

# Package ‘BiocWorkflowTools’

October 16, 2024

**Title** Tools to aid the development of Bioconductor Workflow packages

**Version** 1.30.0

**Encoding** UTF-8

**Description** Provides functions to ease the transition between RMarkdown and LaTeX documents when authoring a Bioconductor Workflow.

**License** MIT + file LICENSE

**Depends** R (>= 3.4)

**Imports** BiocStyle, bookdown, git2r, httr, knitr, rmarkdown, rstudioapi, stringr, tools, utils, usethis

**NeedsCompilation** no

**VignetteBuilder** knitr

**biocViews** Software, ReportWriting

**RoxxygenNote** 7.1.0

**Collate** 'createBiocWorkflow.R' 'f1000\_article.R' 'uploadToOverleaf.R'  
'utils.R'

**git\_url** <https://git.bioconductor.org/packages/BiocWorkflowTools>

**git\_branch** RELEASE\_3\_19

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createBioWorkflow	<i>Create a New Bioconductor Workflow Package</i>
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**Description**

Uses [create\\_package](#) to set up a skeleton for a new Bioconductor workflow package.

**Usage**

```
createBioWorkflow(
  path,
  description = getOption("devtools.desc"),
  rstudio = TRUE,
  open = rstudio
)
```

**Arguments**

<code>path</code>	location to create new package. The last component of the path will be used as the package name.
<code>description</code>	list of description values to override default values or add additional values.
<code>rstudio</code>	if TRUE, creates an RStudio project file.
<code>open</code>	if TRUE, opens the project in a new RStudio session.

**Value**

File path to the R Markdown vignette (invisibly).

**Examples**

```
createBioWorkflow(file.path(tempdir(), "MyWorkflow"), open = FALSE)
```

f1000_article	<i>F1000Research article format</i>
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**Description**

Format for creating F1000Research software tool articles.

## Usage

```
f1000_article(
  toc = FALSE,
  number_sections = FALSE,
  fig_width = 5.67,
  fig_height = fig_width,
  fig_align = "center",
  keep_tex = TRUE,
  citation_package = "natbib",
  md_extensions = "+link_attributes",
  pandoc_args = "--wrap=preserve",
  ...
)
```

## Arguments

<code>toc</code>	TRUE to include a table of contents in the output
<code>number_sections</code>	TRUE to number section headings
<code>fig_width</code>	Default width (in inches) for figures
<code>fig_height</code>	Default height (in inches) for figures
<code>fig_align</code>	Default alignment of figures. Possible values are "center" (default) "left" and "right".
<code>keep_tex</code>	Keep the intermediate tex file used in the conversion to PDF
<code>citation_package</code>	The LaTeX package to process citations, <code>natbib</code> or <code>biblatex</code> . Use <code>none</code> if neither package is to be used.
<code>md_extensions</code>	Markdown extensions to be added or removed from the default definition or R Markdown. See the <a href="#">rmarkdown_format</a> for additional details.
<code>pandoc_args</code>	Additional command line options to pass to pandoc
<code>...</code>	Arguments to <a href="#">pdf_document</a>

## Details

Creates LaTeX sources which can be submitted to F1000Research through Overleaf.

## Value

R Markdown output format to pass to [render](#)

## Citations

R Markdown supports automatic generation of citations. You can find more information on the markdown citation syntax in the [Bibliographies and Citations](#) article in the R Markdown online documentation.

A bibliography file can be specified using the `bibliography` metadata field in the document's YAML header. Metadata variables for customizing citation style include:

**biblio-style** Bibliography style (e.g. "unsrtnat", "plainnat")  
**natbiboptions** Options to natbib LaTeX package (e.g. "number", "super", "round")  
**biblatexoptions** Options to biblatex LaTeX package

## Examples

```
## Not run:  
  
rmarkdown::draft("MyArticle.Rmd", template="f1000_article", package="BiocWorkflowTools")  
  
## End(Not run)
```

**uploadToOverleaf**      *Upload a LaTeX project to Overleaf*

## Description

Upload a LaTeX project to Overleaf

## Usage

```
uploadToOverleaf(path)
```

## Arguments

path	File path to a directory or a single zip file to be uploaded.
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## Value

Does not return any value. The Overleaf project page will automatically open in the default browser.

## Examples

```
## Not run:  
## don't run this code chunk in the example as we don't want to spam Overleaf  
uploadToOverleaf(files = 'MyWorkflow', openInBrowser = TRUE)  
  
## End(Not run)
```

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