Package: beeswarm (via r-universe)

August 22, 2024	
Title The Bee Swarm Plot, an Alternative to Stripchart	
Description The bee swarm plot is a one-dimensional scatter plot like ``stripchart", but with closely-packed, non-overlapping points.	
Version 0.4.0	
Date 2021-05-07	
Imports stats, graphics, grDevices, utils	
NeedsCompilation yes	
License Artistic-2.0	
<pre>URL https://github.com/aroneklund/beeswarm</pre>	
BugReports https://github.com/aroneklund/beeswarm/issues	
Repository https://aroneklund.r-universe.dev	
RemoteUrl https://github.com/aroneklund/beeswarm	
RemoteRef HEAD	
RemoteSha d641db509e7a3f9c8e76147bf779fcb141cdba51	
Contents	
beeswarm	2
	6
bxplot	7
swarmx	9
Index 1	2

beeswarm

Bee swarm plot

Description

Create a bee swarm plot. A bee swarm plot is a one-dimensional scatter plot similar to stripchart, but with various methods to separate coincident points such that each point is visible. Also, beeswarm introduces additional features unavailable in stripchart, such as the ability to control the color and plotting character of each point.

Usage

```
beeswarm(x, ...)
## S3 method for class 'formula'
beeswarm(formula, data = NULL, subset, na.action = NULL,
        pwpch = NULL, pwcol = NULL, pwbg = NULL, pwcex = NULL, dlab, glab, ...)
## Default S3 method:
beeswarm(x,
    method = c("swarm", "compactswarm", "center", "hex", "square"),
    vertical = TRUE, horizontal = !vertical,
    cex = 1, spacing = 1, breaks = NULL,
    labels, at = NULL,
    corral = c("none", "gutter", "wrap", "random", "omit"),
    corralWidth, side = 0L,
    priority = c("ascending", "descending", "density", "random", "none"),
    fast = TRUE,
    pch = par("pch"), col = par("col"), bg = NA,
    pwpch = NULL, pwcol = NULL, pwbg = NULL, pwcex = NULL,
    do.plot = TRUE, add = FALSE, axes = TRUE, log = FALSE,
    xlim = NULL, ylim = NULL, dlim = NULL, glim = NULL,
    xlab = NULL, ylab = NULL, dlab = "", glab = "",
    ...)
```

Arguments

formula	A formula, such as y ~ grp, where y is a numeric vector of data values to be split into groups according to the grouping variable grp (usually a factor).
data	A data.frame (or list) from which the variables in formula should be taken.
subset	An optional vector specifying a subset of observations to be used.
na.action	A function which indicates what should happen when the data contain NAs. The default is to quietly ignore missing values in either the response or the group.
X	A numeric vector, or a data frame or list of numeric vectors, each of which is plotted as an individual swarm.

method Method for arranging points (see Details).

vertical, horizontal

Orientation of the plot. horizontal takes precedence if both are specified.

cex Size of points relative to the default given by par("cex"). Unlike other plotting

functions, this must be a single value. (But see also the pwcex argument)

spacing Relative spacing between points.

breaks Breakpoints for data discretization (optional). Used only if method is "square",

"hex", or "center". If NULL, breakpoints are chosen automatically. If NA, data

is not discretized at all (similar to stripchart with method = "stack").

labels Labels for each group. Recycled if necessary. By default, these are inferred

from the data.

at Numeric vector giving the locations where the swarms should be drawn; defaults

to 1:n where n is the number of groups.

corral Method to adjust points that would be placed outside their own group region

(see Details).

corralWidth Width of the "corral" in user coordinates. If missing, a sensible value will be

chosen.

side Direction to perform jittering: 0: both directions; 1: to the right or upwards; -1:

to the left or downwards.

priority Order used to perform point layout when method is "swarm" or "compactswarm";

ignored otherwise (see Details).

fast Use compiled version of algorithm? This option is ignored for all methods ex-

cept "swarm" and "compactswarm".

pch, col, bg Plotting characters and colors, specified by group. Recycled if necessary (see

Details).

pwpch, pwcol, pwbg, pwcex

"Point-wise" plotting characteristics, specified for each data point (see Details).

do.plot Draw a plot?

add Add to an existing plot?

axes Draw axes and box?

log Use a logarithmic scale on the data axis?

xlim, ylim Limits of the plot.

dlim, glim An alternative way to specify limits (see Details).

xlab, ylab Axis labels.

dlab, glab An alternative way to specify axis labels (see Details).

... Further arguments passed to plot.

Details

Several methods for placing the points are available; each method uses a different algorithm to avoid overlapping points.

The default method, swarm, places points in increasing order. If a point would overlap an existing point, it is shifted sideways (along the group axis) by a minimal amount sufficient to avoid overlap. With this method breaks is ignored.

The methods square, hex, and center do the same thing, but they first discretize the values along the continuous data axis, in order to enable more efficient packing: square places the points on a square grid, hex uses a hexagonal grid, and center uses a centered square grid. By default, the number of breakpoints for discretization is determined by a combination of the available plotting area and the plotting character size. The discretization of the data can be explicitly controlled using breaks. If breaks is set to NA, the data will not be grouped into intervals; this may be a sensible option if the data is already discrete. NOTE that these three methods adjust the data to fit into a grid, and therefore the resulting plots should be interpreted with this in mind.

In contrast to most other plotting functions, changing the size of the graphics device will often change the position of the points.

The plotting characters and colors can be controlled in two ways. First, the arguments pch, col and bg can specify plotting characters and colors in the same way as stripchart and boxplot: in short, the arguments apply to each group as a whole (and are recycled if necessary).

Alternatively, the "point-wise" characteristics of each individual data point can be controlled using pwpch, pwcol, and pwbg, which override pch, col and bg if these are also specified. Likewise, pwcex controls the size of each point relative to the default (which may be adjusted by cex). Notably, the point layout algorithm is applied without considering the point-wise arguments; thus setting pwcex larger than 1 will usually result in partially overlapping points. These arguments can be specified as a list or vector. If supplied using the formula method, the arguments can be specified as part of the formula interface; i.e. they are affected by data and subset.

The dlab and glab labels may be used instead of xlab and ylab if those are not specified. dlab applies to the continuous data axis (the Y axis unless horizontal is TRUE); glab to the group axis. Likewise, dlim and glim can be used to specify limits of the axes instead of xlim or ylim.

This function is intended to be mostly compatible with calls to stripchart or boxplot. Thus, code that works with these functions should work with beeswarm with minimal modification.

By default, swarms from different groups are not prevented from overlapping. Thus, large data sets, or data sets with uneven distributions, may produce somewhat unpleasing beeswarms. If this is a problem, consider reducing cex. Another approach is to control runaway points (those that would be plotted outside a region allotted to each group) with the corral argument: The default, "none", does not control runaway points. "gutter" collects runaway points along the boundary between groups. "wrap" implements periodic boundaries. "random" places runaway points randomly in the region. "omit" omits runaway points. See Examples below.

When using the "swarm" method, priority controls the order in which the points are placed; this generally has a noticeable effect on the resulting appearance. "ascending" gives the "traditional" beeswarm plot in which the points are placed in an ascending order. "descending" is the opposite. "density" prioritizes points with higher local density. "random" places points in a random order. "none" places points in the order provided.

Whereas the "swarm" method places points in a predetermined order, the "compactswarm" method uses a greedy strategy to determine which point will be placed next. This often leads to a more tightly-packed layout. The strategy is very simple: on each iteration, a point that can be placed as close as possible to the non-data axis is chosen and placed. If there are two or more equally good points, priority is used to break ties.

Value

A data frame with plotting information, invisibly.

See Also

```
stripchart, boxplot
```

Examples

```
## One of the examples from 'stripchart'
beeswarm(decrease ~ treatment,
  data = OrchardSprays, log = TRUE,
  pch = 16, col = rainbow(8))
## One of the examples from 'boxplot', with a beeswarm overlay
 boxplot(len ~ dose, data = ToothGrowth,
            main = "Guinea Pigs' Tooth Growth",
            xlab = "Vitamin C dose mg",
            ylab = "Tooth length")
 beeswarm(len ~ dose, data = ToothGrowth, col = 2, add = TRUE)
## Compare the 5 methods
op \leftarrow par(mfrow = c(2,3))
for (m in c("swarm", "compactswarm", "center", "hex", "square")) {
  beeswarm(len ~ dose, data = ToothGrowth, method = m, main = m)
}
par(op)
## Demonstrate the use of 'pwcol'
data(breast)
beeswarm(time_survival ~ ER, data = breast,
  pch = 16, pwcol = 1 + as.numeric(event_survival),
  xlab = "", ylab = "Follow-up time (months)",
  labels = c("ER neg", "ER pos"))
legend("topright", legend = c("Yes", "No"),
  title = "Censored", pch = 16, col = 1:2)
## The list interface
distributions <- list(runif = runif(200, min = -3, max = 3),
                      rnorm = rnorm(200),
                      rlnorm = rlnorm(200, sdlog = 0.5))
beeswarm(distributions, col = 2:4)
## Demonstrate 'pwcol' with the list interface
myCol <- lapply(distributions, function(x) cut(x, breaks = quantile(x), labels = FALSE))</pre>
beeswarm(distributions, pch = 16, pwcol = myCol)
legend("bottomright", legend = 1:4, pch = 16, col = 1:4, title = "Quartile")
## Demonstrate the 'corral' methods
par(mfrow = c(2,3))
beeswarm(distributions, col = 2:4,
```

6 breast

```
main = 'corral = "none" (default)')
beeswarm(distributions, col = 2:4, corral = "gutter",
  main = 'corral = "gutter"')
beeswarm(distributions, col = 2:4, corral = "wrap",
 main = 'corral = "wrap"')
beeswarm(distributions, col = 2:4, corral = "random",
  main = 'corral = "random"')
beeswarm(distributions, col = 2:4, corral = "omit",
  main = 'corral = "omit"')
## Demonstrate 'side' and 'priority'
par(mfrow = c(2,3))
beeswarm(distributions, col = 2:4,
  main = 'Default')
beeswarm(distributions, col = 2:4, side = -1,
  main = 'side = -1')
beeswarm(distributions, col = 2:4, side = 1,
  main = 'side = 1')
beeswarm(distributions, col = 2:4, priority = "descending",
  main = 'priority = "descending"')
beeswarm(distributions, col = 2:4, priority = "random",
 main = 'priority = "random"')
beeswarm(distributions, col = 2:4, priority = "density",
 main = 'priority = "density"')
## Demonstrate 'side' and 'priority' for compact method
par(mfrow = c(2,3))
beeswarm(distributions, col = 2:4, method = "compactswarm",
 main = 'Default')
beeswarm(distributions, col = 2:4, method = "compactswarm", side = -1,
 main = 'side = -1')
beeswarm(distributions, col = 2:4, method = "compactswarm", side = 1,
 main = 'side = 1')
beeswarm(distributions, col = 2:4, method = "compactswarm",
  priority = "descending", main = 'priority = "descending"')
beeswarm(distributions, col = 2:4, method = "compactswarm",
  priority = "random", main = 'priority = "random"')
beeswarm(distributions, col = 2:4, method = "compactswarm",
  priority = "density", main = 'priority = "density"')
## Demonstrate pwcol, pwpch, pwbg, and pwcex
beeswarm(mpg ~ cyl, data = mtcars, cex = 3,
  pwcol = gear, pwbg = am + 1, pwpch = gear + 18, pwcex = hp / 335)
```

breast

Lymph-node-negative primary breast tumors

Description

Tumor molecular measurements and outcome from breast cancer patients.

bxplot 7

Usage

```
data(breast)
```

Format

A data frame with 286 observations on the following 5 variables.

```
ER Estrogen receptor status (factor with levels neg, pos)
```

ESR1 Expression of the ESR1 gene (numeric)

ERBB2 Expression of the ERBB2 gene (numeric)

time_survival Time in months (numeric)

event_survival Coded event: 0 = censored, 1 = metastasis (numeric)

Details

ER, ESR1, and ERBB2 were measured on a tumor specimen taken at surgery (time = 0).

ESR1 and ERBB2 expression values were determined by microarray probe sets 205225_at and 216836_s_at using RMA-normalized data.

Source

Wang Y, Klijn JG, Zhang Y, Sieuwerts AM, Look MP, Yang F, Talantov D, Timmermans M, Meijervan Gelder ME, Yu J, Jatkoe T, Berns EM, Atkins D, Foekens JA. Gene-expression profiles to predict distant metastasis of lymph-node-negative primary breast cancer. Lancet. 2005 Feb 19-25;365(9460):671-9.

Examples

```
data(breast)
with(breast,
  plot(ESR1, ERBB2, col = as.numeric(ER))
)
```

bxplot

Plot quantile lines

Description

Plot lines indicating the specified quantiles for each group. This function is intended as a simplified interpretation of boxplot, which can be combined with a beeswarm (or stripchart) plot.

8 bxplot

Usage

```
bxplot(x, ...)
## S3 method for class 'formula'
bxplot(formula, data = NULL, ..., subset, na.action = NULL)
## Default S3 method:
bxplot(x, probs = c(0.25, 0.5, 0.75),
    vertical = TRUE, horizontal = !vertical, add = FALSE,
    col = par("col"), lty = par("lty"), lwd = NULL,
    at = NULL, width = 0.75, ...)
```

Arguments

formula	A formula, such as y ~ grp, where y is a numeric vector of data values to be split
	into groups according to the grouping variable grp (usually a factor).

data A data frame (or list) from which the variables in formula should be taken.

subset An optional vector specifying a subset of observations to be used.

A function which indicates what should happen when the data contain NAs. The na.action

default is to quietly ignore missing values in either the response or the group.

A numeric vector, or a data frame or list of numeric vectors, each of which is Х

considered as a group.

probs A numeric vector of probabilities with values in [0,1]

vertical, horizontal

Orientation of the plot. horizontal takes precedence if both are specified.

Add to an existing plot? add

col, lty Color and line type for each probability. lwd

Line width for each probability (see below).

Numeric vector giving the locations where the swarms should be drawn; defaults at

to 1:n where n is the number of groups.

width Width of the lines.

Further arguments passed to boxplot.

Details

This function is intended as a minimalistic interpration of boxplot; however, the quantiles plotted by bxplot are not necessarily the same as the hinges plotted by a boxplot.

Notice that specifying a vector of graphical parameters such as 1wd or co1 will refer to each of probs, not to each group in the data (as one might expect by analogy with boxplot).

If 1wd is NULL, and if the probs includes 0.5, 1wd will be set to 3 times par("1wd") for probs=0.5, and par ("lwd") for the others. (Thus something resembling the median line and hinges of a boxplot is produced by default.)

swarmx 9

Value

None.

Examples

```
## bxplot on bottom
beeswarm(len ~ dose, data = ToothGrowth)
bxplot(len ~ dose, data = ToothGrowth, add = TRUE)

## bxplot on top
bxplot(decrease ~ treatment, data = OrchardSprays, probs = 0.5, col = 2)
beeswarm(decrease ~ treatment, data = OrchardSprays, add = TRUE)

## Show deciles
data(breast)
bxplot(time_survival ~ event_survival, data = breast,
    probs = seq(0, 1, by = 0.1), col = rainbow(10))
beeswarm(time_survival ~ event_survival, data = breast,
    pch = 21, bg = "gray75", add = TRUE)
```

swarmx

Adjust 1-d data to separate coincident points

Description

Take a series of points lying in a horizontal or vertical line, and jitter them in the other dimension such that no points are overlapping.

Usage

```
swarmx(x, y,
    xsize = xinch(0.08, warn.log = FALSE),
    ysize = yinch(0.08, warn.log = FALSE),
    log = NULL, cex = par("cex"), side = 0L,
    priority = c("ascending", "descending", "density", "random", "none"),
    fast = TRUE, compact = FALSE)
swarmy(x, y,
    xsize = xinch(0.08, warn.log = FALSE),
    ysize = yinch(0.08, warn.log = FALSE),
    log = NULL, cex = par("cex"), side = 0L,
    priority = c("ascending", "descending", "density", "random", "none"),
    fast = TRUE, compact = FALSE)
```

Arguments

x, y Coordinate vectors in any format supported by xy.coords. xsize, ysize Width and height of the plotting character in user coordinates. 10 swarmx

log	Character string indicating which axes are logarithmic, as in plot.default, or NULL to figure it out automatically.
cex	Relative plotting character size.
side	Direction to perform jittering: 0: both directions; 1: to the right or upwards; -1: to the left or downwards.
priority	Method used to perform point layout (see below).
fast	Use compiled version of algorithm? This option is ignored for all methods except "swarm" and "compactswarm".
compact	Use compact layout? (see below)

Details

For swarmx, the input coordinates must lie in a vertical line. For swarmy, the input coordinates must lie in a horizontal line.

swarmx adjusts coordinates to the left or right; swarmy adjusts coordinates up or down.

priority controls the order in which the points are placed; this has generally has a noticeable effect on the resulting appearance. "ascending" gives the "traditional" beeswarm plot in which the points are placed in an ascending order. "descending" is the opposite. "density" prioritizes points with higher local density. "random" places points in a random order. "none" places points in the order provided.

When compact is FALSE, points are placed in a predetermined order. When compact is TRUE, a greedy strategy is used to determine which point will be placed next. This often leads to a more tightly-packed layout. The strategy is very simple: on each iteration, a point that can be placed as close as possible to the non-data axis is chosen and placed. If there are two or more equally good points, priority is used to break ties.

Usually it makes sense to call this function after a plotting device has already been set up (e.g. when adding points to an existing plot), so that the default values for xsize, ysize, and log will be appropriate.

Value

A data frame with columns x and y with the new coordinates.

See Also

```
beeswarm, jitter
```

Examples

```
## Plot points in one dimension
index <- rep(0, 100)
values <- rnorm(100)
plot(index, values, xlim = c(-0.5, 2.5))
points(swarmx(index + 1, values), col = 2)
points(swarmx(index + 2, values, cex = 1.5), col = 3, cex = 1.5)</pre>
```

swarmx 11

```
## Try the horizontal direction, with a log scale
plot(values, index, log = "x", ylim = c(-1, 2))
points(swarmy(values, index + 1), col = 2)

## Newer examples using "side", "priority", and "compact"
plot(c(-0.5, 3.5), range(values), type = 'n')
points(swarmx(index + 0, values), col = 1)
points(swarmx(index + 0.9, values, side = -1), col = 2)
points(swarmx(index + 1.1, values, side = 1, priority = "descending"), col = 3)
points(swarmx(index + 2 , values, priority = 'density'), col = 4)
points(swarmx(index + 3 , values, priority = 'random'), col = 5)
points(swarmx(index + 3 , values, priority = 'random', compact = TRUE), col = 5)
```

Index

```
\ast datasets
    breast, 6
* dplot
     swarmx, 9
* hplot
    beeswarm, 2
    bxplot, 7
beeswarm, 2, 7, 10
boxplot, 4, 5, 7, 8
breast, 6
bxplot, 7
jitter, 10
plot, 3
{\tt plot.default}, {\color{red} 10}
stripchart, 2, 4, 5, 7
swarmx, 9
swarmy (swarmx), 9
xy.coords, 9
```