

# Package ‘PCAtools’

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**Type** Package

**Title** PCAtools: everything Principal Components Analysis

**Version** 1.0.0

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**Description** Principal Components Analysis (PCA) is a very powerful technique that has wide applicability in data science, bioinformatics, and further afield. It was initially developed to analyse large volumes of data in order to tease out the differences/relationships between the logical entities being analysed. It extracts the fundamental structure of the data without the need to build any model to represent it. This 'summary' of the data is arrived at through a process of reduction that can transform the large number of variables into a lesser number that are uncorrelated, i.e., the principal components, whilst at the same time being capable of easy interpretation on the original data.

**License** GPL-3

**Depends** stats, ggplot2, ggrepel, reshape2, lattice, grDevices, cowplot

**Imports**

**Suggests** RUnit, BiocGenerics, knitr, Biobase, GEOquery, biomaRt, ggplotify

**URL** <https://github.com/kevinblighe/PCAtools>

**biocViews** RNASeq, GeneExpression, Transcription

**VignetteBuilder** knitr

**git\_url** <https://git.bioconductor.org/packages/PCAtools>

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PCAtools-package	<i>PCAtools: everything Principal Components Analysis</i>
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### Description

Principal Components Analysis (PCA) is a very powerful technique that has wide applicability in data science, bioinformatics, and further afield. It was initially developed to analyse large volumes of data in order to tease out the differences/relationships between the logical entities being analysed. It extracts the fundamental structure of the data without the need to build any model to represent it. This 'summary' of the data is arrived at through a process of reduction that can transform the large number of variables into a lesser number that are uncorrelated, i.e., 'principal components', whilst at the same time being capable of easy interpretation on the original data.

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biplot	<i>biplot</i>
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### Description

Draw a bi-plot, comparing 2 selected principal components / eigenvectors.

### Usage

```
biplot(pcaobj,
x = 'PC1',
y = 'PC2',
colby = NULL,
colkey = NULL,
singlecol = NULL,
shape = NULL,
shapekey = NULL,
pointSize = 3.0,
legendPosition = 'none',
legendLabSize = 12,
legendIconSize = 5.0,
xlim = NULL,
ylim = NULL,
lab = TRUE,
labSize = 3.0,
labhjust = 1.5,
labvjust = 0,
```

```

selectLab = NULL,
drawConnectors = TRUE,
widthConnectors = 0.5,
colConnectors = 'grey50',
xlab = paste0(x, ', ',
  round(pcaobj$variance[x], digits=2),
  '% variation'),
xlabAngle = 0,
xlabhjust = 0.5,
xlabvjust = 0.5,
ylab = paste0(y, ', ',
  round(pcaobj$variance[y], digits=2),
  '% variation'),
ylabAngle = 0,
ylabhjust = 0.5,
ylabvjust = 0.5,
axisLabSize = 16,
title = '',
subtitle = '',
caption = '',
titleLabSize = 16,
subtitleLabSize = 12,
captionLabSize = 12,
hline = NULL,
hlineType = 'longdash',
hlineCol = 'black',
hlineWidth = 0.4,
vline = NULL,
vlineType = 'longdash',
vlineCol = 'black',
vlineWidth = 0.4,
gridlines.major = TRUE,
gridlines.minor = TRUE,
borderWidth = 0.8,
borderColour = 'black',
returnPlot = TRUE)

```

### Arguments

pcaobj	Object of class 'pca' created by <code>pca()</code> . <b>REQUIRED.</b>
x	A principal component to plot on x-axis. All principal component names are stored in <code>pcaobj\$label</code> . <b>DEFAULT = 'PC1'. REQUIRED.</b>
y	A principal component to plot on y-axis. All principal component names are stored in <code>pcaobj\$label</code> . <b>DEFAULT = 'PC2'. REQUIRED.</b>
colby	If <code>NULL</code> , all points will be coloured differently. If not <code>NULL</code> , value is assumed to be a column name in <code>pcaobj\$metadata</code> relating to some grouping/categorical variable. <b>DEFAULT = NULL. OPTIONAL.</b>
colkey	Vector of name-value pairs relating to value passed to 'col', e.g., <code>c(A='forestgreen', B='gold')</code> . <b>DEFAULT = NULL. OPTIONAL.</b>
singlecol	If specified, all points will be shaded by this colour. Overrides 'col'. <b>DEFAULT = NULL. OPTIONAL.</b>

shape	If NULL, all points will be have the same shape. If not NULL, value is assumed to be a column name in <code>pcaobj\$metadata</code> relating to some grouping/categorical variable. DEFAULT = NULL. OPTIONAL.
shapekey	Vector of name-value pairs relating to value passed to 'shape', e.g., <code>c(A=10, B=21)</code> . DEFAULT = NULL. OPTIONAL.
pointSize	Size of plotted points. DEFAULT = 3.0. OPTIONAL.
legendPosition	Position of legend ('top', 'bottom', 'left', 'right', 'none'). DEFAULT = 'none'. OPTIONAL.
legendLabSize	Size of plot legend text. DEFAULT = 10. OPTIONAL.
legendIconSize	Size of plot legend icons / symbols. DEFAULT = 3.0. OPTIONAL.
xlim	Limits of the x-axis. DEFAULT = NULL. OPTIONAL.
ylim	Limits of the y-axis. DEFAULT = NULL. OPTIONAL.
lab	Logical, indicating whether or not to label the points in the plot space. Labels will be taken as the original colnames of the input object, usually sample IDs. DEFAULT = TRUE. OPTIONAL.
labSize	Size of labels. DEFAULT = 3.0. OPTIONAL.
labhjust	Horizontal adjustment of label. DEFAULT = 1.5. OPTIONAL.
labvjust	Vertical adjustment of label. DEFAULT = 0. OPTIONAL.
selectLab	A vector containing a subset of lab to plot. DEFAULT = NULL. OPTIONAL.
drawConnectors	Logical, indicating whether or not to connect plot labels to their corresponding points by line connectors. DEFAULT = TRUE. OPTIONAL.
widthConnectors	Line width of connectors. DEFAULT = 0.5. OPTIONAL.
colConnectors	Line colour of connectors. DEFAULT = 'grey50'. OPTIONAL.
xlab	Label for x-axis. DEFAULT = <code>paste0(x, ', ', round(pcaobj\$variance[x], digits=2), '% variation')</code> . OPTIONAL.
xlabAngle	Rotation angle of x-axis labels. DEFAULT = 0. OPTIONAL.
xlabhjust	Horizontal adjustment of x-axis labels. DEFAULT = 0.5. OPTIONAL.
xlabvjust	Vertical adjustment of x-axis labels. DEFAULT = 0.5. OPTIONAL.
ylab	Label for y-axis. DEFAULT = <code>paste0(y, ', ', round(pcaobj\$variance[y], digits=2), '% variation')</code> . OPTIONAL.
ylabAngle	Rotation angle of y-axis labels. DEFAULT = 0. OPTIONAL.
ylabhjust	Horizontal adjustment of y-axis labels. DEFAULT = 0.5. OPTIONAL.
ylabvjust	Vertical adjustment of y-axis labels. DEFAULT = 0.5. OPTIONAL.
axisLabSize	Size of x- and y-axis labels. DEFAULT = 16. OPTIONAL.
title	Plot title. DEFAULT = ''. OPTIONAL.
subtitle	Plot subtitle. DEFAULT = ''. OPTIONAL.
caption	Plot caption. DEFAULT = ''. OPTIONAL.
titleLabSize	Size of plot title. DEFAULT = 16. OPTIONAL.
subtitleLabSize	Size of plot subtitle. DEFAULT = 12. OPTIONAL.
captionLabSize	Size of plot caption. DEFAULT = 12. OPTIONAL.

<code>hline</code>	Draw one or more horizontal lines passing through this/these values on y-axis. For single values, only a single numerical value is necessary. For multiple lines, pass these as a vector, e.g., <code>c(60,90)</code> . DEFAULT = NULL. OPTIONAL.
<code>hlineType</code>	Line type for <code>hline</code> ('blank', 'solid', 'dashed', 'dotted', 'dotdash', 'longdash', 'twodash'). DEFAULT = 'longdash'. OPTIONAL.
<code>hlineCol</code>	Colour of <code>hline</code> . DEFAULT = 'black'. OPTIONAL.
<code>hlineWidth</code>	Width of <code>hline</code> . DEFAULT = 0.4. OPTIONAL.
<code>vline</code>	Draw one or more vertical lines passing through this/these values on x-axis. For single values, only a single numerical value is necessary. For multiple lines, pass these as a vector, e.g., <code>c(60,90)</code> . DEFAULT = NULL. OPTIONAL.
<code>vlineType</code>	Line type for <code>vline</code> ('blank', 'solid', 'dashed', 'dotted', 'dotdash', 'longdash', 'twodash'). DEFAULT = 'longdash'. OPTIONAL.
<code>vlineCol</code>	Colour of <code>vline</code> . DEFAULT = 'black'. OPTIONAL.
<code>vlineWidth</code>	Width of <code>vline</code> . DEFAULT = 0.4. OPTIONAL.
<code>gridlines.major</code>	Logical, indicating whether or not to draw major gridlines. DEFAULT = TRUE. OPTIONAL.
<code>gridlines.minor</code>	Logical, indicating whether or not to draw minor gridlines. DEFAULT = TRUE. OPTIONAL.
<code>borderWidth</code>	Width of the border on the x and y axes. DEFAULT = 0.8. OPTIONAL.
<code>borderColour</code>	Colour of the border on the x and y axes. DEFAULT = 'black'. OPTIONAL.
<code>returnPlot</code>	Logical, indicating whether or not to return the plot object. DEFAULT = TRUE. OPTIONAL.

**Value**

A `ggplot2` object.

**Author(s)**

Kevin Blighe <kevin@clinicalbioinformatics.co.uk>, Myles Lewis <myles.lewis@qmul.ac.uk>

**Examples**

```
options(scipen=10)
options(digits=6)

col <- 20
row <- 20000
mat1 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat1) <- paste0('gene', 1:nrow(mat1))
colnames(mat1) <- paste0('sample', 1:ncol(mat1))

mat2 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat2) <- paste0('gene', 1:nrow(mat2))
colnames(mat2) <- paste0('sample', (ncol(mat1)+1):(ncol(mat1)+ncol(mat2)))
```

```

mat <- cbind(mat1, mat2)

metadata <- data.frame(row.names = colnames(mat))
metadata$Group <- rep(NA, ncol(mat))
metadata$Group[seq(1,40,2)] <- 'A'
metadata$Group[seq(2,40,2)] <- 'B'
metadata$CRP <- sample.int(100, size=ncol(mat), replace=TRUE)
metadata$ESR <- sample.int(100, size=ncol(mat), replace=TRUE)

p <- pca(mat, metadata = metadata, removeVar = 0.1)

biplot(p)

biplot(p, colby = 'Group', shape = 'Group')

biplot(p, colby = 'Group', colkey = c(A = 'forestgreen', B = 'gold'),
  legendPosition = 'right')

biplot(p, colby = 'Group', colkey = c(A='forestgreen', B='gold'),
  shape = 'Group', shapekey = c(A=10, B=21), legendPosition = 'bottom')

```

---

eigencorplot

*eigencorplot*


---

## Description

Correlate principal components to continuous variable metadata and test significancies of these.

## Usage

```

eigencorplot(pcaobj,
  components = getComponents(pcaobj, seq_len(10)),
  metavar,
  titleX = '',
  cexTitleX = 1.0,
  rotTitleX = 0,
  colTitleX = 'black',
  fontTitleX = 2,
  titleY = '',
  cexTitleY = 1.0,
  rotTitleY = 0,
  colTitleY = 'black',
  fontTitleY = 2,
  cexLabX = 1.0,
  rotLabX = 0,
  colLabX = 'black',
  fontLabX = 2,
  cexLabY = 1.0,
  rotLabY = 0,
  colLabY = 'black',
  fontLabY = 2,

```

```

posLab = 'bottomleft',
col = c('blue4', 'blue3', 'blue2', 'blue1', 'white',
        'red1', 'red2', 'red3', 'red4'),
posColKey = 'right',
cexLabColKey = 1.0,
cexCorval = 1.0,
colCorval = 'black',
fontCorval = 1,
scale = TRUE,
main = '',
cexMain = 2,
rotMain = 0,
colMain = 'black',
fontMain = 2,
corFUN = 'pearson',
corUSE = 'pairwise.complete.obs',
signifSymbols = c('***', '**', '*', ''),
signifCutpoints = c(0, 0.001, 0.01, 0.05, 1),
colFrame = 'white',
plotRsquared = FALSE,
returnPlot = TRUE)

```

### Arguments

pcaobj	Object of class 'pca' created by <code>pca()</code> . REQUIRED.
components	The principal components to be included in the plot. DEFAULT = <code>getComponents(pcaobj, seq_len(10))</code> . OPTIONAL.
metavars	A vector of column names in metadata representing continuous variables. REQUIRED.
titleX	X-axis title. DEFAULT = "". OPTIONAL.
cexTitleX	X-axis title cex. DEFAULT = 1.0. OPTIONAL.
rotTitleX	X-axis title rotation in degrees. DEFAULT = 0. OPTIONAL.
colTitleX	X-axis title colour. DEFAULT = 'black'. OPTIONAL.
fontTitleX	X-axis title font style. 1, plain; 2, bold; 3, italic; 4, bold-italic. DEFAULT = 2. OPTIONAL.
titleY	Y-axis title. DEFAULT = "". OPTIONAL.
cexTitleY	Y-axis title cex. DEFAULT = 1.0. OPTIONAL.
rotTitleY	Y-axis title rotation in degrees. DEFAULT = 0. OPTIONAL.
colTitleY	Y-axis title colour. DEFAULT = 'black'. OPTIONAL.
fontTitleY	Y-axis title font style. 1, plain; 2, bold; 3, italic; 4, bold-italic. DEFAULT = 2. OPTIONAL.
cexLabX	X-axis labels cex. DEFAULT = 1.0. OPTIONAL.
rotLabX	X-axis labels rotation in degrees. DEFAULT = 0. OPTIONAL.
colLabX	X-axis labels colour. DEFAULT = 'black'. OPTIONAL.
fontLabX	X-axis labels font style. 1, plain; 2, bold; 3, italic; 4, bold-italic. DEFAULT = 2. OPTIONAL.
cexLabY	Y-axis labels cex. DEFAULT = 1.0. OPTIONAL.

rotLabY	Y-axis labels rotation in degrees. DEFAULT = 0. OPTIONAL.
colLabY	Y-axis labels colour. DEFAULT = 'black'. OPTIONAL.
fontLabY	Y-axis labels font style. 1, plain; 2, bold; 3, italic; 4, bold-italic. DEFAULT = 2. OPTIONAL.
posLab	Positioning of the X- and Y-axis labels. 'bottomleft', bottom and left; 'topright', top and right; 'all', bottom / top and left /right; 'none', no labels. DEFAULT = 'bottomleft'. OPTIONAL.
col	Colour shade gradient for RColorBrewer. DEFAULT = c('blue4', 'blue3', 'blue2', 'blue1', 'white', 'red1', 'red2', 'red3', 'red4'). OPTIONAL.
posColKey	Position of colour key. 'bottom', 'left', 'top', 'right'. DEFAULT = 'right'. OPTIONAL.
cexLabColKey	Colour key labels cex. DEFAULT = 1.0. OPTIONAL.
cexCorval	Correlation values cex. DEFAULT = 1.0. OPTIONAL.
colCorval	Correlation values colour. DEFAULT = 'black'. OPTIONAL.
fontCorval	Correlation values font style. 1, plain; 2, bold; 3, italic; 4, bold-italic. DEFAULT = 1. OPTIONAL.
scale	Logical, indicating whether or not to scale the colour range to max and min cor values. DEFAULT = TRUE. OPTIONAL.
main	Plot title. DEFAULT = ''. OPTIONAL.
cexMain	Plot title cex. DEFAULT = 2. OPTIONAL.
rotMain	Plot title rotation in degrees. DEFAULT = 0. OPTIONAL.
colMain	Plot title colour. DEFAULT = 'black'. OPTIONAL.
fontMain	Plot title font style. 1, plain; 2, bold; 3, italic; 4, bold-italic. DEFAULT = 2. OPTIONAL.
corFUN	Correlation method: 'pearson', 'spearman', or 'kendall'. DEFAULT = 'pearson'. OPTIONAL.
corUSE	Method for handling missing values (see documentation for cor function via ?cor). 'everything', 'all.obs', 'complete.obs', 'na.or.complete', or 'pairwise.complete.obs'. DEFAULT = 'pairwise.complete.obs'. OPTIONAL.
signifSymbols	Statistical significance symbols to display beside correlation values. DEFAULT = c('****', '**', '*', ''). OPTIONAL.
signifCutpoints	Cut-points for statistical significance. DEFAULT = c(0, 0.001, 0.01, 0.05, 1). OPTIONAL.
colFrame	Frame colour. DEFAULT = 'white'. OPTIONAL.
plotRsquared	Logical, indicating whether or not to plot R-squared values. DEFAULT = FALSE. OPTIONAL.
returnPlot	Logical, indicating whether or not to return the plot object. DEFAULT = TRUE. OPTIONAL.

**Value**

A [lattice](#) object.

**Author(s)**

Kevin Blighe <kevin@clinicalbioinformatics.co.uk>, Myles Lewis <myles.lewis@qmul.ac.uk>



**Examples**

```

options(scipen=10)
options(digits=6)

col <- 20
row <- 20000
mat1 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat1) <- paste0('gene', 1:nrow(mat1))
colnames(mat1) <- paste0('sample', 1:ncol(mat1))

mat2 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat2) <- paste0('gene', 1:nrow(mat2))
colnames(mat2) <- paste0('sample', (ncol(mat1)+1):(ncol(mat1)+ncol(mat2)))

mat <- cbind(mat1, mat2)

metadata <- data.frame(row.names = colnames(mat))
metadata$Group <- rep(NA, ncol(mat))
metadata$Group[seq(1,40,2)] <- 'A'
metadata$Group[seq(2,40,2)] <- 'B'
metadata$CRP <- sample.int(100, size=ncol(mat), replace=TRUE)
metadata$ESR <- sample.int(100, size=ncol(mat), replace=TRUE)

p <- pca(mat, metadata = metadata, removeVar = 0.1)

eigencorplot(p, components = getComponents(p, 1:10),
  metavars = c('ESR', 'CRP'))

```

getComponents

*getComponents***Description**

Return the principal component labels for an object of class 'pca'.

**Usage**

```

getComponents(
  pcaobj,
  components = NULL)

```

**Arguments**

pcaobj	Object of class 'pca' created by <code>pca()</code> . <b>REQUIRED.</b>
components	Indices of the principal components whose names will be returned. If <code>NULL</code> , all PC names will be returned. <b>DEFAULT = NULL. OPTIONAL.</b>

**Value**

A [character](#) object.

**Author(s)**

Kevin Blighe <kevin@clinicalbioinformatics.co.uk>, Myles Lewis <myles.lewis@qmul.ac.uk>

**Examples**

```
options(scipen=10)
options(digits=6)

col <- 20
row <- 20000
mat1 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat1) <- paste0('gene', 1:nrow(mat1))
colnames(mat1) <- paste0('sample', 1:ncol(mat1))

mat2 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat2) <- paste0('gene', 1:nrow(mat2))
colnames(mat2) <- paste0('sample', (ncol(mat1)+1):(ncol(mat1)+ncol(mat2)))

mat <- cbind(mat1, mat2)

metadata <- data.frame(row.names = colnames(mat))
metadata$Group <- rep(NA, ncol(mat))
metadata$Group[seq(1,40,2)] <- 'A'
metadata$Group[seq(2,40,2)] <- 'B'
metadata$CRP <- sample.int(100, size=ncol(mat), replace=TRUE)
metadata$ESR <- sample.int(100, size=ncol(mat), replace=TRUE)

p <- pca(mat, metadata = metadata, removeVar = 0.1)

getComponents(p)
```

---

getLoadings

*getLoadings*

---

**Description**

Return component loadings for principal components from an object of class 'pca'.

**Usage**

```
getLoadings(
  pcaobj,
  components = NULL)
```

**Arguments**

pcaobj	Object of class 'pca' created by <code>pca()</code> . <b>REQUIRED.</b>
components	Indices of the principal components whose component loadings will be returned. If NULL, all PC names will be returned. <b>DEFAULT = NULL. OPTIONAL.</b>

**Value**

A `data.frame` object.

**Author(s)**

Kevin Blighe <kevin@clinicalbioinformatics.co.uk>, Myles Lewis <myles.lewis@qmul.ac.uk>

**Examples**

```
options(scipen=10)
options(digits=6)

col <- 20
row <- 20000
mat1 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat1) <- paste0('gene', 1:nrow(mat1))
colnames(mat1) <- paste0('sample', 1:ncol(mat1))

mat2 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat2) <- paste0('gene', 1:nrow(mat2))
colnames(mat2) <- paste0('sample', (ncol(mat1)+1):(ncol(mat1)+ncol(mat2)))

mat <- cbind(mat1, mat2)

metadata <- data.frame(row.names = colnames(mat))
metadata$Group <- rep(NA, ncol(mat))
metadata$Group[seq(1,40,2)] <- 'A'
metadata$Group[seq(2,40,2)] <- 'B'
metadata$CRP <- sample.int(100, size=ncol(mat), replace=TRUE)
metadata$ESR <- sample.int(100, size=ncol(mat), replace=TRUE)

p <- pca(mat, metadata = metadata, removeVar = 0.1)

getLoadings(p)
```

---

getVars

*getVars*

---

**Description**

Return the explained variance for each principal component for an object of class 'pca'.

**Usage**

```
getVars(
  pcaobj,
  components = NULL)
```

**Arguments**

`pcaobj` Object of class 'pca' created by `pca()`. REQUIRED.

`components` Indices of the principal components whose explained variances will be returned. If NULL, all values will be returned. DEFAULT = NULL. OPTIONAL.

**Value**

A numeric object.

**Author(s)**

Kevin Blighe <kevin@clinicalbioinformatics.co.uk>, Myles Lewis <myles.lewis@qmul.ac.uk>

**Examples**

```
options(scipen=10)
options(digits=6)

col <- 20
row <- 20000
mat1 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat1) <- paste0('gene', 1:nrow(mat1))
colnames(mat1) <- paste0('sample', 1:ncol(mat1))

mat2 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat2) <- paste0('gene', 1:nrow(mat2))
colnames(mat2) <- paste0('sample', (ncol(mat1)+1):(ncol(mat1)+ncol(mat2)))

mat <- cbind(mat1, mat2)

metadata <- data.frame(row.names = colnames(mat))
metadata$Group <- rep(NA, ncol(mat))
metadata$Group[seq(1,40,2)] <- 'A'
metadata$Group[seq(2,40,2)] <- 'B'
metadata$CRP <- sample.int(100, size=ncol(mat), replace=TRUE)
metadata$ESR <- sample.int(100, size=ncol(mat), replace=TRUE)

p <- pca(mat, metadata = metadata, removeVar = 0.1)

getVars(p)
```

---

pairsplot

*pairsplot*

---

**Description**

Draw multiple bi-plots.

**Usage**

```
pairsplot(pcaobj,  
  components = getComponents(pcaobj, seq_len(5)),  
  triangle = TRUE,  
  trianglelabSize = 18,  
  plotaxes = TRUE,  
  margingaps = unit(c(0.1, 0.1, 0.1, 0.1), 'cm'),  
  ncol = NULL,  
  nrow = NULL,  
  x = NULL,  
  y = NULL,  
  colby = NULL,  
  colkey = NULL,  
  singlecol = NULL,  
  shape = NULL,  
  shapekey = NULL,  
  pointSize = 1.0,  
  legendPosition = 'none',  
  legendLabSize = 6,  
  legendIconSize = 1.5,  
  xlim = NULL,  
  ylim = NULL,  
  lab = FALSE,  
  labSize = 1.5,  
  labhjust = 1.5,  
  labvjust = 0,  
  selectLab = NULL,  
  drawConnectors = FALSE,  
  widthConnectors = 0.5,  
  colConnectors = 'grey50',  
  xlab = NULL,  
  xlabAngle = 0,  
  xlabhjust = 0.5,  
  xlabvjust = 0.5,  
  ylab = NULL,  
  ylabAngle = 0,  
  ylabhjust = 0.5,  
  ylabvjust = 0.5,  
  axisLabSize = 10,  
  title = NULL,  
  titleLabSize = 32,  
  hline = NULL,  
  hlineType = 'longdash',  
  hlineCol = 'black',  
  hlineWidth = 0.4,  
  vline = NULL,  
  vlineType = 'longdash',  
  vlineCol = 'black',  
  vlineWidth = 0.4,  
  gridlines.major = TRUE,  
  gridlines.minor = TRUE,  
  borderWidth = 0.8,
```

```
borderColour = 'black',
returnPlot = TRUE)
```

### Arguments

pcaobj	Object of class 'pca' created by <code>pca()</code> . REQUIRED.
components	The principal components to be included in the plot. These will be compared in a pairwise fashion via multiple calls to <code>biplot()</code> . DEFAULT = <code>getComponents(pcaobj, seq_len(5))</code> . OPTIONAL.
triangle	Logical, indicating whether or not to draw the plots in the upper panel in a triangular arrangement? Principal component names will be labeled along the diagonal. DEFAULT = TRUE. OPTIONAL.
trianglelabSize	Size of p rincipal component label (when <code>triangle = TRUE</code> ). DEFAULT = 18. OPTIONAL.
plotaxes	Logical, indicating whether or not to draw the axis tick, labels, and titles. DEFAULT = TRUE. OPTIONAL.
marginaps	The margins between plots in the plot space. Takes the form of a 'unit()' variable. DEFAULT = <code>unit(c(0.1, 0.1, 0.1, 0.1), 'cm')</code> . OPTIONAL.
ncol	If <code>triangle = FALSE</code> , the number of columns in the final merged plot. DEFAULT = NULL. OPTIONAL.
nrow	If <code>triangle = FALSE</code> , the number of rows in the final merged plot. DEFAULT = NULL. OPTIONAL.
x	A principal component to plot on x-axis. All principal component names are stored in <code>pcaobj\$label</code> . DEFAULT = NULL. OPTIONAL.
y	A principal component to plot on y-axis. All principal component names are stored in <code>pcaobj\$label</code> . DEFAULT = NULL. OPTIONAL.
colby	If NULL, all points will be coloured differently. If not NULL, value is assumed to be a column name in <code>pcaobj\$metadata</code> relating to some grouping/categorical variable. DEFAULT = NULL. OPTIONAL.
colkey	Vector of name-value pairs relating to value passed to 'col', e.g., <code>c(A='forestgreen', B='gold')</code> . DEFAULT = NULL. OPTIONAL.
singlecol	If specified, all points will be shaded by this colour. Overrides 'col'. DEFAULT = NULL. OPTIONAL.
shape	If NULL, all points will be have the same shape. If not NULL, value is assumed to be a column name in <code>pcaobj\$metadata</code> relating to some grouping/categorical variable. DEFAULT = NULL. OPTIONAL.
shapekey	Vector of name-value pairs relating to value passed to 'shape', e.g., <code>c(A=10, B=21)</code> . DEFAULT = NULL. OPTIONAL.
pointSize	Size of plotted points. DEFAULT = 1.0. OPTIONAL.
legendPosition	Position of legend ('top', 'bottom', 'left', 'right', 'none'). DEFAULT = 'none'. OPTIONAL.
legendLabSize	Size of plot legend text. DEFAULT = 6. OPTIONAL.
legendIconSize	Size of plot legend icons / symbols. DEFAULT = 1.5. OPTIONAL.
xlim	Limits of the x-axis. DEFAULT = NULL. OPTIONAL.
ylim	Limits of the y-axis. DEFAULT = NULL. OPTIONAL.

lab	Logical, indicating whether or not to label the points in the plot space. Labels will be taken as the original colnames of the input object, usually sample IDs. DEFAULT = FALSE. OPTIONAL.
labSize	Size of labels. DEFAULT = 1.5. OPTIONAL.
labhjust	Horizontal adjustment of label. DEFAULT = 1.5. OPTIONAL.
labvjust	Vertical adjustment of label. DEFAULT = 0. OPTIONAL.
selectLab	A vector containing a subset of lab to plot. DEFAULT = NULL. OPTIONAL.
drawConnectors	Logical, indicating whether or not to connect plot labels to their corresponding points by line connectors. DEFAULT = FALSE. OPTIONAL.
widthConnectors	Line width of connectors. DEFAULT = 0.5. OPTIONAL.
colConnectors	Line colour of connectors. DEFAULT = 'grey50'. OPTIONAL.
xlab	Label for x-axis. DEFAULT = NULL. OPTIONAL.
xlabAngle	Rotation angle of x-axis labels. DEFAULT = 0. OPTIONAL.
xlabhjust	Horizontal adjustment of x-axis labels. DEFAULT = 0.5. OPTIONAL.
xlabvjust	Vertical adjustment of x-axis labels. DEFAULT = 0.5. OPTIONAL.
ylab	Label for y-axis. DEFAULT = NULL. OPTIONAL.
ylabAngle	Rotation angle of y-axis labels. DEFAULT = 0. OPTIONAL.
ylabhjust	Horizontal adjustment of y-axis labels. DEFAULT = 0.5. OPTIONAL.
ylabvjust	Vertical adjustment of y-axis labels. DEFAULT = 0.5. OPTIONAL.
axisLabSize	Size of x- and y-axis labels. DEFAULT = 10. OPTIONAL.
title	Plot title. DEFAULT = NULL. OPTIONAL.
titleLabSize	Size of plot title. DEFAULT = 32. OPTIONAL.
hline	Draw one or more horizontal lines passing through this/these values on y-axis. For single values, only a single numerical value is necessary. For multiple lines, pass these as a vector, e.g., c(60,90). DEFAULT = NULL. OPTIONAL.
hlineType	Line type for hline ('blank', 'solid', 'dashed', 'dotted', 'dotdash', 'longdash', 'twodash'). DEFAULT = 'longdash'. OPTIONAL.
hlineCol	Colour of hline. DEFAULT = 'black'. OPTIONAL.
hlineWidth	Width of hline. DEFAULT = 0.4. OPTIONAL.
vline	Draw one or more vertical lines passing through this/these values on x-axis. For single values, only a single numerical value is necessary. For multiple lines, pass these as a vector, e.g., c(60,90). DEFAULT = NULL. OPTIONAL.
vlineType	Line type for vline ('blank', 'solid', 'dashed', 'dotted', 'dotdash', 'longdash', 'twodash'). DEFAULT = 'longdash'. OPTIONAL.
vlineCol	Colour of vline. DEFAULT = 'black'. OPTIONAL.
vlineWidth	Width of vline. DEFAULT = 0.4. OPTIONAL.
gridlines.major	Logical, indicating whether or not to draw major gridlines. DEFAULT = TRUE. OPTIONAL.
gridlines.minor	Logical, indicating whether or not to draw minor gridlines. DEFAULT = TRUE. OPTIONAL.
borderWidth	Width of the border on the x and y axes. DEFAULT = 0.8. OPTIONAL.
borderColour	Colour of the border on the x and y axes. DEFAULT = 'black'. OPTIONAL.
returnPlot	Logical, indicating whether or not to return the plot object. DEFAULT = TRUE. OPTIONAL.

**Value**

A `cowplot` object.

**Author(s)**

Kevin Blighe <kevin@clinicalbioinformatics.co.uk>, Myles Lewis <myles.lewis@qmul.ac.uk>

**Examples**

```
options(scipen=10)
options(digits=6)

col <- 20
row <- 20000
mat1 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat1) <- paste0('gene', 1:nrow(mat1))
colnames(mat1) <- paste0('sample', 1:ncol(mat1))

mat2 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat2) <- paste0('gene', 1:nrow(mat2))
colnames(mat2) <- paste0('sample', (ncol(mat1)+1):(ncol(mat1)+ncol(mat2)))

mat <- cbind(mat1, mat2)

metadata <- data.frame(row.names = colnames(mat))
metadata$Group <- rep(NA, ncol(mat))
metadata$Group[seq(1,40,2)] <- 'A'
metadata$Group[seq(2,40,2)] <- 'B'
metadata$CRP <- sample.int(100, size=ncol(mat), replace=TRUE)
metadata$ESR <- sample.int(100, size=ncol(mat), replace=TRUE)

p <- pca(mat, metadata = metadata, removeVar = 0.1)

pairsplot(p, triangle = TRUE)
```

---

pca

*pca*

---

**Description**

Principal Components Analysis (PCA) is a very powerful technique that has wide applicability in data science, bioinformatics, and further afield. It was initially developed to analyse large volumes of data in order to tease out the differences/relationships between the logical entities being analysed. It extracts the fundamental structure of the data without the need to build any model to represent it. This 'summary' of the data is arrived at through a process of reduction that can transform the large number of variables into a lesser number that are uncorrelated, i.e., the principal components', whilst at the same time being capable of easy interpretation on the original data.



**Usage**

```
pca(
  mat,
  metadata = NULL,
  center = TRUE,
  scale = FALSE,
  removeVar = NULL)
```

**Arguments**

mat	A data-matrix or data-frame containing numerical data only. <b>REQUIRED.</b>
metadata	A data-matrix or data-frame containing metadata. This will be stored in the resulting <code>pca</code> object. Strictly enforced that <code>rownames(metadata) == colnames(mat)</code> . <b>DEFAULT = NULL. OPTIONAL.</b>
center	Center the data before performing PCA? Same as <code>prcomp()</code> 'center' parameter. <b>DEFAULT = TRUE. OPTIONAL.</b>
scale	Scale the data? Same as <code>prcomp()</code> 'scale' parameter. <b>DEFAULT = FALSE. OPTIONAL.</b>
removeVar	Remove this <b>DEFAULT = NULL. OPTIONAL.</b>

**Value**

A `pca` object.

**Author(s)**

Kevin Blighe <kevin@clinicalbioinformatics.co.uk>, Myles Lewis <myles.lewis@qmul.ac.uk>

**Examples**

```
options(scipen=10)
options(digits=6)

col <- 20
row <- 20000
mat1 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat1) <- paste0('gene', 1:nrow(mat1))
colnames(mat1) <- paste0('sample', 1:ncol(mat1))

mat2 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat2) <- paste0('gene', 1:nrow(mat2))
colnames(mat2) <- paste0('sample', (ncol(mat1)+1):(ncol(mat1)+ncol(mat2)))

mat <- cbind(mat1, mat2)

metadata <- data.frame(row.names = colnames(mat))
metadata$Group <- rep(NA, ncol(mat))
metadata$Group[seq(1,40,2)] <- 'A'
metadata$Group[seq(2,40,2)] <- 'B'
metadata$CRP <- sample.int(100, size=ncol(mat), replace=TRUE)
```

```

metadata$ESR <- sample.int(100, size=ncol(mat), replace=TRUE)

p <- pca(mat, metadata = metadata, removeVar = 0.1)

getComponents(p)

getVars(p)

getLoadings(p)

screeplot(p)

screeplot(p, hline = 80)

biplot(p)

biplot(p, colby = 'Group', shape = 'Group')

biplot(p, colby = 'Group', colkey = c(A = 'forestgreen', B = 'gold'),
  legendPosition = 'right')

biplot(p, colby = 'Group', colkey = c(A='forestgreen', B='gold'),
  shape = 'Group', shapekey = c(A=10, B=21), legendPosition = 'bottom')

pairsplot(p, triangle = TRUE)

plotloadings(p, drawConnectors=TRUE)

eigencorplot(p, components = getComponents(p, 1:10),
  metavars = c('ESR', 'CRP'))

```

---

plotloadings

*plotloadings*


---

### Description

Plot the component loadings for selected principal components / eigenvectors and label variables driving variation along these.

### Usage

```

plotloadings(pcaobj,
  components = getComponents(pcaobj, seq_len(5)),
  rangeRetain = 0.05,
  absolute = FALSE,
  col = c('gold', 'white', 'royalblue'),
  colMidpoint = 0,
  shape = 21,
  shapeSizeRange = c(10, 10),
  legendPosition = 'top',
  legendLabSize = 10,
  legendIconSize = 3.0,
  xlim = NULL,

```

```

ylim = NULL,
labSize = 2.0,
labhjust = 1.5,
labvjust = 0,
drawConnectors = TRUE,
positionConnectors = 'right',
widthConnectors = 0.5,
typeConnectors = 'closed',
endsConnectors = 'first',
lengthConnectors = unit(0.01, 'npc'),
colConnectors = 'grey50',
xlab = 'Principal component',
xlabAngle = 0,
xlabhjust = 0.5,
xlabvjust = 0.5,
ylab = 'Component loading',
ylabAngle = 0,
ylabhjust = 0.5,
ylabvjust = 0.5,
axisLabSize = 16,
title = '',
subtitle = '',
caption = '',
titleLabSize = 16,
subtitleLabSize = 12,
captionLabSize = 12,
hline = c(0),
hlineType = 'longdash',
hlineCol = 'black',
hlineWidth = 0.4,
vline = NULL,
vlineType = 'longdash',
vlineCol = 'black',
vlineWidth = 0.4,
gridlines.major = TRUE,
gridlines.minor = TRUE,
borderWidth = 0.8,
borderColour = 'black',
returnPlot = TRUE)

```

### Arguments

pcaobj	Object of class 'pca' created by <code>pca()</code> . REQUIRED.
components	The principal components to be included in the plot. DEFAULT = <code>getComponents(pcaobj, seq_len(5))</code> . OPTIONAL.
rangeRetain	Cut-off value for retaining variables. The function will look across each specified principal component and retain the variables that fall within this top/bottom fraction of the loadings range. DEFAULT = 0.05. OPTIONAL.
absolute	Logical, indicating whether or not to plot absolute loadings. DEFAULT = FALSE. OPTIONAL.
col	Colours used for generation of fill gradient according to loadings values. Can be 2 or 3 colours. DEFAULT = <code>c('gold', 'white', 'royalblue')</code> . OPTIONAL.

colMidpoint	Mid-point (loading) for the colour range. DEFAULT = 0. OPTIONAL.
shape	Shape of the plotted points. DEFAULT = 21. OPTIONAL.
shapeSizeRange	Size range for the plotted points (min, max). DEFAULT = c(10, 10). OPTIONAL.
legendPosition	Position of legend ('top', 'bottom', 'left', 'right', 'none'). DEFAULT = 'top'. OPTIONAL.
legendLabSize	Size of plot legend text. DEFAULT = 10. OPTIONAL.
legendIconSize	Size of plot legend icons / symbols. DEFAULT = 3.0. OPTIONAL.
xlim	Limits of the x-axis. DEFAULT = NULL. OPTIONAL.
ylim	Limits of the y-axis. DEFAULT = NULL. OPTIONAL.
labSize	Size of labels. DEFAULT = 2.0. OPTIONAL.
labhjust	Horizontal adjustment of label. DEFAULT = 1.5. OPTIONAL.
labvjust	Vertical adjustment of label. DEFAULT = 0. OPTIONAL.
drawConnectors	Logical, indicating whether or not to connect plot labels to their corresponding points by line connectors. DEFAULT = TRUE. OPTIONAL.
positionConnectors	Position of the connectors and their labels with respect to the plotted points ('left', 'right'). DEFAULT = 'right'. OPTIONAL.
widthConnectors	Line width of connectors. DEFAULT = 0.5. OPTIONAL.
typeConnectors	Have the arrow head open or filled ('closed')? ('open', 'closed'). DEFAULT = 'closed'. OPTIONAL.
endsConnectors	Which end of connectors to draw arrow head? ('last', 'first', 'both'). DEFAULT = 'first'. OPTIONAL.
lengthConnectors	Length of the connectors. DEFAULT = unit(0.01, 'npc'). OPTIONAL.
colConnectors	Line colour of connectors. DEFAULT = 'grey50'. OPTIONAL.
xlab	Label for x-axis. DEFAULT = 'Principal component'. OPTIONAL.
xlabAngle	Rotation angle of x-axis labels. DEFAULT = 0. OPTIONAL.
xlabhjust	Horizontal adjustment of x-axis labels. DEFAULT = 0.5. OPTIONAL.
xlabvjust	Vertical adjustment of x-axis labels. DEFAULT = 0.5. OPTIONAL.
ylab	Label for y-axis. DEFAULT = 'Component loading'. OPTIONAL.
ylabAngle	Rotation angle of y-axis labels. DEFAULT = 0. OPTIONAL.
ylabhjust	Horizontal adjustment of y-axis labels. DEFAULT = 0.5. OPTIONAL.
ylabvjust	Vertical adjustment of y-axis labels. DEFAULT = 0.5. OPTIONAL.
axisLabSize	Size of x- and y-axis labels. DEFAULT = 16. OPTIONAL.
title	Plot title. DEFAULT = ''. OPTIONAL.
subtitle	Plot subtitle. DEFAULT = ''. OPTIONAL.
caption	Plot caption. DEFAULT = ''. OPTIONAL.
titleLabSize	Size of plot title. DEFAULT = 16. OPTIONAL.
subtitleLabSize	Size of plot subtitle. DEFAULT = 12. OPTIONAL.
captionLabSize	Size of plot caption. DEFAULT = 12. OPTIONAL.

<code>hline</code>	Draw one or more horizontal lines passing through this/these values on y-axis. For single values, only a single numerical value is necessary. For multiple lines, pass these as a vector, e.g., <code>c(60,90)</code> . DEFAULT = <code>c(0)</code> . OPTIONAL.
<code>hlineType</code>	Line type for <code>hline</code> ('blank', 'solid', 'dashed', 'dotted', 'dotdash', 'longdash', 'twodash'). DEFAULT = 'longdash'. OPTIONAL.
<code>hlineCol</code>	Colour of <code>hline</code> . DEFAULT = 'black'. OPTIONAL.
<code>hlineWidth</code>	Width of <code>hline</code> . DEFAULT = 0.4. OPTIONAL.
<code>vline</code>	Draw one or more vertical lines passing through this/these values on x-axis. For single values, only a single numerical value is necessary. For multiple lines, pass these as a vector, e.g., <code>c(60,90)</code> . DEFAULT = NULL. OPTIONAL.
<code>vlineType</code>	Line type for <code>vline</code> ('blank', 'solid', 'dashed', 'dotted', 'dotdash', 'longdash', 'twodash'). DEFAULT = 'longdash'. OPTIONAL.
<code>vlineCol</code>	Colour of <code>vline</code> . DEFAULT = 'black'. OPTIONAL.
<code>vlineWidth</code>	Width of <code>vline</code> . DEFAULT = 0.4. OPTIONAL.
<code>gridlines.major</code>	Logical, indicating whether or not to draw major gridlines. DEFAULT = TRUE. OPTIONAL.
<code>gridlines.minor</code>	Logical, indicating whether or not to draw minor gridlines. DEFAULT = TRUE. OPTIONAL.
<code>borderWidth</code>	Width of the border on the x and y axes. DEFAULT = 0.8. OPTIONAL.
<code>borderColour</code>	Colour of the border on the x and y axes. DEFAULT = 'black'. OPTIONAL.
<code>returnPlot</code>	Logical, indicating whether or not to return the plot object. DEFAULT = TRUE. OPTIONAL.

**Value**

A `ggplot2` object.

**Author(s)**

Kevin Blighe <kevin@clinicalbioinformatics.co.uk>, Myles Lewis <myles.lewis@qmul.ac.uk>

**Examples**

```
options(scipen=10)
options(digits=6)

col <- 20
row <- 20000
mat1 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat1) <- paste0('gene', 1:nrow(mat1))
colnames(mat1) <- paste0('sample', 1:ncol(mat1))

mat2 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat2) <- paste0('gene', 1:nrow(mat2))
colnames(mat2) <- paste0('sample', (ncol(mat1)+1):(ncol(mat1)+ncol(mat2)))
```

```

mat <- cbind(mat1, mat2)

metadata <- data.frame(row.names = colnames(mat))
metadata$Group <- rep(NA, ncol(mat))
metadata$Group[seq(1,40,2)] <- 'A'
metadata$Group[seq(2,40,2)] <- 'B'
metadata$CRP <- sample.int(100, size=ncol(mat), replace=TRUE)
metadata$ESR <- sample.int(100, size=ncol(mat), replace=TRUE)

p <- pca(mat, metadata = metadata, removeVar = 0.1)

plotloadings(p, drawConnectors=TRUE)

```

---

screeplot

*screeplot*


---

### Description

Draw a SCREE plot, showing the distribution of explained variance across all or select principal components / eigenvectors.

### Usage

```

screeplot(pcaobj,
  components = getComponents(pcaobj),
  xlim = NULL,
  ylim = c(0, 100),
  xlab = 'Principal component',
  xlabAngle = 90,
  xlabhjust = 0.5,
  xlabvjust = 0.5,
  ylab = 'Explained variation (%)',
  ylabAngle = 0,
  ylabhjust = 0.5,
  ylabvjust = 0.5,
  axisLabSize = 16,
  title = 'SCREE plot',
  subtitle = '',
  caption = '',
  titleLabSize = 16,
  subtitleLabSize = 12,
  captionLabSize = 12,
  colBar = 'dodgerblue',
  drawCumulativeSumLine = TRUE,
  colCumulativeSumLine = 'red2',
  sizeCumulativeSumLine = 1.5,
  drawCumulativeSumPoints = TRUE,
  colCumulativeSumPoints = 'red2',
  sizeCumulativeSumPoints = 2.0,
  hline = NULL,
  hlineType = 'longdash',

```

```

hlineCol = 'black',
hlineWidth = 0.4,
vline = NULL,
vlineType = 'longdash',
vlineCol = 'black',
vlineWidth = 0.4,
gridlines.major = TRUE,
gridlines.minor = TRUE,
borderWidth = 0.8,
borderColour = 'black',
returnPlot = TRUE)

```

### Arguments

pcaobj	Object of class 'pca' created by <code>pca()</code> . REQUIRED.
components	The principal components to be included in the plot. DEFAULT = <code>getComponents(pcaobj)</code> . OPTIONAL.
xlim	Limits of the x-axis. DEFAULT = NULL. OPTIONAL.
ylim	Limits of the y-axis. DEFAULT = <code>c(0, 100)</code> . OPTIONAL.
xlab	Label for x-axis. DEFAULT = 'Principal component'. OPTIONAL.
xlabAngle	Rotation angle of x-axis labels. DEFAULT = 90. OPTIONAL.
xlabhjust	Horizontal adjustment of x-axis labels. DEFAULT = 0.5. OPTIONAL.
xlabvjust	Vertical adjustment of x-axis labels. DEFAULT = 0.5. OPTIONAL.
ylab	Label for y-axis. DEFAULT = 'Explained variation (%)'. OPTIONAL.
ylabAngle	Rotation angle of y-axis labels. DEFAULT = 0. OPTIONAL.
ylabhjust	Horizontal adjustment of y-axis labels. DEFAULT = 0.5. OPTIONAL.
ylabvjust	Vertical adjustment of y-axis labels. DEFAULT = 0.5. OPTIONAL.
axisLabSize	Size of x- and y-axis labels. DEFAULT = 16. OPTIONAL.
title	Plot title. DEFAULT = 'SCREE plot'. OPTIONAL.
subtitle	Plot subtitle. DEFAULT = ''. OPTIONAL.
caption	Plot caption. DEFAULT = ''. OPTIONAL.
titleLabSize	Size of plot title. DEFAULT = 16. OPTIONAL.
subtitleLabSize	Size of plot subtitle. DEFAULT = 12. OPTIONAL.
captionLabSize	Size of plot caption. DEFAULT = 12. OPTIONAL.
colBar	DEFAULT = 'dodgerblue'. OPTIONAL.
drawCumulativeSumLine	Logical, indicating whether or not to overlay plot with a cumulative explained variance line. DEFAULT = TRUE. OPTIONAL.
colCumulativeSumLine	Colour of cumulative explained variance line. DEFAULT = 'red2'. OPTIONAL.
sizeCumulativeSumLine	Size of cumulative explained variance line. DEFAULT = 1.5. OPTIONAL.
drawCumulativeSumPoints	Logical, indicating whether or not to draw the cumulative explained variance points. DEFAULT = TRUE. OPTIONAL.

<code>colCumulativeSumPoints</code>	Colour of cumulative explained variance points. DEFAULT = 'red2'. OPTIONAL.
<code>sizeCumulativeSumPoints</code>	Size of cumulative explained variance points. DEFAULT = 2.0. OPTIONAL.
<code>hline</code>	Draw one or more horizontal lines passing through this/these values on y-axis. For single values, only a single numerical value is necessary. For multiple lines, pass these as a vector, e.g., <code>c(60,90)</code> . DEFAULT = NULL. OPTIONAL.
<code>hlineType</code>	Line type for <code>hline</code> ('blank', 'solid', 'dashed', 'dotted', 'dotdash', 'longdash', 'twodash'). DEFAULT = 'longdash'. OPTIONAL.
<code>hlineCol</code>	Colour of <code>hline</code> . DEFAULT = 'black'. OPTIONAL.
<code>hlineWidth</code>	Width of <code>hline</code> . DEFAULT = 0.4. OPTIONAL.
<code>vline</code>	Draw one or more vertical lines passing through this/these values on x-axis. For single values, only a single numerical value is necessary. For multiple lines, pass these as a vector, e.g., <code>c(60,90)</code> . DEFAULT = NULL. OPTIONAL.
<code>vlineType</code>	Line type for <code>vline</code> ('blank', 'solid', 'dashed', 'dotted', 'dotdash', 'longdash', 'twodash'). DEFAULT = 'longdash'. OPTIONAL.
<code>vlineCol</code>	Colour of <code>vline</code> . DEFAULT = 'black'. OPTIONAL.
<code>vlineWidth</code>	Width of <code>vline</code> . DEFAULT = 0.4. OPTIONAL.
<code>gridlines.major</code>	Logical, indicating whether or not to draw major gridlines. DEFAULT = TRUE. OPTIONAL.
<code>gridlines.minor</code>	Logical, indicating whether or not to draw minor gridlines. DEFAULT = TRUE. OPTIONAL.
<code>borderWidth</code>	Width of the border on the x and y axes. DEFAULT = 0.8. OPTIONAL.
<code>borderColour</code>	Colour of the border on the x and y axes. DEFAULT = 'black'. OPTIONAL.
<code>returnPlot</code>	Logical, indicating whether or not to return the plot object. DEFAULT = TRUE. OPTIONAL.

**Value**

A `ggplot2` object.

**Author(s)**

Kevin Blighe <kevin@clinicalbioinformatics.co.uk>, Myles Lewis <myles.lewis@qmul.ac.uk>

**Examples**

```
options(scipen=10)
options(digits=6)

col <- 20
row <- 20000
mat1 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat1) <- paste0('gene', 1:nrow(mat1))
colnames(mat1) <- paste0('sample', 1:ncol(mat1))
```



```
mat2 <- matrix(
  rexp(col*row, rate = 0.1),
  ncol = col)
rownames(mat2) <- paste0('gene', 1:nrow(mat2))
colnames(mat2) <- paste0('sample', (ncol(mat1)+1):(ncol(mat1)+ncol(mat2)))

mat <- cbind(mat1, mat2)

metadata <- data.frame(row.names = colnames(mat))
metadata$Group <- rep(NA, ncol(mat))
metadata$Group[seq(1,40,2)] <- 'A'
metadata$Group[seq(2,40,2)] <- 'B'
metadata$CRP <- sample.int(100, size=ncol(mat), replace=TRUE)
metadata$ESR <- sample.int(100, size=ncol(mat), replace=TRUE)

p <- pca(mat, metadata = metadata, removeVar = 0.1)

screepilot(p)

screepilot(p, hline = 80)
```

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