Package 'BiocManager'

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Title Access the Bioconductor Project Package Repository

Description A convenient tool to install and update Bioconductor packages.	
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R topics documented:	
available BiocManager-pkg install repositories valid version	2 2 2 3 6 8
Index	1(

2 BiocManager-pkg

available

Discover packages available for installation.

Description

Discover packages available for installation.

Usage

```
available(pattern = "", include_installed = TRUE)
```

Arguments

pattern

 $character(1)\ pattern\ to\ filter\ (via\ grep(\texttt{pattern=...}))\ available\ packages;\ the$

filter is not case sensitive.

include_installed

logical(1) When TRUE, include installed packages in list of available packages; when FALSE, exclude installed packages.

Value

character() vector of package names available for installation.

Examples

```
avail <- BiocManager::available()
length(avail)
BiocManager::available("bs.*hsapiens")</pre>
```

BiocManager-pkg

Install or update Bioconductor, CRAN, or GitHub packages

Description

This package provides tools for managing *Bioconductor* and other packages in a manner consistent with *Bioconductor*'s package versioning and release system.

BiocManager-pkg 3

Details

Main functions are as follows; additional help is available for each function, e.g., ?BiocManager::version.

BiocManager::install() Install or update packages from Bioconductor, CRAN, and GitHub.

BiocManager::version() Report the version of *Bioconductor* in use.

BiocManager::available() Return a character() vector of package names available (at BiocManager::repositories(for installation.

BiocManager::valid() Determine whether installed packages are from the same version of *Bioconductor*.

BiocManager::repositories() *Bioconductor* and other repository URLs to discover packages for installation.

The version of _Bioconductor_ in use is determined by the installed version of a second package, BiocVersion. BiocVersion is installed automatically during first use of `BiocManager::install()`. If BiocVersion has not yet been installed, the version is determined by code in base R.

Author(s)

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See Also

Useful links:

• Report bugs at https://github.com/Bioconductor/BiocManager/issues

Examples

```
R.version.string
packageVersion("BiocManager")
if ("BiocVersion" %in% rownames(installed.packages()))
    packageVersion("BiocVersion")
BiocManager::version()
```

4 install

install

Install or update Bioconductor, CRAN, and GitHub packages

Description

The BiocManager::install() function installs or updates *Bioconductor* and CRAN packages in a *Bioconductor* release. Upgrading to a new *Bioconductor* release may require additional steps; see https://bioconductor.org/install.

Usage

```
install(pkgs = character(), ..., site_repository = character(),
  update = TRUE, ask = TRUE, checkBuilt = FALSE,
  version = BiocManager::version())
```

Arguments

pkgs	character() vector of package names to install or update. A missing value updates installed packages according to update = and ask =. Package names containing a '/' are treated as GitHub repositories and installed using remotes::install_github().	
• • •	Additional arguments used by install.packages().	
site_repository		
	(Optional) character(1) vector representing an additional repository in which to look for packages to install. This repository will be prepended to the default repositories (which you can see with BiocManager::repositories()).	
update	logical(1). When FALSE, BiocManager::install() does not attempt to update old packages. When TRUE, update old packages according to ask.	
ask	logical(1) indicating whether to prompt user before installed packages are updated. If TRUE, user can choose whether to update all outdated packages without further prompting, to pick packages to update, or to cancel updating (in a non-interactive session, no packages will be updated unless ask = FALSE).	
checkBuilt	logical(1). If TRUE a package built under an earlier major.minor version of R (e.g., 3.4) is considered to be old.	
version	character(1) <i>Bioconductor</i> version to install, e.g., version = "3.8". The special symbol version = "devel" installs the current 'development' version.	

Details

Installation of *Bioconductor* and CRAN packages use R's standard functions for library management – install.packages(), available.packages(), update.packages(). Installation of GitHub packages uses the remotes::install_github().

When installing CRAN or *Bioconductor* packages, typical arguments include: lib.loc, passed to old.packages() and used to determine the library location of installed packages to be updated; and lib, passed to install.packages() to determine the library location where pkgs are to be installed.

repositories 5

When installing GitHub packages, ... is passed to the **remotes** package functions <code>install_github()</code> and <code>remotes:::install()</code>. A typical use is to build vignettes, via dependencies=TRUE, build_vignettes=TRUE. BIOCONDUCTOR_ONLINE_VERSION_DIAGNOSIS is an environment variable or global options() which, when set to FALSE, avoids the R and <code>Bioconductor</code> version checks that are done by querying an online configuration file. Setting <code>BIOCONDUCTOR_ONLINE_VERSION_DIAGNOSIS</code> to FALSE can speed package loading when internet access is slow or non-existent, but may result in out-of-date information about the current release and development versions of <code>Bioconductor</code>.

Value

```
BiocManager::install() returns the pkgs argument, invisibly.
```

See Also

```
BiocManager::repositories() returns the Bioconductor and CRAN repositories used by install(). install.packages() installs the packages themselves (used by BiocManager::install internally). update.packages() updates all installed packages (used by BiocManager::install internally). chooseBioCmirror() allows choice of a mirror from all public Bioconductor mirrors. chooseCRANmirror() allows choice of a mirror from all public CRAN mirrors.
```

Examples

```
## Not run:
## update previously installed packages
BiocManager::install()

## install Bioconductor packages, and prompt to update all
## installed packages
BiocManager::install(c("GenomicRanges", "edgeR"))

## install a CRAN and Bioconductor packages:
BiocManager::install(c("survival", "SummarizedExperiment"))

## install a package from source:
BiocManager::install("IRanges", type="source")

## End(Not run)
```

repositories

Display current Bioconductor and CRAN repositories.

Description

repositories() reports the URLs from which to install *Bioconductor* and CRAN packages. It is used by BiocManager::install() and other functions.

6 valid

Usage

```
repositories(site_repository = character(),
 version = BiocManager::version())
```

Arguments

site_repository

(Optional) character(1) representing an additional repository (e.g., a URL to an organization's internally maintained repository) in which to look for packages to install. This repository will be prepended to the default repositories returned

by the function.

version (Optional) character(1) or package_version indicating the *Bioconductor*

version (e.g., "3.8") for which repositories are required.

Value

Named character() of repositories.

See Also

```
BiocManager::install() Installs or updates Bioconductor, CRAN, and GitHub packages.
chooseBioCmirror() choose an alternative Bioconductor mirror; not usually necessary.
chooseCRANmirror() choose an alternative CRAN mirror; not usually necessary.
setRepositories() Select additional repositories for searching.
```

Examples

```
BiocManager::repositories()
## Not run:
BiocManager::repositories(version="3.8")
## End(Not run)
```

valid

Validate installed package versions against correct versions.

Description

Check that installed packages are consistent (neither out-of-date nor too new) with the version of R and Bioconductor in use.

valid 7

Usage

```
valid(pkgs = installed.packages(lib.loc, priority = priority),
  lib.loc = NULL, priority = "NA", type = getOption("pkgType"),
  filters = NULL, ..., checkBuilt = FALSE,
  site_repository = character())

## S3 method for class 'biocValid'
print(x, ...)
```

Arguments

pkgs	A character() vector of package names for checking, or a matrix as returned by installed.packages.	
lib.loc	A character() vector of library location(s) of packages to be validated; see installed.packages().	
priority	character(1) Check validity of all, "base", or "recommended" packages; see installed.packages().	
type	character(1) The type of available package (e.g., binary, source) to check validity against; see available.packages().	
filters	character(1) Filter available packages to check validity against; see available.packages().	
	Additional arguments, passed to BiocManager::install() when fix=TRUE.	
checkBuilt	logical(1). If TRUE a package built under an earlier major.minor version of R (e.g., 3.4) is considered to be old.	
site_repository		
	character(1). See ?install.	
Х	A 'biocValid' object returned by 'BiocManager::valid()'.	

Details

This function compares the version of installed packages to the version of packages associated with the version of R and Bioconductor currently in use.

```
Packages are reported as 'out-of-date' if a more recent version is available at the repositories specified by 'BiocManager::repositories()'. Usually, 'BiocManager::install()' is sufficient to update packages to their most recent version.

Packages are reported as 'too new' if the installed version is more recent than the most recent available in the 'BiocManager::repositories()'. It is possible to down-grade by re-installing a too new package "PkgA" with 'BiocManger::install("PkgA")'. It is important for the user to understand how their installation became too new, and to avoid this in the future.
```

8 version

Value

biocValid list object with elements too_new and out_of_date containing data.frames with packages and their installed locations that are too new or out-of-date for the current version of *Bioconductor*.

'print()' is invoked for its side effect.

Author(s)

Martin Morgan <martin.morgan@roswellpark.org>

See Also

```
BiocManager::install() to update installed packages.
```

Examples

```
BiocManager::valid()
```

version

Version of Bioconductor currently in use.

Description

version() reports the version of *Bioconductor* appropriate for this version of R, or the version of *Bioconductor* requested by the user.

Usage

```
version()
## S3 method for class 'version_sentinel'
print(x, ...)
```

Arguments

x An unknown_version instance used to represent the situation when the version of Bioconductor in use cannot be determined.

. . . Additional arguments, ignored.

Details

version() (and all functions requiring version information) fails when version cannot be validated e.g., because internet access is not available.

Value

A two-digit version, e.g., 3.8, of class package_version describing the version of *Bioconductor* in use.

version 9

Examples

BiocManager::version()

Index

```
*Topic environment
    install, 4
    repositories, 5
    valid, 6
_PACKAGE (BiocManager-pkg), 2
available, 2
available.packages, 7
BiocManager (BiocManager-pkg), 2
BiocManager-pkg, 2
BiocManager-pkg-package
        (BiocManager-pkg), 2
BIOCONDUCTOR_ONLINE_VERSION_DIAGNOSIS
        (install), 4
chooseBioCmirror, 5, 6
chooseCRANmirror, 5, 6
install, 4, 6-8
install.packages, 4, 5
install_github, 5
installed.packages, 7
old.packages, 4
print.biocValid(valid), 6
print.version_sentinel (version), 8
repositories, 4, 5, 5
setRepositories, 6
update.packages, 5
valid, 6
{\tt version}, {\color{red} 8}
```