

# Package ‘alevinQC’

October 14, 2021

**Type** Package

**Title** Generate QC Reports For Alevin Output

**Version** 1.8.0

**Description** Generate QC reports summarizing the output from an alevin run.  
Reports can be generated as html or pdf files, or as shiny applications.

**Encoding** UTF-8

**Depends** R (>= 4.0)

**Imports** rmarkdown (>= 2.5), tools, methods, ggplot2, GGally, dplyr,  
rjson, shiny, shinydashboard, DT, stats, utils, tximport (>=  
1.17.4), cowplot, rlang

**RoxxygenNote** 7.1.1

**Suggests** knitr, BiocStyle, testthat

**VignetteBuilder** knitr

**biocViews** QualityControl, SingleCell

**URL** <https://github.com/csoneson/alevinQC>

**BugReports** <https://github.com/csoneson/alevinQC/issues>

**License** MIT + file LICENSE

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**Author** Charlotte Soneson [aut, cre] (<<https://orcid.org/0000-0003-3833-2169>>),  
Avi Srivastava [aut]

**Maintainer** Charlotte Soneson <charlottesoneson@gmail.com>

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alevinQC-pkg	<i>alevinQC</i>
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### Description

`alevinQC`

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alevinQCReport	<i>Generate alevin summary report</i>
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### Description

Generate a report summarizing the main aspects of an alevin quantification run. The report generation assumes that alevin has been run with the `--dumpFeatures` flag to generate the necessary output files.

### Usage

```
alevinQCReport(
  baseDir,
  sampleId,
  outputFile,
  outputDir = "./",
  outputFormat = NULL,
  showCode = FALSE,
  forceOverwrite = FALSE,
  knitrProgress = FALSE,
  quiet = FALSE,
  ignorePandoc = FALSE,
  customCBLList = list(),
  ...
)
```

## Arguments

baseDir	Path to the output directory from the alevin run (should be the directory containing the alevin directory).
sampleId	Sample ID, will be used to set the title for the report.
outputFile	File name of the output report. The file name extension must be either .html or .pdf, and consistent with the value of outputFormat.
outputDir	Path to the output directory where the report will be generated.
outputFormat	The format of the output report. Either "html_document" or "pdf_document". The file name extension of outputFile must be consistent with this choice.
showCode	Logical, whether to display the R code in the report.
forceOverwrite	Logical, whether to force overwrite an existing report with the same name in the output directory.
knitrProgress	Logical, whether to display the progress of knitr when generating the report.
quiet	Logical, whether to show progress messages.
ignorePandoc	Logical, determines what to do if pandoc or pandoc-citeproc is missing (if Sys.which("pandoc") or Sys.which("pandoc-citeproc") returns ""). If ignorePandoc is TRUE, only a warning is given. The figures will be generated, but not the final report. If ignorePandoc is FALSE (default), the execution stops immediately.
customCBLList	Named list with custom set(s) of barcodes to provide summary statistics/plots for, in addition to the whitelists generated by alevin.
...	Other arguments that will be passed to rmarkdown::render.

## Details

When the function is called, a .Rmd template file will be copied into the output directory, and rmarkdown::render will be called to generate the final report. If there is already a .Rmd file with the same name in the output directory, the function will raise an error and stop, to avoid overwriting the existing file. The reason for this behaviour is that the copied template in the output directory will be deleted once the report is generated.

## Value

Generates a summary report in the outputDir directory, and returns (invisibly) the name of the generated report.

## Author(s)

Charlotte Soneson

## Examples

```
alevinQCReport(baseDir = system.file("extdata/alevin_example_v0.14",
                                      package = "alevinQC"),
               sampleId = "example", outputFile = "alevinReport.html",
               outputDir = tempdir(), forceOverwrite = TRUE)
```

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alevinQCShiny	<i>Generate alevin summary shiny app</i>
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## Description

Generate a shiny app summarizing the main aspects of an alevin quantification run. The app generation assumes that alevin has been run with the –dumpFeatures flag to generate the necessary output files.

## Usage

```
alevinQCShiny(baseDir, sampleId, customCBLlist = list())
```

## Arguments

baseDir	Path to the output directory from the alevin run (should be the directory containing the alevin directory).
sampleId	Sample ID, will be used set the title for the app.
customCBLlist	Named list with custom set(s) of barcodes to provide summary statistics/plots for, in addition to the whitelists generated by alevin.

## Value

A shiny app.

## Author(s)

Charlotte Soneson

## Examples

```
app <- alevinQCShiny(baseDir = system.file("extdata/alevin_example_v0.14",
                                              package = "alevinQC"),
                      sampleId = "example")
if (interactive()) {
  shiny::runApp(app)
}
```

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checkAlevinInputFiles *Check that all required input files are available*

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## Description

Check that all required input files are available

## Usage

```
checkAlevinInputFiles(baseDir)
```

## Arguments

baseDir Path to the output directory from the alevin run (should be the directory containing the alevin directory).

## Value

Returns nothing, raises an error if any of the required files are missing.

## Author(s)

Charlotte Soneson

## Examples

```
checkAlevinInputFiles(system.file("extdata/alevin_example_v0.14",
                                 package = "alevinQC"))
```

---

plotAlevinBarcodeCollapse

*Summary plot of cell barcode collapsing*

---

## Description

Plot the original frequency of each cell barcode in the original whitelist against the frequency after collapsing similar cell barcodes.

## Usage

```
plotAlevinBarcodeCollapse(cbTable)
```

## Arguments

cbTable data.frame (such as the cbTable returned by readAlevinQC) with barcode frequencies before and after collapsing.

**Value**

A ggplot object

**Author(s)**

Charlotte Soneson

**Examples**

```
alevin <- readAlevinQC(system.file("extdata/alevin_example_v0.14",
                                    package = "alevinQC"))
plotAlevinBarcodeCollapse(alevin$cbTable)
```

**plotAlevinHistogram**     *Histogram of selected summary statistic*

**Description**

Histogram of selected summary statistic

**Usage**

```
plotAlevinHistogram(
  cbTable,
  plotVar = "dedupRate",
  axisLabel = plotVar,
  colName = "inFinalWhiteList",
  cbName = "final whitelist"
)
```

**Arguments**

cbTable	data.frame (such as the cbTable returned by <code>readAlevinQC</code> ) containing the desired summary statistic in a column.
plotVar	Character scalar giving the name of a numeric column of cbTable to plot.
axisLabel	Character scalar giving the label of the selected statistic (will be displayed as the axis label in the plot).
colName	Character scalar giving the name of a logical column of cbTable to use for filling the bars in the histogram.
cbName	Character scalar giving the name of the set of barcodes defined by colName, used for labelling the plot legend.

**Value**

A ggplot object

**Author(s)**

Charlotte Soneson

**Examples**

```
alevin <- readAlevinQC(system.file("extdata/alevin_example_v0.14",
                                    package = "alevinQC"))
plotAlevinHistogram(alevin$cbTable, plotVar = "dedupRate",
                    axisLabel = "Deduplication rate",
                    colName = "inFinalWhiteList",
                    cbName = "final whitelist")
```

---

**plotAlevinKneeNbrGenes**

*Knee plot of the number of detected genes per cell*

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**Description**

Plot the number of detected genes per cell in decreasing order. Only cells contained in the original whitelist are considered.

**Usage**

```
plotAlevinKneeNbrGenes(cbTable)
```

**Arguments**

cbTable	data.frame (such as the cbTable returned by <code>readAlevinQC</code> ) with the number of detected genes per cell.
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**Value**

A ggplot object

**Author(s)**

Charlotte Soneson

**Examples**

```
alevin <- readAlevinQC(system.file("extdata/alevin_example_v0.14",
                                    package = "alevinQC"))
plotAlevinKneeNbrGenes(alevin$cbTable)
```

`plotAlevinKneeRaw`      *Knee plot of raw cell barcode frequencies*

### Description

Plot the raw cell barcode frequencies in decreasing order, and indicate a predetermined breakpoint (indicating barcodes included in the original whitelist) using color as well as a label.

### Usage

```
plotAlevinKneeRaw(cbTable)
```

### Arguments

<code>cbTable</code>	<code>data.frame</code> with raw barcode frequencies (such as the <code>cbTable</code> returned by <code>readAlevinQC</code> ).
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### Value

A `ggplot` object

### Author(s)

Charlotte Soneson

### Examples

```
alevin <- readAlevinQC(system.file("extdata/alevin_example_v0.14",
                                    package = "alevinQC"))
plotAlevinKneeRaw(alevin$cbTable)
```

`plotAlevinQuant`      *Panel of plots with quantification summary statistics*

### Description

Panel of plots with quantification summary statistics

### Usage

```
plotAlevinQuant(
  cbTable,
  colName = "inFinalWhiteList",
  cbName = "final whitelist"
)
```

**Arguments**

cbTable	data.frame (such as the cbTable returned by readAlevinQC) with collapsed barcode frequencies, the total UMI count and the number of detected genes for each cell.
colName	Character scalar giving the name of a logical column of cbTable to use for coloring the points.
cbName	Character scalar giving the name of the set of barcodes defined by colName, used for labelling the plot legend.

**Value**

A ggplot object

**Author(s)**

Charlotte Soneson

**Examples**

```
alevin <- readAlevinQC(system.file("extdata/alevin_example_v0.14",
                                     package = "alevinQC"))
plotAlevinQuant(alevin$cbTable, colName = "inFinalWhiteList",
                 cbName = "final whitelist")
```

**plotAlevinQuantPairs** *Pairs plot with quantification summary statistics*

**Description**

Pairs plot with quantification summary statistics

**Usage**

```
plotAlevinQuantPairs(cbTable, colName = "inFinalWhiteList")
```

**Arguments**

cbTable	data.frame (such as the cbTable returned by readAlevinQC) with collapsed barcode frequencies, the total UMI count and the number of detected genes for each cell.
colName	Character scalar giving the name of a logical column of cbTable to use for coloring the points.

**Value**

A ggmatrix object

## Author(s)

Charlotte Soneson

## Examples

```
alevin <- readAlevinQC(system.file("extdata/alevin_example_v0.14",
                                    package = "alevinQC"))
plotAlevinQuantPairs(alevin$cbTable, colName = "inFinalWhiteList")
```

`readAlevinQC` *Read alevin data required to generate summary report*

## Description

Read all alevin output files required to generate the summary report or shiny app.

## Usage

```
readAlevinQC(baseDir, customCBLList = list())
```

## Arguments

- |                            |   |
|----------------------------|---|
| <code>baseDir</code>       | Path to the output directory from the alevin run (should be the directory containing the <code>alevin</code> directory).              |
| <code>customCBLList</code> | Named list with custom set(s) of barcodes to provide summary statistics/plots for, in addition to the whitelists generated by alevin. |

## Value

A list collecting all necessary information for generating the summary report/shiny app.

## Author(s)

Charlotte Soneson

## Examples

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