# **R** documentation

of 'blandAltmanLehPlot.Rd'

September 3, 2020

blandAltmanLehPlot Generate Bland-Altman agreement plot between predicted and expected cell type proportions from test samples.

# Description

Generate Bland-Altman agreement plot between predicted and expected cell type proportions from test samples. The Bland-Altman agreement plots can be displayed all mixed or split based on cell type (CellType) or the number of cell types present in the sample (nMix). See facet.by argument and examples for more information.

#### Usage

```
blandAltmanLehPlot(
  object,
  colors,
  color.by,
  facet.by = NULL,
  log.2 = FALSE,
  filter.sc = TRUE,
  density = TRUE,
  color.density = "darkblue",
  size.point = 0.05,
  alpha.point = 1,
  ncol = NULL,
  nrow = NULL
  title = NULL,
  theme = theme_grey(),
)
```

#### **Arguments**

object

DigitalDLSorter object with trained.model slot containing metrics in eval.stats.samples slot.

2 blandAltmanLehPlot

colors	Vector of colors to use. Only vectors with a number of colors equal to or greater than the levels of color.by will be accepted. By default it is used a list of custom colors provided by the package.
color.by	Variable used to color data. The options are nMix and CellType.
facet.by	Variable used to display data in different panels. If it is NULL, the plot is not separated into different panels. The options are nMix (by number of different cell types) and CellType (by cell type).
filter.sc	Boolean indicating if filter single-cell profiles and only display correlations of results associated with bulk samples (TRUE by default).
density	Boolean indicating if show density lines (TRUE by default).
color.density	
	Color of density lines if density argument is equal to TRUE.
size.point	Size of points (0.1 by default).
alpha.point	Alpha of points (0.1 by default).
ncol	Number of columns if facet.by is different than NULL.
nrow	Number of rows if facet.by is different than NULL.
title	Title of the plot.
theme	ggplot theme.
•••	Additional argument for facet_wrap ggplot function if facet.by is not equal to NULL.
log2	If show Bland-Altman agreement plot in log2 space (FALSE by default).

### See Also

calculateEvalMetrics corrExpPredPlot distErrorPlot barErrorPlot

# **Examples**

```
## Bland-Altman plot by cell type
blandAltmanLehPlot(
  object = DDLSChung,
  facet.by = "CellType",
  color.by = "CellType",
  corr = "both"
)
## Bland-Altman plot of all samples mixed
blandAltmanLehPlot(
  object = DDLSChung,
  facet.by = NULL,
  color.by = "CellType",
  alpha.point = 0.3
)
```

# Index

```
barErrorPlot, 2
blandAltmanLehPlot, 1

calculateEvalMetrics, 2
corrExpPredPlot, 2

distErrorPlot, 2
```