

# Package ‘ggQC’

January 30, 2018

**Type** Package

**Title** Quality Control Charts for 'ggplot'

**Version** 0.0.2

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**Description** Plot single and faceted type quality control charts for 'ggplot'.

**Depends** R (>= 2.10)

**License** GPL-3 | file LICENSE

**Encoding** UTF-8

**LazyData** true

**Suggests** testthat, gridExtra, knitr, rmarkdown, reshape2, plyr

**RoxygenNote** 6.0.1

**Imports** ggplot2, stats, dplyr, tidyr

**VignetteBuilder** knitr

**NeedsCompilation** no

**Repository** CRAN

**Date/Publication** 2018-01-30 14:23:59 UTC

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---

cBar\_LCL

*Lower Control Limit: Count Data (c-chart)*

---

**Description**

Calculates lower control limit (LCL) for count data acquired over the same-sized area of opportunity. Negative values are reported as 0.

**Usage**

```
cBar_LCL(y, na.rm = FALSE, ...)
```

**Arguments**

y	Vector of count data. Each observation having the same-area of opportunity.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
...	further arguments passed to or from other methods.

**Value**

A number; 3-sigma lower control limit (LCL). Function returns 0 for negative values.

**Examples**

```
set.seed(5555)
y <- rpois(30, 9)
cBar_LCL(y)
```

---

cBar\_UCL

*Upper Control Limit: Count Data (c-chart)*

---

**Description**

Calculates upper control limit (UCL) for count data acquired over the same-sized area of opportunity.

**Usage**

```
cBar_UCL(y, na.rm = FALSE, ...)
```

**Arguments**

y	Vector of count data. Each observation having the same-area of opportunity.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
...	further arguments passed to or from other methods.

**Value**

A number; 3-sigma upper control limit (UCL)

**Examples**

```
set.seed(5555)
y <- rpois(30, 9)
cBar_UCL(y)
```

---

mR	<i>Mean One-Point Moving Range</i>
----	------------------------------------

---

**Description**

Calculates the mean one-point moving range used when constructing a moving-range chart.

**Usage**

```
mR(y, na.rm = TRUE, ...)
```

**Arguments**

y	Vector of values
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
...	further arguments passed to or from other methods.

**Value**

A number; mean one-point moving range.

**Examples**

```
set.seed(5555)
values <- rnorm(n = 100, mean = 25, sd = 1)
mR(values)
```

---

mR_points	<i>One Point Moving Range of Vector</i>
-----------	---

---

**Description**

Calculates a one-point moving range vector given an input vector of values. Output often used to produce mR-chart.

**Usage**

```
mR_points(y)
```

**Arguments**

y : vector of values

**Value**

Vector of one-point moving range.

**Examples**

```
y <- seq(-5:5)
mR_points(y)
```

---

mR\_UCL

*Mean One-Point Moving Range Upper Control Limit (UCL)*

---

**Description**

Calculates the mean one-point moving range UCL used when constructing a moving-range chart.

**Usage**

```
mR_UCL(y, na.rm = FALSE, ...)
```

**Arguments**

y Vector of values

na.rm a logical value indicating whether NA values should be stripped before the computation proceeds.

... further arguments passed to or from other methods.

**Value**

A number; mean one-point moving range UCL.

**Examples**

```
set.seed(5555)
values <- rnorm(n = 100, mean = 25, sd = 1)
mR_UCL(values)
```

---

npBar	<i>Mean Value: Binomial Data (np-chart)</i>
-------	---

---

**Description**

Calculates the mean value for binomial count data acquired over the same-sized area of opportunity.

**Usage**

```
npBar(y, n, na.rm = FALSE, ...)
```

**Arguments**

y	Vector of binomial count data (not proportions). Each observation having the same-area of opportunity.
n	A number representing the area of opportunity.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
...	further arguments passed to or from other methods.

**Value**

A number; mean value

**Examples**

```
set.seed(5555)
p <- rbinom(n = 100, size = 30, prob = .2)
npBar(y = p, n = 30)
```

---

npBar_LCL	<i>Lower Control Limit: Binomial Data (np-chart)</i>
-----------	--

---

**Description**

Calculates lower control limit (LCL) for binomial count data acquired over the same-sized area of opportunity.

**Usage**

```
npBar_LCL(y, n, na.rm = FALSE, ...)
```

**Arguments**

y	Vector of binomial count data (not proportions). Each observation having the same-area of opportunity.
n	A number representing the area of opportunity.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
...	further arguments passed to or from other methods.

**Value**

A number; 3-sigma upper control limit (LCL)

**Examples**

```
set.seed(5555)
p <- rbinom(n = 100, size = 30, prob = .2)
npBar_LCL(y = p, n = 30)
```

---

npBar\_UCL

*Upper Control Limit: Binomial Data (np-chart)*


---

**Description**

Calculates upper control limit (UCL) for binomial count data acquired over the same-sized area of opportunity.

**Usage**

```
npBar_UCL(y, n, na.rm = FALSE, ...)
```

**Arguments**

y	Vector of binomial count data (not proportions). Each observation having the same-area of opportunity.
n	A number representing the area of opportunity.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
...	further arguments passed to or from other methods.

**Value**

A number; 3-sigma upper control limit (UCL)

## Examples

```
set.seed(5555)
p <- rbinom(n = 100, size = 30, prob = .2)
npBar_UCL(y = p, n = 30)
```

---

pBar

*Mean Proportion: Binomial Data (p-chart)*

---

## Description

Calculates overall mean proportion for binomial proportion data acquired over a variable area of opportunity.

## Usage

```
pBar(y, n, na.rm = FALSE, ...)
```

## Arguments

y	Vector of binomial proportion data (not counts). Observations may have a different area of opportunity, n.
n	A vector representing the area of opportunity.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
...	further arguments passed to or from other methods.

## Value

A vector of mean proportion, length equal to length of parameter y.

## Examples

```
set.seed(5555)
p <- rbinom(n = 100, size = 30, prob = .2)
n <- rpois(100, 100)
pBar(y = p/n, n = n)
```

---

pBar\_LCL                      *Lower Control Limit: Binomial Data (p-chart)*

---

**Description**

Calculates point-wise lower control limit (LCL) for binomial proportion data acquired over a variable area of opportunity.

**Usage**

```
pBar_LCL(y, n, na.rm = FALSE, ...)
```

**Arguments**

**y**                      Vector of binomial proportion data (not counts). Observations may have a different area of opportunity, n.

**n**                      A vector representing the area of opportunity.

**na.rm**                a logical value indicating whether NA values should be stripped before the computation proceeds.

**...**                further arguments passed to or from other methods.

**Value**

A vector; point-wise 3-sigma lower control limit (LCL)

**Examples**

```
set.seed(5555)
p <- rbinom(n = 100, size = 30, prob = .2)
n <- rpois(100, 100)
pBar_LCL(y = p/n, n = n)
```

---

pBar\_UCL                      *Upper Control Limit: Binomial Data (p-chart)*

---

**Description**

Calculates point-wise upper control limit (UCL) for binomial proportion data acquired over a variable area of opportunity.

**Usage**

```
pBar_UCL(y, n, na.rm = FALSE, ...)
```

**Arguments**

<code>y</code>	Vector of binomial proportion data (not counts). Observations may have a different area of opportunity, <code>n</code> .
<code>n</code>	A vector representing the area of opportunity.
<code>na.rm</code>	a logical value indicating whether NA values should be stripped before the computation proceeds.
<code>...</code>	further arguments passed to or from other methods.

**Value**

A vector; point-wise 3-sigma upper control limit (UCL)

**Examples**

```
set.seed(5555)
p <- rbinom(n = 100, size = 30, prob = .2)
n <- rpois(100, 100)
pBar_UCL(y = p/n, n = n)
```

---

QCrange

*Range: Max Min Difference*

---

**Description**

Given a set of numbers, function calculates the difference between the maximum and minimum value.

**Usage**

```
QCrange(y)
```

**Arguments**

`y` : vector of values

**Value**

a number.

**Examples**

```
y <- seq(-5:5)
QCrange(y)
```

**Description**

Calculates QC chart lines for the following chart types and reports in a dataframe:

- **Individuals Charts:** mR, XmR,
- **Attribute Charts:** c, np, p, u,
- **Studentized Charts:** xBar.rBar, xBar.rMedian, xBar.sBar, xMedian.rBar, xMedian.rMedian,
- **Dispersion Charts:** rBar, rMedian, sBar.

**Usage**

```
QC_Lines(data = NULL, value = NULL, grouping = NULL, formula = NULL,
         n = NULL, method = "xBar.rBar", na.rm = FALSE)
```

**Arguments**

data	vector or dataframe, as indicated below for each chart type <ul style="list-style-type: none"> <li>• <b>Individuals &amp; Attribute Charts:</b> vector of values;</li> <li>• <b>Studentized &amp; Dispersion Charts:</b> dataframe</li> </ul>
value	<b>Studentized Charts</b> and <b>Dispersion Charts</b> , numeric vector in dataframe with values of interest
grouping	<b>Studentized Charts</b> and <b>Dispersion Charts:</b> single factor/variable to split the dataframe "values" by
formula	<b>Studentized Charts</b> and <b>Dispersion Charts:</b> a formula, such as $y \sim x1 + x2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables
n	number or vector as indicated below for each chart type. <ul style="list-style-type: none"> <li>• <b>Individuals Charts:</b> No effect</li> <li>• <b>Attribute Charts:</b> (p and u) vector, indicating sample area of opportunity.</li> <li>• <b>Attribute Charts:</b> (np) number, indicating constant sampling area of opportunity.</li> <li>• <b>Studentized Charts:</b> number, user specified subgroup size.</li> <li>• <b>Dispersion Charts:</b> No effect</li> </ul>
method	string, calling the following methods: <ul style="list-style-type: none"> <li>• <b>Individuals Charts:</b> mR, XmR,</li> <li>• <b>Attribute Charts:</b> c, np, p, u,</li> <li>• <b>Studentized Charts:</b> xBar.rBar, xBar.rMedian, xBar.sBar, xMedian.rBar, xMedian.rMedian</li> <li>• <b>Dispersion Charts:</b> rBar, rMedian, sBar.</li> </ul>
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.

**Value**

a dataframe,

- **Attribute Data:** (p and u) Center Line, Upper Control Limit and Lower Control limit for each point.
- **Other Data:** single line dataframe, with relevant control limits noted in column headings.

**Note**

If using the **formula** argument do not use **value** and **group** arguments.

**References**

Wheeler, DJ, and DS Chambers. Understanding Statistical Process Control, 2nd Ed. Knoxville, TN: SPC, 1992. Print.

**Examples**

```
#####
# Example 1: Charts other than "p" or "u" #
#####

# Load Libraries -----
require(ggQC)
require(plyr)
require(ggplot2)

# Setup Data -----
set.seed(5555)
Process1 <- data.frame(processID = as.factor(rep(1,100)),
                      metric_value = rnorm(100,0,1),
                      subgroup_sample=rep(1:20, each=5),
                      Process_run_id = 1:100)

set.seed(5555)
Process2 <- data.frame(processID = as.factor(rep(2,100)),
                      metric_value = rnorm(100,5, 1),
                      subgroup_sample=rep(1:10, each=10),
                      Process_run_id = 101:200)

Both_Processes <- rbind(Process1, Process2)

# QC Values For Individuals -----
# All Together
QC_Lines(data = Both_Processes$metric_value, method = "XmR")

# For Each Process
ddply(Both_Processes, .variables = "processID",
      .fun =function(df){
        QC_Lines(data = df$metric_value, method = "XmR")
      }
    )
```

```

)

# QC Values For Studentized Runs-----
# All Together
  QC_Lines(data = Both_Processes,
           formula = metric_value ~ subgroup_sample)

# For Each Process
  dply(Both_Processes, .variables = "processID",
       .fun =function(df){
         QC_Lines(data = df, formula = metric_value ~ subgroup_sample)
       }
)

#####
# Example 2 "p" data #
#####

# Setup p Data -----
set.seed(5555)
bin_data <- data.frame(
  trial = 1:30,
  Num_Incomplete_Items = rpois(n = 30, lambda = 30),
  Num_Items_in_Set = runif(n = 30, min = 50, max = 100))

bin_data$Proportion_Incomplete <- bin_data$Num_Incomplete_Items/bin_data$Num_Items_in_Set

# QC_Lines for "p" data -----
QC_Lines(data = bin_data$Proportion_Incomplete,
         n = bin_data$Num_Items_in_Set, method="p")

#####
# Example 3 "u" data #
#####

# Setup u Data -----
set.seed(5555)
bin_data <- data.frame(
  trial=1:30,
  Num_of_Blemishes = rpois(n = 30, lambda = 30),
  Num_Items_Inspected = runif(n = 30, min = 50, max = 100))

bin_data$Blemish_Rate <- bin_data$Num_of_Blemishes/bin_data$Num_Items_Inspected

# QC Lines for "u" data -----
QC_Lines(data = bin_data$Blemish_Rate,
         n = bin_data$Num_Items_Inspected, method="u")

```

QC\_Violations

*Calculate QC Violations***Description**

function that calculates QC violations on sequentially ordered data based on the following 4 rules:

- **Violation Same Side:** 8 or more consecutive, same-side points
- **Violation 1 Sigma:** 4 or more consecutive, same-side points exceeding 1 sigma
- **Violation 2 Sigma:** 2 or more consecutive, same-side points exceeding 2 sigma
- **Violation 3 Sigma:** any points exceeding 3 sigma

**Usage**

```
QC_Violations(data, value = NULL, grouping = NULL, formula = NULL,
              method = NULL, ...)
```

**Arguments**

data	vector or dataframe, as indicated below for each chart type <ul style="list-style-type: none"> <li>• <b>Individuals:</b> vector of values;</li> <li>• <b>Studentized Charts:</b> dataframe</li> </ul>
value	<b>Studentized Charts:</b> numeric vector in dataframe with values of interest
grouping	<b>Studentized Charts:</b> single factor/variable to split the dataframe "values" by
formula	<b>Studentized Charts:</b> a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables
method	string, calling the following methods: <ul style="list-style-type: none"> <li>• <b>Individuals Charts:</b> XmR,</li> <li>• <b>Studentized Charts:</b> xBar.rBar, xBar.rMedian, xBar.sBar, xMedian.rBar, xMedian.rMedian</li> </ul>
...	further arguments passed to or from other methods.

**Value**

a dataframe, with the following columns

- **data:** The input data if XmR, mean or median by group for Studentized methods
- **z\_score:** z-score for the data point
- **Index:** number, indicating the order of the input data
- **Violation\_Result:** description of the type of test being run.
  - **Violation Same Side:** 8 or more consecutive, same-side points
  - **Violation 1 Sigma:** 4 or more consecutive, same-side points exceeding 1 sigma
  - **Violation 2 Sigma:** 2 or more consecutive, same-side points exceeding 2 sigma
  - **Violation 3 Sigma:** any points exceeding 3 sigma
- **Index:** boolean, does the data point violate the rule?

**Note**

If using the **formula** argument do not use **value** and **group** arguments.

**References**

Wheeler, DJ, and DS Chambers. Understanding Statistical Process Control, 2nd Ed. Knoxville, TN: SPC, 1992. Print.

**Examples**

```
#####
# Example 1: XmR Check Violations #
#####
# Load Libraries -----
require(ggQC)

# Setup Data -----

set.seed(5555)
QC_XmR <- data.frame(
  data = c(c(-1, 2.3, 2.4, 2.5), #Outlier Data
           sample(c(rnorm(60),5,-5), 62, replace = FALSE), #Normal Data
           c(1,-.3, -2.4,-2.6,-2.5,-2.7, .3)), #Outlier Data
  Run_Order = 1:73 #Run Order
)

QC_Vs <- QC_Violations(data = QC_XmR$data, method = "XmR")

#####
# Example 2: Xbar Check Violations #
#####

# Setup Some Data -----
QC_xBar.rBar <- do.call(rbind, lapply(1:3, function(X){
  set.seed(5555+X) #Loop over 3 seeds
  data.frame(
    sub_group = rep(1:42), #Define Subgroups
    sub_class = letters[X],
    c(
      c(runif(n = 5, min = 2.0,3.2)), #Outlier Data
      sample(c(rnorm(30),5,-4), 32, replace = FALSE), #Normal Data
      c(runif(n = 5, min = -3.2, max = -2.0)) #Outlier Data
    )
  )
})
)

colnames(QC_xBar.rBar) <- c("sub_group","sub_class", "value")
QC_Vs <- QC_Violations(data = QC_xBar.rBar,
  formula = value~sub_group,
  method = "xBar.rBar")
```

---

rBar	<i>Mean Subgroup Range</i>
------	----------------------------

---

**Description**

Calculates the mean subgroup range used when constructing a XbarR chart.

**Usage**

```
rBar(data, value, grouping, formula = NULL, ...)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables
...	further arguments passed to or from other methods.

**Value**

A number; mean subgroup range.

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
rBar(data = df, formula = v~g)
```

---

rBar_LCL	<i>Mean Subgroup Range Lower Control Limit (LCL)</i>
----------	--

---

**Description**

Calculates the mean subgroup range Lower control limit (UCL) used when constructing a XbarR chart.

**Usage**

```
rBar_LCL(data = data, value = value, grouping = grouping,
  formula = NULL, ...)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
formula	a formula, such as $y \sim x1 + x2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables
...	further arguments passed to or from other methods.

**Value**

A number; mean subgroup range lower control limit (LCL).

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
rBar_LCL(data = df, formula = v~g)
```

---

rBar\_UCL

*Mean Subgroup Range Upper Control Limit (UCL)*


---

**Description**

Calculates the mean subgroup range upper control limit (UCL) used when constructing a XbarR chart.

**Usage**

```
rBar_UCL(data = data, value = value, grouping = grouping,
          formula = NULL, ...)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
formula	a formula, such as $y \sim x1 + x2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables
...	further arguments passed to or from other methods.

**Value**

A number; mean subgroup range upper control limit (UCL).

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
rBar_UCL(data = df, formula = v~g)
```

---

rMedian

*Median of Subgroup Ranges*


---

**Description**

Calculates the median of subgroup ranges, used when constructing xBar\_rMedian charts.

**Usage**

```
rMedian(data, value, grouping, formula = NULL, ...)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
formula	a formula, such as $y \sim x1 + x2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables
...	further arguments passed to or from other methods.

**Value**

A number; median subgroup range.

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
rMedian(data = df, formula = v~g)
```

---

rMedian_LCL	<i>Median of Subgroup Ranges Lower Control Limit (LCL)</i>
-------------	--

---

**Description**

Calculates the median of subgroup range Lower control limit (LCL) used when constructing a xBar\_rMedian chart.

**Usage**

```
rMedian_LCL(data = data, value = value, grouping = grouping,
            formula = NULL, ...)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
formula	a formula, such as $y \sim x1 + x2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables
...	further arguments passed to or from other methods.

**Value**

A number; median of subgroup range lower control limit (LCL).

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
rMedian_LCL(data = df, formula = v~g)
```

---

rMedian_UCL	<i>Median of Subgroup Ranges Upper Control Limit (UCL)</i>
-------------	--

---

**Description**

Calculates the median of subgroup range upper control limit (UCL) used when constructing a xBar\_rMedian chart.

**Usage**

```
rMedian_UCL(data = data, value = value, grouping = grouping,
            formula = NULL, ...)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables
...	further arguments passed to or from other methods.

**Value**

A number; median of subgroup range upper control limit (UCL).

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
rMedian_UCL(data = df, formula = v~g)
```

---

sBar

*Mean Subgroup Standard Deviation*


---

**Description**

Calculates the mean subgroup standard deviation used when constructing a XbarS chart.

**Usage**

```
sBar(data, value, grouping, formula = NULL, ...)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables
...	further arguments passed to or from other methods.

**Value**

A number; mean subgroup standard deviation.

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
sBar(data = df, formula = v~g)
```

---

sBar_LCL	<i>Mean Subgroup Standard Deviation Lower Control Limit (LCL)</i>
----------	---

---

**Description**

Calculates the mean subgroup standard deviation Lower control limit (UCL) used when constructing a XbarS chart.

**Usage**

```
sBar_LCL(data = data, value = value, grouping = grouping,
          formula = NULL, ...)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
formula	a formula, such as $y \sim x1 + x2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables
...	further arguments passed to or from other methods.

**Value**

A number; mean subgroup standard deviation lower control limit (LCL).

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
sBar_LCL(data = df, formula = v~g)
```

---

sBar_UCL	<i>Mean Subgroup Standard Deviation Upper Control Limit (UCL)</i>
----------	---

---

**Description**

Calculates the mean subgroup standard deviation upper control limit (UCL) used when constructing a XbarS chart.

**Usage**

```
sBar_UCL(data = data, value = value, grouping = grouping,
          formula = NULL, ...)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables
...	further arguments passed to or from other methods.

**Value**

A number; mean subgroup standard deviation upper control limit (UCL).

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
sBar_UCL(data = df, formula = v~g)
```

---

stat\_mR

*Generate mR chart in ggplot*


---

**Description**

ggplot stat used to create a mR chart in ggplot

**Usage**

```
stat_mR(mapping = NULL, data = NULL, geom = "point",
        position = "identity", show.legend = NA, inherit.aes = TRUE,
        na.rm = FALSE, color.mr_point = "black", color.mr_line = "black",
        color.qc_limits = "red", color.qc_center = "blue", ...)
```

**Arguments**

mapping	Set of aesthetic mappings created by <a href="#">aes</a> or <a href="#">aes_</a> . If specified and <code>inherit.aes = TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.
data	The data to be displayed in this layer. There are three options: If <code>NULL</code> , the default, the data is inherited from the plot data as specified in the call to <a href="#">ggplot</a> . A <code>data.frame</code> , or other object, will override the plot data. All objects will be fortified to produce a data frame. See <a href="#">fortify</a> for which variables will be created. A function will be called with a single argument, the plot data. The return value must be a <code>data.frame</code> , and will be used as the layer data.

<code>geom</code>	The geometric object to use display the data
<code>position</code>	Position adjustment, either as a string, or the result of a call to a position adjustment function.
<code>show.legend</code>	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
<code>inherit.aes</code>	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <code>borders</code> .
<code>na.rm</code>	a logical value indicating whether NA values should be stripped before the computation proceeds.
<code>color.mr_point</code>	color, to be used for the mR points.
<code>color.mr_line</code>	color, to be used for line connecting points.
<code>color.qc_limits</code>	color, used to colorize the plot's upper and lower mR control limits.
<code>color.qc_center</code>	color, used to colorize the plot's center line.
<code>...</code>	other arguments passed on to <code>layer</code> . These are often aesthetics, used to set an aesthetic to a fixed value, like <code>color = "red"</code> or <code>size = 3</code> . They may also be parameters to the paired <code>geom/stat</code> .

## Value

data need to produce the mR plot in ggplot.

## Examples

```
#####
# Example 1: mR Chart #
#####

# Load Libraries -----
require(ggQC)
require(ggplot2)

# Setup Data -----
set.seed(5555)
Process1 <- data.frame(processID = as.factor(rep(1,100)),
                      metric_value = rnorm(100,0,1),
                      subgroup_sample=rep(1:20, each=5),
                      Process_run_id = 1:100)

set.seed(5556)
Process2 <- data.frame(processID = as.factor(rep(2,100)),
                      metric_value = rnorm(100,5, 1),
                      subgroup_sample=rep(1:10, each=10),
                      Process_run_id = 101:200)

Both_Processes <- rbind(Process1, Process2)
```

```
# One Plot Both Processes -----
ggplot(Both_Processes, aes(x=Process_run_id, y = metric_value)) +
  stat_mR() + ylab("Moving Range")

# Facet Plot - Both Processes -----
ggplot(Both_Processes, aes(x=Process_run_id, y = metric_value)) +
  stat_mR() + ylab("Moving Range") +
  facet_grid(.~processID, scales = "free_x")
```

stat\_pareto

*Generate a Pareto Plot with ggplot***Description**

stat function to create ggplot Pareto chart

**Usage**

```
stat_pareto(mapping = NULL, data = NULL, geom = "point",
  position = "identity", show.legend = NA, inherit.aes = TRUE,
  group = 1, na.rm = FALSE, point.color = "black", point.size = 2,
  line.color = "black", line.size = 0.5, bars.fill = c("red", "white"),
  ...)
```

**Arguments**

mapping	Set of aesthetic mappings created by <a href="#">aes</a> or <a href="#">aes_</a> . If specified and <code>inherit.aes = TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.
data	The data to be displayed in this layer. There are three options: If <code>NULL</code> , the default, the data is inherited from the plot data as specified in the call to <a href="#">ggplot</a> . A <code>data.frame</code> , or other object, will override the plot data. All objects will be fortified to produce a data frame. See <a href="#">fortify</a> for which variables will be created. A function will be called with a single argument, the plot data. The return value must be a <code>data.frame</code> , and will be used as the layer data.
geom	The geometric object to use display the data
position	Position adjustment, either as a string, or the result of a call to a position adjustment function.
show.legend	logical. Should this layer be included in the legends? <code>NA</code> , the default, includes if any aesthetics are mapped. <code>FALSE</code> never includes, and <code>TRUE</code> always includes.
inherit.aes	If <code>FALSE</code> , overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <a href="#">borders</a> .

group	defines grouping for variable for pareto plot, default and suggested is 1.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
point.color	color, used to define point color of cumulative percentage line
point.size	number, used to define point size of cumulative percentage line
line.color	color, used to define line color of cumulative percentage line
line.size	color, used to define line weight of cumulative percentage line
bars.fill	character vector length 2, start and end colors for pareto bars.
...	other arguments passed on to <a href="#">layer</a> . These are often aesthetics, used to set an aesthetic to a fixed value, like color = "red" or size = 3. They may also be parameters to the paired geom/stat.

## Value

Pareto plot.

## Examples

```
#####
# Example 1: Pareto Plot #
#####

# Load Libraries -----
require(ggQC)
require(ggplot2)

# Setup Data -----
df <- data.frame(
  x = letters[1:10],
  y = as.integer(runif(n = 10, min = 0, max=100))
)

# Render Pareto Plot -----

ggplot(df, aes(x=x, y=y)) +
  stat_pareto(point.color = "red",
             point.size = 3,
             line.color = "black",
             #size.line = 1,
             bars.fill = c("blue", "orange"),
             )
```

stat\_QC

*Produce QC Charts with ggplot Framework.***Description**

Produce QC charts with ggplot framework. Support for faceting and layering of multiple QC chart lines on a single plot. Charts supported (see method argument for call):

- **Individuals Charts:** mR, XmR,
- **Attribute Charts:** c, np, p, u,
- **Studentized Charts:** xBar.rBar, xBar.rMedian, xBar.sBar, xMedian.rBar, xMedian.rMedian,
- **Dispersion Charts:** rBar, rMedian, sBar.

To label chart lines see [stat\\_QC\\_labels](#)

**Usage**

```
stat_QC(mapping = NULL, data = NULL, geom = "hline",
        position = "identity", na.rm = FALSE, show.legend = NA,
        inherit.aes = TRUE, n = NULL, method = "xBar.rBar",
        color.qc_limits = "red", color.qc_center = "green", ...)
```

**Arguments**

mapping	Set of aesthetic mappings created by <a href="#">aes</a> or <a href="#">aes_</a> . If specified and <code>inherit.aes = TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.
data	The data to be displayed in this layer. There are three options: If <code>NULL</code> , the default, the data is inherited from the plot data as specified in the call to <a href="#">ggplot</a> . A <code>data.frame</code> , or other object, will override the plot data. All objects will be fortified to produce a data frame. See <a href="#">fortify</a> for which variables will be created. A function will be called with a single argument, the plot data. The return value must be a <code>data.frame</code> , and will be used as the layer data.
geom	The geometric object to use display the data
position	Position adjustment, either as a string, or the result of a call to a position adjustment function.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
show.legend	logical. Should this layer be included in the legends? <code>NA</code> , the default, includes if any aesthetics are mapped. <code>FALSE</code> never includes, and <code>TRUE</code> always includes.
inherit.aes	If <code>FALSE</code> , overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <a href="#">borders</a> .

n number, for

- **Studentized Charts**, used for custom or hypothetical subgroup size.
- **np Charts**, used to specify a fixed area of opportunity.

method string, calling the following methods:

- **Individuals Charts**: mR, XmR,
- **Attribute Charts**: c, np, p, u,
- **Studentized Charts**: xBar.rBar, xBar.rMedian, xBar.sBar, xMedian.rBar, xMedian.rMedian
- **Dispersion Charts**: rBar, rMedian, sBar.

color.qc\_limits color, used to colorize the plot's upper and lower mR control limits.

color.qc\_center color, used to colorize the plot's center line.

... other arguments passed on to [layer](#). These are often aesthetics, used to set an aesthetic to a fixed value, like `color = "red"` or `size = 3`. They may also be parameters to the paired geom/stat.

## Value

data need to produce the mR plot in ggplot.

## Examples

```
# Load Libraries -----
require(ggQC)
require(ggplot2)

# Setup Data -----
set.seed(5555)
Process1 <- data.frame(processID = as.factor(rep(1,100)),
  metric_value = rnorm(100,0,1),
  subgroup_sample = rep(1:20, each=5),
  Process_run_id = 1:100)

set.seed(5556)
Process2 <- data.frame(processID = as.factor(rep(2,100)),
  metric_value = rnorm(100,5, 1),
  subgroup_sample = rep(1:10, each=10),
  Process_run_id = 101:200)

Both_Processes <- rbind(Process1, Process2)

#####
# Example 1: XmR Chart #
#####

EX1.1 <- ggplot(Both_Processes, aes(x=Process_run_id, y = metric_value)) +
  geom_point() + geom_line() + stat_QC(method="XmR") +
  stat_QC_labels(method="XmR", digits = 2) +
```

```

  facet_grid(~processID, scales = "free_x")
#EX1.1

EX1.2 <- ggplot(Both_Processes, aes(x=Process_run_id, y = metric_value)) +
  stat_mR() + ylab("Moving Range") +
  stat_QC_labels(method="mR", digits = 2) +
  facet_grid(~processID, scales = "free_x")
#EX1.2

#####
# Example 2: XbarR Chart #
#####

EX2.1 <- ggplot(Both_Processes, aes(x = subgroup_sample,
                                   y = metric_value,
                                   group = processID)) +
  stat_summary(fun.y = "mean", color = "blue", geom = c("point")) +
  stat_summary(fun.y = "mean", color = "blue", geom = c("line")) +
  stat_QC(method = "xBar.rBar") + facet_grid(~processID, scales = "free_x")
#EX2.1

EX2.2 <- ggplot(Both_Processes, aes(x = subgroup_sample,
                                   y = metric_value,
                                   group = processID)) +
  stat_summary(fun.y = "QCrange", color = "blue", geom = "point") +
  stat_summary(fun.y = "QCrange", color = "blue", geom = "line") +
  stat_QC(method = "rBar") +
  ylab("Range") +
  facet_grid(~processID, scales = "free_x")
#EX2.2

#####
# Example 3: p Chart #
#####
# p chart Setup -----
set.seed(5556)
bin_data <- data.frame(
  trial=1:30,
  Num_Incomplete_Items = rpois(30, lambda = 30),
  Num_Items_in_Set = runif(n = 30, min = 50, max = 100))
bin_data$Proportion_Incomplete <- bin_data$Num_Incomplete_Items/bin_data$Num_Items_in_Set

# Plot p chart -----
EX3.1 <- ggplot(data = bin_data, aes(x=trial,
                                   y=Proportion_Incomplete,
                                   n=Num_Items_in_Set)) +
  geom_point() + geom_line() +
  stat_QC(method = "p")
#EX3.1

#####
# Example 4: u Chart #
#####

```

```

# u chart Setup -----
set.seed(5555)
bin_data <- data.frame(
  trial=1:30,
  Num_of_Blemishes = rpois(30, lambda = 30),
  Num_Items_Inspected = runif(n = 30, min = 50, max = 100)
)
bin_data$Blemish_Rate <- bin_data$Num_of_Blemishes/bin_data$Num_Items_Inspected

# Plot u chart -----
EX4.1 <- ggplot(data = bin_data, aes(x=trial,
                                     y=Blemish_Rate,
                                     n=Num_Items_Inspected)) +
  geom_point() + geom_line() +
  stat_QC(method = "u")
#EX4.1

#####
# Example 5: np Chart #
#####
# np chart Setup -----
set.seed(5555)
bin_data <- data.frame(
  trial=1:30,
  NumNonConforming = rbinom(30, 30, prob = .50))
Units_Tested_Per_Batch <- 60

# Plot np chart -----
EX5.1 <- ggplot(data = bin_data, aes(trial, NumNonConforming)) +
  geom_point() +
  stat_QC(method = "np", n = Units_Tested_Per_Batch)
#EX5.1

```

---

stat\_QC\_labels

---

*Write QC Line Labels to ggplot QC Charts.*


---

## Description

Write QC line labels to ggplot QC Charts. Useful if you want to see the value of the center line and QC limits. see method argument for methods supported.

## Usage

```

stat_QC_labels(mapping = NULL, data = NULL, geom = "label",
  position = "identity", na.rm = FALSE, show.legend = NA,
  inherit.aes = TRUE, n = NULL, digits = 1, method = "xBar.rBar",
  color.qc_limits = "red", color.qc_center = "black", text.size = 3, ...)

```

**Arguments**

mapping	Set of aesthetic mappings created by <a href="#">aes</a> or <a href="#">aes_</a> . If specified and <code>inherit.aes = TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.
data	The data to be displayed in this layer. There are three options: If <code>NULL</code> , the default, the data is inherited from the plot data as specified in the call to <a href="#">ggplot</a> . A <code>data.frame</code> , or other object, will override the plot data. All objects will be fortified to produce a data frame. See <a href="#">fortify</a> for which variables will be created. A function will be called with a single argument, the plot data. The return value must be a <code>data.frame</code> , and will be used as the layer data.
geom	The geometric object to use display the data
position	Position adjustment, either as a string, or the result of a call to a position adjustment function.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <a href="#">borders</a> .
n	number, for <ul style="list-style-type: none"> <li>• <b>Studentized Charts</b>, used for custom or hypothetical subgroup size.</li> <li>• <b>np Charts</b>, used to specify a fixed area of opportunity.</li> </ul>
digits	integer, indicating the number of decimal places
method	string, calling the following methods: <ul style="list-style-type: none"> <li>• <b>Individuals Charts</b>: <code>mR</code>, <code>XmR</code>,</li> <li>• <b>Attribute Charts</b>: <code>c</code>, <code>np</code>, <code>p</code>, <code>u</code>,</li> <li>• <b>Studentized Charts</b>: <code>xBar.rBar</code>, <code>xBar.rMedian</code>, <code>xBar.sBar</code>, <code>xMedian.rBar</code>, <code>xMedian.rMedian</code></li> <li>• <b>Dispersion Charts</b>: <code>rBar</code>, <code>rMedian</code>, <code>sBar</code>.</li> </ul>
color.qc_limits	color, used to colorize the plot's upper and lower mR control limits.
color.qc_center	color, used to colorize the plot's center line.
text.size	number, size of the text label
...	other arguments passed on to <a href="#">layer</a> . These are often aesthetics, used to set an aesthetic to a fixed value, like <code>color = "red"</code> or <code>size = 3</code> . They may also be parameters to the paired <code>geom/stat</code> .

**Value**

data need to produce the mR plot in [ggplot](#).

## Examples

```
#####
# Example 1: mR Chart #
#####

# Load Libraries -----
require(ggQC)
require(ggplot2)

# Setup Data -----
set.seed(5555)
Process1 <- data.frame(processID = as.factor(rep(1,100)),
                       metric_value = rnorm(100,0,1),
                       subgroup_sample=rep(1:20, each=5),
                       Process_run_id = 1:100)

set.seed(5556)
Process2 <- data.frame(processID = as.factor(rep(2,100)),
                       metric_value = rnorm(100,5, 1),
                       subgroup_sample=rep(1:10, each=10),
                       Process_run_id = 101:200)

Both_Processes <- rbind(Process1, Process2)

# Facet Plot - Both Processes -----
EX1.1 <- ggplot(Both_Processes, aes(x=Process_run_id, y = metric_value)) +
  geom_point() + geom_line() + stat_QC(method="XmR") +
  stat_QC_labels(method="XmR", digits = 2) +
  facet_grid(~processID, scales = "free_x")
#EX1.1

EX1.2 <- ggplot(Both_Processes, aes(x=Process_run_id, y = metric_value)) +
  stat_mR() + ylab("Moving Range") +
  stat_QC_labels(method="mR", digits = 2) +
  facet_grid(~processID, scales = "free_x")
#EX1.2

#####
# Example 2: XbarR Chart #
#####
# Facet Plot - Studentized Process -----

EX2.1 <- ggplot(Both_Processes, aes(x=subgroup_sample,
                                   y = metric_value,
                                   group = processID)) +
  geom_point(alpha=.2) +
  stat_summary(fun.y = "mean", color="blue", geom=c("point")) +
  stat_summary(fun.y = "mean", color="blue", geom=c("line")) +
  stat_QC() + facet_grid(~processID, scales = "free_x") +
  stat_QC_labels(text.size =3, label.size=.1)
#EX2.1

EX2.2 <- ggplot(Both_Processes, aes(x=subgroup_sample,
```

```

      y = metric_value,
      group = processID)) +
  stat_summary(fun.y = "QCrange", color="blue", geom = "point") +
  stat_summary(fun.y = "QCrange", color="blue", geom = "line") +
  stat_QC(method="rBar") +
  stat_QC_labels(digits=2, method="rBar") +
  ylab("Range") +
  facet_grid(.~processID, scales = "free_x")
#EX2.2

```

---

stat\_qc\_violations      *Inspect QC Violations*

---

## Description

ggplot stat function that renders a faceted plot of QC violations based on the following 4 rules:

- **Violation Same Side:** 8 or more consecutive, same-side points
- **Violation 1 Sigma:** 4 or more consecutive, same-side points exceeding 1 sigma
- **Violation 2 Sigma:** 2 or more consecutive, same-side points exceeding 2 sigma
- **Violation 3 Sigma:** any points exceeding 3 sigma

## Usage

```

stat_qc_violations(mapping = NULL, data = NULL, geom = "point",
  position = "identity", show.legend = NA, inherit.aes = TRUE,
  na.rm = FALSE, method = "xBar.rBar", geom_points = TRUE,
  geom_line = TRUE, point.size = 1.5, point.color = "black",
  violation_point.color = "red", rule.color = "darkgreen",
  line.color = NULL, ...)

```

## Arguments

mapping	Set of aesthetic mappings created by <a href="#">aes</a> or <a href="#">aes_</a> . If specified and <code>inherit.aes = TRUE</code> (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.
data	The data to be displayed in this layer. There are three options: If <code>NULL</code> , the default, the data is inherited from the plot data as specified in the call to <a href="#">ggplot</a> . A <code>data.frame</code> , or other object, will override the plot data. All objects will be fortified to produce a data frame. See <a href="#">fortify</a> for which variables will be created. A function will be called with a single argument, the plot data. The return value must be a <code>data.frame.</code> , and will be used as the layer data.
geom	The geometric object to use display the data

position	Position adjustment, either as a string, or the result of a call to a position adjustment function.
show.legend	logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.
inherit.aes	If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. <code>borders</code> .
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
method	string, calling the following methods: <ul style="list-style-type: none"> <li>• <b>Individuals Charts:</b> <code>XmR</code>,</li> <li>• <b>Studentized Charts:</b> <code>xBar.rBar</code>, <code>xBar.rMedian</code>, <code>xBar.sBar</code>, <code>xMedian.rBar</code>, <code>xMedian.rMedian</code></li> </ul>
geom_points	boolean, draw points
geom_line	boolean, draw line
point.size	number, size of points on chart
point.color	string, color of points on charts (e.g., "black")
violation_point.color	string, color of violation points on charts (e.g., "red")
rule.color	string, color or horizontal rules indicating distribution center and sigma levels
line.color	string, color of lines connecting points
...	other arguments passed on to <code>layer</code> . These are often aesthetics, used to set an aesthetic to a fixed value, like <code>color = "red"</code> or <code>size = 3</code> . They may also be parameters to the paired <code>geom/stat</code> .

## Value

faceted plot.

## Examples

```
#####
# Example 1: XmR Check Violations #
#####
# Load Libraries -----
require(ggQC)
require(ggplot2)

# Setup Data -----

set.seed(5555)
QC_XmR <- data.frame(
  data = c(c(-1, 2.3, 2.4, 2.5), #Outlier Data
           sample(c(rnorm(60),5,-5), 62, replace = FALSE), #Normal Data
           c(1,-.3, -2.4,-2.6,-2.5,-2.7, .3)), #Outlier Data
  Run_Order = 1:73 #Run Order
```

```

)

# Render QC Violation Plot -----

EX1 <- ggplot(QC_XmR, aes(x = Run_Order, y = data)) +
  stat_qc_violations(method = "XmR") #Makes facet graph with violations
#EX1
#####
# Example 2: Xbar Check Violations #
#####

# Setup Some Data -----
QC_xBar.rBar <- do.call(rbind, lapply(1:3, function(X){
  set.seed(5555+X) #Loop over 3 seeds
  data.frame(
    sub_group = rep(1:42), #Define Subgroups
    sub_class = letters[X],
    c(
      c(runif(n = 5, min = 2.0,3.2)), #Outlier Data
      sample(c(rnorm(30),5,-4), 32, replace = FALSE), #Normal Data
      c(runif(n = 5, min = -3.2, max = -2.0)) #Outlier Data
    )
  )
})
)

colnames(QC_xBar.rBar) <- c("sub_group", "sub_class", "value")

# Render QC Violation Plot -----
EX2 <- ggplot(QC_xBar.rBar, aes(x = sub_group, y = value)) +
  stat_qc_violations(method = "xBar.rBar")
  #stat_qc_violations(method="xBar.rMedian")
  #stat_qc_violations(method="xBar.sBar")
  #stat_qc_violations(method="xMedian.rBar")
  #stat_qc_violations(method="xMedian.rMedian")
#EX2
#####
# Complete User Control - Bypass stat_qc_violation #
#####
#### The code below has two options if you are looking for complete
#### control over the look and feel of the graph. Use option 1 or option
#### 2 as appropriate. If you want something quick and easy use examples above.

##### Option 1: Setup for XmR Type Data
# QC_XmR: Defined in Example 1
QC_Vs <- QC_Violations(data = QC_XmR$data, method = "XmR")
QC_Stats <- QC_Lines(data = QC_XmR$data, method = "XmR")
MEAN <- QC_Stats$mean
SIGMA <- QC_Stats$sigma

##### Option 2: Setup for xBar.rBar Type Data

```

```

# QC_xBar.rBar: Defined in Example 2
QC_Vs <- QC_Violations(data = QC_xBar.rBar,
                      formula = value~sub_group,
                      method = "xBar.rBar")
QC_Stats <- QC_Lines(data = QC_xBar.rBar,
                   formula = value~sub_group,
                   method = "xBar.rBar")
MEAN <- QC_Stats$xBar_Bar
SIGMA <- QC_Stats$sigma

##### Setup second table for horizontal rules
FacetNames <- c("Violation Same Side",
               "Violation 1 Sigma",
               "Violation 2 Sigma",
               "Violation 3 Sigma")

QC_Vs$Violation_Result <- ordered(QC_Vs$Violation_Result,
                                 levels=FacetNames)

QC_Stats_df <- data.frame(
  Violation_Result = factor(x = FacetNames, levels = FacetNames),
  SigmaPlus = MEAN+SIGMA*0:3,
  MEAN = MEAN,
  SigmaMinus = MEAN-SIGMA*0:3
)

##### Make the Plot
ggplot(QC_Vs, aes(x=Index, y=data, color=Violation, group=1)) +
  geom_point() + geom_line() +
  facet_grid(.~Violation_Result) +
  geom_hline(data = QC_Stats_df, aes(yintercept = c(SigmaPlus))) +
  geom_hline(data = QC_Stats_df, aes(yintercept = c(SigmaMinus))) +
  geom_hline(data = QC_Stats_df, aes(yintercept = c(MEAN)))

```

---

uBar

*Mean Rate: Count Data (u-chart)*


---

### Description

Calculates overall mean rate for count data acquired over a variable area of opportunity.

### Usage

```
uBar(y, n, na.rm = FALSE, ...)
```

### Arguments

**y** Vector of counts per unit opportunity (rate). Observations may have a different area of opportunity, n.

**n** A vector representing the area of opportunity.

na.rm            a logical value indicating whether NA values should be stripped before the computation proceeds.  
 ...            further arguments passed to or from other methods.

**Value**

A vector of mean rate, length equal to length of parameter y.

**Examples**

```
set.seed(5555)
counts <- rpois(100, 25)
n <- rpois(100, 15)
uBar(y = counts / n, n = n)
```

---

uBar\_LCL

*Lower Control Limit: Count Data (u-chart)*


---

**Description**

Calculates point-wise lower control limit (LCL) for count data acquired over a variable area of opportunity.

**Usage**

```
uBar_LCL(y, n, na.rm = FALSE, ...)
```

**Arguments**

y            Vector of counts per unit opportunity (rate). Observations may have a different area of opportunity, n.  
 n            A vector representing the area of opportunity.  
 na.rm       a logical value indicating whether NA values should be stripped before the computation proceeds.  
 ...        further arguments passed to or from other methods.

**Value**

A vector; point-wise 3-sigma lower control limit (LCL)

**Examples**

```
set.seed(5555)
counts <- rpois(100, 25)
n <- rpois(100, 15)
uBar_LCL(y = counts / n, n = n)
```

---

uBar\_UCL

*Upper Control Limit: Count Data (u-chart)*

---

### Description

Calculates point-wise upper control limit (UCL) for count data acquired over a variable area of opportunity.

### Usage

```
uBar_UCL(y, n, na.rm = FALSE, ...)
```

### Arguments

y	Vector of counts per unit opportunity (rate). Observations may have a different area of opportunity, n.
n	A vector representing the area of opportunity.
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
...	further arguments passed to or from other methods.

### Value

A vector; point-wise 3-sigma upper control limit (UCL)

### Examples

```
set.seed(5555)
counts <- rpois(100, 25)
n <- rpois(100, 15)
uBar_UCL(y = counts / n, n = n)
```

---

xBar\_Bar

*Mean of Subgroup Means*

---

### Description

Calculates the mean subgroup means used when constructing a xBar-R or xBar-S charts.

### Usage

```
xBar_Bar(data, value, grouping, formula = NULL, ...)
```

**Arguments**

<code>data</code>	data frame to be processed
<code>value</code>	numeric vector in a data frame with values of interest.
<code>grouping</code>	single factor/variable to split the data frame "values" by.
<code>formula</code>	a formula, such as $y \sim x_1 + x_2$ , where the $y$ variable is numeric data to be split into groups according to the grouping $x$ factors/variables
<code>...</code>	further arguments passed to or from other methods.

**Value**

A number; mean of subgroup means.

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
xBar_Bar(data = df, formula = v~g)
```

---

<code>xBar_one_LCL</code>	<i>xBar_One Lower Control Limit (LCL)</i>
---------------------------	---

---

**Description**

Calculates the `xBar_One` LCL used when constructing a `xBar-One` chart.

**Usage**

```
xBar_one_LCL(y, na.rm = FALSE, ...)
```

**Arguments**

<code>y</code>	Vector of values
<code>na.rm</code>	a logical value indicating whether NA values should be stripped before the computation proceeds.
<code>...</code>	further arguments passed to or from other methods.

**Value**

A number; `xBar_One` Lower Control Limit (LCL)

**Examples**

```
set.seed(5555)
values <- rnorm(n = 100, mean = 25, sd = 1)
xBar_one_LCL(values)
```

---

xBar_one_UCL	<i>xBar_One Upper Control Limit (UCL)</i>
--------------	---

---

**Description**

Calculates the xBar\_One UCL used when constructing a xBar-One chart.

**Usage**

```
xBar_one_UCL(y, na.rm = FALSE, ...)
```

**Arguments**

y	Vector of values
na.rm	a logical value indicating whether NA values should be stripped before the computation proceeds.
...	further arguments passed to or from other methods.

**Value**

A number; xBar\_One Upper Control Limit (UCL)

**Examples**

```
set.seed(5555)
values <- rnorm(n = 100, mean = 25, sd = 1)
xBar_one_UCL(values)
```

---

xBar_rBar_LCL	<i>Mean of Subgroup Means Lower Control Limit (LCL)</i>
---------------	---

---

**Description**

Calculates the mean of subgroup means lower control limit used when constructing a xBar-R charts.

**Usage**

```
xBar_rBar_LCL(data, value, grouping, n = NULL, natural = F,
  formula = NULL)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
n	a number indicating a hypothetical subgroup size other than, function determined subgroup n determined by the floor length of subgroup values.
natural	logical, if TRUE calculate limits for individuals (n=1) else calculate for n determined by the floor length of subgroup values
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables

**Value**

A number; mean of subgroup means lower control limit.

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
xBar_rBar_LCL(data = df, formula = v~g)
```

---

xBar\_rBar\_UCL

---

*Mean of Subgroup Means Upper Control Limit (UCL)*


---

**Description**

Calculates the mean of subgroup means upper control limit used when constructing a xBar-R charts.

**Usage**

```
xBar_rBar_UCL(data, value, grouping, n = NULL, natural = F,
  formula = NULL)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
n	a number indicating a hypothetical subgroup size other than, function determined subgroup n determined by the floor length of subgroup values.
natural	logical, if TRUE calculate limits for individuals (n=1) else calculate for n determined by the floor length of subgroup values
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables

**Value**

A number; mean of subgroup means upper control limit.

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
xBar_rBar_UCL(data = df, formula = v~g)
```

---

xBar_rMedian_LCL	<i>Mean of Subgroup Means Lower Control Limit (LCL) based on Median Range</i>
------------------	---

---

**Description**

Calculates the mean of subgroup means lower control limit based on the median range. The result is used when constructing a xBar-rMedian charts.

**Usage**

```
xBar_rMedian_LCL(data, value, grouping, n = NULL, natural = F,
  formula = NULL)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
n	a number indicating a hypothetical subgroup size other than, function determined subgroup n determined by the floor length of subgroup values.
natural	logical, if TRUE calculate limits for individuals (n=1) else calculate for n determined by the floor length of subgroup values
formula	a formula, such as $y \sim x1 + x2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables

**Value**

A number; mean of subgroup means Lower Control Limit (LCL) based on Median Range

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
xBar_rMedian_LCL(data = df, formula = v~g)
```

---

xBar_rMedian_UCL	<i>Mean of Subgroup Means Upper Control Limit (UCL) based on Median Range</i>
------------------	---

---

### Description

Calculates the mean of subgroup means upper control limit based on the median range. The result is used when constructing a xBar-rMedian charts.

### Usage

```
xBar_rMedian_UCL(data, value, grouping, n = NULL, natural = F,
  formula = NULL)
```

### Arguments

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
n	a number indicating a hypothetical subgroup size other than, function determined subgroup n determined by the floor length of subgroup values.
natural	logical, if TRUE calculate limits for individuals (n=1) else calculate for n determined by the floor length of subgroup values
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables

### Value

A number; mean of subgroup means Upper Control Limit (UCL) based on Median Range

### Examples

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
xBar_rMedian_UCL(data = df, formula = v~g)
```

---

xBar_sBar_LCL	<i>Mean of Subgroup Means Lower Control Limit (LCL) based on Standard Deviation</i>
---------------	---

---

### Description

Calculates the mean of subgroup means lower control limit based on the standard deviation. The result is used when constructing a xBar-S charts.

### Usage

```
xBar_sBar_LCL(data, value, grouping, n = NULL, natural = F,
  formula = NULL)
```

### Arguments

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
n	a number indicating a hypothetical subgroup size other than, function determined subgroup n determined by the floor length of subgroup values.
natural	logical, if TRUE calculate limits for individuals (n=1) else calculate for n determined by the floor length of subgroup values
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables

### Value

A number; mean of subgroup means Lower Control Limit (LCL) based on standard deviation

### Examples

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
xBar_sBar_LCL(data = df, formula = v~g)
```

---

xBar_sBar_UCL	<i>Mean of Subgroup Means Upper Control Limit (UCL) based on Standard Deviation</i>
---------------	---

---

### Description

Calculates the mean of subgroup means upper control limit based on the standard deviation. The result is used when constructing a xBar-S charts.

### Usage

```
xBar_sBar_UCL(data, value, grouping, n = NULL, natural = F,
  formula = NULL)
```

### Arguments

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
n	a number indicating a hypothetical subgroup size other than, function determined subgroup n determined by the floor length of subgroup values.
natural	logical, if TRUE calculate limits for individuals (n=1) else calculate for n determined by the floor length of subgroup values
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables

### Value

A number; mean of subgroup means Upper Control Limit (UCL) based on standard deviation

### Examples

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
xBar_sBar_UCL(data = df, formula = v~g)
```

---

xMedian_Bar	<i>Mean of Subgroup Medians</i>
-------------	---------------------------------

---

**Description**

Calculates the mean of subgroup medians used when constructing a xMedian-R charts.

**Usage**

```
xMedian_Bar(data, value, grouping, formula = NULL, ...)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables
...	further arguments passed to or from other methods.

**Value**

A number; mean of subgroup medians.

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
xMedian_Bar(data = df, formula = v~g)
```

---

xMedian_rBar_LCL	<i>Mean of Subgroup Medians Lower Control Limit (LCL) based on Mean Range</i>
------------------	---

---

**Description**

Calculates the mean of subgroup medians lower control limit based on the mean range. The result is used when constructing a xMedian-R charts.

**Usage**

```
xMedian_rBar_LCL(data, value, grouping, n = NULL, natural = F,
  formula = NULL)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
n	a number indicating a hypothetical subgroup size other than, function determined subgroup n determined by the floor length of subgroup values.
natural	logical, if TRUE calculate limits for individuals (n=1) else calculate for n determined by the floor length of subgroup values
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables

**Value**

A number; mean of subgroup medians Lower Control Limit (LCL) based on mean range

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
xBar_rMedian_LCL(data = df, formula = v~g)
```

---

xMedian_rBar_UCL	<i>Mean of Subgroup Medians Upper Control Limit (UCL) based on mean Range</i>
------------------	---

---

**Description**

Calculates the mean of subgroup medians upper control limit based on the mean subgroup range. The result is used when constructing a xMedian-R charts.

**Usage**

```
xMedian_rBar_UCL(data, value, grouping, n = NULL, natural = F,
  formula = NULL)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
n	a number indicating a hypothetical subgroup size other than, function determined subgroup n determined by the floor length of subgroup values.
natural	logical, if TRUE calculate limits for individuals (n=1) else calculate for n determined by the floor length of subgroup values
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables

**Value**

A number; mean of subgroup means Upper Control Limit (UCL) based on Median Range

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
xMedian_rBar_UCL(data = df, formula = v~g)
```

---

xMedian_rMedian_LCL	<i>Mean of Subgroup Medians Lower Control Limit (LCL) based on Median Range</i>
---------------------	---

---

**Description**

Calculates the mean of subgroup medians lower control limit based on the median subgroup range. The result is used when constructing a xMedian-rMedian charts.

**Usage**

```
xMedian_rMedian_LCL(data, value, grouping, n = NULL, natural = F,
  formula = NULL)
```

**Arguments**

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
n	a number indicating a hypothetical subgroup size other than, function determined subgroup n determined by the floor length of subgroup values.
natural	logical, if TRUE calculate limits for individuals (n=1) else calculate for n determined by the floor length of subgroup values
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables

**Value**

A number; mean of subgroup median Lower Control Limit (LCL) based on Median Range

**Examples**

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
xMedian_rMedian_LCL(data = df, formula = v~g)
```

---

xMedian\_rMedian\_UCL    *Mean of Subgroup Medians Upper Control Limit (UCL) based on Median Range*

---

### Description

Calculates the mean of subgroup medians upper control limit based on the median subgroup range. The result is used when constructing a xMedian-rMedian charts.

### Usage

```
xMedian_rMedian_UCL(data, value, grouping, n = NULL, natural = F,
  formula = NULL)
```

### Arguments

data	data frame to be processed
value	numeric vector in a data frame with values of interest.
grouping	single factor/variable to split the data frame "values" by.
n	a number indicating a hypothetical subgroup size other than, function determined subgroup n determined by the floor length of subgroup values.
natural	logical, if TRUE calculate limits for individuals (n=1) else calculate for n determined by the floor length of subgroup values
formula	a formula, such as $y \sim x_1 + x_2$ , where the y variable is numeric data to be split into groups according to the grouping x factors/variables

### Value

A number; mean of subgroup median upper Control Limit (UCL) based on Median Range

### Examples

```
set.seed(5555)
df <- data.frame(v=rnorm(60, 0, 1), g=rep(c("A","B","C","D","E"), each=12))
xMedian_rMedian_UCL(data = df, formula = v~g)
```

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