## Package 'BiocCheck'

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**Version** 1.40.0

Title Bioconductor-specific package checks

**Description** BiocCheck guides maintainers through Bioconductor best practicies. It runs Bioconductor-specific package checks by searching through package code, examples, and vignettes. Maintainers are required to address all errors, warnings, and most notes produced.

**Depends** R (>= 4.3.0)

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VignetteBuilder knitr

- **Imports** BiocBaseUtils, BiocFileCache, BiocManager, biocViews (>= 1.33.7), callr, codetools, graph, httr2, knitr, methods, rvest, stringdist, tools, utils
- Suggests RUnit, BiocGenerics, Biobase, jsonlite, rmarkdown, downloader, devtools (>= 1.4.1), usethis, BiocStyle, GenomicRanges, gert

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biocViews Infrastructure

URL https://github.com/Bioconductor/BiocCheck

BugReports https://github.com/Bioconductor/BiocCheck/issues

RoxygenNote 7.3.1

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BiocCheck

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BiocCheck	Check a package's adherence with the Bioconductor Package Guide-
	lines

## Description

Analyzes an R package for adherence with Bioconductor package guidelines and best practices. The check outputs are categorized into ERROR, WARNING, and NOTE. See the vignette for more details. BiocCheck is complementary to R CMD check, which should always be run first.

#### Usage

```
BiocCheck(
   package = getwd(),
   checkDir = dirname(package),
   debug = FALSE,
   callr = FALSE,
   ...
)
```

#### Arguments

package

The path to an R package directory or tarball (.tar.gz). The BiocCheck function is intended to be run from the package directory; therefore, the current working directory (given by getwd()) is the default.

## BiocCheck

checkDir	The directory where the BiocCheck output directory will be stored. By default, it will be placed in the same directory as the package directory i.e., dirname(pkg_dir).
debug	Whether to append the names of functions that correspond to each condition raised by BiocCheck in the written log (i.e., in the ' <package_name>.BiocCheck' folder). This option is only relevant to developers and contributors to BiocCheck.</package_name>
callr	logical(1) Whether to use the callr package to run BiocCheck in an isolated R session to prevent namespace collisions.
	See the details section for available options. When running BiocCheck, options can be specified as:
	BiocCheck(package, `no-check-vignettes`=TRUE)

#### Details

BiocCheck() reviews R packages for adherence with Bioconductor package guidelines and best practices. See https://contributions.bioconductor.org for the latest guidance for writing Bioconductor software. Some rationale behind these best practices can be seen in the vignette and pages in the references section. The vignette also provides detailed explanations of all the checks performed by BiocCheck.

BiocCheck is called within R with

BiocCheck(<package>)

where package points to the source directory or the .tar.gz tarball that was created using R CMD build.

*Note* that BiocCheck is complementary to R CMD check. R CMD check should always be run first for best results.

#### Value

BiocCheck() is chiefly called for the side effect of the check reporting. The function also creates a <package\_name>.BiocCheck folder and returns a BiocCheck reference class with three main list elements:

error	Items to address before the package can be accepted
warning	Strongly suggested items that may require attention
note	Items to consider, though not required, before acceptance

## dot-options

new-package enable checks specific to new packages

no-check-dependencies disable check for bad dependencies

no-check-deprecated disable check for usage of deprecated packages

**no-check-remotes** disable check for usage of remote packages other than those hosted on CRAN or Bioconductor

no-check-version-num disable check for valid version number

no-check-R-ver disable check for valid R version

no-check-pkg-size disable check for package tarball size no-check-file-size disable check for individual file size no-check-bioc-views disable biocViews-specific checks (for non-BioC packages) no-check-bbs disable BBS-specific checks (for non-BioC packages). Valid DESCRIPTION no-check-description disable DESCRIPTION file checks no-check-vignettes disable vignette checks no-check-library-calls disable check usage of functions that install or update packages no-check-install-self disable check for require or library of itself no-check-coding-practices disable check for some common best coding practices no-check-function-len disable check for function length **no-check-man-doc** disable checks for man page documentation no-check-news disable checks for NEWS file no-check-unit-tests disable checks for unit tests no-check-skip-bioc-tests disable check for tests that skip when on bioc no-check-formatting disable checks for file formatting no-check-CRAN disable check for if package exists in CRAN no-check-bioc-help disable check for registration on Bioconductor mailing list and support site build-output-file file containing R CMD build output, for additional analysis quit-with-status enable exit code option when performing check

#### Author(s)

Dan Tenenbaum, Lori Shepherd, and Marcel Ramos

#### References

https://contributions.bioconductor.org

## See Also

BiocCheck-class, Message-class

#### Examples

```
packageDir <- system.file("testpackages", "testpkg0", package="BiocCheck")
BiocCheck(packageDir, `quit-with-status`=FALSE)</pre>
```

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#### Description

The BiocCheck class provides a framework for reporting checks based on Bioconductor guidelines. The class has several methods for working with the provided checks that handle and display messages and the display of the metadata. These methods also record the output of the BiocCheck() report in both plain text and JSON formats.

**Note** that currently, multiple BiocCheck runs will interfere with each other given that they are implemented via a reference class semantic. When running multiple checks in the same session, you can separate these instances by running them in separate processes (e.g., via BiocParallel).

#### Details

The metadata includes a number of standard fields to allow easier troubleshooting and display of potentially relevant information. Currently, the fields included are

BiocCheckVersion The version of the BiocCheck package

BiocVersion The version of Bioconductor

Package The name of the package in check

PackageVersion The version of the package in check

sourceDir The directory of the package source or tarball in check

installDir The directory where the package is installed for testing, a temporary location by default

**BiocCheckDir** The directory where the <package>.BiocCheck folder is saved. Usually the same folder as the package in check

platform The platform/OS where the check is taking place

isTarBall Whether the package in check is a source directory or a tarball

#### Value

A BiocCheck instance

#### Fields

log list() A running list of all conditions raised (i.e., notes, warnings, errors)

check character(1) The title of the last check used for logging purposes.

error, warning, note list() Finer extraction of each condition type

metadata list() A list of additional information relevant to the package and its state. See details.

verbose logical(1) Whether to show the full information pertaining to the checks. A FALSE value will only show the condition messages and not any relevant files or additional information. The defaults are FALSE and TRUE for BiocCheck and BiocCheckGitClone, respectively.

## See Also

Message-class

## Examples

bc <- BiocCheck:::.BiocCheck</pre>

bc\$verbose

BiocCheck-methods A list of methods for the BiocCheck reference class

## Description

A list of methods for the BiocCheck reference class

## Arguments

	character() A vector that makes up the BiocCheck exception message (e.g., 'Vignette must be built by R CMD build'). The character vector is handled with paste0 and made into a list and appended with help_text and messages.
help_text	character(1) Additional text prompting a list of files (e.g., "Found in files:")
condition	character(1) One of the three conditions handled: error, warning, or note
messages	character() Often a vector of file names where the check was triggered.
verbose	logical(1) Whether or not to output both the help_text and messages as part of the report
debug	logical(1) Whether to append the name of the originating check name into for trace-ability
checkName	character(1) The title of the current group of checks. It can be set with handleCheck, e.g., handleCheck("Checking for version number mismatch"). Internally, it 'is saved with setCheck and obtained with getLastCheck.
isOnBBS	logical(1) Indicates whether the checks are being run on the Bioconductor Build System (BBS). This is helpful for avoiding the creation of folders in the BBS.
file	character(1) A path to a JSON file for writing or reading as created by toJSON and fromJSON BiocCheck methods.

## Value

An internal BiocCheck R5 Reference Class used to document conditions such as errors, warnings, and notes

#### methods

add Include a condition to the BiocCheck report getLastCheck Obtain the name of the last check run setCheck Create a new element in the internal list for a check get Extract the list of conditions raised by BiocCheck getNum Tally the number of condition provided by the input zero Reset the internal log of the condition provided getBiocCheckDir Report and create the <package>.BiocCheck directory as obtained from the metadata composeReport Simplify the list structure from the log and provide a character vector of conditions raised report Write the 00BiocCheck.log report into the BiocCheck folder toJSON Write a JSON file to the location indicated with the conditions raised from JSON Read a JSON file from the location indicated with the output of previous conditions raised in the check **show** Display the information in the class. Currently empty. show\_meta Display the metadata information stored in the metadata field

BiocCheckGitClone Checks specific to a Git clone of a package repository

#### Description

Analyzes an R package for adherence with Bioconductor package guidelines and best practices. The check outputs are categorized into ERROR, WARNING, and NOTE. This function is typically used in the Bioconductor Build System (BBS) and not intended for general use.

## Usage

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

#### Arguments

package	A directory containing an R source package. Not a package tar ball.
	Currently, only quit-with-status is available. See BiocCheck

#### Details

BiocCheckGitClone() reviews R packages for adherence with Bioconductor package guidelines and best practices. See <a href="https://contributions.bioconductor.org">https://contributions.bioconductor.org</a> for the latest guidance for writing Bioconductor software. This function should only be run on a source directory and not on a tarball.

BiocCheckGitClone is called within R with, as

BiocCheckGitClone(<package>)

where package is the source directory containing the R package.

## Value

BiocCheckGitClone() is chiefly called for the side effect of the check reporting. The function returns a BiocCheck reference class with three main list elements:

error	Items to address before the package can be accepted
warning	Strongly suggested items that may require attention
note	Items to consider, though not required, before acceptance

## Author(s)

Lori Shepherd

## References

https://contributions.bioconductor.org

#### See Also

BiocCheck-class

## Examples

```
packageDir <- system.file("testpackages", "testpkg0", package="BiocCheck")
BiocCheckGitClone(packageDir, `quit-with-status`=FALSE)</pre>
```

Context	Report context of events to user with a data.frame of events and loca-
	tions

## Description

Report context of events to user with a data.frame of events and locations

## Usage

```
Context(file = "", lines = character(), idx = logical(), offset = 0L)
```

## Arguments

file	character(1) full path (including package name) of file being summarized.
lines	character() vector of text lines in file
idx	logical() same length as lines indicating lines in which event occurs
offset	'integer(1)' The number of lines to add to the 'Line' column calculation. It is mainly used to account for the number of lines that the YAML header occupies.

## Value

Context: a data.frame() with columns File, Line, and Context

Message-class

## Description

A lower level Message helper class for BiocCheck

#### Value

A Message class instance

#### Fields

msg list() A list of character messages usually grown with append with conditions raised by a check

condition character(1) One of the three conditions handled: error, warning, or note

## See Also

BiocCheck-class

Message-methods A list of methods for the Message reference class

#### Description

A list of methods for the Message reference class

#### Arguments

condition	character(1) One of the three conditions handled: error, warning, or note
verbose	logical(1) Whether to output the full text in the check or only the check name itself in the report
	list() A nested list with the check name as the top level layer. Second level lists include any help_text and messages that are part of the check.

#### Value

An internal R5 Reference Class to handle messages and their conditions, e.g., for errors, warnings, or notes.

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