

# Package ‘ngsReports’

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**Title** Load FastqQC reports and other NGS related files

**Description** This package provides methods and object classes for parsing FastQC reports and output summaries from other NGS tools into R. As well as parsing files, multiple plotting methods have been implemented for visualising the parsed data. Plots can be generated as static ggplot objects or interactive plotly objects.

**URL** <https://github.com/smped/ngsReports>

**BugReports** <https://github.com/smped/ngsReports/issues>

**License** LGPL-3

**Encoding** UTF-8

**Depends** R (>= 4.2.0), BiocGenerics, ggplot2 (>= 3.5.0), patchwork (>= 1.1.1), tibble (>= 1.3.1)

**Imports** Biostrings, checkmate, dplyr (>= 1.1.0), forcats, ggdendro, grDevices (>= 3.6.0), grid, jsonlite, lifecycle, lubridate, methods, plotly (>= 4.9.4), reshape2, rlang, rmarkdown, scales, stats, stringr, tidyr, tidyselect (>= 0.2.3), utils, zoo

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**Collate** 'AllGenerics.R' 'validationFunctions.R' 'FastpData.R' 'FastpDataList.R' 'FastpFile.R' 'FastqcData.R' 'FastqcDataList.R' 'FastqcFile.R' 'PwfCols.R' 'S4coercion.R' 'TheoreticalGC.R' 'aaa.R' 'data.R' 'errMsg.R' 'estGcDistn.R' 'extract.R' 'fqName.R' 'fqVersion.R' 'getColours.R' 'getGC.R' 'getModule.R' 'getSummary.R' 'helpers.R' 'importNgsLogs.R' 'importSJ.R' 'isCompressed.R' 'maxAdapterContent.R' 'ngsReports-package.R' 'overRep2Fasta.R' 'path.R' 'plotAdapterContent.R' 'plotAlignmentSummary.R' 'plotAssemblyStats.R' 'plotBaseQuals.R' 'plotDupLevels.R' 'plotFastqcPCA.R' 'plotGcContent.R' 'plotInsertSize.R'

'plotKmers.R' 'plotNContent.R' 'plotOverrep.R'  
 'plotReadTotals.R' 'plotSeqContent.R' 'plotSeqLengthDistn.R'  
 'plotSeqQuals.R' 'plotSummary.R' 'pwf.R' 'readTotals.R'  
 'summariseOverrep.R' 'writeHtmlReport.R'

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---

<code>.addPercent</code>	<i>Add a percentage sign</i>
--------------------------	------------------------------

---

### Description

Add a percentage sign to the end of a string

### Usage

```
.addPercent(x)
```

### Arguments

x	Any vector
---	------------

### Value

character vector

### Examples

```
x <- 1:10
ngsReports:::.addPercent(x)
```

---

<code>.emptyPlot</code>	<i>Create an empty plot with supplied text</i>
-------------------------	--

---

### Description

Create an empty plot with supplied text

### Usage

```
.emptyPlot(x)
```

### Details

Create plot using `theme_void` and only with the supplied text

**Value**

A ggplot2 object

**Examples**

```
ngsReports:::.emptyPlot("This is an empty plot")
```

---

*.FastpFile-class*      *The .FastpFile Object Class*

---

**Description**

The *.FastpFile* Object Class defines a path to the output from the standalone tool fastp. **[Experimental]**

**Arguments**

x                      character(1) denoting a file.path

**Details**

This class simply refers to a fastp output file after checking for existence and validity (i.e. the correct internal structure). Underlying files are expected to be in json format

The helper function *.FastpFile()* is a simple constructor which checks validity and enables construction of other dependent classes.

**Value**

An object of class *.FastqcFile*

**Slots**

path Character vector of length 1 which contains a valid file path.

---

*.FastqcFile-class*      *The .FastqcFile Object Class*

---

### Description

The *.FastqcFile* Object Class defines a path to the output from the standalone tool FastQC. **[Stable]**

### Arguments

x                      character(1) denoting a file.path

### Details

This class simply refers to a fastqc output file after checking for existence and validity (i.e. the correct internal structure). Underlying files can be zipped (*\*\_fastqc.zip*) or extracted directories

The helper function *.FastqcFile()* is a simple constructor which checks validity and enables construction of other dependent classes.

### Value

An object of class *.FastqcFile*

### Slots

path Character vector of length 1 which contains a valid file path.

### Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)[1]

# As this is the root structure, we can only call this
# function with an individual file
ff <- ngsReports:::.FastqcFile(fl)
```

---

*.getToolName*                      *Identify tool name*

---

### Description

Identify tool name for log files after reading in using *readLines*.

### Usage

```
.getToolName(x, possTypes)
```

**Arguments**

x                      Character vector as output when readLines to a supplied log file

**Details**

Checks for all the required fields in the lines provided

**Value**

logical(1)

---

.hidePWFRects	<i>Hide PWF tooltips from line plots</i>
---------------	--

---

**Description**

Hide tooltips from PWF rectangles in line plots

**Usage**

.hidePWFRects(x)

**Arguments**

x                      plotlyObject\$x\$data

**Value**

plotlyObject\$x\$data

---

.makeDendro	<i>Make the dendrogram for heatmap-style plots</i>
-------------	--

---

**Description**

Set the clusters for heatmap-style interactive plots

**Usage**

.makeDendro(df, rowVal, colVal, value)

**Arguments**

df                      The data frame to be clustered  
rowVal                  The rows to be clustered  
colVal                  The value which will become column names  
value                    The value to use for the clustering

**Value**

A dendrogram

**Examples**

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fileList <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)
cols <- c("Filename", "Position", "Illumina_Universal_Adapter")
ac <- getModule(fileList, "Adapter_Content")[cols]
ngsReports:::makeDendro(df = ac,
                        rowVal = "Filename",
                        colVal = "Position",
                        value = "Illumina_Universal_Adapter")
```

---

.makeLabels

*Perform the checks and return the labels*

---

**Description**

Checks for the presence of labels and returns defaults

**Usage**

```
.makeLabels(
  x,
  labels,
  pattern = ".(fast|fq|bam|sam|cram).*",
  col = "Filename",
  ...
)
```

**Arguments**

x	A data.frame with a column titled "Filename"
labels	Named vector of labels for plotting
pattern	character Regular expression to remove from filenames
col	character Column to use for generating labels
...	Not used

**Details**

Takes a named vector of labels and checks for the correct fields. If no vector is supplied, returns the file names missing the specified pattern, which defaults to removing the suffixes fastq(.gz), fq(.gz), bam, sam or cram.



**Value**

Named character vector

**Examples**

```
f <- paste0(c("File1", "File2"), ".fastq")
df <- data.frame(FileName = f, stringsAsFactors = FALSE)
ngsReports:::makeLabels(df)
```

---

<code>.makePwfGradient</code>	<i>Construct a gradient using PwfCols</i>
-------------------------------	---

---

**Description**

Construct a gradient using PwfCols

**Usage**

```
.makePwfGradient(  
  vals,  
  pwfCols,  
  breaks = c(0, 5, 10, 100),  
  passLow = TRUE,  
  na.value = "white"  
)
```

**Arguments**

<code>vals</code>	The values which need to have the scale generated
<code>pwfCols</code>	An object of class PwfCols
<code>breaks</code>	The breaks for the PWF bins
<code>passLow</code>	Is the PASS category at the low or high end of the numeric range
<code>na.value</code>	The colour to plot for missing values

**Details**

This constructs a list of arguments for passing to `scale_fill_gradientn()` using the values contained in the fill aesthetic and the supplied breaks for PASS/WARN/FAIL criteria.

**Value**

Returns a ggplot list

---

<code>.makeSidebar</code>	<i>Shortcut for making the status sidebar</i>
---------------------------	---

---

**Description**

Shortcut for making the status sidebar

**Usage**

```
.makeSidebar(status, key, pwfCols, usePlotly = TRUE)
```

**Arguments**

<code>status</code>	A data.frame with columns 'Filename' & 'Status'
<code>key</code>	A vector of values corresponding to the Filename column
<code>pwfCols</code>	An object of class PwfCols
<code>usePlotly</code>	If TRUE, output is a plotly panel

**Value**

if `usePlotly = TRUE`, a plotly object. The sidebar for an interactive plot showing PASS/WARN/FAIL status for each file. If `usePlotly = FALSE` the underlying ggplot object will be returned.

---

<code>.parseAdapterRemovalLogs</code>	<i>Parse data from Adapter Removal log files</i>
---------------------------------------	--

---

**Description**

Parse data from Adapter Removal log files

**Usage**

```
.parseAdapterRemovalLogs(data, which = 3)
```

**Arguments**

<code>data</code>	List of lines read using <code>readLines</code> on one or more files
<code>which</code>	which element of the log file to return. Can be 1:4, "sequences", "settings", "statistics" or "distribution"

**Details**

Checks for structure will have been performed

**Value**

tibble

---

`.parseBowtieLogs`      *Parse data from Bowtie log files*

---

**Description**

Parse data from Bowtie log files

**Usage**

```
.parseBowtieLogs(data, ...)
```

**Arguments**

<code>data</code>	List of lines read using <code>readLines</code> on one or more files
<code>...</code>	Not used

**Details**

Checks for structure will have been performed

**Value**

data.frame

---

`.parseBuscoLogs`      *Parse data from BUSCO log files*

---

**Description**

Parse data from BUSCO log files

**Usage**

```
.parseBuscoLogs(data, ...)
```

**Arguments**

<code>data</code>	List of lines read using <code>readLines</code> on one or more files
<code>...</code>	Not used

**Details**

Checks for structure will have been performed

**Value**

data.frame

---

*.parseCutadaptLogs*      *Parse data from cutadapt log files*


---

**Description**

Parse data from cutadapt log files

**Usage**`.parseCutadaptLogs(data, which = 1)`**Arguments**

<code>data</code>	List of lines read using <code>readLines</code> on one or more files
<code>which</code>	which element of the log file to return. Can be <code>summary</code> , <code>adapter1</code> , <code>adapter2</code> , <code>adapter3</code> or <code>overview</code> , or any integer in 1:5

**Details**

Checks for structure will have been performed

**Value**

tibble

---

*.parseDuplicationMetricsLogs*  
*Parse data from Picard duplicationMetrics log files*


---

**Description**

Parse data from Picard duplicationMetrics log files

**Usage**`.parseDuplicationMetricsLogs(data, which = 1)`**Arguments**

<code>data</code>	List of lines read using <code>readLines</code> on one or more files
<code>which</code>	which element of the log file to return. Can be 1:2, "metrics" or "histogram"

**Details**

Checks for structure will have been performed

**Value**

tibble

---

`.parseFeatureCountsLogs` *Parse data from featureCounts summary files*

---

**Description**

Parse data from featureCounts summary files

**Usage**

`.parseFeatureCountsLogs(data, ...)`

**Arguments**

<code>data</code>	List of lines read using <code>readLines</code> on one or more files
<code>...</code>	Not used

**Details**

Checks for structure will have been performed

**Value**

tibble

---

`.parseFlagstatLogs` *Parse data from samtools flagstat files*

---

**Description**

Parse data from samtools flagstat files

**Usage**

`.parseFlagstatLogs(data, ...)`

**Arguments**

<code>data</code>	List of lines read using <code>readLines</code> on one or more files
<code>...</code>	Not used

**Details**

Checks for structure will have been performed

**Value**

`data.frame`

---

<code>.parseHisat2Logs</code>	<i>Parse data from HISAT2 log files</i>
-------------------------------	---

---

**Description**

Parse data from HISAT2 log files

**Usage**

```
.parseHisat2Logs(data, ...)
```

**Arguments**

<code>data</code>	List of lines read using <code>readLines</code> on one or more files
<code>...</code>	Not used

**Details**

Checks for structure will have been performed

**Value**

`data.frame`

---

`.parseMacs2CallpeakLogs`

*Parse data from macs2 callpeak log files*

---

**Description**

Parse data from macs2 callpeak log files

**Usage**

`.parseMacs2CallpeakLogs(data, ...)`

**Arguments**

<code>data</code>	List of lines read using <code>readLines</code> on one or more files
<code>...</code>	Not used

**Details**

Checks for structure will have been performed

**Value**

`data.frame`

---

`.parseQuastLogs`

*Parse data from BUSCO log files*

---

**Description**

Parse data from BUSCO log files

**Usage**

`.parseQuastLogs(data, ...)`

**Arguments**

<code>data</code>	List of lines read using <code>readLines</code> on one or more files
<code>...</code>	Not used

**Details**

Checks for structure will have been performed

**Value**

`data.frame`

---

`.parseRnaseqcMetricsLogs`*Parse data from rnaseqc metrics files*

---

**Description**

Parse data from rnaseqc metrics files

**Usage**

```
.parseRnaseqcMetricsLogs(data, ...)
```

**Arguments**

<code>data</code>	List of lines read using <code>readLines</code> on one or more files
<code>...</code>	Not used

**Details**

Checks for structure will have been performed

**Value**

`data.frame`

---

`.parseSalmonLibFormatCountsLogs`*Parse data from salmon lib\_format\_counts.json files*

---

**Description**

Parse data from salmon lib\_format\_counts.json files

**Usage**

```
.parseSalmonLibFormatCountsLogs(data, ...)
```

**Arguments**

<code>data</code>	List of lines read using <code>readLines</code> on one or more files
<code>...</code>	Not used

**Details**

Checks for structure will have been performed



**Value**

data.frame

---

`.parseSalmonMetaInfoLogs`

*Parse data from salmon meta\_info.json files*

---

**Description**

Parse data from salmon meta\_info.json files

**Usage**

`.parseSalmonMetaInfoLogs(data, ...)`

**Arguments**

<code>data</code>	List of lines read using readLines on one or more files
<code>...</code>	Not used

**Details**

Checks for structure will have been performed

**Value**

data.frame

---

`.parseStarLogs`

*Parse data from STAR log files*

---

**Description**

Parse data from STAR log files

**Usage**

`.parseStarLogs(data, ...)`

**Arguments**

<code>data</code>	List of lines read using readLines on one or more files
<code>...</code>	Not used

**Details**

Checks for structure will have been performed

**Value**

tibble

---

*.parseTrimmomaticLogs* *Parse data from trimmomatic log files*

---

**Description**

Parse data from trimmomatic log files

**Usage**

```
.parseTrimmomaticLogs(data, ...)
```

**Arguments**

<code>data</code>	List of lines read using <code>readLines</code> on one or more files
<code>...</code>	not used

**Details**

Checks for structure will have been performed

**Value**

tibble

---

*.parseUmitoolsDedupLogs*  
*Parse data from umi\_tools dedup files*

---

**Description**

Parse data from umi\_tools dedup files

**Usage**

```
.parseUmitoolsDedupLogs(data, ...)
```

**Arguments**

data            List of lines read using readLines on one or more files  
...            Not used

**Details**

Checks for structure will have been performed

**Value**

data.frame

---

.prepHeatmap	<i>Prepare the final heatmap for plotting</i>
--------------	---

---

**Description**

Add dendrogram & status bar to ggplot2 heatmap

**Usage**

```
.prepHeatmap(x, status, segments, usePlotly, heat_w = 8, pwf, hv = NULL)
```

**Arguments**

x                a ggplot2 heatmap produced by ngsReports  
status           a tibble with the columns Filename and Status  
segments        a dendrogram produced during clustering of samples  
usePlotly       logical(1)  
hv               character vector of fields to include in hoverinfo

**Value**

Either a ggplot2 object assembled using patchwork, or an interactive plotly object

---

<code>.renderDendro</code>	<i>Set up dendrograms for interactive plots</i>
----------------------------	---

---

### Description

A commonly used (hidden) function for setting up dendrograms for interactive plots. based on code found at <https://plot.ly/ggplot2/ggdendro-dendrograms/>

### Usage

```
.renderDendro(df)
```

### Arguments

<code>df</code>	A data.frame as required
-----------------	--------------------------

### Details

Create plot using `theme_dendro`

### Value

A plotly object

---

<code>.splitByTab</code>	<i>Split elements of a vector into a data.frame</i>
--------------------------	---

---

### Description

Split elements of a character vector by the tab separator

### Usage

```
.splitByTab(x, firstRowToNames = TRUE, tab = "\\t")
```

### Arguments

<code>x</code>	A character vector
<code>firstRowToNames</code>	logical Should the first element be used for column names
<code>tab</code>	character The string used to represent the tab symbol

### Details

This will split a vector into a data.frame checking that every line has the same number of separators. By default the first element will be set as the column names.

This is designed to take input from `readLines()`

**Value**

A data frame

**Examples**

```
x <- c("ColA\tColB", "Value1\tValue2")
ngsReports:::splitByTab(x, firstRowToNames = TRUE)
ngsReports:::splitByTab(x, firstRowToNames = FALSE)
```

---

.updateThemeFromDots *Add custom theme elements from dotArgs*

---

**Description**

Add custom theme elements from dotArgs

**Usage**

```
.updateThemeFromDots(p, ...)
```

**Arguments**

p	ggplot object
...	Standard dot arguments

**Value**

ggplot2 object

---

estGcDistn *Estimate a GC Content Distribution From Sequences*

---

**Description**

Generate a GC content distribution from sequences for a given read length and fragment length

**Usage**

```
estGcDistn(x, n = 1e+06, r1 = 100, fl = 200, fragSd = 30, bins = 101, ...)

## S4 method for signature 'ANY'
estGcDistn(x, n = 1e+06, r1 = 100, fl = 200, fragSd = 30, bins = 101, ...)

## S4 method for signature 'character'
estGcDistn(x, n = 1e+06, r1 = 100, fl = 200, fragSd = 30, bins = 101, ...)

## S4 method for signature 'DNAStrngSet'
estGcDistn(x, n = 1e+06, r1 = 100, fl = 200, fragSd = 30, bins = 101, ...)
```

**Arguments**

x	DNAStr ingSet or path to a fasta file
n	The number of reads to sample
r1	Read Lengths to sample
f1	The mean of the fragment lengths sequenced
fragSd	The standard deviation of the fragment lengths being sequenced
bins	The number of bins to estimate
...	Not used

**Details**

The function takes the supplied object and returns the theoretical GC content distribution. Using a fixed read length essentially leads to a discrete distribution so the bins argument is used to define the number of bins returned. This defaults to 101 for 0 to 100% inclusive.

The returned values are obtained by interpolating the values obtained during sampling. This avoids returned distributions with gaps and jumps as would be obtained setting readLengths at values not in multiples of 100.

Based heavily on <https://github.com/mikelove/fastqcTheoreticalGC>

**Value**

A tibble with two columns: GC\_Content and Freq denoting the proportion of GC and frequency of occurrence respectively

**Examples**

```
faDir <- system.file("extdata", package = "ngsReports")
faFile <- list.files(faDir, pattern = "fasta", full.names = TRUE)
df <- estGcDistn(faFile, n = 200)
```

---

FastpData-class

*The FastpData Object Class*


---

**Description**

The FastpData Object Class **[Experimental]**

**Usage**

```
FastpData(x)
```

**Arguments**

x	Path to a single zip archive or extracted folder for a individual fastp report.
---	---

**Details**

This object class is the main object required for generating plots and tables. Instantiation will first check for a .json file with the correct data structure, and will then parse all the data into R as a FastpData object. Fastp modules are contained as individual slots, which can be viewed using slotNames. Sub-modules are also contained within many larger modules with modules being based on the sections within a fastp html report

Individual modules can be returned using the function `getModule()` and specifying which module/sub-module is required. See `getModule()` for more details.

**Value**

An object of class FastpData

**Slots**

**Summary** Contains three submodules 1) Before\_filtering, 2) After\_filtering and 3) Filtering\_result.

All values presented in the initial table for individual fastp reports are contained in other sections of the report

**Adapters** Contains a tibble with all data from this module

**Duplication** Contains a tibble with all duplication results

**Insert\_size** Contains a tibble with all insert size estimates

**Before\_filtering, After\_filtering** The modules can be selected for either Read1 or Read2 paired logical(1) indicating whether the file is from paired-end sequencing

**command** character(1) with the executed command

**version** character(1) with the fastp version being used

**path** Path to the Fastp report

---

FastpDataList-class    *The FastpDataList Object Class*

---

**Description**

The FastpDataList Object Class [**Stable**]

**Usage**

```
FastpDataList(x)
```

**Arguments**

x                      Character vector of file paths specifying paths to fastp.json.gz output

**Value**

An object of class FastpDataList

**Slots**

... this can either be a single character vector of paths to fastp files, or several instances of .Fastp-File objects

---

FastqcData-class      *The FastqcData Object Class*

---

**Description**

The FastqcData Object Class [**Stable**]

**Usage**

```
FastqcData(x)
```

**Arguments**

x                      Path to a single zip archive or extracted folder for a individual FastQC report.

**Details**

This object class is the main object required for generating plots and tables. Instantiation will first test for a compressed file (or extracted directory) with the correct data structure, and will then parse all the data into R as a FastqcData object. FastQC modules are contained as individual slots, which can be viewed using `slotNames`.

Individual modules can be returned using the function `getModule()` and specifying which module is required. See [getModule\(\)](#) for more details.

**Value**

An object of class FastqcData

**Slots**

Summary      Summary of PASS/WARN/FAIL status for each module

Basic\_Statistics      The Basic\_Statistics table from the top of a FastQC html report

Per\_base\_sequence\_quality      The underlying data from the Per\_base\_sequence\_quality module

Per\_sequence\_quality\_scores      The underlying data from the Per\_sequence\_quality\_scores module

Per\_base\_sequence\_content      The underlying data from the Per\_base\_sequence\_content module

Per\_sequence\_GC\_content      The underlying data from the Per\_sequence\_GC\_content module

Per\_base\_N\_content      The underlying data from the Per\_base\_N\_content module

Sequence\_Length\_Distribution      The underlying data from the Sequence\_Length\_Distribution module



Sequence\_Duplication\_Levels The underlying data from the Sequence\_Duplication\_Levels module

Overrepresented\_sequences The underlying data from the Overrepresented\_sequences module

Adapter\_Content The underlying data from the Adapter\_Content module

Kmer\_Content The underlying data from the Kmer\_Content module

Total\_Deduplicated\_Percentage Estimate taken from the plot data for Sequence\_Duplication\_Levels. Only included in later versions of FastQC

version The version of FastQC used for generation of the report (if available)

path Path to the FastQC report

### Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
f1 <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)[1]

# Load the FASTQC data as a FastqcData object
fd <- FastqcData(f1)
fd
```

---

FastqcDataList-class *The FastqcDataList Object Class*

---

### Description

The FastqcDataList Object Class **[Stable]**

### Usage

```
FastqcDataList(x)
```

### Arguments

x Character vector of file paths specifying paths to FastQC reports

### Value

An object of class FastqcDataList

### Slots

... this can either be a single character vector of paths to FASTQC files, or several instances of .FastqcFile objects

## Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)
fdl
```

---

fqcVersion, FastqcData-method  
*Get the FASTQC version*

---

## Description

Get the FASTQC version used to generate the initial files

## Usage

```
## S4 method for signature 'FastqcData'
fqcVersion(object)

## S4 method for signature 'FastqcDataList'
fqcVersion(object)

## S4 method for signature 'ANY'
fqcVersion(object)
```

## Arguments

object            An object of class FastqcData or FastqcDataList

## Value

A character vector (FastqcData), or tibble (FastqcDataList)

## Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)

# Get the FASTQC version
fqcVersion(fdl)
```

---

fqName	<i>Return the Underlying Fastq File Names from Fastqc/Fastp Objects</i>
--------	---

---

## Description

Return the Underlying Fastq File Names from Fastqc/Fastp Objects

## Usage

```
fqName(object)

## S4 method for signature 'ANY'
fqName(object)

## S4 method for signature 'FastqcData'
fqName(object)

## S4 method for signature 'FastqcDataList'
fqName(object)

fqName(object) <- value

## S4 replacement method for signature 'FastqcData'
fqName(object) <- value

## S4 replacement method for signature 'FastqcDataList'
fqName(object) <- value

## S4 method for signature 'FastpData'
fqName(object)

## S4 method for signature 'FastpDataList'
fqName(object)
```

## Arguments

object	An object able to extract an Fastq name from
value	Replacement value for fqName

## Value

Returns the names of the Fastq files the FastQC report was generated from, without any preceding directories.

## Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)
fqName(fdl)

nm <- paste0(letters[seq_along(fdl)], ".fq")
fqName(fdl) <- nm
fqName(fdl)
```

---

gcAvail

*List Genomes or Transcriptomes with Theoretical GC Content*

---

## Description

List available genomes or transcriptomes in a TheoreticalGC object

## Usage

```
gcAvail(object, type)

## S4 method for signature 'TheoreticalGC'
gcAvail(object, type)
```

## Arguments

object	An object of class TheoreticalGC
type	character indicating either Genome or Transcriptome

## Details

An object of class TheoreticalGC can hold the theoretical GC content for one or more species, for either the genome or transcriptome. This function checks which species are available in the given object, for either the genome or transcriptome, as supplied to the parameter type.

## Value

A tibble object

## Examples

```
gcAvail(gcTheoretical, "Genome")
```

---

gcTheoretical	<i>Theoretical GC content</i>
---------------	-------------------------------

---

## Description

This object contains the theoretical GC content for each provided species, for both the genome and transcriptome, where available.

## Usage

```
gcTheoretical
```

## Format

An object of class `TheoreticalGC` of length 1.

## Details

The object is defined with the S4 class `TheoreticalGC`. Species for which information is available can be found using the command `gcAvail(gcTheoretical)` and selecting the appropriate type.

Metadata is accessible using `mData(gcTheoretical)`.

All GC content was calculated using code from <https://github.com/mikelove/fastqcTheoreticalGC> using `BSgenome` packages. This provides a default set of GC content data for common organisms generated using 100bp reads/fragments and 1e6 reads.

## See Also

```
gcAvail
```

## Examples

```
## Check which genomes are included
gcAvail(gcTheoretical, "Genome")

## Check which transcriptomes are included
gcAvail(gcTheoretical, "Transcriptome")
```

---

`getColours`*Work with objects of class PwfCols*

---

**Description**

Get and modify colours from objects of class PwfCols

**Usage**

```
## S4 method for signature 'PwfCols'
getColours(object)

## S4 method for signature 'PwfCols'
setColours(object, PASS, WARN, FAIL, MAX)

## S4 method for signature 'PwfCols'
setAlpha(object, alpha)
```

**Arguments**

<code>object</code>	An object of class PwfCols
<code>PASS</code>	The colour denoting PASS on all plots, in rgb format
<code>WARN</code>	The colour denoting WARN on all plots, in rgb format
<code>FAIL</code>	The colour denoting FAIL on all plots, in rgb format
<code>MAX</code>	The colour denoting the limit of values in rgb format
<code>alpha</code>	Numeric(1). Ranges from 0 to 1 by default, but can also be on the range 0 to 255.

**Details**

Use `getColours` to obtain the colours in an object of class PwfCols.  
These can be modified using the functions `setColours` and `setAlpha`

**Value**

`getColours` will return a character vector of colours corresponding to PASS/WARN/FAIL  
`setColours` will return an object of class PwfCols  
`setAlpha` will return an object of class PwfCols

**Examples**

```
getColours(pwf)

# How to add transparency
pwf2 <- setAlpha(pwf, 0.1)
getColours(pwf2)
```

---

getGC	<i>Get Theoretical GC content</i>
-------	-----------------------------------

---

**Description**

Get the GC content data from a TheoreticalGC object

**Usage**

```
getGC(object, name, type)

## S4 method for signature 'ANY'
getGC(object, type)

## S4 method for signature 'TheoreticalGC'
getGC(object, name, type)
```

**Arguments**

object	An object of class Theoretical GC
name	The Name of the species in 'Gspecies' format, e.g. Hsapiens
type	The type of GC content. Can only be either "Genome" or "Transcriptome"

**Value**

A tibble object

**Examples**

```
getGC(gcTheoretical, name = "Hsapiens", type = "Genome")
```

---

getModule, FastqcData-method	<i>Retrieve a given module from a Fastqc* Object</i>
------------------------------	--

---

**Description**

Retrieve a specific module from a Fastqc\* object as a data.frame

**Usage**

```
## S4 method for signature 'FastqcData'
getModule(object, module)

## S4 method for signature 'FastqcDataList'
getModule(object, module)

## S4 method for signature 'ANY'
getModule(object, module)

## S4 method for signature 'FastpData'
getModule(object, module)

## S4 method for signature 'FastpDataList'
getModule(object, module)
```

**Arguments**

object	Can be a <code>FastqcData</code> , <code>fastqcDataList</code> , or simply a character vector of paths
module	The requested module as contained in a FastQC report. Possible values are <code>Summary</code> , <code>Basic_Statistics</code> , <code>Per_base_sequence_quality</code> , <code>Per_tile_sequence_quality</code> , <code>Per_sequence_quality_scores</code> , <code>Per_base_sequence_content</code> , <code>Per_sequence_GC_content</code> , <code>Per_base_N_content</code> , <code>Sequence_Length_Distribution</code> , <code>Sequence_Duplication_Levels</code> , <code>Overrepresented_sequences</code> , <code>Adapter_Content</code> , <code>Kmer_Content</code> , <code>Total_Deduplicated_Percentage</code> . Note that spelling and capitalisation is exactly as contained within a FastQC report, with the exception that spaces have been converted to underscores. Partial matching is implemented for this argument.

**Details**

This function will return a given module from a `Fastqc*` object as a `data.frame`. Note that each module will be it's own unique structure, although all will return a `data.frame`

**Value**

A single tibble containing module-level information from all FastQC reports contained in the `Fastqc*` object.

**Examples**

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)

# Extract the Summary module, which corresponds to the PASS/WARN/FAIL flags
getModule(fdl, "Summary")
```



```
# The Basic_Statistics module corresponds to the table at the top of each
# FastQC report
getModule(fdl, "Basic_Statistics")
```

---

```
getSummary,.FastqcFile-method
```

*Get the summary information from Fastqc Files*

---

## Description

Read the information from the summary.txt files in each .FastqcFile

## Usage

```
## S4 method for signature '.FastqcFile'
getSummary(object)

## S4 method for signature 'ANY'
getSummary(object)

## S4 method for signature 'FastqcData'
getSummary(object)

## S4 method for signature 'FastqcDataList'
getSummary(object)
```

## Arguments

object	Can be a FastqcData, FastqcDataList object or a vector of paths to unparsed FastQC reports.
--------	---

## Details

This simply extracts the summary of PASS/WARN/FAIL status for every module as defined by the tool FastQC for each supplied file.

## Value

A tibble containing the PASS/WARN/FAIL status for each module, as defined in a FastQC report.

## Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)
```

```
# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(f1)

# Return a tibble/tibble with the raw information
getSummary(fdl)
```

---

importNgsLogs

*Import Various NGS-related log files*


---

## Description

**[Maturing]** Imports NGS-related log files such as those generated from stderr.

## Usage

```
importNgsLogs(x, type = "auto", which, stripPaths = TRUE)
```

## Arguments

x	character. Vector of filenames. All log files must be of the same type. Duplicate file paths will be silently ignored.
type	character. The type of file being imported. Can be one of bowtie, bowtie2, hisat2, star, flagstat, featureCounts, duplicationMetrics, cutadapt, umitoolsDedup, macs2Callpeak, adapterRemoval, rnaseqMetrics, quast, salmonLibFormatCounts, salmonMetaInfo or busco. Defaults to type = "auto" which will automatically detect the file type for all implemented types.
which	Which element of the parsed object to return. Ignored in all file types except when type is set to duplicationMetrics, cutadapt or adapterRemoval. See details for possible values. To return all elements, set this value to 'all'
stripPaths	logical(1). Remove paths from the Filename column

## Details

Imports one or more log files as output by tools such as: bowtie, bowtie2, featureCounts, Hisat2, STAR, salmon picard MarkDuplicates, cutadapt, flagstat, macs2Callpeak, Adapter Removal, trimmomatic, rnaseqMetrics, quast or busco. autoDetect can be used to detect the log type by parsing the file.

The featureCounts log file corresponds to the counts.out.summary, not the main counts.out file.

Whilst most log files return a single tibble, some are more complex with multiple modules.

adapterRemoval can return one of four modules (which = 1:4). When calling by name, the possible values are sequences, settings, statistics or distribution. Partial matching is implemented.

cutadapt can return one of five modules (which = 1:5). When calling by name the possible modules are summary, adapter1, adapter2, adapter3 or overview. Note that adapter2/3 may be missing from these files depending on the nature of your data. If cutadapt log files are obtained using report=minimal, all supplied log files must be of this format and no modules can be returned.

duplicationMetrics will return either the metrics of histogram. These can be requested by setting which as 1 or 2, or naming either module.

### Value

A tibble. Column names are broadly similar to the text in supplied files, but have been modified for easier handling under R naming conventions.

### Examples

```
f <- c("bowtiePE.txt", "bowtieSE.txt")
bowtieLogs <- system.file("extdata", f, package = "ngsReports")
df <- importNgsLogs(bowtieLogs, type = "bowtie")
```

---

importSJ

*Import STAR Splice Junctions*

---

### Description

Import the SJ.out.tab files produced by STAR

### Usage

```
importSJ(x, stripPaths = TRUE)
```

### Arguments

x                    vector of file paths to SJ.out.tab files  
stripPaths        logical(1) Remove directory prefixes from the file paths in x

### Details

Imports one or more splice-junction output files as produced by STAR. If all are located in separated directories with identical names, be sure to set the argument stripPaths = FALSE

All co-ordinates are 1-based, in keeping with the STAR manual

### Value

A tibble

### Author(s)

Stephen Pederson [stephen.pederson@adelaide.edu.au](mailto:stephen.pederson@adelaide.edu.au)

**Examples**

```
sjFiles <- system.file("extdata", "SJ.out.tab", package = "ngsReports")
# Import leaving the complete file path in the column Filename
# The argument stripPaths is set as TRUE by default
df <- importSJ(sjFiles, stripPaths = FALSE)
```

---

isCompressed	<i>Check to see if a file is compressed</i>
--------------	---

---

**Description**

Check to see if a file, or vector of files is compressed

**Usage**

```
isCompressed(path, type = c("zip", "gzip"), verbose = FALSE)
```

**Arguments**

path	The path to one or more files
type	The type of compression to check for. Currently only ZIP/GZIP files have been implemented.
verbose	logical/integer Determine the level of output to show as messages

**Details**

Reads the first four bytes from the local file header. If the file is a .ZIP file, this should match the magic number PK\003\004.

This function assumes that the first thing in a zip archive is the .ZIP entry with the local file header signature. ZIP files containing a self-extracting archive may not exhibit this structure and will return FALSE

**Value**

A logical vector

**Examples**

```
# Get the files included with the package
fileDir <- system.file("extdata", package = "ngsReports")
allFiles <- list.files(fileDir, pattern = "zip$", full.names = TRUE)
isCompressed(allFiles)
```

---

maxAdapterContent	<i>Get the maximum Adapter Content</i>
-------------------	--

---

## Description

Get the maximum Adapter Content across one or more FASTQC reports

## Usage

```
maxAdapterContent(x, asPercent = TRUE)
```

## Arguments

x	Can be a <code>.FastqcFile</code> , <code>FastqcData</code> , <code>FastqcDataList</code> or path
asPercent	logical. Format the values as percentages with the added <code>%</code> symbol

## Details

This will extract the `Adapter_Content` module from the supplied object, and provide a tibble with the final value for each file.

## Value

A tibble object containing the percent of reads with each adapter type at the final position

## Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)

# Get the maxAdapterContent
maxAdapterContent(fdl)
```

---

mData	<i>Extract Metadata for TheoreticalGC objects</i>
-------	---

---

**Description**

Extract Metadata for TheoreticalGC objects

**Usage**

```
mData(object)

## S4 method for signature 'TheoreticalGC'
mData(object)
```

**Arguments**

object            An object of class Theoretical GC

**Value**

A tibble object

**Examples**

```
mData(gcTheoretical)
```

---

overRep2Fasta	<i>Write fasta of Over-Represented sequences.</i>
---------------	---

---

**Description**

Output overrepresented sequences to disk in fasta format.

**Usage**

```
overRep2Fasta(x, path, n = 10, labels, noAdapters = TRUE, ...)

## S4 method for signature 'ANY'
overRep2Fasta(x, path, n = 10, labels, noAdapters = TRUE, ...)

## S4 method for signature 'FastqcData'
overRep2Fasta(x, path, n = 10, labels, noAdapters = TRUE, ...)

## S4 method for signature 'FastqcDataList'
overRep2Fasta(x, path, n = 10, labels, noAdapters = TRUE, ...)
```

**Arguments**

x	Can be a FastqcData or FastqcDataList
path	Path to export the fasta file to. Reverts to a default in the working directory if not supplied
n	The number of sequences to output
labels	An optional named factor of labels for the file names. All filenames must be present in the names. File extensions are dropped by default.
noAdapters	logical. Remove any sequences identified as possible adapters or primers by FastQC
...	Used to pass any alternative patterns to remove from the end of filenames

**Details**

Fasta will contain Filename, Possible Source, Percent of total reads

**Value**

Exports to a fasta file, and returns the fasta information invisibly

**Examples**

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)

# Export the top10 Overrepresented Sequences as a single fasta file
faOut <- file.path(tempdir(), "top10.fa")
overRep2Fasta(fdl, path = faOut)
```

---

path

*Return the File Paths from an object*

---

**Description**

Return the File Paths from an object

**Usage**

```
## S4 method for signature '.FastqcFile'  
path(object)  
  
## S4 method for signature 'FastqcData'  
path(object)  
  
## S4 method for signature 'FastqcDataList'  
path(object)  
  
## S4 method for signature '.FastpFile'  
path(object)  
  
## S4 method for signature 'FastpData'  
path(object)  
  
## S4 method for signature 'FastpDataList'  
path(object)
```

**Arguments**

object            An object of class .FastqcFile

**Details**

Obtains the file.path for objects of multiple classes

**Value**

A character vector of the file paths to the underlying FastQC reports

**Examples**

```
# Get the files included with the package  
packageDir <- system.file("extdata", package = "ngsReports")  
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)  
  
# Load the FASTQC data as a FastqcDataList object  
fdl <- FastqcDataList(fl)  
path(fdl)
```

---

plotAdapterContent     *Draw an Adapter Content Plot*

---

**Description**

Draw an Adapter Content Plot across one or more FASTQC reports



**Usage**

```
plotAdapterContent(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  ...  
)  
  
## S4 method for signature 'ANY'  
plotAdapterContent(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  ...  
)  
  
## S4 method for signature 'FastqcData'  
plotAdapterContent(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  pwfCols,  
  showPwf = TRUE,  
  warn = 5,  
  fail = 10,  
  scaleColour = NULL,  
  plotlyLegend = FALSE,  
  ...  
)  
  
## S4 method for signature 'FastqcDataList'  
plotAdapterContent(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  pwfCols,  
  showPwf = TRUE,  
  warn = 5,  
  fail = 10,  
  plotType = c("heatmap", "line"),  
  adapterType = "Total",  
  cluster = FALSE,  
  dendrogram = FALSE,  
  heat_w = 8L,  
  ...  
)
```

```

    scaleFill = NULL,
    scaleColour = NULL,
    plotlyLegend = FALSE,
    ...
)

## S4 method for signature 'FastpData'
plotAdapterContent(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  scaleFill = NULL,
  plotlyLegend = FALSE,
  plotTheme = theme_get(),
  ...
)

## S4 method for signature 'FastpDataList'
plotAdapterContent(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  pwfCols,
  showPwf = FALSE,
  warn = 5,
  fail = 10,
  cluster = FALSE,
  dendrogram = FALSE,
  scaleFill = NULL,
  plotTheme = theme_get(),
  heat_w = 8L,
  ...
)

```

### Arguments

x	Can be a FastqcData, a FastqcDataList or character vector of file paths
usePlotly	logical. Output as ggplot2 (default) or plotly object.
labels	An optional named vector of labels for the file names. All filenames must be present in the names.
pattern	regex used to trim the ends of all filenames for plotting
...	Used to pass additional attributes to theme() for FastQC objects and geoms for Fastp objects
pwfCols	Object of class PwfCols() containing the colours for PASS/WARN/FAIL
showPwf	logical(1) Show PASS/WARN/FAIL status as would be included in a standard FastQC report

warn, fail	The default values for warn and fail are 5 and 10 respectively (i.e. percentages)
plotlyLegend	logical(1) Show legend when choosing interactive plots. Ignored for heatmaps
plotType	character. Can only take the values plotType = "heatmap" or plotType = "line"
adapterType	A regular expression matching the adapter(s) to be plotted. To plot all adapters summed, specify adapterType = "Total". This is the default behaviour.
cluster	logical default FALSE. If set to TRUE, fastqc data will be clustered using hierarchical clustering
dendrogram	logical redundant if cluster is FALSE if both cluster and dendrogram are specified as TRUE then the dendrogram will be displayed.
heat_w	Width of the heatmap relative to other plot components
scaleFill, scaleColour	scale_fill\* and scale_colour_\* objects
plotTheme	Set theme elements by passing a <a href="#">theme</a>

### Details

This extracts the Adapter\_Content module from the supplied object and generates a ggplot2 object, with a set of minimal defaults. The output of this function can be further modified using the standard ggplot2 methods.

When x is a single or FastqcData object line plots will always be drawn for all adapters. Otherwise, users can select line plots or heatmaps. When plotting more than one fastqc file, any undetected adapters will not be shown.

An interactive version of the plot can be made by setting usePlotly as TRUE

### Value

A standard ggplot2 object, or an interactive plotly object

### Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
f1 <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(f1)

# The default plot
plotAdapterContent(fdl)

# Also subset the reads to just the R1 files
r1 <- grepl("R1", fqName(fdl))
plotAdapterContent(fdl[r1])

# Plot just the Universal Adapter
# and change the y-axis using ggplot2::scale_y_continuous
plotAdapterContent(fdl, adapterType = "Illumina_Universal", plotType = "line") +
```

```

facet_wrap(~Filename) +
guides(colour = "none")

# For FastpData object, the plots are slightly different
fp <- FastpData(system.file("extdata/fastp.json.gz", package = "ngsReports"))
plotAdapterContent(fp, scaleFill = scale_fill_brewer(palette = "Set1"))

```

---

plotAlignmentSummary *Plot a summary of alignments*

---

### Description

Plot a summary of alignments from a set of log files

### Usage

```

plotAlignmentSummary(
  x,
  type = c("star", "bowtie", "bowtie2", "hisat2"),
  usePlotly = FALSE,
  stripPaths = TRUE,
  asPercent = FALSE,
  ...,
  fill = c("red", "yellow", "blue", rgb(0, 0.5, 1))
)

```

### Arguments

x	Paths to one or more alignment log files
type	The aligner used. Can be one of star, bowtie, bowtie2 or hisat2
usePlotly	logical. If TRUE an interactive plot will be generated.
stripPaths	logical(1). Remove paths from the Filename column
asPercent	Show alignments as percentages, with the alternative (FALSE) being the total number of reads If FALSE a ggplot object will be output
...	Used to pass additional attributes to theme() and between methods
fill	Colours used to fill the bars. Passed to scale_fill_manual.

### Details

Loads a set of alignment log files and creates a default plot. Implemented aligners are bowtie, bowtie2, Hisat2 and STAR.

### Value

A ggplot2 object, or a plotly object

## Examples

```
f <- c("bowtie2PE.txt", "bowtie2SE.txt")
bowtie2Logs <- system.file("extdata", f, package = "ngsReports")
plotAlignmentSummary(bowtie2Logs, "bowtie2")
```

---

plotAssemblyStats      *Plot a summary of assembly logs*

---

## Description

Plot a summary of assembly stats from a set of log files

## Usage

```
plotAssemblyStats(
  x,
  type = c("quast", "busco"),
  usePlotly = FALSE,
  plotType = c("bar", "paracoord"),
  ...
)
```

## Arguments

x	Paths to one or more log files
type	The tool used. Can be one of quast or busco
usePlotly	logical. If TRUE an interactive plot will be generated. If FALSE a ggplot object will be output
plotType	character. Plot type to output, one of bar or paracoord.
...	Used to pass additional attributes to theme() and between methods

## Details

Loads a set of assembly log files and creates a default plot. Implemented tools are quast and BUSCO. quast will plot a parallel coordinate plot of some assembly statistics BUSCO will plot a stacked barplot of completeness statistics

## Value

A ggplot2 object, or a plotly object

## Examples

```
#getquast log filenames
quastFiles <- system.file("extdata",
c("quast1.tsv", "quast2.tsv"), package = "ngsReports")

# The default plot
plotAssemblyStats(quastFiles)
```

---

plotBaseQuals

*Plot the Base Qualities for each file*

---

## Description

Plot the Base Qualities for each file as separate plots

## Usage

```
plotBaseQuals(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)

## S4 method for signature 'ANY'
plotBaseQuals(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)

## S4 method for signature 'FastqcData'
plotBaseQuals(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  pwfCols,
  warn = 25,
  fail = 20,
  boxWidth = 0.8,
  showPwf = TRUE,
  plotlyLegend = FALSE,
  ...
)

## S4 method for signature 'FastqcDataList'
plotBaseQuals(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  pwfCols,
  warn = 25,
  fail = 20,
```

```
    showPwf = TRUE,
    boxWidth = 0.8,
    plotType = c("heatmap", "boxplot"),
    plotValue = "Mean",
    cluster = FALSE,
    dendrogram = FALSE,
    nc = 2,
    heat_w = 8L,
    ...
)

## S4 method for signature 'FastpData'
plotBaseQuals(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  pwfCols,
  warn = 25,
  fail = 20,
  showPwf = FALSE,
  module = c("Before_filtering", "After_filtering"),
  reads = c("read1", "read2"),
  readsBy = c("facet", "linetype"),
  bases = c("A", "T", "C", "G", "mean"),
  scaleColour = NULL,
  plotTheme = theme_get(),
  plotlyLegend = FALSE,
  ...
)

## S4 method for signature 'FastpDataList'
plotBaseQuals(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  pwfCols,
  warn = 25,
  fail = 20,
  showPwf = FALSE,
  module = c("Before_filtering", "After_filtering"),
  plotType = "heatmap",
  plotValue = c("mean", "A", "T", "C", "G"),
  scaleFill = NULL,
  plotTheme = theme_get(),
  cluster = FALSE,
  dendrogram = FALSE,
```

```

    heat_w = 8L,
    ...
)

```

### Arguments

x	Can be a FastqcData, FastqcDataList or character vector of file paths
usePlotly	logical Default FALSE will render using ggplot. If TRUE plot will be rendered with plotly
labels	An optional named vector of labels for the file names. All filenames must be present in the names.
pattern	Regex to remove from the end of the Fastp report and Fastq file names
...	Used to pass additional attributes to theme() and between methods
pwfCols	Object of class PwfCols() to give colours for pass, warning, and fail values in plot
warn, fail	The default values for warn and fail are 30 and 20 respectively (i.e. percentages)
boxWidth	set the width of boxes when using a boxplot
showPwf	Include the Pwf status colours
plotlyLegend	logical(1) Show legend for interactive plots. Only called when drawing line plots
plotType	character Can be either "boxplot" or "heatmap"
plotValue	character Type of data to be presented. Can be any of the columns returned by the appropriate call to getModule()
cluster	logical default FALSE. If set to TRUE, fastqc data will be clustered using hierarchical clustering
dendrogram	logical redundant if cluster is FALSE if both cluster and dendrogram are specified as TRUE then the dendrogram will be displayed.
nc	numeric. The number of columns to create in the plot layout. Only used if drawing boxplots for multiple files in a FastqcDataList
heat_w	Relative width of any heatmap plot components
module	Select Before and After filtering when using a FastpDataList
reads	Create plots for read1, read2 or all when using a FastpDataList
readsBy	If paired reads are present, separate using either linetype or by facet
bases	Which bases to include on the plot
scaleColour	ggplot discrete colour scale, passed to lines
plotTheme	theme object
scaleFill	ggplot2 continuous scale. Passed to heatmap cells



**Details**

When acting on a `FastqcDataList`, this defaults to a heatmap using the mean `Per_base_sequence_quality` score. A set of plots which replicate those obtained through a standard FastQC html report can be obtained by setting `plotType = "boxplot"`, which uses `facet_wrap` to provide the layout as a single `ggplot` object.

When acting on a `FastqcData` object, this replicates the `Per_base_sequence_quality` plots from FastQC with no faceting.

For large datasets, subsetting by R1 or R2 reads may be helpful.

An interactive plot can be obtained by setting `usePlotly = TRUE`.

**Value**

A standard `ggplot2` object or an interactive `plotly` object

**Examples**

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)

# The default plot for multiple libraries is a heatmap
plotBaseQuals(fdl)

# The default plot for a single library is the standard boxplot
plotBaseQuals(fdl[[1]])

# FastpData objects have qualities by base
fp <- FastpData(system.file("extdata/fastp.json.gz", package = "ngsReports"))
plotBaseQuals(
  fp, plotTheme = theme(plot.title = element_text(hjust = 0.5))
)
```

---

plotDupLevels

*Plot the combined Sequence\_Duplication\_Levels information*

---

**Description**

Plot the `Sequence_Duplication_Levels` information for a set of FASTQC reports

**Usage**

```
plotDupLevels(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)  
  
## S4 method for signature 'ANY'  
plotDupLevels(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)  
  
## S4 method for signature 'FastqcData'  
plotDupLevels(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  pwfCols,  
  warn = 20,  
  fail = 50,  
  showPwf = TRUE,  
  plotlyLegend = FALSE,  
  lineCol = c("red", "blue"),  
  lineWidth = 1,  
  ...  
)  
  
## S4 method for signature 'FastqcDataList'  
plotDupLevels(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  pwfCols,  
  warn = 20,  
  fail = 50,  
  showPwf = TRUE,  
  plotlyLegend = FALSE,  
  deduplication = c("pre", "post"),  
  plotType = c("heatmap", "line"),  
  cluster = FALSE,  
  dendrogram = FALSE,  
  heatCol = hcl.colors(50, "inferno"),  
  heat_w = 8,  
  ...  
)  
  
## S4 method for signature 'FastpData'  
plotDupLevels(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",
```

```

    pwfCols,
    warn = 20,
    fail = 50,
    showPwf = FALSE,
    maxLevel = 10,
    lineCol = "red",
    barFill = "dodgerblue4",
    barCol = barFill,
    plotlyLegend = FALSE,
    plotTheme = theme_get(),
    ...
)

## S4 method for signature 'FastpDataList'
plotDupLevels(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  pwfCols,
  warn = 20,
  fail = 50,
  showPwf = FALSE,
  plotlyLegend = FALSE,
  plotType = c("bar", "heatmap"),
  barFill = "blue",
  barCol = "blue",
  cluster = FALSE,
  dendrogram = FALSE,
  scaleFill = NULL,
  plotTheme = theme_get(),
  heat_w = 8,
  maxLevel = 10,
  ...
)

```

### Arguments

x	Can be a FastqcData, FastqcDataList or file path
usePlotly	logical Default FALSE will render using ggplot. If TRUE plot will be rendered with plotly
labels	An optional named vector of labels for the file names. All filenames must be present in the names. File extensions are dropped by default.
pattern	regex to remove from the end of fastp & fastq file names
...	Used to pass additional attributes to theme() and between methods
pwfCols	Object of class PwfCols() to give colours for pass, warning, and fail values in the plot

warn, fail	The default values for warn and fail are 20 and 50 respectively (i.e. percentages)
showPwf	logical(1) Show PWF rectangles in the background
plotlyLegend	logical(1) Show legend for line plots when using interactive plots
lineCol, lineWidth	Colours and width of lines drawn
deduplication	Plot Duplication levels 'pre' or 'post' deduplication. Can only take values "pre" and "post"
plotType	Choose between "heatmap" and "line"
cluster	logical default FALSE. If set to TRUE, fastqc data will be clustered using hierarchical clustering
dendrogram	logical Plot will automatically be clustered if TRUE.
heatCol	Colour palette used for the heatmap
heat_w	Relative width of the heatmap relative to other plot components
maxLevel	The maximum duplication level to plot. Beyond this level, all values will be summed
barFill, barCol	Colours for bars when calling geom_col()
plotTheme	<a href="#">theme</a> object. Applied after a call to theme_bw()
scaleFill	Discrete scale used to fill heatmap cells

### Details

This extracts the Sequence\_Duplication\_Levels from the supplied object and generates a ggplot2 object, with a set of minimal defaults. For multiple reports, this defaults to a heatmap with block sizes proportional to the percentage of reads belonging to that duplication category.

If setting usePlotly = FALSE, the output of this function can be further modified using standard ggplot2 syntax. If setting usePlotly = TRUE an interactive plotly object will be produced.

### Value

A standard ggplot2 or plotly object

### Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)

# Draw the default plot for a single file
plotDupLevels(fdl[[1]])

plotDupLevels(fdl)
```

---

plotFastqcPCA                      *Draw a PCA plot for Fast QC modules*

---

### Description

Draw a PCA plot for Fast QC modules across multiple samples **[Experimental]**

### Usage

```
plotFastqcPCA(  
  x,  
  module = "Per_sequence_GC_content",  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  sz = 4,  
  groups,  
  ...  
)  
  
## S4 method for signature 'ANY'  
plotFastqcPCA(  
  x,  
  module = "Per_sequence_GC_content",  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  sz = 4,  
  groups,  
  ...  
)  
  
## S4 method for signature 'character'  
plotFastqcPCA(  
  x,  
  module = "Per_sequence_GC_content",  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  sz = 4,  
  groups,  
  ...  
)  
  
## S4 method for signature 'FastqcDataList'  
plotFastqcPCA(  
  x,
```

```

    module = "Per_sequence_GC_content",
    usePlotly = FALSE,
    labels,
    pattern = ".(fast|fq|bam).*",
    sz = 4,
    groups,
    pc = c(1, 2),
    ...
  )

```

### Arguments

x	Can be a FastqcDataList or character vector of file paths
module	character vector containing the desired FastQC module (eg. c("Per_base_sequence_quality", "Per_base_sequence_content"))
usePlotly	logical. Output as ggplot2 (default) or plotly object.
labels	An optional named vector of labels for the file names. All file names must be present in the names of the vector.
pattern	Regex to remove from the end of any filenames
sz	The size of the text labels
groups	Optional factor of the same length as x. If provided, the plot will be coloured using this factor as the defined groups. Ellipses will also be added to the final plot.
...	Used to pass additional attributes to theme() and between methods
pc	The two components to be plotted

### Details

This carries out PCA on a single FastQC module and plots the output using either ggplot or plotly. Current modules for PCA are Per\_base\_sequence\_quality, Per\_sequence\_quality\_scores, Per\_sequence\_GC\_content, Per\_base\_sequence\_content, and Sequence\_Length\_Distribution.

If a factor is provided in the groups argument, this will be applied to the plotting colours and ellipses will be drawn using these groups. Only the labels will be plotted using geom\_text()

### Value

A standard ggplot2 object, or an interactive plotly object

### Examples

```

# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)
grp <- as.factor(gsub(".+(R[12]).*", "\\1", fqName(fdl)))

```

```
plotFastqcPCA(fdl, module = "Per_sequence_GC_content", groups = grp)
```

---

```
plotGcContent          Plot the Per Sequence GC Content
```

---

### Description

Plot the Per Sequence GC Content for a set of FASTQC files

### Usage

```
plotGcContent(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)
```

```
## S4 method for signature 'ANY'
```

```
plotGcContent(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)
```

```
## S4 method for signature 'FastqcData'
```

```
plotGcContent(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  theoreticalGC = TRUE,
  gcType = c("Genome", "Transcriptome"),
  species = "Hsapiens",
  GCobject,
  plotlyLegend = FALSE,
  Fastafile,
  n = 1e+06,
  counts = FALSE,
  scaleColour = NULL,
  lineCols = c("red3", "black"),
  linetype = 1,
  linewidth = 0.5,
  ...
)
```

```
## S4 method for signature 'FastqcDataList'
```

```
plotGcContent(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  theoreticalGC = TRUE,
  gcType = c("Genome", "Transcriptome"),
```

```

    species = "Hsapiens",
    GCobject,
    Fastafilename,
    n = 1e+06,
    plotType = c("heatmap", "line", "cdf"),
    cluster = FALSE,
    dendrogram = FALSE,
    heat_w = 8,
    pwfCols,
    showPwf = TRUE,
    scaleFill = NULL,
    scaleColour = NULL,
    plotlyLegend = FALSE,
    lineCols = RColorBrewer::brewer.pal(12, "Paired"),
    linetype = 1,
    linewidth = 0.5,
    ...
)

## S4 method for signature 'FastpData'
plotGcContent(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  theoreticalGC = TRUE,
  gcType = c("Genome", "Transcriptome"),
  species = "Hsapiens",
  GCobject,
  Fastafilename,
  n = 1e+06,
  plotType = "bar",
  scaleFill = NULL,
  plotlyLegend = FALSE,
  plotTheme = theme_get(),
  ...
)

## S4 method for signature 'FastpDataList'
plotGcContent(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  theoreticalGC = TRUE,
  gcType = c("Genome", "Transcriptome"),
  species = "Hsapiens",
  GCobject,

```



```

Fastafilename,
n = 1e+06,
plotType = "bar",
scaleFill = NULL,
plotTheme = theme_get(),
plotlyLegend = FALSE,
...
)

```

### Arguments

x	Can be a FastqcData, FastqcDataList or character vector of file paths
usePlotly	logical Default FALSE will render using ggplot. If TRUE plot will be rendered with plotly
labels	An optional named vector of labels for the file names.
pattern	Pattern to remove from the end of filenames
...	Used to pass various potting parameters to themes and geoms.
theoreticalGC	logical default is FALSE to give the true GC content, set to TRUE to normalize values of GC_Content by the theoretical values using <code>gcTheoretical()</code> . species must be specified. For Fastqc* objects, the entire distributions will be used, whereas for the Fastp* objects, only the expected mean value is shown as a horizontal line
gcType	character Select type of data to normalize GC content against. Accepts either "Genome" (default) or "Transcriptome".
species	character if gcTheory is TRUE it must be accompanied by a species. Species currently supported can be obtained using <code>mData(gcTheoretical)</code>
GCobject	an object of class GCTheoretical. Defaults to the gcTheoretical object supplied with the package
plotlyLegend	logical(1) Show legend on interactive line plots
Fastafilename	a fasta file contains DNA sequences to generate theoretical GC content
n	number of simulated reads to generate theoretical GC content from Fastafilename
counts	logical. Plot the counts from each file if counts = TRUE, otherwise frequencies will be plotted. Ignored if calling the function on a FastqcDataList.
scaleColour	ggplot2 scale for line colours
lineCols, linetype, linewidth	Line colour type and width for observed and theoretical GC lines
plotType	Takes values "line", "heatmap" or "cdf"
cluster	logical default FALSE. If set to TRUE, fastqc data will be clustered using hierarchical clustering
dendrogram	logical redundant if cluster is FALSE if both cluster and dendrogram are specified as TRUE then the dendrogram will be displayed.
heat_w	Relative width of any heatmap plot components

pwfCols	Object of class <code>PwfCols()</code> to give colours for pass, warning, and fail values in plot
showPwf	logical(1) Show Pwf Status on the plot
scaleFill	ggplot2 scale for filling heatmap cells or bars
plotTheme	<code>theme</code> object

### Details

Makes plots for GC\_Content. When applied to a single FastqcData object a simple line plot will be drawn, with Theoretical GC content overlaid if desired.

When applied to multiple FastQC reports, the density at each GC content bin can be shown as a heatmap by setting `theoreticalGC = FALSE`. By default the difference in observed and expected theoretical GC is shown. Species and genome/transcriptome should also be set if utilising the theoretical GC content.

As an alternative to a heatmap, a series of overlaid distributions can be shown by setting `plotType = "line"`.

Can produce a static ggplot2 object or an interactive plotly object.

### Value

A ggplot2 or plotly object

### Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)

# The default plot for a FastqcDataList
plotGcContent(fdl)

# Plot a single FastqcData object
plotGcContent(fdl[[1]])
```

---

plotInsertSize	<i>Plot Insert Size Distributions</i>
----------------	---------------------------------------

---

### Description

Plot the insert size distribution from one of Fastp reports

**Usage**

```
plotInsertSize(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)
```

```
## S4 method for signature 'FastpData'
```

```
plotInsertSize(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  plotType = c("histogram", "cumulative"),
  counts = FALSE,
  plotTheme = theme_get(),
  expand.x = 0.01,
  expand.y = c(0, 0.05),
  ...
)
```

```
## S4 method for signature 'FastpDataList'
```

```
plotInsertSize(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  plotType = c("heatmap", "line", "cumulative"),
  plotTheme = theme_get(),
  scaleFill = NULL,
  scaleColour = NULL,
  cluster = FALSE,
  dendrogram = FALSE,
  heat_w = 8,
  ...
)
```

**Arguments**

<code>x</code>	A <code>FastpData</code> or <code>FastpDataList</code> object
<code>usePlotly</code>	logical. Generate an interactive plot using plotly
<code>labels</code>	An optional named vector of labels for the file names. All file names must be present in the names of the vector.
<code>pattern</code>	Regex to remove from the end of any filenames
<code>...</code>	Passed to <code>geom*</code> functions during plotting
<code>plotType</code>	Determine the plot type. Options vary with the input structure
<code>counts</code>	logical(1) Plot read counts, or percentages (default)
<code>plotTheme</code>	a <a href="#">theme</a> object
<code>expand.x</code> , <code>expand.y</code>	Axis expansions

scaleFill	Continuous scale used to fill heatmap cells. Defaults to the "inferno" palette
scaleColour	Discrete scale for adding line colours
cluster	logical default FALSE. If set to TRUE, data will be clustered using hierarchical clustering
dendrogram	logical redundant if cluster is FALSE if both cluster and dendrogram are specified as TRUE the dendrogram will be displayed.
heat_w	Width of the heatmap relative to other plot components

### Details

Takes a Fastp as a set of Fastp reports and plot insert size distributions. Plots can be drawn as cumulative totals or the default histograms for a single report, and as boxplots or heatmaps for a set of reports

### Value

A ggplot or plotly object

### Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastp.json.gz", full.names = TRUE)
fp <- FastpData(fl)
plotInsertSize(
  fp, counts = TRUE, fill = "steelblue4",
  plotTheme = theme(plot.title = element_text(hjust = 0.5))
)
plotInsertSize(fp, plotType = "cumulative")
```

---

plotKmers

*Plot Overrepresented Kmers*

---

### Description

Plot Overrepresented Kmers

### Usage

```
plotKmers(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)

## S4 method for signature 'ANY'
plotKmers(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)

## S4 method for signature 'FastqcData'
plotKmers(
```

```

    x,
    usePlotly = FALSE,
    labels,
    pattern = ".(fast|fq|bam).*",
    n = 6,
    linewidth = 0.5,
    plotlyLegend = FALSE,
    scaleColour = NULL,
    pal = c("red", "blue", "green", "black", "magenta", "yellow"),
    ...
)

## S4 method for signature 'FastqcDataList'
plotKmers(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  cluster = FALSE,
  dendrogram = FALSE,
  pwfCols,
  showPwf = TRUE,
  scaleFill = NULL,
  heatCol = hcl.colors(50, "inferno"),
  heat_w = 8,
  ...
)

## S4 method for signature 'FastpData'
plotKmers(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  module = c("Before_filtering", "After_filtering"),
  reads = c("read1", "read2"),
  readsBy = c("facet", "mean", "diff"),
  trans = "log2",
  scaleFill = NULL,
  plotTheme = theme_get(),
  plotlyLegend = FALSE,
  ...
)

```

## Arguments

x                    Can be a FastqcData, FastqcDataList or file paths

usePlotly	logical Default FALSE will render using ggplot. If TRUE plot will be rendered with plotly
labels	An optional named vector of labels for the file names. All filenames must be present in the names.
pattern	regex to drop from the end of filenames
...	Used to pass parameters to theme for FastqcData objects and to geoms for FastpData objects
n	numeric. The number of Kmers to show.
linewidth	Passed to geom_line()
plotlyLegend	Show legend for interactive plots
pal	The colour palette. If the vector supplied is less than n, grDevices::colorRampPalette() will be used
cluster	logical default FALSE. If set to TRUE, fastqc data will be clustered using hierarchical clustering
dendrogram	logical redundant if cluster is FALSE if both cluster and dendrogram are specified as TRUE then the dendrogram will be displayed.
pwfCols	Object of class PwfCols() to give colours for pass, warning, and fail values in the plot
showPwf	Show the PASS/WARN/FAIL status
scaleFill, scaleColour	ggplot2 scales to be used for colour palettes
heatCol	Colour palette used for the heatmap. Default is inferno from the viridis set of palettes
heat_w	Relative width of any heatmap plot components
module	The module to obtain data from when using a FastpData object
reads	Either read1 or read2. Only used when using a FastpData object
readsBy	Strategy for visualising both read1 and read2. Can be set to show each set of reads by facet, or within the same plot taking the mean of the enrichment above mean, or the difference in the enrichment above mean
trans	Function for transforming the count/mean ratio. Set as NULL to use the ratio without transformation
plotTheme	<a href="#">theme</a> object

### Details

As the Kmer Content module present in FastQC reports is relatively uninformative, and omitted by default in later versions of FastQC, these are rudimentary plots.

Plots for FastqcData objects replicate those contained in a FastQC report, whilst the heatmap generated from FastqcDataList objects simply show the location and abundance of over-represented Kmers.

### Value

A standard ggplot2 object or an interactive plotly object

**Examples**

```

# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)
plotKmers(fdl[[1]])

# Use a FastpData object
fl <- system.file("extdata", "fastp.json.gz", package = "ngsReports")
fp <- FastpData(fl)
plotKmers(fp, size = 2)
plotKmers(
  fp, reads = "read1", size = 2, trans = NULL,
  scaleFill = scale_fill_gradient(low = "white", high = "black")
)

```

---

plotNContent

*Draw an N Content Plot*


---

**Description**

Draw an N Content Plot across one or more FastQC reports

**Usage**

```

plotNContent(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)

## S4 method for signature 'ANY'
plotNContent(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)

## S4 method for signature 'FastqcData'
plotNContent(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  pwfCols,
  warn = 5,
  fail = 20,
  showPwf = TRUE,
  ...,
  lineCol = "red"
)

## S4 method for signature 'FastqcDataList'

```

```
plotNContent(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  pwfCols,  
  warn = 5,  
  fail = 20,  
  showPwf = TRUE,  
  cluster = FALSE,  
  dendrogram = FALSE,  
  heat_w = 8,  
  scaleFill = NULL,  
  ...  
)  
  
## S4 method for signature 'FastpData'  
plotNContent(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  module = c("Before_filtering", "After_filtering"),  
  moduleBy = c("facet", "colour", "linetype"),  
  reads = c("read1", "read2"),  
  readsBy = c("facet", "colour", "linetype"),  
  scaleColour = NULL,  
  scaleLine = NULL,  
  plotTheme = theme_get(),  
  plotlyLegend = FALSE,  
  ...  
)  
  
## S4 method for signature 'FastpDataList'  
plotNContent(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  module = c("Before_filtering", "After_filtering"),  
  reads = c("read1", "read2"),  
  scaleFill = NULL,  
  plotTheme = theme_get(),  
  cluster = FALSE,  
  dendrogram = FALSE,  
  heat_w = 8,  
  ...  
)
```



**Arguments**

x	Can be a FastqcData, FastqcDataList or file paths
usePlotly	logical. Output as ggplot2 (default) or plotly object.
labels	An optional named vector of labels for the file names. All filenames must be present in the names.
pattern	Regex used to trim the end of filenames
...	Used to pass additional attributes to theme() for FastqcData objects and to geom* calls for FastpData-based objects
pwfCols	Object of class PwfCols() containing the colours for PASS/WARN/FAIL
warn, fail	The default values for warn and fail are 5 and 10 respectively (i.e. percentages)
showPwf	logical(1) Show the PASS/WARN/FAIL status
lineCol	Line colours
cluster	logical default FALSE. If set to TRUE, fastqc data will be clustered using hierarchical clustering
dendrogram	logical redundant if cluster is FALSE if both cluster and dendrogram are specified as TRUE then the dendrogram will be displayed.
heat_w	Relative width of any heatmap plot components
scaleFill, scaleColour, scaleLine	ggplot2 scale objects
module	Used for Fastp* structures to show results before or after filtering
moduleBy, readsBy	How to show each module or set of reads on the plot
reads	Show plots for read1, read2 or both.
plotTheme	theme object
plotlyLegend	logical(1) Show legend on interactive plots

**Details**

This extracts the N\_Content from the supplied object and generates a ggplot2 object, with a set of minimal defaults. The output of this function can be further modified using the standard ggplot2 methods.

When x is a single FastqcData object line plots will always be drawn for all Ns. Otherwise, users can select line plots or heatmaps.

**Value**

A standard ggplot2 object, or an interactive plotly object

**Examples**

```
## Using a Fastp Data object
fl <- system.file("extdata/fastp.json.gz", package = "ngsReports")
fp <- FastpData(fl)
plotNContent(fp)
plotNContent(
  fp, pattern = "_001.+",
  moduleBy = "colour", scaleColour = scale_colour_brewer(palette = "Set1"),
  plotTheme = theme(
    legend.position = 'inside', legend.position.inside = c(0.99, 0.99),
    legend.justification = c(1, 1), plot.title = element_text(hjust = 0.5)
  )
)
```

---

plotOverrep

*Plot a summary of Over-represented Sequences*


---

**Description**

Plot a summary of Over-represented Sequences for a set of FASTQC reports

**Usage**

```
plotOverrep(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  pwfCols,
  ...
)

## S4 method for signature 'ANY'
plotOverrep(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  pwfCols,
  ...
)

## S4 method for signature 'character'
plotOverrep(
  x,
  usePlotly = FALSE,
```

```

    labels,
    pattern = ".(fast|fq|bam).*",
    pwfCols,
    ...
)

## S4 method for signature 'FastqcData'
plotOverrep(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  pwfCols,
  n = 10,
  expand.x = c(0, 0, 0.05, 0),
  expand.y = c(0, 0.6, 0, 0.6),
  plotlyLegend = FALSE,
  ...
)

## S4 method for signature 'FastqcDataList'
plotOverrep(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  pwfCols,
  showPwf = TRUE,
  cluster = FALSE,
  dendrogram = FALSE,
  scaleFill = NULL,
  paletteName = "Set1",
  panel_w = 8,
  expand.x = c(0, 0, 0.05, 0),
  expand.y = rep(0, 4),
  ...
)

```

### Arguments

x	Can be a FastqcData, FastqcDataList or file paths
usePlotly	logical Default FALSE will render using ggplot. If TRUE plot will be rendered with plotly
labels	An optional named factor of labels for the file names. All filenames must be present in the names.
pattern	Regex to remove from the end of any filenames
pwfCols	Object of class <code>PwfCols()</code> containing the colours for PASS/WARN/FAIL

...	Used to pass additional attributes to theme() and between methods
n	The number of sequences to plot from an individual file
expand.x, expand.y	Output from expansion() or numeric vectors of length 4. Passed to scale_*_continuous()
plotlyLegend	Show legend on interactive plots
showPwf	Show PASS/WARN/FAIL status on the plot
cluster	logical default FALSE. If set to TRUE, fastqc data will be clustered using hierarchical clustering
dendrogram	logical redundant if cluster is FALSE if both cluster and dendrogram are specified as TRUE then the dendrogram will be displayed.
scaleFill	ggplot scale object
paletteName	Name of the palette for colouring the possible sources of the overrepresented sequences. Must be a palette name from RColorBrewer. Ignored if specifying the scaleFill separately
panel_w	Width of main panel on output

### Details

Percentages are obtained by simply summing those within a report. Any possible double counting by FastQC is ignored for the purposes of a simple approximation.

Plots generated from a FastqcData object will show the top n sequences grouped by their predicted source & coloured by whether the individual sequence would cause a WARN/FAIL.

Plots generated from a FastqcDataList group sequences by predicted source and summarise as a percentage of the total reads.

### Value

A standard ggplot2 object

### Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)

# A brief summary across all FastQC reports
plotOverrep(fdl)
```

---

plotReadTotals	<i>Draw a barplot of read totals</i>
----------------	--------------------------------------

---

**Description**

Draw a barplot of read totals

**Usage**

```
plotReadTotals(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)
```

```
## S4 method for signature 'ANY'
```

```
plotReadTotals(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)
```

```
## S4 method for signature 'FastqcDataList'
```

```
plotReadTotals(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  duplicated = TRUE,
  bars = c("stacked", "adjacent"),
  vertBars = TRUE,
  divBy = 1,
  barCols = c("red", "blue"),
  expand.y = c(0, 0.02),
  plotlyLegend = FALSE,
  ...
)
```

```
## S4 method for signature 'FastpDataList'
```

```
plotReadTotals(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  adjPaired = TRUE,
  divBy = 1e+06,
  scaleFill = NULL,
  labMin = 0.05,
  status = TRUE,
  labelVJ = 0.5,
  labelFill = "white",
  plotTheme = theme_get(),
  vertBars = FALSE,
  plotlyLegend = FALSE,
  expand.y = c(0, 0.05),
)
```

```
    ...
  )
```

### Arguments

x	Can be a FastqcData, FastqcDataList or file paths
usePlotly	logical Default FALSE will render using ggplot. If TRUE plot will be rendered with plotly
labels	An optional named vector of labels for the file names. All filenames must be present in the names.
pattern	Regex used to trim the end of filenames
...	Used to pass additional attributes to theme()
duplicated	logical(1). Include deduplicated read total estimates to plot charts
bars	If duplicated = TRUE, show unique and deduplicated reads as "stacked" or "adjacent".
vertBars	logical(1) Show bars as vertical or horizontal
divBy	Scale read totals by this value. The default shows the y-axis in millions for FastpDataList objects, but does not scale FastQC objects, for the sake of backwards compatability
barCols	Colours for duplicated and unique reads.
expand.y	Passed to <code>ggplot2::expansion</code> for the axis showing totals
plotlyLegend	logical(1) Show legend on interactive plots
adjPaired	Scale read totals by 0.5 when paired
scaleFill	ScaleDiscrete function to be applied to the plot
labMin	Only show labels for filtering categories higher than this values as a proportion of reads. Set to any number > 1 to turn off labels
status	logical(1) Include read status in the plot
labelVJ	Relative vertical position to labels within each bar.
labelFill	Passed to <code>geom_label</code>
plotTheme	<code>theme</code> to be added to the plot

### Details

Draw a barplot of read totals using the standard ggplot2 syntax. The raw data from `readTotals()` can otherwise be used to manually create a plot.

Duplication levels are based on the value shown on FASTQC reports at the top of the DeDuplicated-Totals plot, which is known to be inaccurate. As it still gives a good guide as to sequence diversity it is included as the default. This can be turned off by setting `duplicated = FALSE`.

For FastpDataList objects, duplication statistics are not part of the default module containing Read-Totals. However, the status of reads and the reason for being retained or filtered is, and as such these are shown instead of duplication statistics.

**Value**

Returns a ggplot or plotly object

**Examples**

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)

# Plot the Read Totals showing estimated duplicates
plotReadTotals(fdl)

# Plot the Read Totals without estimated duplicates
plotReadTotals(fdl, duplicated = FALSE)
```

---

plotSeqContent	<i>Plot the per base content as a heatmap</i>
----------------	---

---

**Description**

Plot the Per Base content for a set of FASTQC files.

**Usage**

```
plotSeqContent(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)

## S4 method for signature 'ANY'
plotSeqContent(x, usePlotly = FALSE, labels, pattern = ".(fast|fq|bam).*", ...)

## S4 method for signature 'FastqcData'
plotSeqContent(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  bases = c("A", "T", "C", "G"),
  scaleColour = NULL,
  plotTheme = theme_get(),
  plotlyLegend = FALSE,
  expand.x = 0.02,
  expand.y = c(0, 0.05),
  ...
)
```

```
## S4 method for signature 'FastqcDataList'
plotSeqContent(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  pwfCols,
  showPwf = TRUE,
  plotType = c("heatmap", "line", "residuals"),
  scaleColour = NULL,
  plotTheme = theme_get(),
  cluster = FALSE,
  dendrogram = FALSE,
  heat_w = 8,
  plotlyLegend = FALSE,
  nc = 2,
  ...
)

## S4 method for signature 'FastpData'
plotSeqContent(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  module = c("Before_filtering", "After_filtering"),
  reads = c("read1", "read2"),
  readsBy = c("facet", "linetype"),
  moduleBy = c("facet", "linetype"),
  bases = c("A", "T", "C", "G", "N", "GC"),
  scaleColour = NULL,
  scaleLine = NULL,
  plotlyLegend = FALSE,
  plotTheme = theme_get(),
  expand.x = 0.02,
  expand.y = c(0, 0.05),
  ...
)

## S4 method for signature 'FastpDataList'
plotSeqContent(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
  module = c("Before_filtering", "After_filtering"),
  moduleBy = c("facet", "linetype"),
  reads = c("read1", "read2"),
```



```

readsBy = c("facet", "linetype"),
bases = c("A", "T", "C", "G", "N", "GC"),
showPwf = FALSE,
pwfCols,
warn = 10,
fail = 20,
plotType = c("heatmap", "line", "residuals"),
plotlyLegend = FALSE,
scaleColour = NULL,
scaleLine = NULL,
plotTheme = theme_get(),
cluster = FALSE,
dendrogram = FALSE,
heat_w = 8,
expand.x = c(0.01),
expand.y = c(0, 0.05),
nc = 2,
...
)

```

### Arguments

x	Can be a FastqcData, FastqcDataList or file paths
usePlotly	logical. Generate an interactive plot using plotly
labels	An optional named vector of labels for the file names. All file names must be present in the names of the vector.
pattern	Regex to remove from the end of any filenames
...	Used to pass additional attributes to plotting geoms
bases	Which bases to draw on the plot. Also becomes the default plotting order by setting these as factor levels
scaleColour	Discrete colour scale as a ggplot ScaleDiscrete object If not provided, will default to <a href="#">scale_colour_manual</a>
plotTheme	<a href="#">theme</a> object to be applied. Note that all plots will have <a href="#">theme_bw</a> theme applied by default, as well as any additional themes supplied here
plotlyLegend	logical(1) Show legends for interactive plots. Ignored for heatmaps
expand.x, expand.y	Passed to <a href="#">expansion</a> in the x- and y-axis scales respectively
pwfCols	Object of class <a href="#">PwfCols()</a> to give colours for pass, warning, and fail values in plot
showPwf	Show PASS/WARN/FAIL categories as would be defined in a FastQC report
plotType	character. Type of plot to generate. Must be "line", "heatmap" or "residuals"
cluster	logical default FALSE. If set to TRUE, fastqc data will be clustered using hierarchical clustering
dendrogram	logical redundant if cluster is FALSE if both cluster and dendrogram are specified as TRUE then the dendrogram will be displayed.

heat_w	Relative width of any heatmap plot components
nc	Specify the number of columns if plotting a FastqcDataList as line plots. Passed to <code>facet_wrap</code> .
module	Fastp Module to show. Can only be Before/After_filtering
reads	Which set of reads to show
readsBy, moduleBy	When plotting both R1 & R2 and both modules, separate by either linetype or linetype
scaleLine	Discrete scale_linetype object. Only relevant if plotting values by linetype
warn, fail	Default values for WARN and FAIL based on FastQC reports. Only applied to heatmaps for FastpDataList objects

### Details

Per base sequence content (%A, %T, %G, %C), is shown as four overlaid heatmap colours when plotting from multiple reports. The individual line plots are able to be generated by setting `plotType = "line"`, and the layout is determined by `facet_wrap` from `ggplot2`.

Individual line plots are also generated when plotting from a single `FastqcData` object.

If setting `usePlotly = TRUE` for a large number of reports, the plot can be slow to render. An alternative may be to produce a plot of residuals for each base, produced by taking the position-specific mean for each base.

### Value

A `ggplot2` object or an interactive `plotly` object

### Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)

# The default plot
plotSeqContent(fdl)

fp <- FastpData(system.file("extdata/fastp.json.gz", package = "ngsReports"))
plotSeqContent(fp)
plotSeqContent(fp, moduleBy = "linetype", bases = c("A", "C", "G", "T"))
```

---

plotSeqLengthDistn      *Plot the Sequence Length Distribution*

---

### Description

Plot the Sequence Length Distribution across one or more FASTQC reports

### Usage

```
plotSeqLengthDistn(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  ...  
)  
  
## S4 method for signature 'ANY'  
plotSeqLengthDistn(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  ...  
)  
  
## S4 method for signature 'character'  
plotSeqLengthDistn(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  ...  
)  
  
## S4 method for signature 'FastqcData'  
plotSeqLengthDistn(  
  x,  
  usePlotly = FALSE,  
  labels,  
  pattern = ".(fast|fq|bam).*",  
  counts = TRUE,  
  plotType = c("line", "cdf"),  
  expand.x = c(0, 0.2, 0, 0.2),  
  plotlyLegend = FALSE,  
  colour = "red",  
  ...  
)
```

```

)

## S4 method for signature 'FastqcDataList'
plotSeqLengthDistn(
  x,
  usePlotly = FALSE,
  labels,
  pattern = "(fast|fq|bam).*",
  counts = FALSE,
  plotType = c("heatmap", "line", "cdf"),
  cluster = FALSE,
  dendrogram = FALSE,
  heat_w = 8,
  pwfCols,
  showPwf = TRUE,
  scaleFill = NULL,
  scaleColour = NULL,
  heatCol = hcl.colors(50, "inferno"),
  plotlyLegend = FALSE,
  ...
)

```

### Arguments

x	Can be a FastqcData, FastqcDataList or file paths
usePlotly	logical. Output as ggplot2 or plotly object.
labels	An optional named vector of labels for the file names. All filenames must be present in the names.
pattern	Regex to remove from the end of any filenames
...	Used to pass additional attributes to theme()
counts	logical Should distributions be shown as counts or frequencies (percentages)
plotType	character. Can only take the values plotType = "heatmap" plotType = "line" or plotType = "cdf"
expand.x	Output from expansion() or numeric vector of length 4. Passed to scale_x_discrete
plotlyLegend	logical(1) Show legend for interactive line plots
colour	Line colour
cluster	logical default FALSE. If set to TRUE, fastqc data will be clustered using hierarchical clustering
dendrogram	logical redundant if cluster and usePlotly are FALSE. If both cluster and dendrogram are specified as TRUE then the dendrogram will be displayed.
heat_w	Relative width of any heatmap plot components
pwfCols	Object of class PwfCols() to give colours for pass, warning, and fail values in plot
showPwf	logical(1) Show PASS/WARN/FAIL status

scaleFill, scaleColour  
 Optional ggplot scale objects

heatCol            The colour scheme for the heatmap

### Details

This extracts the Sequence Length Distribution from the supplied object and generates a ggplot2 object, with a set of minimal defaults. The output of this function can be further modified using the standard ggplot2 methods.

A cdf plot can also be generated to provide guidance for minimum read length in some NGS workflows, by setting plotType = "cdf". If all libraries have reads of identical lengths, these plots may be less informative.

An alternative interactive plot is available by setting the argument usePlotly = TRUE.

### Value

A standard ggplot2 object, or an interactive plotly object

### Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)

# Plot as a frequency plot using lines
plotSeqLengthDistn(fdl)

# Or plot the cdf
plotSeqLengthDistn(fdl, plotType = "cdf")
```

---

plotSeqQuals

*Plot the Per Sequence Quality Scores*

---

### Description

Plot the Per Sequence Quality Scores for a set of FASTQC reports

### Usage

```
plotSeqQuals(
  x,
  usePlotly = FALSE,
  labels,
  pattern = ".(fast|fq|bam).*",
```

```
    pwfCols,  
    ...  
  )  
  
  ## S4 method for signature 'ANY'  
  plotSeqQuals(  
    x,  
    usePlotly = FALSE,  
    labels,  
    pattern = ".(fast|fq|bam).*",  
    pwfCols,  
    ...  
  )  
  
  ## S4 method for signature 'character'  
  plotSeqQuals(  
    x,  
    usePlotly = FALSE,  
    labels,  
    pattern = ".(fast|fq|bam).*",  
    pwfCols,  
    ...  
  )  
  
  ## S4 method for signature 'FastqcData'  
  plotSeqQuals(  
    x,  
    usePlotly = FALSE,  
    labels,  
    pattern = ".(fast|fq|bam).*",  
    pwfCols,  
    showPwf = TRUE,  
    counts = FALSE,  
    alpha = 0.1,  
    warn = 30,  
    fail = 20,  
    colour = "red",  
    plotlyLegend = FALSE,  
    ...  
  )  
  
  ## S4 method for signature 'FastqcDataList'  
  plotSeqQuals(  
    x,  
    usePlotly = FALSE,  
    labels,  
    pattern = ".(fast|fq|bam).*",  
    pwfCols,
```

```

counts = FALSE,
alpha = 0.1,
warn = 30,
fail = 20,
showPwf = TRUE,
plotType = c("heatmap", "line"),
dendrogram = FALSE,
cluster = FALSE,
scaleFill = NULL,
heatCols = hcl.colors(100, "inferno"),
heat_w = 8,
scaleColour = NULL,
plotlyLegend = FALSE,
...
)

```

### Arguments

x	Can be a FastqcData, FastqcDataList or path
usePlotly	logical Default FALSE will render using ggplot. If TRUE plot will be rendered with plotly
labels	An optional named vector of labels for the file names. All file names must be present in the names of the vector.
pattern	Regex to remove from the end of any filenames
pwfCols	Object of class PwfCols() containing the colours for PASS/WARN/FAIL
...	Used to pass various potting parameters to theme. Can also be used to set size and colour for box outlines.
showPwf	logical(1) Show PASS/WARN/FAIL status
counts	logical. Plot the counts from each file if counts = TRUE, otherwise the frequencies will be plotted
alpha	set alpha for line graph bounds
warn, fail	The default values for warn and fail are 5 and 10 respectively (i.e. percentages)
colour	Colour for single line plots
plotlyLegend	logical(1) Show legend for interactive line plots
plotType	character. Can only take the values plotType = "heatmap" or plotType = "line"
dendrogram	logical redundant if cluster is FALSE if both cluster and dendrogram are specified as TRUE then the dendrogram will be displayed.
cluster	logical default FALSE. If set to TRUE, fastqc data will be clustered using hierarchical clustering
scaleFill, scaleColour	ggplot2 scales
heatCols	Colour palette for the heatmap
heat_w	Relative width of any heatmap plot components

### Details

Plots the distribution of average sequence quality scores across the set of files. Values can be plotted either as counts (`counts = TRUE`) or as frequencies (`counts = FALSE`).

Any faceting or scale adjustment can be performed after generation of the initial plot, using the standard methods of `ggplot2` as desired.

### Value

A standard `ggplot2` object, or an interactive `plotly` object

### Examples

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
f1 <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(f1)

# The default plot
plotSeqQuals(fdl)

# Also subset the reads to just the R1 files
r1 <- grepl("R1", fqName(fdl))
plotSeqQuals(fdl[r1])
```

---

plotSummary

*Plot the PASS/WARN/FAIL information*

---

### Description

Extract the PASS/WARN/FAIL summaries and plot them

### Usage

```
plotSummary(
  x,
  usePlotly = FALSE,
  labels,
  pwfCols,
  cluster = FALSE,
  dendrogram = FALSE,
  ...
)

## S4 method for signature 'ANY'
plotSummary(
```



```

    x,
    usePlotly = FALSE,
    labels,
    pwfCols,
    cluster = FALSE,
    dendrogram = FALSE,
    ...
)

## S4 method for signature 'character'
plotSummary(
  x,
  usePlotly = FALSE,
  labels,
  pwfCols,
  cluster = FALSE,
  dendrogram = FALSE,
  ...
)

## S4 method for signature 'FastqcDataList'
plotSummary(
  x,
  usePlotly = FALSE,
  labels,
  pwfCols,
  cluster = FALSE,
  dendrogram = FALSE,
  ...,
  gridlineWidth = 0.2,
  gridlineCol = "grey20"
)

```

### Arguments

x	Can be a FastqcData, FastqcDataList or character vector of file paths
usePlotly	logical. Generate an interactive plot using plotly
labels	An optional named vector of labels for the file names. All filenames must be present in the names. File extensions are dropped by default.
pwfCols	Object of class <code>PwfCols()</code> containing the colours for PASS/WARN/FAIL
cluster	logical default FALSE. If set to TRUE, fastqc data will be clustered using hierarchical clustering
dendrogram	logical redundant if cluster is FALSE if both cluster and dendrogram are specified as TRUE then the dendrogram will be displayed.
...	Used to pass various potting parameters to theme.
gridlineWidth, gridlineCol	Passed to <code>geom_hline</code> and <code>geom_vline</code> to determine width and colour of gridlines

**Details**

This uses the standard ggplot2 syntax to create a three colour plot. The output of this function can be further modified using the standard ggplot2 methods if required.

**Value**

A ggplot2 object (usePlotly = FALSE) or an interactive plotly object (usePlotly = TRUE)

**Examples**

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)

# Check the overall PASS/WARN/FAIL status
plotSummary(fdl)
```

---

pwf

*Colours for PASS/WARN/FAIL*

---

**Description**

Default colours for PASS/WARN/FAIL

**Usage**

pwf

**Format**

An object of class PwfCols of length 1.

**Details**

pwf is an object of class PwfCols supplied with the package and used as the default colouring. Colours correspond approximately to PASS, WARN and FAIL from the FASTQC reports, with the additional colour (MAX) included to indicate an extreme FAIL. In order, these colours in the default vector are green (rgb(0, 0.8, 0)), yellow (rgb(0.9, 0.9, 0.2)), red (rgb(0.8, 0.2, 0.2)) and white (rgb(1, 1, 1))

**Examples**

```
# Make a pie chart showing the default colours
pie(rep(1,4), labels = names(pwf), col = getColours(pwf))
```

---

PwfCols-class

*The PwfCols class and associated methods*


---

**Description**

Define the PwfCols class and associated methods

**Details**

This is an object of with four colours in components named PASS, WARN, FAIL and MAX. Used to indicate these categories as defined on the standard plots from fastqc.

**Value**

An S4 object of class PwfCols

**Slots**

PASS A vector of length 1, defining the colour for PASS in rgb format. Defaults to rgb(0, 0.8, 0)

WARN A vector of length 1, defining the colour for WARN in rgb format. Defaults to rgb(0.9, 0.9, 0.2)

FAIL A vector of length 1, defining the colour for FAIL in rgb format. Defaults to rgb(0.8, 0.2, 0.2)

MAX A vector of length 1, defining the colour for an extreme FAIL or NA in rgb format. Defaults to rgb(1, 1, 1)

---

readTotals

*Get the read totals*


---

**Description**

Get the read totals from one or more FASTQC reports

**Usage**

```
readTotals(x)
```

**Arguments**

x Can be a FastqcData, FastqcDataList, FastpData, FastpDataList or file paths

**Value**

A tibble with the columns Filename and Total\_Sequences

**Examples**

```
# Get the files included with the package
packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)

# Load the FASTQC data as a FastqcDataList object
fdl <- FastqcDataList(fl)

# Print the read totals
readTotals(fdl)
```

---

summariseOverrep	<i>Summarise Overrepresented Sequences</i>
------------------	--

---

**Description**

Summarise the Overrepresented sequences found in one or more QC files

**Usage**

```
summariseOverrep(x, ...)

## S4 method for signature 'FastpData'
summariseOverrep(x, step = c("Before", "After"), min_count = 0, ...)

## S4 method for signature 'FastpDataList'
summariseOverrep(
  x,
  min_count = 0,
  step = c("Before", "After"),
  vals = c("count", "rate"),
  fn = c("mean", "sum", "max"),
  by = c("reads", "sequence"),
  ...
)

## S4 method for signature 'FastqcDataList'
summariseOverrep(
  x,
  min_count = 0,
  vals = c("Count", "Percentage"),
  fn = c("mean", "sum", "max"),
  pattern = ".*",
  ...
)
```

```
## S4 method for signature 'FastqcData'
summariseOverrep(
  x,
  min_count = 0,
  vals = c("Count", "Percentage"),
  fn = c("mean", "sum", "max"),
  pattern = ".*",
  by = "Filename",
  ...
)
```

### Arguments

x	An object of a suitable class
...	Not used
step	Can be 'Before', 'After' or both to obtain data from the Before_filtering or After_filtering modules
min_count	Filter sequences with counts less than this value, both before and after filtering
vals	Values to use for creating summaries across multiple files. For FastpDataList objects these can be "count" and/or "rate", whilst for FastqcDataList objects these values can be "Count" and/or "Percentage"
fn	Functions to use when summarising values across multiple files
by	character vector of columns to summarise by. See <a href="#">dplyr::summarise</a>
pattern	Regular expression to filter the Possible_Source column by

### Details

This function prepares a useful summary of all over-represented sequences as reported by either fastp or FastQC

### Value

A tibble

Tibble columns will vary between Fastp\*, FastqcDataList and FastqcData objects. Calling this function on list-type objects will attempt to summarise the presence each over-represented sequence across all files.

In particular, FastqcData objects will provide the requested summary statistics across all sequences within a file

### Examples

```
## For operations on a FastpData object
f <- system.file("extdata/fastp.json.gz", package = "ngsReports")
fp <- FastpData(f)
summariseOverrep(fp, min_count = 100)

## Applying the function to a FastqcDataList
```

```

packageDir <- system.file("extdata", package = "ngsReports")
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)
fdl <- FastqcDataList(fl)
summariseOverrep(fdl)

# An alternative viewpoint can be obtained using
fdl |> lapply(summariseOverrep) |> dplyr::bind_rows()

```

---

TheoreticalGC-class    *The TheoreticalGC Object Class*

---

### Description

Contains Theoretical GC content for a selection of species

### Details

Estimates are able to be retained for genomic and transcriptomic sequences. Values are stored as frequencies.

### Value

An object of class TheoreticalGC

### Slots

Genome A data.frame containing theoretical GC content for genomic sequences

Transcriptome A data.frame containing theoretical GC content for transcriptomic sequences

mData A data.frame containing metadata about all species in the object

### Examples

```

## How to form an object using your own fasta file
faDir <- system.file("extdata", package = "ngsReports")
faFile <- list.files(faDir, pattern = "fasta", full.names = TRUE)
gen_df <- estGcDistn(faFile, n = 200)
gen_df <- dplyr::rename(gen_df, Athaliana = Freq)
mData_df <-
  data.frame(Name = "Athaliana", Genome = TRUE, Transcriptome = FALSE)
tr_df <- data.frame()
myGC <- new(
  "TheoreticalGC", Genome = gen_df, Transcriptome = tr_df, mData = mData_df)

```

---

writeHtmlReport	<i>Write an HTML Summary Report</i>
-----------------	-------------------------------------

---

## Description

Compiles an HTML report using a supplied template

## Usage

```
writeHtmlReport(
  fastqcDir,
  template,
  outDir,
  usePlotly = TRUE,
  species = "Hsapiens",
  gcType = c("Genome", "Transcriptome"),
  nOver = 30,
  targetsDF,
  overwrite = FALSE,
  quiet = TRUE
)
```

## Arguments

fastqcDir	A directory containing zipped, or extracted FastQC reports
template	The template file which will be copied into fastqcDir
outDir	The directory to write the compiled document to
usePlotly	Generate interactive plots?
species	Species/closely related species of sequenced samples
gcType	Is the data "Transcriptomic" or "Genomic" in nature?
nOver	The maximum number of Overrepresented Sequences to show
targetsDF	A data.frame with at least two columns named Filename and Label. The file-names should match the original fastq files, and the labels should be simply alternative labels for these files for convenience.
overwrite	logical. Overwrite any previous copies of the template file in the destination directory
quiet	logical. Show or hide markdown output in the Console.

## Details

This will take a user supplied template, or the file supplied with the package and create an HTML summary of all standard FASTQC plots for all files in the supplied directory.

**Value**

Silently returns TRUE and will output a compiled HTML file from the supplied Rmarkdown template file

**Examples**

```
## Not run:
packageDir <- system.file("extdata", package = "ngsReports")
fileList <- list.files(packageDir, pattern = "fastqc.zip", full.names= TRUE)
# Copy these files to tempdir() to avoid overwriting
# any files in the package directory
file.copy(fileList, tempdir(), overwrite = TRUE)
writeHtmlReport(fastqcDir = tempdir())

## End(Not run)
```

---

[,FastqcDataList,numeric,missing-method  
*Extract Elements*

---

**Description**

Extract elements from FastqcDataList Object

**Usage**

```
## S4 method for signature 'FastqcDataList,numeric,missing'
x[i, j, ..., drop = TRUE]

## S4 method for signature 'FastqcDataList,character,missing'
x[i, j, ..., drop = TRUE]

## S4 method for signature 'FastqcDataList,logical,missing'
x[i, j, ..., drop = TRUE]

## S4 method for signature 'FastqcDataList,ANY,missing'
x[i, j, ..., drop = TRUE]

## S4 method for signature 'FastpDataList,numeric,missing'
x[i, j, ..., drop = TRUE]

## S4 method for signature 'FastpDataList,character,missing'
x[i, j, ..., drop = TRUE]

## S4 method for signature 'FastpDataList,logical,missing'
x[i, j, ..., drop = TRUE]
```



```
## S4 method for signature 'FastpDataList,ANY,missing'  
x[i, j, ..., drop = TRUE]
```

### Arguments

x	A FastqcDataList or FastpDataList
i	character, logical or integer vector
j	not used
...	not used
drop	not used

### Details

Extract elements in a consistent manner with R conventions

### Value

Will return a subset of the original object following the standard rules for subsetting objects

### Examples

```
# Get the files included with the package  
packageDir <- system.file("extdata", package = "ngsReports")  
fl <- list.files(packageDir, pattern = "fastqc.zip", full.names = TRUE)  
  
# Load the FASTQC data as a FastqcDataList object  
fdl <- FastqcDataList(fl)  
  
# Subsetting using the standard methods  
fdl[1]  
fdl[[1]]
```

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