>
<tr>
<td>color.legend</td>
<td>the name to assign to the legend created by palette when its levels are not numeric values. Defaults to NA (no name).</td>
</tr>
<tr>
<td>shape.palette</td>
<td>the palette to control the shapes of the nodes set by shape when the shapes are not numeric values. Defaults to NULL, which will create an array of shape values if shape is not a numeric value or a vector of numeric values.</td>
</tr>
<tr>
<td>shape.legend</td>
<td>the name to assign to the legend created by shape when its levels are not numeric values. Defaults to NA (no name).</td>
</tr>
<tr>
<td>size.palette</td>
<td>the palette to control the sizes of the nodes set by size when the sizes are not numeric values.</td>
</tr>
<tr>
<td>size.legend</td>
<td>the name to assign to the legend created by size. Defaults to NA (no name).</td>
</tr>
<tr>
<td>size.zero</td>
<td>whether to accept zero-sized nodes based on the value(s) of size. Defaults to FALSE, which ensures that zero-sized nodes are still shown in the plot and its size legend.</td>
</tr>
<tr>
<td>size.cut</td>
<td>whether to cut the size of the nodes into a certain number of quantiles. Accepts TRUE, which tries to cut the sizes into quartiles, or any positive numeric value, which tries to cut the sizes into that many quantiles. If the size of the nodes do not contain the specified number of distinct quantiles, the largest possible number is used. See quantile and cut for details. Defaults to FALSE (does nothing).</td>
</tr>
<tr>
<td>size.min</td>
<td>whether to subset the network to nodes with a minimum size, based on the values of size. Defaults to NA (preserves all nodes).</td>
</tr>
<tr>
<td>size.max</td>
<td>whether to subset the network to nodes with a maximum size, based on the values of size. Defaults to NA (preserves all nodes).</td>
</tr>
<tr>
<td>label</td>
<td>whether to label the nodes. If set to TRUE, nodes are labeled with their vertex names. If set to a vector that contains as many elements as there are nodes in net, nodes are labeled with these. If set to any other vector of values, the nodes are labeled only when their vertex name matches one of these values. Defaults to FALSE (no labels).</td>
</tr>
<tr>
<td>label.alpha</td>
<td>the level of transparency of the node labels, as a numeric value, a vector of numeric values, or as a vertex attribute containing numeric values. Defaults to 1 (no transparency).</td>
</tr>
<tr>
<td>label.color</td>
<td>the color of the node labels, as a color value, a vector of color values, or as a vertex attribute containing color values. Defaults to &quot;black&quot;.</td>
</tr>
<tr>
<td>label.size</td>
<td>the size of the node labels, in points, as a numeric value, a vector of numeric values, or as a vertex attribute containing numeric values. Defaults to max_size / 2 (half the maximum node size), which defaults to 4.5.</td>
</tr>
<tr>
<td>label.trim</td>
<td>whether to apply some trimming to the node labels. Accepts any function that can process a character vector, or a strictly positive numeric value, in which case the labels are trimmed to a fixed-length substring of that length: see substr for details. Defaults to FALSE (does nothing).</td>
</tr>
<tr>
<td>node.alpha</td>
<td>see alpha</td>
</tr>
<tr>
<td>node.color</td>
<td>see color</td>
</tr>
<tr>
<td>node.label</td>
<td>see label</td>
</tr>
<tr>
<td>Argument</td>
<td>Description</td>
</tr>
<tr>
<td>------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>node.shape</td>
<td>see shape</td>
</tr>
<tr>
<td>node.size</td>
<td>see size</td>
</tr>
<tr>
<td>edge.alpha</td>
<td>the level of transparency of the edges. Defaults to the value of alpha, which</td>
</tr>
<tr>
<td></td>
<td>defaults to 1.</td>
</tr>
<tr>
<td>edge.color</td>
<td>the color of the edges, as a color value, a vector of color values, or as an</td>
</tr>
<tr>
<td></td>
<td>edge attribute containing color values. Defaults to &quot;grey50&quot;.</td>
</tr>
<tr>
<td>edge.lty</td>
<td>the linetype of the edges, as a linetype value, a vector of linetype values,</td>
</tr>
<tr>
<td></td>
<td>or as an edge attribute containing linetype values. Defaults to &quot;solid&quot;.</td>
</tr>
<tr>
<td>edge.size</td>
<td>the size of the edges, in points, as a numeric value, a vector of numeric</td>
</tr>
<tr>
<td></td>
<td>values, or as an edge attribute containing numeric values. All edge sizes must</td>
</tr>
<tr>
<td></td>
<td>be strictly positive. Defaults to 0.25.</td>
</tr>
<tr>
<td>edge.label</td>
<td>the labels to plot at the middle of the edges, as a single value, a vector of</td>
</tr>
<tr>
<td></td>
<td>values, or as an edge attribute. Defaults to NULL (no edge labels).</td>
</tr>
<tr>
<td>edge.label.alpha</td>
<td>the level of transparency of the edge labels, as a numeric value, a vector of</td>
</tr>
<tr>
<td></td>
<td>numeric values, or as an edge attribute containing numeric values. Defaults to</td>
</tr>
<tr>
<td></td>
<td>1 (no transparency).</td>
</tr>
<tr>
<td>edge.label.color</td>
<td>the color of the edge labels, as a color value, a vector of color values, or</td>
</tr>
<tr>
<td></td>
<td>as an edge attribute containing color values. Defaults to label.color, which</td>
</tr>
<tr>
<td></td>
<td>defaults to &quot;black&quot;.</td>
</tr>
<tr>
<td>edge.label.fill</td>
<td>the background color of the edge labels. Defaults to &quot;white&quot;.</td>
</tr>
<tr>
<td>edge.label.size</td>
<td>the size of the edge labels, in points, as a numeric value, a vector of numeric</td>
</tr>
<tr>
<td></td>
<td>values, or as an edge attribute containing numeric values. All edge label sizes</td>
</tr>
<tr>
<td></td>
<td>must be strictly positive. Defaults to max.size / 2(half the maximum node size), which defaults to 4.5.</td>
</tr>
<tr>
<td>arrow.size</td>
<td>the size of the arrows for directed network edges, in points. See arrow for</td>
</tr>
<tr>
<td></td>
<td>details. Defaults to 0 (no arrows).</td>
</tr>
<tr>
<td>arrow.gap</td>
<td>a setting aimed at improving the display of edge arrows by plotting slightly</td>
</tr>
<tr>
<td></td>
<td>shorter edges. Accepts any value between 0 and 1, where a value of 0.05 will</td>
</tr>
<tr>
<td></td>
<td>generally achieve good results when the size of the nodes is reasonably small.</td>
</tr>
<tr>
<td></td>
<td>Defaults to 0 (no shortening).</td>
</tr>
<tr>
<td>arrow.type</td>
<td>the type of the arrows for directed network edges. See arrow for details.</td>
</tr>
<tr>
<td></td>
<td>Defaults to &quot;closed&quot;.</td>
</tr>
<tr>
<td>legend.size</td>
<td>the size of the legend symbols and text, in points. Defaults to 9.</td>
</tr>
<tr>
<td>legend.position</td>
<td>the location of the plot legend(s). Accepts all legend.position values sup-</td>
</tr>
<tr>
<td></td>
<td>ported by theme. Defaults to &quot;right&quot;.</td>
</tr>
<tr>
<td>...</td>
<td>other arguments passed to the geom_text object that sets the node labels: see</td>
</tr>
<tr>
<td></td>
<td>geom_text for details.</td>
</tr>
</tbody>
</table>
Details

The degree centrality measures that can be produced through the size argument will take the directedness of the network into account, but will be unweighted. To compute weighted network measures, see the tnet package by Tore Opsahl (help("tnet", package = "tnet").

The nodes of bipartite networks can be mapped to their mode by passing the "mode" argument to any of alpha, color, shape and size, in which case the nodes of the primary mode will be mapped as "actor", and the nodes of the secondary mode will be mapped as "event".

Author(s)

Moritz Marbach and Francois Briatte, with help from Heike Hoffmann, Pedro Jordano and Ming-Yu Liu

See Also

ggnet in this package, gplot in the sna package, and plot.network in the network package

Examples

library(network)

# random adjacency matrix
x <- 10
ndyads <- x * (x - 1)
density <- x / ndyads
m <- matrix(0, nrow = x, ncol = x)
dimnames(m) <- list(letters[1:x], letters[1:x])
m[row(m) != col(m)] <- runif(ndyads) < density
m

# random undirected network
n <- network::network(m, directed = FALSE)
n
ggnet2(n, label = TRUE)
ggnet2(n, label = TRUE, shape = 15)
ggnet2(n, label = TRUE, shape = 15, color = "black", label.color = "white")

# add vertex attribute
x = network.vertex.names(n)
x = ifelse(x %in% c("a", "e", "i"), "vowel", "consonant")
n %v% "phono" = x

ggnet2(n, color = "phono")
ggnet2(n, color = "phono", palette = c("vowel" = "gold", "consonant" = "grey"))
ggnet2(n, shape = "phono", color = "phono")

if (require(RColorBrewer)) {

    # random groups
    n %v% "group" <- sample(LETTERS[1:3], 10, replace = TRUE)
ggnetworkmap

- Plot a network with ggplot2 suitable for overlay on a ggmap:: map ggplot, or other ggplot

Description

This is a descendent of the original ggnet function. ggnet added the innovation of plotting the network geographically. However, ggnet needed to be the first object in the ggplot chain. ggnetworkmap does not. If passed a ggplot object as its first argument, such as output from ggmap, ggnetworkmap will plot on top of that chart, looking for vertex attributes lon and lat as coordinates. Otherwise, ggnetworkmap will generate coordinates using the Fruchterman-Reingold algorithm.

Usage

ggnetworkmap(gg, net, size = 3, alpha = 0.75, weight, node.group, node.color = NULL, node.alpha = NULL, ring.group, segment.alpha = NULL, segment.color = "grey", great.circles = FALSE, segment.size = 0.25, arrow.size = 0, label.nodes = FALSE, label.size = size/2, ...)

Arguments

- **gg**: an object of class ggplot.
- **net**: an object of class network, or any object that can be coerced to this class, such as an adjacency or incidence matrix, or an edge list: see edgeset.constructors and network for details. If the object is of class igraph and the intergraph package is installed, it will be used to convert the object: see asNetwork for details.
size  size of the network nodes. Defaults to 3. If the nodes are weighted, their area is proportionally scaled up to the size set by size.

alpha  a level of transparency for nodes, vertices and arrows. Defaults to 0.75.

weight  if present, the unquoted name of a vertex attribute in data. Otherwise nodes are unweighted.

node.group  NULL, the default, or the unquoted name of a vertex attribute that will be used to determine the color of each node.

node.color  If node.group is null, a character string specifying a color.

node.alpha  transparency of the nodes. Inherits from alpha.

ring.group  if not NULL, the default, the unquoted name of a vertex attribute that will be used to determine the color of each node border.

segment.alpha  transparency of the vertex links. Inherits from alpha

segment.color  color of the vertex links. Defaults to "grey".

great.circles  whether to draw edges as great circles using the geosphere package. Defaults to FALSE

segment.size  size of the vertex links, as a vector of values or as a single value. Defaults to 0.25.

arrow.size  size of the vertex arrows for directed network plotting, in centimeters. Defaults to 0.

label.nodes  label nodes with their vertex names attribute. If set to TRUE, all nodes are labelled. Also accepts a vector of character strings to match with vertex names.

label.size  size of the labels. Defaults to size / 2.

...  other arguments supplied to geom_text for the node labels. Arguments pertaining to the title or other items can be achieved through ggplot2 methods.

Details

This is a function for plotting graphs generated by network or igraph in a more flexible and elegant manner than permitted by gnet. The function does not need to be the first plot in the ggplot chain, so the graph can be plotted on top of a map or other chart. Segments can be straight lines, or plotted as great circles. Note that the great circles feature can produce odd results with arrows and with vertices beyond the plot edges; this is a ggplot2 limitation and cannot yet be fixed. Nodes can have two color schemes, which are then plotted as the center and ring around the node. The color schemes are selected by adding scale_fill_ or scale_color_ just like any other ggplot2 plot. If there are no rings, scale_color sets the color of the nodes. If there are rings, scale_color sets the color of the rings, and scale_fill sets the color of the centers. Note that additional arguments in the ... are passed to geom_text for plotting labels.

Author(s)

Amos Elberg <amos.elberg@gmail.com>. Original by Moritz Marbach <mmarbach@mail.uni-mannheim.de>, Francois Briatte <f.briatte@gmail.com>
Examples

# small function to display plots only if it's interactive
p_ <- GGally:::print_if_interactive
invisible(lapply(c("ggplot2", "maps", "network", "sna"), base::library, character.only = TRUE))

## Example showing great circles on a simple map of the USA

airports <- read.csv("http://datasets.flowingdata.com/tuts/maparcs/airports.csv", header = TRUE)
rownames(airports) <- airports$iata

# select some random flights
set.seed(1234)
flights <- data.frame(
  origin = sample(airports[200:400, ]$iata, 200, replace = TRUE),
  destination = sample(airports[200:400, ]$iata, 200, replace = TRUE)
)

# convert to network
flights <- network(flights, directed = TRUE)

# add geographic coordinates
flights %>% mutate(lat = airports[ network.vertex.names(flights), "lat" ])
flights %>% mutate(lon = airports[ network.vertex.names(flights), "long" ])

# drop isolated airports
delete.vertices(flights, which(degree(flights) < 2))

# compute degree centrality
flights %>% mutate("degree" = degree(flights, gmode = "digraph")

# add random groups
flights %>% mutate("mygroup" = sample(letters[1:4], network.size(flights), replace = TRUE)

# create a map of the USA
usa <- ggplot(map_data("usa"), aes(x = long, y = lat)) +
  geom_polygon(aes(group = -group), color = "grey65",
  fill = "#f9f9f9", size = 0.2)

# overlay network data to map
p <- ggnetworkmap(
  usa, flights, size = 4, great.circles = TRUE,
  node.group = mygroup, segment.color = "steelblue",
  ring.group = degree, weight = degree
)
p(p)

## Exploring a community of spambots found on Twitter
## Data by Amos Elberg: see ?twitter_spambots for details

data(twitter_spambots)
# create a world map
world <- fortify(map("world", plot = FALSE, fill = TRUE))
world <- ggplot(world, aes(x = long, y = lat)) +
  geom_polygon(aes(group = group), color = "grey65",
               fill = "#f9f9f9", size = 0.2)

# view global structure
p <- ggnetworkmap(world, twitter_spambots)
p_(p)

# domestic distribution
p <- ggnetworkmap(net = twitter_spambots)
p_(p)

# topology
p <- ggnetworkmap(net = twitter_spambots, arrow.size = 0.5)
p_(p)

# compute indegree and outdegree centrality
twitter_spambots %>% "indegree" <- degree(twitter_spambots, cmode = "indegree")
twitter_spambots %>% "outdegree" <- degree(twitter_spambots, cmode = "outdegree")

p <- ggnetworkmap(
  net = twitter_spambots,
  arrow.size = 0.5,
  node.group = indegree,
  ring.group = outdegree, size = 4
) +
  scale_fill_continuous("Indegree", high = "red", low = "yellow") +
  labs(color = "Outdegree")
p_(p)

# show some vertex attributes associated with each account
p <- ggnetworkmap(
  net = twitter_spambots,
  arrow.size = 0.5,
  node.group = followers,
  ring.group = friends,
  size = 4,
  weight = indegree,
  label.nodes = TRUE, vjust = -1.5
) +
  scale_fill_continuous("Followers", high = "red", low = "yellow") +
  labs(color = "Friends") +
  scale_color_continuous(low = "lightgreen", high = "darkgreen")
p_(p)
Description

gnostic - Plot matrix of statistical model diagnostics

Usage

gnostic(model, ..., columnsX = attr(data, "var_x"), columnsY = c(".resid", "sigma", ".hat", ".cooks")
, columnLabelsX = attr(data, "var_x_label"),
, columnLabelsY = gsub("\", ",", gsub("\", "", columnsY)),
, xlab = "explanatory variables", ylab = "diagnostics",
, title = paste(deparse(model$call, width.cutoff = 500L), collapse = "\n"),
, continuous = list(default = ggally_points, .fitted = ggally_points, .se.fit =
  ggally_nostic_se_fit, .resid = ggally_nostic_resid, .hat =
  ggally_nostic_hat, .sigma = ggally_nostic_sigma, .cooks =
  ggally_nostic_cooksd)
, combo = list(default = ggally_box_no_facet, fitted = ggally_box_no_facet,
  .se.fit = ggally_nostic_se_fit, .resid = ggally_nostic_resid, .hat =
  ggally_nostic_hat, .sigma = ggally_nostic_sigma, .cooks =
  ggally_nostic_cooksd)
, discrete = list(default = ggally_ratio, .fitted = ggally_ratio, .se.fit =
  ggally_ratio, .resid = ggally_ratio, .hat = ggally_ratio, .sigma =
  ggally_ratio, .cooks = ggally_ratio, .std.resid = ggally_ratio)
, data = broomify(model))

Arguments

model statistical model object such as output from stats::lm or stats::glm

... arguments passed directly to ggduo

columnsX columns to be displayed in the plot matrix. Defaults to the predictor columns of the model

columnsY rows to be displayed in the plot matrix. Defaults to residuals, leave one out sigma value, diagonal of the hat matrix, and Cook’s Distance. The possible values are the response variables in the model and the added columns provided by broom::augment(model). See details for more information.

columnLabelsX, columnLabelsY column and row labels to display in the plot matrix

xlab, ylab, title plot matrix labels passed directly to ggmatrix

continuous, combo, discrete list of functions for each y variable. See details for more information.

data data defaults to a ‘broomify’ed model object. This object will contain information about the X variables, Y variables, and multiple broom outputs. See broomify(model) for more information

‘columnsY’ broom::augment() collects data from the supplied model and returns a data.frame with the following columns (taken directly from broom documentation). These columns are the only allowed values in the columnsY parameter to gnostic.
.resid  Residuals
.hat    Diagonal of the hat matrix
.sigma  Estimate of residual standard deviation when corresponding observation is dropped from
         model
.cooksd  Cooks distance, cooks.distance
.fitted  Fitted values of model
.se.fit  Standard errors of fitted values
.std.resid Standardized residuals

**response variable name** The response variable in the model may be added. Such as "mpg" in the
model `lm(mpg ~ ., data = mtcars)`

### 'continuous', 'combo', 'discrete' types

Similar to **ggduo** and **ggpairs**, functions may be supplied to display the different column types. However, since the Y rows are fixed, each row has its own corresponding function in each of the
plot types: continuous, combo, and discrete. Each plot type list can have keys that correspond
to the broom::augment() output: ".fitted", ".resid", ".std.resid", ".sigma", ".se.fit", ".hat", ".cooks". An extra key, "default", is used to plot the response variables of the model
if they are included. Having a function for each diagnostic allows for very fine control over the
diagnostics plot matrix. The functions for each type list are wrapped into a switch function that
calls the function corresponding to the y variable being plotted. These switch functions are then
passed directly to the types parameter in **ggduo**.

### Examples

```r
# small function to display plots only if it's interactive
p_ <- GGally::print_if_interactive
data(mtcars)

# use mtcars dataset and alter the 'am' column to display actual name values
mtc <- mtcars
mtc$am <- c("0" = "automatic", "1" = "manual")[as.character(mtc$am)]

# step the complete model down to a smaller model
mod <- stats::step(stats::lm(mpg ~ ., data = mtc), trace = FALSE)

# display using defaults
pm <- ggnostic(mod)
p_(pm)

# color by am value
pm <- ggnostic(mod, mapping = ggplot2::aes(color = am))
p_(pm)

# turn resid smooth error ribbon off
pm <- ggnostic(mod, continuous = list(.resid = wrap("nostic_resid", se = FALSE)))
p_(pm)
```

## ggpairs

### Description

Make a matrix of plots with a given data set

### Usage

```r
ggpairs(data, mapping = NULL, columns = 1:ncol(data), title = NULL,
upper = list(continuous = "cor", combo = "box_no_facet", discrete = 
"facetbar", na = "na"), lower = list(continuous = "points", combo = 
"facethist", discrete = "facetbar", na = "na"), diag = list(continuous = 
"densityDiag", discrete = "barDiag", na = "naDiag"), params = NULL, ..., 
xlab = NULL, ylab = NULL, axisLabels = c("show", "internal", "none"),
columnLabels = colnames(data[columns]), labeller = "label_value",
switch = NULL, showStrips = NULL, legend = NULL,
cardinality_threshold = 15, legends = stop("deprecated"))
```

### Arguments

- **data**: data set using. Can have both numerical and categorical data.
- **mapping**: aesthetic mapping (besides x and y). See `aes()`. If mapping is numeric, columns will be set to the mapping value and mapping will be set to NULL.
- **columns**: which columns are used to make plots. Defaults to all columns.
- **title, xlab, ylab**: title, x label, and y label for the graph
- **upper**: see Details
- **lower**: see Details
- **diag**: see Details
- **params**: deprecated. Please see `wrap_fn_with_param_arg`
- **...**: deprecated. Please use mapping
- **axisLabels**: either "show" to display axisLabels, "internal" for labels in the diagonal plots, or "none" for no axis labels
- **columnLabels**: label names to be displayed. Defaults to names of columns being used.
### ggplot2: ggpairs

**labeller**
- labeller for facets. See **labellers**. Common values are "label_value" (default) and "label_parsed".

**switch**
- switch parameter for facet_grid. See ggplot2::facet_grid. By default, the labels are displayed on the top and right of the plot. If "x", the top labels will be displayed to the bottom. If "y", the right-hand side labels will be displayed to the left. Can also be set to "both"

**showStrips**
- boolean to determine if each plot’s strips should be displayed. NULL will default to the top and right side plots only. TRUE or FALSE will turn all strips on or off respectively.

**legend**
- May be the two objects described below or the default NULL value. The legend position can be moved by using ggplot2’s theme element `theme(legend.position = "bottom")`
  - **a numeric vector of length 2** provides the location of the plot to use the legend for the plot matrix’s legend. Such as `legend = c(3, 5)` which will use the legend from the plot in the third row and fifth column
  - **a single numeric value** provides the location of a plot according to the display order. Such as `legend = 3` in a plot matrix with 2 rows and 5 columns displayed by column will return the plot in position `c(1, 2)`
  - **a object from grab_legend()** a predetermined plot legend that will be displayed directly

**cardinality_threshold**
- maximum number of levels allowed in a character / factor column. Set this value to NULL to not check factor columns. Defaults to 15

**legends**
- deprecated

### Details

**upper** and **lower** are lists that may contain the variables ‘continuous’, ‘combo’, ‘discrete’, and ‘na’. Each element of the list may be a function or a string. If a string is supplied, it must implement one of the following options:

- **continuous** exactly one of (`points`, `smooth`, `smooth_loess`, `density`, `cor`, `blank`). This option is used for continuous X and Y data.
- **combo** exactly one of (`box`, `box_no_facet`, `dot`, `dot_no_facet`, `facethist`, `facetdensity`, `denstrip`, `blank`). This option is used for either continuous X and categorical Y data or categorical X and continuous Y data.
- **discrete** exactly one of (`facetbar`, `ratio`, `blank`). This option is used for categorical X and Y data.
- **na** exactly one of (`na`, `blank`). This option is used when all X data is NA, all Y data is NA, or either all X or Y data is NA.

**diag** is a list that may only contain the variables ‘continuous’, ‘discrete’, and ‘na’. Each element of the diag list is a string implementing the following options:

- **continuous** exactly one of (`densityDiag`, `barDiag`, `blankDiag`). This option is used for continuous X data.
- **discrete** exactly one of (`barDiag`, `blankDiag`). This option is used for categorical X and Y data.
na exactly one of ('naDiag', 'blankDiag'). This option is used when all X data is NA.

If 'blank' is ever chosen as an option, then ggpairs will produce an empty plot.

If a function is supplied as an option to upper, lower, or diag, it should implement the function api of function(data, mapping, ...){#make ggplot2 plot}. If a specific function needs its parameters set, wrap(fn, param1 = val1, param2 = val2) the function with its parameters.

Value

ggmatrix object that if called, will print

Author(s)

Barret Schloerke <schloerke@gmail.com>, Jason Crowley <crowley.jason.s@gmail.com>, Di Cook <dicook@iastate.edu>, Heike Hofmann <hofmann@iastate.edu>, Hadley Wickham <h.wickham@gmail.com>

References


See Also

wrap v1_ggmatrix_theme

Examples

# small function to display plots only if it's interactive
p_ <- GGally::print_if_interactive

## Quick example, with and without colour
data(flea)
ggpairs(flea, columns = 2:4)  
pm <- ggpairs(flea, columns = 2:4, ggplot2::aes(colour=species))  
p_(pm)
# Note: colour should be categorical, else you will need to reset
# the upper triangle to use points instead of trying to compute corr

data(tips, package = "reshape")
p_ <- ggpairs(tips[, 1:3])  
p_(pm)

pm <- ggpairs(tips[, 1:3], columnLabels = c("Total Bill", "Tip", "Sex"))  
p_(pm)

pm <- ggpairs(tips, upper = "blank")  
p_(pm)

## Plot Types
# Change default plot behavior
pm <- ggpairs(
    tips[, c(1, 3, 4, 2)]},

```r
# small function to display plots only if it's interactive
p_ <- GGally::print_if_interactive

## Quick example, with and without colour
data(flea)
ggpairs(flea, columns = 2:4)  
pm <- ggpairs(flea, columns = 2:4, ggplot2::aes(colour=species))  
p_(pm)
# Note: colour should be categorical, else you will need to reset
# the upper triangle to use points instead of trying to compute corr

data(tips, package = "reshape")
p_ <- ggpairs(tips[, 1:3])  
p_(pm)

pm <- ggpairs(tips[, 1:3], columnLabels = c("Total Bill", "Tip", "Sex"))  
p_(pm)

pm <- ggpairs(tips, upper = "blank")  
p_(pm)

## Plot Types
# Change default plot behavior
pm <- ggpairs(
    tips[, c(1, 3, 4, 2)]},
```

```r
# small function to display plots only if it's interactive
p_ <- GGally::print_if_interactive

## Quick example, with and without colour
data(flea)
ggpairs(flea, columns = 2:4)  
pm <- ggpairs(flea, columns = 2:4, ggplot2::aes(colour=species))  
p_(pm)
# Note: colour should be categorical, else you will need to reset
# the upper triangle to use points instead of trying to compute corr

data(tips, package = "reshape")
p_ <- ggpairs(tips[, 1:3])  
p_(pm)

pm <- ggpairs(tips[, 1:3], columnLabels = c("Total Bill", "Tip", "Sex"))  
p_(pm)

pm <- ggpairs(tips, upper = "blank")  
p_(pm)

## Plot Types
# Change default plot behavior
pm <- ggpairs(
    tips[, c(1, 3, 4, 2)]},
```
upper = list(continuous = "density", combo = "box_no_facet"),
lower = list(continuous = "points", combo = "dot_no_facet")
)
p_(p)
# Supply Raw Functions (may be user defined functions!)
pm <- ggpairs( 
tips[, c(1, 3, 4, 2)],
upper = list(continuous = ggally_density, combo = ggally_box_no_facet),
lower = list(continuous = ggally_points, combo = ggally_dot_no_facet)
)
p_(p)

# Use sample of the diamonds data
data(diamonds, package="ggplot2")
diamonds.samp <- diamonds[sample(1:dim(diamonds)[1], 1000), ]

# Different aesthetics for different plot sections and plot types
pm <- ggpairs( 
diamonds.samp[, 1:5],
mapping = ggplot2::aes(color = cut),
upper = list(continuous = wrap("density", alpha = 0.5), combo = "box_no_facet"),
lower = list(continuous = wrap("points", alpha = 0.3), combo = wrap("dot_no_facet", alpha = 0.4)),
title = "Diamonds"
)
p_(p)

## Axis Label Variations
## Only Variable Labels on the diagonal (no axis labels)
pm <- ggpairs(tips[, 1:3], axisLabels="internal")
p_(p)
## Only Variable Labels on the outside (no axis labels)
pm <- ggpairs(tips[, 1:3], axisLabels="none")
p_(p)

## Facet Label Variations
# Default:
df_x <- rnorm(100)
df_y <- df_x + rnorm(100, 0, 0.1)
df <- data.frame(x = df_x, y = df_y, c = sqrt(df_x^2 + df_y^2))
pm <- ggpairs( 
df,
columnLabels = c("alpha[foo]", "alpha[bar]", "sqrt(alpha[foo]^2 + alpha[bar]^2]")
)
p_(p)
## Parsed labels:
pm <- ggpairs( 
df,
columnLabels = c("alpha[foo]", "alpha[bar]", "sqrt(alpha[foo]^2 + alpha[bar]^2]"),
labeller = "label_parsed"
)
p_(p)

## Plot Insertion Example
ggparcoord

A function for plotting static parallel coordinate plots, utilizing the ggplot2 graphics package.

Usage

ggparcoord(data, columns = 1:ncol(data), groupColumn = NULL, scale = "std", scaleSummary = "mean", centerObsID = 1, missing = "exclude", order = columns, showPoints = FALSE, splineFactor = FALSE, alphaLines = 1, boxplot = FALSE, shadeBox = NULL, mapping = NULL, title = "")

Arguments

data the dataset to plot

columns a vector of variables (either names or indices) to be axes in the plot
groupColumn a single variable to group (color) by

scale method used to scale the variables (see Details)
scaleSummary if scale="center", summary statistic to univariately center each variable by
centerObsID if scale="centerObs", row number of case plot should univariately be centered on
missing method used to handle missing values (see Details)
order method used to order the axes (see Details)
showPoints logical operator indicating whether points should be plotted or not
splineFactor logical or numeric operator indicating whether spline interpolation should be
used. Numeric values will multiplied by the number of columns, TRUE will
default to cubic interpolation, AsIs to set the knot count directly and 0, FALSE,
or non-numeric values will not use spline interpolation.
alphaLines value of alpha scaler for the lines of the parcoord plot or a column name of the
data
boxplot logical operator indicating whether or not boxplots should underlay the distri-
bution of each variable
shadeBox color of underlaying box which extends from the min to the max for each vari-
able (no box is plotted if shadeBox == NULL)
mapping aes string to pass to ggplot object
title character string denoting the title of the plot

Details

scale is a character string that denotes how to scale the variables in the parallel coordinate plot. Options:
  • std: univariately, subtract mean and divide by standard deviation
  • robust: univariately, subtract median and divide by median absolute deviation
  • unimiMax: univariately, scale so the minimum of the variable is zero, and the maximum is
    one
  • globalMinMax: no scaling is done; the range of the graphs is defined by the global minimum
    and the global maximum
  • center: use unimiMax to standardize vertical height, then center each variable at a value
    specified by the scaleSummary param
  • centerObs: use unimiMax to standardize vertical height, then center each variable at the
    value of the observation specified by the centerObsID param

missing is a character string that denotes how to handle missing missing values. Options:
  • exclude: remove all cases with missing values
  • mean: set missing values to the mean of the variable
  • median: set missing values to the median of the variable
  • min10: set missing values to 10% below the minimum of the variable
  • random: set missing values to value of randomly chosen observation on that variable

order is either a vector of indices or a character string that denotes how to order the axes (variables)
of the parallel coordinate plot. Options:
  • (default): order by the vector denoted by columns
  • (given vector): order by the vector specified
• anyClass: order variables by their separation between any one class and the rest (as opposed to their overall variation between classes). This is accomplished by calculating the F-statistic for each class vs. the rest, for each axis variable. The axis variables are then ordered (decreasing) by their maximum of k F-statistics, where k is the number of classes.

• allClass: order variables by their overall F statistic (decreasing) from an ANOVA with groupColumn as the explanatory variable (note: it is required to specify a groupColumn with this ordering method). Basically, this method orders the variables by their variation between classes (most to least).

• skewness: order variables by their sample skewness (most skewed to least skewed)

• Outlying: order by the scagnostic measure, Outlying, as calculated by the package scagnostics. Other scagnostic measures available to order by are Skewed, Clumpy, Sparse, Striated, Convex, Skinny, Stringy, and Monotonic. Note: To use these methods of ordering, you must have the scagnostics package loaded.

Value

ggplot object that if called, will print

Author(s)

Jason Crowley <crowley.jason.s@gmail.com>, Barret Schloerke <schloerke@gmail.com>, Di Cook <dicook@iastate.edu>, Heike Hofmann <hofmann@iastate.edu>, Hadley Wickham <h.wickham@gmail.com>

Examples

# small function to display plots only if it's interactive
p_ <- GGally::print_if_interactive

# use sample of the diamonds data for illustrative purposes
data(diamonds, package="ggplot2")
diamonds.samp <- diamonds[sample(1:dim(diamonds)[1], 100), ]

# basic parallel coordinate plot, using default settings
p <- ggparcoord(data = diamonds.samp, columns = c(1, 5:10))
p_(p)

# this time, color by diamond cut
p <- ggparcoord(data = diamonds.samp, columns = c(1, 5:10), groupColumn = 2)
p_(p)

# underlay univariate boxplots, add title, use uniminmax scaling
p <- ggparcoord(data = diamonds.samp, columns = c(1, 5:10), groupColumn = 2,
                scale = "uniminmax", boxplot = TRUE, title = "Parallel Coord. Plot of Diamonds Data")
p_(p)

# utilize ggplot2 aes to switch to thicker lines
p <- ggparcoord(data = diamonds.samp, columns = c(1, 5:10), groupColumn = 2,
                title ="Parallel Coord. Plot of Diamonds Data", mapping = ggplot2::aes(size = 1)) +
ggplot2::scale_size_identity()
p_(p)
# basic parallel coord plot of the msleep data, using 'random' imputation and # coloring by diet (can also use variable names in the columns and groupColumn # arguments)
data(msleep, package="ggplot2")
p <- ggparcoord(data = msleep, columns = 6:11, groupColumn = "vore", missing = # 'random', scale = "uniminmax")
  p_(p)

# center each variable by its median, using the default missing value handler, # 'exclude'
p <- ggparcoord(data = msleep, columns = 6:11, groupColumn = "vore", scale = # "center", scaleSummary = "median")
  p_(p)

# with the iris data, order the axes by overall class (Species) separation using # the anyClass option
p <- ggparcoord(data = iris, columns = 1:4, groupColumn = 5, order = "anyClass")
  p_(p)

# add points to the plot, add a title, and use an alpha scalar to make the lines # transparent
p <- ggparcoord(data = iris, columns = 1:4, groupColumn = 5, order = "anyClass",
  showPoints = TRUE, title = "Parallel Coordinate Plot for the Iris Data",
  alaphalines = 0.3)
  p_(p)

# color according to a column
iris2 <- iris
iris2$alphaLevel <- c("setosa" = 0.2, "versicolor" = 0.3, "virginica" = 0)[iris2$Species]
p <- ggparcoord(data = iris2, columns = 1:4, groupColumn = 5, order = "anyClass",
  showPoints = TRUE, title = "Parallel Coordinate Plot for the Iris Data",
  alaphalines = "alphaLevel")
  p_(p)

## Use splines on values, rather than lines (all produce the same result)
columns <- c(1, 5:10)
p <- ggparcoord(diamonds.samp, columns, groupColumn = 2, splineFactor = TRUE)
  p_(p)
p <- ggparcoord(diamonds.samp, columns, groupColumn = 2, splineFactor = 3)
  p_(p)
splineFactor <- length(columns) * 3
p <- ggparcoord(diamonds.samp, columns, groupColumn = 2, splineFactor = I(splineFactor))
  p_(p)

---

**ggscatmat**

*ggscatmat - a traditional scatterplot matrix for purely quantitative variables*
**Description**

This function makes a scatterplot matrix for quantitative variables with density plots on the diagonal and correlation printed in the upper triangle.

**Usage**

```r
ggscatmat(data, columns = 1:ncol(data), color = NULL, alpha = 1, corMethod = "pearson")
```

**Arguments**

- **data**: a data matrix. Should contain numerical (continuous) data.
- **columns**: an option to choose the column to be used in the raw dataset. Defaults to `1:ncol(data)`.
- **color**: an option to group the dataset by the factor variable and color them by different colors. Defaults to `NULL`.
- **alpha**: an option to set the transparency in scatterplots for large data. Defaults to `1`.
- **corMethod**: method argument supplied to `cor`.

**Author(s)**

Mengjia Ni, Di Cook <dicook@monash.edu>

**Examples**

```r
data(flea)
ggscatmat(flea, columns = 2:4)
ggscatmat(flea, columns = 2:4, color = "species")
```

---

**ggsurv**

*Survival curves with ggplot2*

**Description**

This function produces Kaplan-Meier plots using ggplot2. As a first argument it needs a `survfit` object, created by the `survival` package. Default settings differ for single stratum and multiple strata objects.

**Usage**

```r
ggsurv(s, CI = "def", plot.cens = TRUE, surv.col = "gg.def", 
cens.col = "gg.def", lty.est = 1, lty.ci = 2, size.est = 0.5, 
size.ci = size.est, cens.size = 2, cens.shape = 3, back.white = FALSE, 
xlab = "Time", ylab = "Survival", main = ", order.legend = TRUE)```
Arguments

- **s**: an object of class `survfit`
- **CI**: should a confidence interval be plotted? Defaults to TRUE for single stratum objects and FALSE for multiple strata objects.
- **plot.cens**: mark the censored observations?
- **surv.col**: colour of the survival estimate. Defaults to black for one stratum, and to the default ggplot2 colours for multiple strata. Length of vector with colour names should be either 1 or equal to the number of strata.
- **cens.col**: colour of the points that mark censored observations.
- **lty.est**: linetype of the survival curve(s). Vector length should be either 1 or equal to the number of strata.
- **lty.ci**: linetype of the bounds that mark the 95% CI.
- **size.est**: line width of the survival curve
- **size.ci**: line width of the 95% CI
- **cens.size**: point size of the censoring points
- **cens.shape**: shape of the points that mark censored observations.
- **back.white**: if TRUE the background will not be the default grey of ggplot2 but will be white with borders around the plot.
- **xlab**: the label of the x-axis.
- **ylab**: the label of the y-axis.
- **main**: the plot label.
- **order.legend**: boolean to determine if the legend display should be ordered by final survival time

Value

An object of class `ggplot`

Author(s)

Edwin Thoen <edwinthoen@gmail.com>

Examples

```r
if (require(survival) && require(scales)) {
  data(lung, package = "survival")
  sf.lung <- survival::survfit(Surv(time, status) ~ 1, data = lung)
  ggsurv(sf.lung)

  # Multiple strata examples
  sf.sex <- survival::survfit(Surv(time, status) ~ sex, data = lung)
  pl.sex <- ggsurv(sf.sex)
  pl.sex
```
# Adjusting the legend of the ggsurv fit

pl.sex +  
ggplot2::guides(linetype = FALSE) +  
ggplot2::scale_colour_discrete(  
  name = 'Sex',  
  breaks = c(1,2),  
  labels = c('Male', 'Female'))

# We can still adjust the plot after fitting
data(kidney, package = "survival")
sf.kid <- survival::survfit(Surv(time, status) ~ disease, data = kidney)
pl.kid <- ggsurv(sf.kid, plot.cens = FALSE)
pl.kid

# Zoom in to first 80 days
pl.kid + ggplot2::coord_cartesian(xlim = c(0, 80), ylim = c(0.45, 1))

# Add the diseases names to the plot and remove legend
pl.kid +  
ggplot2::annotate("text",  
  label = c("PKD", "Other", "GN", "AN"),  
  x = c(90, 125, 5, 60),  
  y = c(0.8, 0.65, 0.55, 0.30),  
  size = 5,  
  colour = scales::hue_pal(
    h = c(0, 360) + 15,  
    c = 100,  
    l = 65,  
    h.start = 0,  
    direction = 1
  )(4)  
) +  
ggplot2::guides(color = FALSE, linetype = FALSE)

---

**ggts**

*Multiple Time Series*

**Description**

GGally implementation of ts.plot. Wraps around the ggduo function and removes the column strips

**Usage**

`ggts(..., columnLabelsX = NULL, xlab = "time")`
**glyphplot**

**Arguments**

- ... supplied directly to `ggduo`
- columnLabelsX remove top strips for the X axis by default
- xlab defaults to "time"

**Value**

ggmatrix object

**Examples**

```r
ggts(pigs, "time", c("gilts", "profit", "s_per_herdsz", "production", "herdsz"))
```

---

**Description**

Glyph plot class

**Usage**

```r
glyphplot(data, width, height, polar, x_major, y_major)

is.glyphplot(x)

## S3 method for class 'glyphplot'
x[

## S3 method for class 'glyphplot'
print(x, ...)
```

**Arguments**

- `data` A data frame containing variables named in `x_major`, `x_minor`, `y_major` and `y_minor`.
- `height, width` The height and width of each glyph. Defaults to 95% of the `resolution` of the data. Specify the width absolutely by supplying a numeric vector of length 1, or relative to the
- `polar` A logical of length 1, specifying whether the glyphs should be drawn in polar coordinates. Defaults to `FALSE`.
- `x_major, y_major` The name of the variable (as a string) for the major x and y axes. Together, the
- `x` glyphplot to be printed
- `...` ignored
Create the data needed to generate a glyph plot.

Usage

glyphs(data, x_major, x_minor, y_major, y_minor, polar = FALSE,
      height = ggplot2::rel(0.95), width = ggplot2::rel(0.95),
      y_scale = identity, x_scale = identity)

Arguments

data A data frame containing variables named in x_major, x_minor, y_major and y_minor.
x_major, x_minor, y_major, y_minor
The name of the variable (as a string) for the major and minor x and y axes. Together, each unique
polar A logical of length 1, specifying whether the glyphs should be drawn in polar coordinates. Defaults to FALSE.
height, width The height and width of each glyph. Defaults to 95% of the resolution of the data. Specify the width absolutely by supplying a numeric vector of length 1, or relative to the
y_scale, x_scale The scaling function to be applied to each set of minor values within a grid cell. Defaults to identity so that no scaling is performed.

Examples

data(nasa)
nasaLate <- nasa[
  nasa$date >= as.POSIXct("1998-01-01") &
  nasa$lat >= 20 &
  nasa$lat <= 40 &
  nasa$long >= -80 &
  nasa$long <= -60
, ]
temp.gly <- glyphs(nasaLate, "long", "day", "lat", "surftemp", height=2.5)
Grab the legend and print it as a plot

**Description**

Grab the legend and print it as a plot

**Usage**

```r
grab_legend(p)
```

## S3 method for class 'legend_guide_box'

### Examples

```r
library(ggplot2)
histPlot <- qplot(  
  x = Sepal.Length,  
  data = iris,  
  fill = Species,  
  geom = "histogram",  
  binwidth = 1/4
)
(right <- histPlot)  
(bottom <- histPlot + theme(legend.position = "bottom"))  
(top <- histPlot + theme(legend.position = "top"))  
(left <- histPlot + theme(legend.position = "left"))

grab_legend(right)  
grab_legend(bottom)  
grab_legend(top)  
grab_legend(left)
```
Data related to happiness from the General Social Survey, 1972-2006.

Description

This data extract is taken from Hadley Wickham’s productplots package. The original description follows, with minor edits.

Usage

data(happy)

Format

A data frame with 51020 rows and 10 variables

Details

The data is a small sample of variables related to happiness from the General Social Survey (GSS). The GSS is a yearly cross-sectional survey of Americans, run from 1972. We combine data for 25 years to yield 51,020 observations, and of the over 5,000 variables, we select nine related to happiness:

• age. age in years: 18–89.
• degree. highest education: lt high school, high school, junior college, bachelor, graduate.
• finrela. relative financial status: far above, above average, average, below average, far below.
• happy. happiness: very happy, pretty happy, not too happy.
• health. health: excellent, good, fair, poor.
• marital. marital status: married, never married, divorced, widowed, separated.
• sex. sex: female, male.
• wtsall. probability weight. 0.43–6.43.

References

lowertriangle

lowertriangle - rearrange dataset as the preparation of ggscatmat function

Description

function for making the melted dataset used to plot the lowertriangle scatterplots.

Usage

lowertriangle(data, columns = 1:ncol(data), color = NULL)

Arguments

data a data matrix. Should contain numerical (continuous) data.
columns an option to choose the column to be used in the raw dataset. Defaults to 1:ncol(data)
color an option to choose a factor variable to be grouped with. Defaults to (NULL)

Author(s)

Mengjia Ni, Di Cook <dicook@monash.edu>

Examples

data(flea)
head(lowertriangle(flea, columns= 2:4))
head(lowertriangle(flea))
head(lowertriangle(flea, color="species"))

model_response_variables

Model term names

Description

Retrieve either the response variable names, the beta variable names, or beta variable names. If the model is an object of class 'lm', by default, the beta variable names will include anova significance stars.

Usage

model_response_variables(model, data = broom::augment(model))
model_beta_variables(model, data = broom::augment(model))
model_beta_label(model, data = broom::augment(model), lmStars = TRUE)
Arguments

model  model in question
data  equivalent to broom::augment(model)
lmStars  boolean that determines if stars are added to labels

Value

character vector of names

Description

This data was provided by NASA for the competition.

Usage

data(nasa)

Format

A data frame with 41472 rows and 17 variables

Details

The data shows 6 years of monthly measurements of a 24x24 spatial grid from Central America:

- time integer specifying temporal order of measurements
- x, y, lat, long spatial location of measurements.
- cloudhigh, cloudlow, cloudmid, ozone, pressure, surftemp, temperature are the various satellite measurements.
- date, day, month, year specifying the time of measurements.
- id unique id for each spatial position.

References

This data contains about the United Kingdom Pig Production from the book 'Data' by Andrews and Herzberg. The original data can be on Statlib: http://lib.stat.cmu.edu/datasets/Andrews/T62.1

Usage
data(pigs)

Format
A data frame with 48 rows and 8 variables

Details
The time variable has been added from a combination of year and quarter

- time year + (quarter - 1) / 4
- year year of production
- quarter quarter of the year of production
- gilts number of sows giving birth for the first time
- profit ratio of price to an index of feed price
- s_per_herdsz ratio of the number of breeding pigs slaughtered to the total breeding herd size
- production number of pigs slaughtered that were reared for meat
- herdsz breeding herd size

References
print.ggmatrix  
*Print ggmatrix object*

Description

Print method taken from ggplot2::print.ggplot and altered for a ggmatrix object

Usage

```r
## S3 method for class 'ggmatrix'
print(x, newpage = is.null(vp), vp = NULL, ...)
```

Arguments

- `x`  
  plot to display
- `newpage`  
  draw new (empty) page first?
- `vp`  
  viewport to draw plot in
- `...`  
  arguments passed onto ggmatrix_gtable

Author(s)

Barret Schloerke

Examples

```r
data(tips, package = "reshape")
pMat <- ggpairs(tips, c(1,3,2), mapping = ggplot2::aes_string(color = "sex"))
pMat # calls print(pMat), which calls print.ggmatrix(pMat)
```

print_if_interactive  
*Print if not CRAN*

Description

Small function to print a plot if the R session is interactive or in a travis build

Usage

```r
print_if_interactive(p)
```

Arguments

- `p`  
  plot to be displayed
psychademic

UCLA canonical correlation analysis data

Description

This data contains 600 observations on eight variables

Usage

data(psychademic)

Format

A data frame with 600 rows and 8 variables

Details

- locus_of_control - psychological
- self_concept - psychological
- motivation - psychological. Converted to four character groups
- read - academic
- write - academic
- math - academic
- science - academic
- female - academic. Dropped from original source
- sex - academic. Added as a character version of female column

References

**putPlot**

**Put Plot**

**Description**

Function to place your own plot in the layout.

**Usage**

```r
putPlot(pm, value, i, j)
```

```r
## S3 replacement method for class 'ggmatrix'
pm[i, j, ...] <- value
```

**Arguments**

- `pm` ggally object to be altered
- `value` ggplot object to be placed
- `i` row from the top
- `j` column from the left
- `...` ignored

**Author(s)**

Barret Schloerke <schloerke@gmail.com>

**Examples**

```r
custom_car <- ggpairs(mtcars[, c("mpg", "wt", "cyl")], upper = "blank", title = "Custom Example")
# ggplot example taken from example(geom_text)
plot <- ggplot2::ggplot(mtcars, ggplot2::aes(x=wt, y=mpg, label=rownames(mtcars)))
plot <- plot +
    ggplot2::geom_text(ggplot2::aes(colour=factor(cyl)), size = 3) +
    ggplot2::scale_colour_discrete(l=40)
custom_car[1, 2] <- plot
personal_plot <- ggally_text(
    "ggpairs allows you\nto put in your\nown plot.\nLike that one.\n" <---
)
custom_car[1, 3] <- personal_plot
# custom_car

# remove plots after creating a plot matrix
custom_car[2,1] <- NULL
custom_car[3,1] <- "blank" # the same as storing null
custom_car[3,2] <- NULL
custom_car
```
Rescaling functions

Usage
range01(x)
max1(x)
mean0(x)
min0(x)
rescale01(x, xlim = NULL)
rescale11(x, xlim = NULL)

Arguments
x numeric vector
xlim value used in range

Find order of variables

Usage
scag_order(scag, vars, measure)

Arguments
scag scagnostics object
vars character vector of the variables to be ordered
measure scagnostics measure to order according to
Value

character vector of variable ordered according to the given scagnostic measure

Author(s)

Barret Schloerke

scatmat

scatmat - plot the lower triangle plots and density plots of the scatter plot matrix.

Description

function for making scatterplots in the lower triangle and diagonal density plots.

Usage

scatmat(data, columns = 1:ncol(data), color = NULL, alpha = 1)

Arguments

data a data matrix. Should contain numerical (continuous) data.
columns an option to choose the column to be used in the raw dataset. Defaults to 1:ncol(data)
color an option to group the dataset by the factor variable and color them by different colors. Defaults to NULL
alpha an option to set the transparency in scatterplots for large data. Defaults to 1.

Author(s)

Mengjia Ni, Di Cook <dicook@monash.edu>

Examples

data(flea)
scatmat(flea, columns=2:4)
scatmat(flea, columns= 2:4, color="species")
**singleClassOrder**

*Order axis variables*

**Description**

Order axis variables by separation between one class and the rest (most separation to least).

**Usage**

```r
singleClassOrder(classVar, axisVars, specClass = NULL)
```

**Arguments**

- `classVar` class variable (vector from original dataset)
- `axisVars` variables to be plotted as axes (data frame)
- `specClass` character string matching to level of `classVar`; instead of looking for separation between any class and the rest, will only look for separation between this class and the rest

**Value**

character vector of names of `axisVars` ordered such that the first variable has the most separation between one of the classes and the rest, and the last variable has the least (as measured by F-statistics from an ANOVA)

**Author(s)**

Jason Crowley <crowley.jason.s@gmail.com>

---

**skewness**

*Sample skewness*

**Description**

Calculate the sample skewness of a vector while ignoring missing values.

**Usage**

```r
skewness(x)
```

**Arguments**

- `x` numeric vector
**Value**

sample skewness of x

**Author(s)**

Jason Crowley <crowley.jason.s@gmail.com>

---

### str.ggmatrix

**ggmatrix structure**

---

**Description**

View the condensed version of the ggmatrix object. The attribute "class" is ALWAYS altered to "_class" to avoid recursion.

**Usage**

```r
## S3 method for class 'ggmatrix'
str(object, ..., raw = FALSE)
```

**Arguments**

- `object` ggmatrix object to be viewed
- `...` passed on to the default str method
- `raw` boolean to determine if the plots should be converted to text or kept as original objects

---

### twitter_spambots

**Twitter spambots**

---

**Description**

A network of spambots found on Twitter as part of a data mining project.

**Usage**

```r
data(twitter_spambots)
```

**Format**

An object of class `network` with 120 edges and 94 vertices.
Details

Each node of the network is identified by the Twitter screen name of the account and further carries five vertex attributes:

- location user’s location, as provided by the user
- lat latitude, based on the user’s location
- lon longitude, based on the user’s location
- followers number of Twitter accounts that follow this account
- friends number of Twitter accounts followed by the account

Author(s)

Amos Elberg

---

uppertriangle  

uppertriangle - rearrange dataset as the preparation of ggscatmat function

Description

function for making the dataset used to plot the uppertriangle plots.

Usage

uppertriangle(data, columns = 1:ncol(data), color = NULL, corMethod = "pearson")

Arguments

data a data matrix. Should contain numerical (continuous) data.
columns an option to choose the column to be used in the raw dataset. Defaults to 1:ncol(data)
color an option to choose a factor variable to be grouped with. Defaults to (NULL)
corMethod method argument supplied to cor

Author(s)

Mengjia Ni, Di Cook <dicook@monash.edu>

Examples

data(flea)  
head(uppertriangle(flea, columns=2:4))  
head(uppertriangle(flea))  
head(uppertriangle(flea, color="species"))
v1_ggmatrix_theme

Modify a ggmatrix object by adding an ggplot2 object to all plots

Description

Modify a ggmatrix object by adding an ggplot2 object to all plots

Usage

v1_ggmatrix_theme()

Examples

ggpairs(iris, 1:2) + v1_ggmatrix_theme()
# move the column names to the left and bottom
ggpairs(iris, 1:2, switch = "both") + v1_ggmatrix_theme()

wrap_fn_with_param_arg

Wrap a function with different parameter values

Description

Wraps a function with the supplied parameters to force different default behavior. This is useful for functions that are supplied to ggpairs. It allows you to change the behavior of one function, rather than creating multiple functions with different parameter settings.

Usage

wrap_fn_with_param_arg(funcVal, params = NULL,
        funcArgName = deparse(substitute(funcVal)))

wrapp(funcVal, params = NULL, funcArgName = deparse(substitute(funcVal)))

wrap(funcVal, ..., funcArgName = deparse(substitute(funcVal)))

wrap_fn_with_params(funcVal, ..., funcArgName = deparse(substitute(funcVal)))

Arguments

funcVal function that the params will be applied to. The function should follow the api of function(data, mapping, ...){}. funcVal is allowed to be a string of one of the ggally_NAME functions, such as “points” for ggally_points or “facetdensity” for ggally_facetdensity.
wrap_fn_with_param_arg

params named vector or list of parameters to be applied to the funcVal
funcArgName name of function to be displayed
... named parameters to be supplied to wrap_fn_with_param_arg

Details

wrap is identical to wrap_fn_with_params. These function take the new parameters as arguments.
wrapp is identical to wrap_fn_with_param_arg. These functions take the new parameters as a single list.

The params and fn attributes are there for debugging purposes. If either attribute is altered, the function must be re-wrapped to have the changes take effect.

Value

a function(data, mapping, ...){} that will wrap the original function with the parameters applied as arguments

Examples

# small function to display plots only if it's interactive
p_ <- GGally::print_if_interactive

# example function that prints 'val'
fn <- function(data, mapping, val = 2) {
  print(val)
}
fn(data = NULL, mapping = NULL) # 2

# wrap function to change default value 'val' to 5 instead of 2
wrapped_fn1 <- wrap(fn, val = 5)
wrapped_fn1(data = NULL, mapping = NULL) # 5
# you may still supply regular values
wrapped_fn1(data = NULL, mapping = NULL, val = 3) # 3

# wrap function to change 'val' to 5 using the arg list
wrapped_fn2 <- wrap_fn_with_param_arg(fn, params = list(val = 5))
wrapped_fn2(data = NULL, mapping = NULL) # 5

# change parameter settings in ggpairs for a particular function
## Goal output:
regularPlot <- ggally_points(
  iris,
  ggplot2::aes(Sepal.Length, Sepal.Width),
  size = 5, color = "red"
)
p_(regularPlot)

# Wrap ggally_points to have parameter values size = 5 and color = 'red'
w_ggally_points <- wrap(ggally_points, size = 5, color = "red")
wrappedPlot <- w_ggally_points(
  iris,


```r
ggplot2::aes(Sepal.Length, Sepal.Width) )

p_(wrappedPlot)

# Double check the aes parameters are the same for the geom_point layer
identical(regularPlot$layers[[1]]$aes_params, wrappedPlot$layers[[1]]$aes_params)

# Use a wrapped function in ggpairs
pm <- ggpairs(iris, 1:3, lower = list(continuous = wrap(ggally_points, size = 5, color = "red")))
p_(pm)

pm <- ggpairs(iris, 1:3, lower = list(continuous = w_ggally_points))
p_(pm)
```
## Index

*Topic **datasets**
- australia_PISA2012, 6
- flea, 9
- happy, 74
- nasa, 76
- pigs, 77
- psychademic, 79
- twitter_spambots, 84

*Topic **hplot**
- getPlot, 11
- ggally_barDiag, 11
- ggally_blank, 12
- ggally_box, 13
- ggally_cor, 13
- ggally_density, 14
- ggally_densityDiag, 15
- ggally_denstrip, 16
- ggally_dot, 18
- ggally_dot_and_box, 19
- ggally_facetbar, 19
- ggally_facetdensity, 20
- ggally_facetdensitystrip, 21
- ggally_facethist, 21
- ggally_na, 22
- ggally_points, 29
- ggally_ratio, 30
- ggally_smooth, 31
- ggally_text, 32
- ggmatrix, 43
- ggpairs, 60
- putPlot, 80

+ gg, 4, 4
[ .ggmatrix (getPlot), 11
[ .glyphplot (glyphplot), 71
[ <- .ggmatrix (putPlot), 80

add_ref_boxes, 5
add_ref_lines, 5
aes, 36, 41, 60
arrow, 47, 52

AsIs, 65
asNetwork, 46, 50, 54
augment, 8, 58, 59
australia_PISA2012, 6

brew_colors, 7
brewer_pal, 50
broomify, 8, 58

cooks.distance, 23, 59
cor, 34, 35, 68, 85
cut, 34, 47, 51

degree, 47, 50

edgeset_constructors, 46, 50, 54
expand_range, 46, 50

facetgrid, 37, 41, 43, 61
find_plot_type, 8
flea, 9
fn_switch, 10

geom_line, 5, 24–26
geom_point, 33
geom_rect, 5
geom_smooth, 26, 31
geom_text, 35, 48, 52
geom_tile, 30
getPlot, 11
ggally_barDiag, 11
ggally_blank, 12
ggally_blankDiag (ggally_blank), 12
ggally_box, 13
ggally_box_no_facet (ggally_box), 13
ggally_cor, 13
ggally_density, 14
ggally_densityDiag, 15
ggally_denstrip, 16
ggally_diagonalAxis, 17
ggally_dot, 18
ggally_dot_and_box, 19
ggally_dot_no_facet (ggally_dot), 18
ggally_facetbar, 19
ggally_facetdensity, 20
ggally_facetdensitystrip, 21
ggally_facethist, 21
ggally_na, 22
ggally_naDiag (ggally_na), 22
ggally_nostic_cooksd, 23
ggally_nostic_hat, 24
ggally_nostic_line, 23, 24, 25, 26–28
ggally_nostic_resid, 26, 29
ggally_nostic_se_fit, 27
ggally_nostic_sigma, 28
ggally_nostic_std_resid, 29
ggally_points, 29
ggally_ratio, 30
ggally_smooth, 31
ggally_smooth_lm (ggally_smooth), 31
ggally_smooth_loess (ggally_smooth), 31
ggally_text, 32
ggcoef, 32
ggcorr, 33
ggduo, 36, 41, 58, 59, 71
ggfacet, 40
gglegend, 41
ggmatrix, 43, 58
ggmatrix_gtable, 45
gnet, 46, 49, 53
gnet2, 46, 48, 49
gnetnetworkmap, 54
ggnostic, 57
ggpairs, 41, 59, 60
ggparcoord, 64
ggplot, 25
ggscattermat, 67
ggsurv, 68
ggts, 70
glance, 8
glm, 58
glyphplot, 71
glyphs, 72
gplot, 48, 53
gplot.layout, 46, 50
grab_legend, 37, 44, 61, 73
happy, 74
identity, 72
igraph, 46, 50, 54
influence, 24, 27, 28
intergraph, 46, 50, 54
is.glyphplot (glyphplot), 71
labellers, 37, 43, 61
lm, 58
lowertriangle, 75
max1 (rescale01), 81
mean0 (rescale01), 81
min0 (rescale01), 81
model_beta_label
(model_response_variables), 75
model_beta_variables
(model_response_variables), 75
model_response_variables, 75
nasa, 76
network, 46, 48, 50, 53, 54
pigs, 77
plot.network, 48, 53
predict, 27
print.ggmatrix, 78
print.glyphplot (glyphplot), 71
print.legend_guide_box (grab_legend), 73
print_if_interactive, 78
progress_bar, 45
psychademic, 79
putPlot, 80
quantile, 47, 51
range01 (rescale01), 81
RCColorBrewer, 50
rescale01, 81
rescale11 (rescale01), 81
residuals, 26
resolution, 71, 72
rstandard, 29
scag_order, 81
scatmat, 82
singleClassOrder, 83
skewness, 83
sna, 46, 48, 50, 53
str.ggmatrix, 84
substr, 48, 51
INDEX

theme, 4, 35, 48, 52
tidy, 8, 33
twitter_spambots, 84

unit, 43
uppertriangle, 85

vl_ggmatrix_theme, 86

wrap, 38, 41, 42, 62
wrap(wrap_fn_with_param_arg), 86
wrap_fn_with_param_arg, 60, 86
wrap_fn_with_params
    (wrap_fn_with_param_arg), 86
wrapp (wrap_fn_with_param_arg), 86