# Package 'BiocCheck'

October 16, 2019

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<b>Version</b> 1.20.0
Title Bioconductor-specific package checks
<b>Description</b> Executes Bioconductor-specific package checks.
<b>Depends</b> R (>= 3.5.0)
License Artistic-2.0
LazyData true
VignetteBuilder knitr
<b>Imports</b> biocViews (>= 1.33.7), BiocManager, stringdist, graph, httr, tools, optparse, codetools, methods, utils, knitr
<b>Suggests</b> RUnit, BiocGenerics, Biobase, RJSONIO, rmarkdown, devtools (>= 1.4.1), usethis, BiocStyle
Enhances codetoolsBioC
biocViews Infrastructure
<pre>URL https://github.com/Bioconductor/BiocCheck/issues</pre>
git_url https://git.bioconductor.org/packages/BiocCheck
git_branch RELEASE_3_9
git_last_commit 68d9869
git_last_commit_date 2019-05-02
Date/Publication 2019-10-15
Author Bioconductor Package Maintainer [aut, cre], Lori Shepherd [ctb], Daniel von Twisk [ctb], Kevin Rue [ctb]
Maintainer Bioconductor Package Maintainer <maintainer@bioconductor.org></maintainer@bioconductor.org>
R topics documented:
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BiocCheck	Check a package for compliance with Bioconductor package guide- lines

#### **Description**

Analyzes R packages for compliance with Bioconductor package guidelines and best practices. Divides output into three categories: ERROR, WARNING, and NOTE. This function is mostly meant to be called from the operating system's command line (via R CMD BiocCheck; see the vignette). Not meant to replace R CMD check, which should always be run first.

## Usage

```
BiocCheck(package = ".", ...)
usage()
```

### **Arguments**

package A directory or tarball (.tar.gz file) containing an R source package.

Run usage() to see the available options. When running BiocCheck interactively, options can be passed like so: BiocCheck(package, `no-check-vignettes`=TRUE)

#### **Details**

BiocCheck() analyzes R packages for compliance with Bioconductor package guidelines and best practices. For the rationale behind these guidelines and best practices, see the vignette and pages in the references section. usage() displays the options that can be passed to BiocCheck().

BiocCheck is typically called from the operating system's command line, as R CMD BiocCheck package

where package is a directory or .tar.gz file containing an R source package.

BiocCheck is not meant as a replacement for R CMD check, which should always be run first for best results.

Installing BiocCheck will attempt to install the BiocCheck script, which could fail; details in the vignette.

See the vignette for detailed explanations of all the checks performed by BiocCheck.

#### Value

Mostly called for the side effect of the information displayed. When called interactively, returns a list with three components:

ERROR Items that must be fixed before the package can be accepted into Bioconductor.

WARNING We strongly suggest fixing these items.

NOTE Fixing these items is not required, there is no expectation that considerations

will escalate in severity.

### Author(s)

Dan Tenenbaum

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#### References

http://www.bioconductor.org/developers/how-to/coding-style/http://www.bioconductor.org/developers/package-guidelines/http://www.bioconductor.org/developers/how-to/version-numbering/http://www.bioconductor.org/developers/how-to/unitTesting-guidelines/

### **Examples**

```
packageDir <- system.file("testpackages", "testpkg0", package="BiocCheck")
BiocCheck(packageDir, `quit-with-status`=FALSE)
## or from the operating system's command line as:
## R CMD BiocCheck myPackage</pre>
```

BiocCheckGitClone

Additional checks for a package for compliance with Bioconductor package guidelines

### **Description**

Analyzes R packages for compliance with Bioconductor package guidelines and best practices. Divides output into three categories: ERROR, WARNING, and NOTE. This function is mostly meant to be called from the operating system's command line (via R CMD BiocCheckGitClone). Not meant to replace R CMD check or R CMD BiocCheck, which should always be run first.

## Usage

```
BiocCheckGitClone(package=".", ...)
usage2()
```

# **Arguments**

package A directory containing an R source package. Not a package tar ball.

Only available option currently is quit-with-status. See BiocCheck

# **Details**

BiocCheckGitClone() analyzes R packages for compliance with Bioconductor package guidelines and best practices. This function should only be run on a open source directory not a tarball.

BiocCheckGitClone is typically called from the operating system's command line, as R CMD BiocCheckGitClone package

where package is a directory containing an R source package.

BiocCheckGitClone is not meant as a replacement for R CMD check or R CMD BiocCheck, which should always be run first for best results.

Installing BiocCheck will attempt to install the BiocCheckGitClone script, which could fail; details in the vignette.

See the vignette for detailed explanations of all the checks performed by BiocCheckGitClone.

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#### Value

Mostly called for the side effect of the information displayed. When called interactively, returns a list with three components:

ERROR Items that must be fixed before the package can be accepted into Bioconductor.

WARNING We strongly suggest fixing these items.

NOTE Fixing these items is not required, there is no expectation that considerations

will escalate in severity.

### Author(s)

Lori Shepherd

#### References

```
http://www.bioconductor.org/developers/how-to/coding-style/http://www.bioconductor.org/developers/package-guidelines/http://www.bioconductor.org/developers/how-to/version-numbering/http://www.bioconductor.org/developers/how-to/unitTesting-guidelines/
```

## **Examples**

```
packageDir <- system.file("testpackages", "testpkg0", package="BiocCheck")
BiocCheckGitClone(packageDir, `quit-with-status`=FALSE)
## or from the operating system's command line as:
## R CMD BiocCheckGitClone myPackage</pre>
```

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