

# Package ‘pacs’

February 27, 2023

**Title** Supplementary Tools for R Packages Developers

**Version** 0.4.10

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**Description** Supplementary utils for CRAN maintainers and R packages developers.

Validating the library, packages and lock files.

Exploring a complexity of a specific package like evaluating its size in bytes with all dependencies.

The shiny app complexity could be explored too.

Assessing the life duration of a specific package version.

Checking a CRAN package check page status for any errors and warnings.

Retrieving a DESCRIPTION or NAMESPACE file for any package version.

Comparing DESCRIPTION or NAMESPACE files between different package versions.

Getting a list of all releases for a specific package.

The Bioconductor is partly supported.

**License** GPL (>= 3)

**URL** <https://github.com/Polkas/pacs>, <https://polkas.github.io/pacs/>

**BugReports** <https://github.com/Polkas/pacs/issues>

**Encoding** UTF-8

**RoxygenNote** 7.2.1

**Depends** R (>= 3.5.0)

**Imports** curl, memoise, jsonlite, xml2, stringi

**Suggests** remotes, renv, withr, pkgsearch, mockery, testthat (>= 3.0.0), knitr, rmarkdown

**Config/testthat/edition** 3

**VignetteBuilder** knitr

**NeedsCompilation** no

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**Repository** CRAN

**Date/Publication** 2023-02-27 19:42:32 UTC

**R topics documented:**

app_deps	2
app_size	4
biocran_repos	5
bio_releases	5
checked_packages	6
compareVersionsMax	7
compareVersionsMin	7
cran_flavors	8
dir_size	8
lib_validate	9
lock_validate	11
match_flavors	13
pacs_base	13
pacs_lifeduration	14
pac_checkpage	15
pac_checkred	16
pac_compare_namespace	17
pac_compare_versions	18
pac_deps	19
pac_deps_dev	20
pac_deps_heavy	21
pac_deps_timemachine	22
pac_deps_user	23
pac_description	24
pac_health	25
pac_isin	27
pac_islast	28
pac_last	29
pac_lifeduration	29
pac_namespace	31
pac_size	32
pac_timemachine	32
pac_true_size	34
pac_validate	35
<b>Index</b>	<b>37</b>

---

app\_deps

*The shiny app dependencies*


---

**Description**

the shiny app dependencies packages are checked recursively. The c("Depends", "Imports", "LinkingTo") DESCRIPTION files fields are checked recursively. The required dependencies have to be installed in the local repository. The default arguments setup is recommended.

**Usage**

```
app_deps(
  path = ".",
  fields = c("Depends", "Imports", "LinkingTo"),
  lib.loc = .libPaths(),
  local = TRUE,
  base = FALSE,
  description_v = FALSE,
  recursive = TRUE,
  repos = biocran_repos()
)
```

**Arguments**

path	character path to the shiny app. Default: "."
fields	character vector listing the types of dependencies, a subset of c("Depends", "Imports", "LinkingTo", "Suggests", "Enhances"). Character string "all" is shorthand for that vector, character string "most" for the same vector without "Enhances", character string "strong" (default) for the first three elements of that vector. Default: c("Depends", "Imports", "LinkingTo")
lib.loc	character vector of search paths with local packages. Default: .libPaths()
local	logical if to use local repository (or newest remote packages). Default: TRUE
base	logical if to add base packages too. If TRUE then pacs::pacs_base() are taken into account. Default: FALSE
description_v	logical if the dependencies version should be taken from description files, minimal required. By default installed versions are taken. Default: FALSE
recursive	logical if to assess the dependencies recursively. Default: TRUE
repos	character vector base URLs of the repositories to use. By default checking CRAN and newest Bioconductor per R version. Default pacs::biocran_repos()

**Value**

character vector with dependency packages or data.frame when checking recursively.

**Note**

renv package has to be installed.

**Examples**

```
## Not run:
library(renv)
# Please update the path to the custom shiny app
app_path <- system.file("examples/04_mpg", package = "shiny")
pacs::app_deps(app_path)
pacs::app_deps(app_path, recursive = FALSE)

## End(Not run)
```

---

app_size	<i>Size of the shiny app</i>
----------	------------------------------

---

### Description

The size of shiny app is a sum of dependencies packages and the app directory. The app dependencies packages are checked recursively, and only in local repository. The default arguments setup is recommended.

### Usage

```
app_size(
  path = ".",
  fields = c("Depends", "Imports", "LinkingTo"),
  lib.loc = .libPaths(),
  recursive = TRUE
)
```

### Arguments

path	character path to the shiny app. Default: "."
fields	character vector listing the types of dependencies, a subset of c("Depends", "Imports", "LinkingTo", "Suggests", "Enhances"). Character string "all" is shorthand for that vector, character string "most" for the same vector without "Enhances", character string "strong" (default) for the first three elements of that vector. Default: c("Depends", "Imports", "LinkingTo")
lib.loc	character vector of search paths with local packages. Default: .libPaths()
recursive	logical if to assess the dependencies recursively. Default: TRUE

### Value

numeric size in bytes, to get MB ten divide by  $10^{*6}$ .

### Note

renv package has to be installed. base packages (pacs: : pacs\_base()) are not taken into account.

### Examples

```
## Not run:
library(renv)
# Please update the path to the shiny app
cat(pacs::app_size(system.file("examples/04_mpg", package = "shiny")) / 10**6, "MB")

## End(Not run)
```

---

biocran_repos	<i>CRAN and Bioconductor repositories</i>
---------------	---

---

**Description**

CRAN and Bioconductor repositories. The newest Bioconductor release for the specific R version is assumed.

**Usage**

```
biocran_repos(version = NULL)
```

**Arguments**

**version** character the Bioconductor release. By default the newest Bioconductor release for the specific R version is assumed, if not available only CRAN repository is returned. Available Bioconductor versions for your R version could be checked with `pacs::bio_releases()`. Default NULL

**Value**

named character vector of repositories.

**Note**

The Internet connection is needed to get Bioconductor repositories.

**Examples**

```
## Not run:  
pacs::biocran_repos()  
  
## End(Not run)
```

---

bio_releases	<i>Retrieving all Bioconductor releases</i>
--------------	---

---

**Description**

Retrieving all Bioconductor releases. The data is downloaded from <https://www.bioconductor.org/about/release-ann>

**Usage**

```
bio_releases()
```

**Value**

data.frame with the same structure as the html table on <https://www.bioconductor.org/about/release-announcements/>

**Note**

Results are cached for 30 minutes with memoise package.

**Examples**

```
## Not run:  
pacs::bio_releases()  
  
## End(Not run)
```

---

checked_packages	<i>Retrieving all R CRAN packages check pages statuses.</i>
------------------	---

---

**Description**

Retrieving all R CRAN packages check pages statuses. The data is downloaded from <https://cran.r-project.org/web/checks/>

**Usage**

```
checked_packages()
```

**Value**

data.frame with the same structure as the html table on [https://cran.r-project.org/web/checks/check\\_summary\\_by\\_](https://cran.r-project.org/web/checks/check_summary_by_)

**Note**

Results are cached for 30 minutes with memoise package. Some packages could be duplicated as not all tests are performed for a new version so two versions still coexists. Checks with asterisks (\*) indicate that checking was not fully performed, this is a case for less than 1% of all packages.

**Examples**

```
## Not run:  
pacs::checked_packages()  
  
## End(Not run)
```

---

compareVersionsMax      *Maximum version across the vector*

---

**Description**

Reduce function over the `utils::compareVersion`

**Usage**

```
compareVersionsMax(vec, na.rm = TRUE)
```

**Arguments**

`vec`                      character vector.  
`na.rm`                    logical if to remove NA values.

**Value**

character maximum version

**Examples**

```
compareVersionsMax(c("1.1.1", "0.2.0"))
```

---

compareVersionsMin      *Minimum version across the vector*

---

**Description**

Reduce function over the `utils::compareVersion`

**Usage**

```
compareVersionsMin(vec, na.rm = TRUE)
```

**Arguments**

`vec`                      character vector.  
`na.rm`                    logical if to remove NA values.

**Value**

character minimal version

**Examples**

```
compareVersionsMin(c("1.1.1", "0.2.0"))
```

---

cran_flavors	<i>Retrieving all R CRAN servers flavors</i>
--------------	--

---

**Description**

Retrieving all R CRAN servers flavors. The data is downloaded from [https://cran.r-project.org/web/checks/check\\_flavors.html](https://cran.r-project.org/web/checks/check_flavors.html)

**Usage**

```
cran_flavors()
```

**Value**

data.frame with the same structure as the html table on [https://cran.r-project.org/web/checks/check\\_flavors.html](https://cran.r-project.org/web/checks/check_flavors.html)

**Note**

Results are cached for 30 minutes with memoise package.

**Examples**

```
## Not run:
pacs::cran_flavors()

## End(Not run)
```

---

dir_size	<i>Size of the package</i>
----------	----------------------------

---

**Description**

size of package.

**Usage**

```
dir_size(path = ".", recursive = TRUE)
```

**Arguments**

path	character path to the shiny app. Default: "."
recursive	logical if to assess the dependencies recursively. Default: TRUE

**Value**

numeric size in bytes, to get MB ten divide by  $10^{*6}$ .



**Examples**

```
## Not run:
cat(pacs::dir_size(system.file(package = "stats")) / 10**6, "MB")

## End(Not run)
```

---

lib_validate	<i>Validate the local library</i>
--------------	-----------------------------------

---

**Description**

Checking if installed packages have correct versions taking into account all DESCRIPTION files requirements. Moreover identifying which packages are newest releases. Optionally we could add life duration and CRAN check page status for each package.

**Usage**

```
lib_validate(
  lib.loc = .libPaths(),
  fields = c("Depends", "Imports", "LinkingTo"),
  lifeduration = FALSE,
  checked = list(scope = character(0), flavors = NULL),
  built = FALSE,
  repos = biocran_repos()
)
```

**Arguments**

lib.loc	character vector of search paths with local packages. Default: <code>.libPaths()</code>
fields	character vector listing the types of dependencies, a subset of <code>c("Depends", "Imports", "LinkingTo", "Suggests", "Enhances")</code> . Character string "all" is shorthand for that vector, character string "most" for the same vector without "Enhances", character string "strong" (default) for the first three elements of that vector. Default: <code>c("Depends", "Imports", "LinkingTo")</code>
lifeduration	logical if to assess life duration for each package in the library. For installed newest releases of packages, a local evaluation is used. MEATCRAN CRANDB is used for libraries with less than 500 packages. Otherwise the direct web page download from CRAN is used. Default: 'FALSE'
checked	list with two named fields, scope and flavor. scope of R CRAN check pages statuses to consider, any of <code>c("ERROR", "FAIL", "WARN", "NOTE")</code> . flavor is a vector of CRAN machines to consider, which might be retrieved with <code>pacs::cran_flavors()\$flavor</code> . By default an empty scope field deactivated assessment for checked column, and NULL flavor will results in checking all machines. Default: <code>list(scope = character(0), flavor = NULL)</code>

<b>built</b>	logical if to add an R version under which each package was installed. Useful mainly for a local usage. Packages installed with a previous version of R could not work correctly with the new version of R. Default: FALSE
<b>repos</b>	character vector of repositories URLs to use. By default checking CRAN and newest Bioconductor per R version. Default <code>pac::biocran_repos()</code>

## Value

data.frame with 4/6/8/9/10 columns.

**Package** character a package name.

**Version.expected.min** character expected by DESCRIPTION files minimal version. "" means not specified so the newest version.

**Version.have** character installed package version.

**version\_status** numeric -1/0/1 which comes from `utils::compareVersion` function. 0 means that we have the same version as required by DESCRIPTION files. -1 means we have too low version installed, this is an error. 1 means we have higher version.

**built** character package was built under this R version

**built\_status** integer if the package was built under the current R version, then 1 (good) and for older R versions 0 (possibly bad). A package built under older R version or mix of packages built under different versions could bring possible failures.

**newest** logical (Internet needed) if the installed version is the newest one. For Bioconductor if is the newest one per R version.

**cran** logical (Internet needed) if the package is on CRAN, version is not taken into account here.

**checked** (Optional) (Internet needed) logical if the NEWEST package contains any specified statuses on CRAN check page. `pac::checked_packages` is used to quickly retrieve all statuses at once.

**lifeduration** (Optional) (Internet needed) integer number of days a package was released.

## Note

Version.expected.min column not count packages which are not a dependency for any package, so could not be find in DESCRIPTION files. When turn on the lifeduration options, calculations might be time consuming for libraries bigger than 500 packages. Results are cached for 30 minutes with memoise package. BioConductor packages are tested only in available scope, checked is not assessed for them. The crandb R packages database is a part of METACRAN project, source: Csárdi G, Salmon M (2022). pkgsearch: Search and Query CRAN R Packages. <https://github.com/r-hub/pkgsearch>, <https://r-hub.github.io/pkgsearch/>.

## Examples

```
## Not run:
pac::lib_validate()
pac::lib_validate(checkedred = list(scope = c("ERROR", "FAIL", "WARN")))
pac::lib_validate(checkedred = list(
  scope = c("ERROR", "FAIL"),
  flavors = pac::match_flavors()
```

```

))
# activate lifeduration argument, could be time consuming for bigger libraries.
pacs::lib_validate(
  lifeduration = TRUE,
  checkred = list(scope = c("ERROR", "FAIL"))
)
# only R CRAN repository
pacs::lib_validate(repos = "https://cran.rstudio.com/")

## End(Not run)

```

---

lock\_validate                      *Validate a specific renv lock file*

---

### Description

This function will be especially useful when renv lock file is built manually. Checking if packages in the lock file have correct versions taking into account their DESCRIPTION files requirements (c("Depends", "Imports", "LinkingTo")). Moreover identifying which packages are newest releases. Optionally we could add life duration and CRAN check page status for each dependency.

### Usage

```

lock_validate(
  path,
  lifeduration = FALSE,
  checkred = list(scope = character(0), flavors = NULL),
  lib.loc = .libPaths(),
  repos = biocran_repos()
)

```

### Arguments

path	character path to the shiny app. Default: "."
lifeduration	logical if to assess life duration for each package in the library. For installed newest releases of packages, a local evaluation is used. MEATCRAN CRANDB is used for libraries with less than 500 packages. Otherwise the direct web page download from CRAN is used. Default: 'FALSE'
checkred	list with two named fields, scope and flavor. scope of R CRAN check pages statuses to consider, any of c("ERROR", "FAIL", "WARN", "NOTE"). flavor is a vector of CRAN machines to consider, which might be retrieved with pacs::cran_flavors()\$flavor. By default an empty scope field deactivated assessment for checkred column, and NULL flavor will results in checking all machines. Default: list(scope = character(0), flavor = NULL)
lib.loc	character vector of search paths with local packages. Default: .libPaths()
repos	character vector of repositories URLs to use. By default checking CRAN and newest Bioconductor per R version. Default pacs::biocran_repos()

**Value**

data.frame with 2/6/7/8 columns.

**Package** character a package name.

**Version.expected.min** (conditional) (Internet needed) character expected by DESCRIPTION files minimal version. "" means not specified so the newest version.

**Version.expected** character package version in the renv lock file.

**version\_status** (conditional) numeric -1/0/1 which comes from `utils::compareVersion` function. 0 means that we have the same version as required by DESCRIPTION files. -1 means we have too low version installed, this is an error. 1 means we have higher version.

**newest** logical (Internet needed) if the installed version is the newest one.

**cran** logical (Internet needed) if the package is on CRAN, version is not taken into account here.

**checked** (Optional) (Internet needed) logical if the NEWEST package contains any specified statuses on CRAN check page.

**lifeduration** (Optional) (Internet needed) integer number of days a package was released.

**Note**

Version.expected.min column not count packages which are not a dependency for any package, so could not be find in DESCRIPTION files. Version.expected.min and version\_status are assessed only if there are less than 500 packages in the lock file. When turn on the lifeduration option, calculations might be time consuming when there is more than 500 packages. The cranR packages database is a part of METACRAN project, source: Csárdi G, Salmon M (2022). pkgsearch: Search and Query CRAN R Packages. <https://github.com/r-hub/pkgsearch>, <https://r-hub.github.io/pkgsearch/>

**Examples**

```
## Not run:
# path or url
url <- "https://raw.githubusercontent.com/Polkas/pacs/master/tests/testthat/files/renv_test.lock"
pacs::lock_validate(url)

pacs::lock_validate(
  url,
  checkred = list(scope = c("ERROR", "FAIL"), flavors = pacs::match_flavors())
)

pacs::lock_validate(
  url,
  lifeduration = TRUE,
  checkred = list(scope = c("ERROR", "FAIL"), flavors = NULL)
)

## End(Not run)
```

---

match_flavors	<i>Get all matched CRAN servers to the local OS</i>
---------------	---

---

**Description**

CRAN servers matched to the local OS.

**Usage**

```
match_flavors()
```

**Value**

character vector matched server names.

**Note**

The Internet connection is needed to use the function.

**Examples**

```
## Not run:  
pacs::match_flavors()  
  
## End(Not run)
```

---

pacs_base	<i>Get base R packages</i>
-----------	----------------------------

---

**Description**

get base packages, all or only startup.

**Usage**

```
pacs_base(startup = FALSE)
```

**Arguments**

startup           logical include only startup packages. Default: FALSE

**Value**

character vector

**Examples**

```
## Not run:
pacs_base()
pacs_base(startup = TRUE)

## End(Not run)
```

---

pacs\_lifeduration      *Packages life duration for a specific version*

---

**Description**

packages life duration for certain versions.

**Usage**

```
pacs_lifeduration(
  pacs,
  versions,
  source = c("crandb", "loop_crandb", "loop_cran"),
  lib.loc = .libPaths(),
  repos = biocran_repos()
)
```

**Arguments**

pacs	character vector of packages names.
versions	character vector of packages versions.
source	character one of c("crandb", "loop_crandb", "loop_cran"). The "crandb" works if less than <code>getOption("pacs.crandb_limit")</code> (currently 500) packages are looked for. Default: "crandb"
lib.loc	character vector of search paths with local packages. Default: <code>.libPaths()</code>
repos	character vector of repositories URLs to use. By default checking CRAN and newest Bioconductor per R version. Default <code>pacs::biocran_repos()</code>

**Value**

data.frame with two columns package name and life duration.

**Note**

Results are cached for 30 minutes with `memoise` package. The `crandb` R packages database is a part of METACRAN project, source: Csárdi G, Salmon M (2022). `pkgsearch`: Search and Query CRAN R Packages. <https://github.com/r-hub/pkgsearch>, <https://r-hub.github.io/pkgsearch/>. For `source = "loop_cran"` the function will scrap two CRAN URLs. Works only with CRAN packages. Please as a courtesy to the R CRAN, don't overload their servers by constantly using this function.

**Examples**

```
## Not run:
pacs::pacs_lifeduration(c("dplyr", "tidyr"), c("1.0.0", "1.2.0"))
pacs::pacs_lifeduration(c("dplyr", "tidyr"), c("1.0.0", "1.2.0"), source = "loop_cran")
# last versions
pacs::pacs_lifeduration(c("dplyr", "tidyr"), sapply(c("dplyr", "tidyr"), pacs::pac_last))

## End(Not run)
```

---

pac\_checkpage

*Retrieving the R CRAN package check page*

---

**Description**

Retrieving the R CRAN package check page.

**Usage**

```
pac_checkpage(pac)
```

**Arguments**

pac                    character a package name.

**Value**

data.frame.

**Note**

Results are cached for 30 minutes with memoise package. If you need to check many packages at once then is recommended usage of pacs::checked\_packages. Please as a courtesy to the R CRAN, don't overload their server by constantly using this function.

**Examples**

```
## Not run:
pacs::pac_checkpage("dplyr")

## End(Not run)
```

---

pac\_checked

*Checking the R CRAN package check page status*

---

## Description

using package R CRAN check page to validate if there are ANY errors and/or fails and/or warnings and/or notes.

## Usage

```
pac_checked(pac, scope = c("ERROR", "FAIL"), flavors = NULL)
```

## Arguments

pac	character a package name.
scope	character vector scope of the check, accepted values c("ERROR", "FAIL", "WARN", "NOTE"). Default: c("ERROR", "FAIL")
flavors	character vector of CRAN server names to consider, possible names could be get with <code>pac::cran_flavors()\$Flavor</code> . The <code>pac::match_flavors()</code> function could be used to get CRAN server names matched for your local OS. By default all CRAN machines are considered NULL value. Default: NULL

## Value

logical if the package fail under specified criteria.

## Note

Results are cached for 30 minutes with memoise package. If you need to check many packages at once then is recommended usage of `pac::checked_packages`. The used repository <https://cran.rstudio.com/>. Please as a courtesy to the R CRAN, don't overload their server by constantly using this function.

## Examples

```
## Not run:
pac::pac_checked("dplyr")
pac::pac_checked("dplyr", scope = c("ERROR"))
pac::pac_checked("dplyr",
  scope = c("ERROR", "FAIL", "WARN"),
  flavors = pac::match_flavors()
)

## End(Not run)
```



---

pac\_compare\_namespace *Compare NAMESPACE exports between specific CRAN packages versions*

---

## Description

using the remote github CRAN mirror to compare NAMESPACE exports between specific packages versions.

## Usage

```
pac_compare_namespace(  
  pac,  
  old = NULL,  
  new = NULL,  
  lib.loc = .libPaths(),  
  repos = "https://cran.rstudio.com/"  
)
```

## Arguments

pac	character a package name.
old	character an old version of package, default local version. Default: NULL
new	character a new version of package, default newest version. Default: NULL
lib.loc	character vector of search paths with local packages. Default: .libPaths()
repos	character vector repositories URLs to use. Used only for the validation. Default https://cran.rstudio.com/

## Value

list with c("imports", "exports", "exportPatterns", "importClasses", "importMethods", "exportClasses", "exportMethods", "exportClassPatterns", "dynlibs", "S3methods") slots, and added and removed ones for each of them.

## Examples

```
## Not run:  
pacs::pac_compare_namespace("shiny", "1.0.0", "1.6.0")  
pacs::pac_compare_namespace("shiny", "1.0.0", "1.6.0")$exports  
# local version to newest one  
pacs::pac_compare_namespace("shiny")  
  
## End(Not run)
```

---

pac\_compare\_versions *Compare DESCRIPTION files dependencies between specific CRAN packages versions*

---

### Description

using the remote github CRAN mirror to compare DESCRIPTION files dependencies between specific packages versions.

### Usage

```
pac_compare_versions(
  pac,
  old = NULL,
  new = NULL,
  fields = c("Imports", "Depends", "LinkingTo"),
  lib.loc = .libPaths(),
  repos = "https://cran.rstudio.com/"
)
```

### Arguments

pac	character a package name.
old	character an old version of package, default local version. Default: NULL
new	character a new version of package, default newest version. Default: NULL
fields	character vector listing the types of dependencies, a subset of c("Depends", "Imports", "LinkingTo", "Suggests", "Enhances"). Character string "all" is shorthand for that vector, character string "most" for the same vector without "Enhances", character string "strong" (default) for the first three elements of that vector. Default: c("Depends", "Imports", "LinkingTo")
lib.loc	character vector of search paths with local packages. Default: .libPaths()
repos	character vector repositories URLs to use. Used only for the validation. Default https://cran.rstudio.com/

### Value

data.frame with 4 columns.

**Package** character package names.

**Version.OLD** character versions of dependencies required by an old package version.

**Version.NEW** character versions of dependencies required by a new package version.

**version\_status** numeric -1/0/1 which comes from `utils::compareVersion` function. 0 means that both versions have the same requirement. -1 means that the new version remove this requirement. 1 means that the new version added a new requirement.

**Examples**

```
## Not run:
pacs::pac_compare_versions("memoise", "0.2.1", "2.0.0")
pacs::pac_compare_versions("memoise", "0.2.1")
# local version to newest one
pacs::pac_compare_versions("memoise")

## End(Not run)
```

pac\_deps

*Package dependencies***Description**

Package dependencies from DESCRIPTION files with installed or expected versions or newest released.

**Usage**

```
pac_deps(
  pac,
  fields = c("Depends", "Imports", "LinkingTo"),
  lib.loc = .libPaths(),
  base = FALSE,
  local = TRUE,
  description_v = FALSE,
  attr = TRUE,
  recursive = TRUE,
  repos = biocran_repos()
)
```

**Arguments**

pac	character a package name.
fields	character vector listing the types of dependencies, a subset of c("Depends", "Imports", "LinkingTo", "Suggests", "Enhances"). Character string "all" is shorthand for that vector, character string "most" for the same vector without "Enhances", character string "strong" (default) for the first three elements of that vector. Default: c("Depends", "Imports", "LinkingTo")
lib.loc	character vector of search paths with local packages. Default: .libPaths()
base	logical if to add base packages too. If TRUE then pacs::pacs_base() are taken into account. Default: FALSE
local	logical if to use local repository (or newest remote packages). Default: TRUE
description_v	logical if the dependencies version should be taken from description files, minimal required. By default installed versions are taken. Default: FALSE

attr	logical if a package and its version should be added as an attribute of data.frame or for FALSE as an additional record. Default: TRUE
recursive	logical if to assess the dependencies recursively. Default: TRUE
repos	character vector of repositories URLs to use. By default checking CRAN and newest Bioconductor per R version. Default pacs::biocran_repos()

**Value**

data.frame with packages and their versions. Versions are taken from installed.packages or newest released.

**Note**

When function is invoked in the loop afterwards results could be aggregated like, stats::aggregate(results[, c("Version"), drop = FALSE], list(Package = results\$Package), pacs::compareVersionsMax).

**Examples**

```
## Not run:
pacs::pac_deps("stats", base = TRUE)$Package
pacs::pac_deps("memoise")$Package
pacs::pac_deps("memoise", description_v = FALSE)
# raw dependencies from DESCRIPTION file
pacs::pac_deps("memoise", description_v = TRUE, recursive = FALSE)
# raw dependencies from DESCRIPTION file - last release
pacs::pac_deps("memoise", description_v = TRUE, local = FALSE, recursive = FALSE)

## End(Not run)
```

---

pac\_deps\_dev

*Package dependencies - developer perspective*

---

**Description**

A higher-level function, build from pacs::pacs\_deps. Package dependencies installed when e.g. R CMD check a package. "Depends", "Imports", "LinkingTo", "Suggests" fields from the DESCRIPTION file and their recursive dependencies taken from "Depends", "Imports", "LinkingTo" fields. Dependencies are taken remotely for the newest version.

**Usage**

```
pac_deps_dev(
  pac,
  base = FALSE,
  local = FALSE,
  attr = TRUE,
  repos = pacs::biocran_repos()
)
```

**Arguments**

pac	character a package name.
base	logical if to add base packages too. If TRUE then pacs::pacs_base() are taken into account. Default: FALSE
local	logical if to use local repository (or newest remote packages). Default: FALSE
attr	logical if a package and its version should be added as an attribute of data.frame or for FALSE as an additional record. Default: TRUE
repos	character vector of repositories URLs to use. By default checking CRAN and newest Bioconductor per R version. Default pacs::biocran_repos()

**Value**

data.frame with packages and their versions. Versions are taken from installed.packages or newest released.

**Examples**

```
## Not run:
pacs::pac_deps_dev("dplyr")
pacs::pac_deps_dev("pacs")
# with the main package in the list
pacs::pac_deps_dev("pacs", attr = FALSE)

## End(Not run)
```

---

pac_deps_heavy	<i>Package direct dependencies and number of dependencies for each of them</i>
----------------	--

---

**Description**

A higher-level function, build from pacs::pacs\_deps and tools::package\_dependencies. A tool to identify a main sources of dependencies, which direct dependencies are the heaviest one.

**Usage**

```
pac_deps_heavy(
  pac,
  fields = c("Depends", "Imports", "LinkingTo"),
  lib.loc = .libPaths(),
  base = FALSE,
  local = FALSE,
  repos = pacs::biocran_repos()
)
```

**Arguments**

pac	character a package name.
fields	character vector listing the types of dependencies, a subset of c("Depends", "Imports", "LinkingTo", "Suggests", "Enhances"). Character string "all" is shorthand for that vector, character string "most" for the same vector without "Enhances", character string "strong" (default) for the first three elements of that vector. Default: c("Depends", "Imports", "LinkingTo")
lib.loc	character vector of search paths with local packages. Default: .libPaths()
base	logical if to add base packages too. If TRUE then pacs::pacs_base() are taken into account. Default: FALSE
local	logical if to use local repository (or newest remote packages). Default: FALSE
repos	character vector of repositories URLs to use. By default checking CRAN and newest Bioconductor per R version. Default pacs::biocran_repos()

**Value**

data.frame with three columns c("Package", "NrDeps", "NrUniqueDeps"): package name, number of dependencies and number of unique dependencies (not shared by other direct dependencies).

**Note**

Please take into account that the sum of the dependencies is not equal to the number of dependencies of the main package, because some dependencies are overlapping.

**Examples**

```
## Not run:
pacs::pac_deps_heavy("caret")
pacs::pac_deps_heavy("dplyr")

## End(Not run)
```

---

pac\_deps\_timemachine *R CRAN package dependencies for a certain version or time point*

---

**Description**

Package dependencies from DESCRIPTION files retrieved recursively for certain version or time point.

**Usage**

```
pac_deps_timemachine(
  pac,
  version = NULL,
  at = NULL,
  fields = c("Depends", "Imports", "LinkingTo"),
  recursive = TRUE
)
```

**Arguments**

pac	character a package name.
version	character version of a package. Default: NULL
at	Date from which to take the version. Default: NULL
fields	character vector listing the types of dependencies, a subset of c("Depends", "Imports", "LinkingTo", "Suggests", "Enhances"). Character string "all" is shorthand for that vector, character string "most" for the same vector without "Enhances", character string "strong" (default) for the first three elements of that vector. Default: c("Depends", "Imports", "LinkingTo")
recursive	logical if to assess the dependencies recursively. Default: TRUE

**Value**

named vector package dependencies and their versions at the release date of main package plus one day.

**Note**

Longer lived version is taken if 2 is available at the same date (switch time).

**Examples**

```
## Not run:
pacs::pac_deps_timemachine("memoise", "0.2.1")
pacs::pac_deps_timemachine("memoise", at = as.Date("2019-01-01"))
pacs::pac_deps_timemachine("dplyr", at = as.Date("2015-01-01"))

## End(Not run)
```

---

pac\_deps\_user

*Package dependencies - user perspective*

---

**Description**

A higher-level function, build from `pacs::pacs_deps`. Package dependencies installed when run `installed.packages`. "Depends", "Imports", "LinkingTo" fields from the DESCRIPTION file and their recursive dependencies taken from the same fields. Dependencies are taken remotely for the newest version.

**Usage**

```
pac_deps_user(
  pac,
  base = FALSE,
  local = FALSE,
  attr = TRUE,
  repos = pacs::biocran_repos()
)
```

**Arguments**

pac	character a package name.
base	logical if to add base packages too. If TRUE then pacs::pacs_base() are taken into account. Default: FALSE
local	logical if to use local repository (or newest remote packages). Default: FALSE
attr	logical if a package and its version should be added as an attribute of data.frame or for FALSE as an additional record. Default: TRUE
repos	character vector of repositories URLs to use. By default checking CRAN and newest Bioconductor per R version. Default pacs::biocran_repos()

**Value**

data.frame with packages and their versions. Versions are taken from installed.packages or newest released.

**Examples**

```
## Not run:
pacs::pac_deps_user("dplyr")
pacs::pac_deps_user("pacs")
# with the main package in the list
pacs::pac_deps_user("pacs", attr = FALSE)

## End(Not run)
```

---

pac\_description      *package DESCRIPTION file*

---

**Description**

CRAN package DESCRIPTION file taken locally or remotely from GITHUB CRAN mirror or CRAN website. By default works for the newest package version.



**Usage**

```
pac_description(  
  pac,  
  version = NULL,  
  at = NULL,  
  local = FALSE,  
  lib.loc = .libPaths(),  
  repos = "https://cran.rstudio.com/"  
)
```

**Arguments**

pac	character a package name.
version	character version of a package. Default: NULL
at	Date from which to take the version. Default: NULL
local	logical if to use local repository (or newest remote packages). Default: FALSE
lib.loc	character vector of search paths with local packages. Default: .libPaths()
repos	character vector repositories URLs to use. Used only for the validation. Default https://cran.rstudio.com/

**Value**

list with names proper for DESCRIPTION file fields.

**Note**

Results are cached for 30 minutes with memoise package.

**Examples**

```
## Not run:  
pacs::pac_description("dplyr", version = "0.8.0")  
pacs::pac_description("dplyr", at = as.Date("2019-02-01"))  
  
## End(Not run)
```

---

pac\_health

*CRAN package health state at a specific Date or for a specific version*

---

**Description**

a package health for a certain version or at a specific Date. By default works for the newest package version. A healthy package was published for more than x days, where default is 14 days. CRAN team gives around one/two week to resolved a package which gave errors under the check page. The newest release is checked for any warnings/errors on the R CRAN package check page.

**Usage**

```
pac_health(
  pac,
  version = NULL,
  at = NULL,
  limit = 14,
  scope = c("ERROR", "FAIL"),
  flavors = NULL,
  lib.loc = .libPaths(),
  repos = "https://cran.rstudio.com/",
  source = c("crandb", "cran")
)
```

**Arguments**

pac	character a package name.
version	character version of a package. Default: NULL
at	Date from which to take the version. Default: NULL
limit	numeric at least days to treat as healthy, " $\geq$ limit". Default: 14
scope	character vector scope of the check, accepted values c("ERROR", "FAIL", "WARN", "NOTE"). Default: c("ERROR", "FAIL")
flavors	character vector of CRAN server names to consider, possible names could be get with <code>pac::cran_flavors()\$Flavor</code> . The <code>pac::match_flavors()</code> function could be used to get CRAN server names matched for your local OS. By default all CRAN machines are considered NULL value. Default: NULL
lib.loc	character vector of search paths with local packages. Default: <code>.libPaths()</code>
repos	character vector repositories URLs to use. Default <code>https://cran.rstudio.com/</code>
source	character one of c("crandb", "cran"). Using the MEATCRAN CRANDB or the direct web page download from CRAN. Default: "crandb"

**Value**

logical if a package is healthy.

**Note**

Results are cached for 30 minutes with memoise package. The crandb R packages database is a part of METACRAN project, source: Csárdi G, Salmon M (2022). pkgsearch: Search and Query CRAN R Packages. <https://github.com/r-hub/pkgsearch>, <https://r-hub.github.io/pkgsearch/>. For source = "cran" the function will scrap two CRAN URLs. Works only with CRAN packages. Please as a courtesy to the R CRAN, don't overload their servers by constantly using this function.

**Examples**

```
## Not run:
pac::pac_health("memoise")
pac::pac_health("dplyr", version = "0.8.0", limit = 14)
```

```
pac::pac_health("dplyr", at = as.Date("2019-02-14"))
pac::pac_health("dplyr", limit = 14, scope = c("ERROR", "FAIL"))

## End(Not run)
```

---

pac\_isin

*Checking if a package is in repositories*

---

### Description

using `utils::available.packages` to check if package is in repositories.

### Usage

```
pac_isin(pac, repos = biocran_repos())
```

### Arguments

pac	character a package name.
repos	character vector base URLs of the repositories to use. By default checking CRAN and newest Bioconductor per R version. Default <code>pac::biocran_repos()</code>

### Value

logical if a package is inside repositories.

### Note

Results are cached for 30 minutes with `memoise` package.

### Examples

```
## Not run:
pac_isin("dplyr")
pac_isin("dplyr", repos = "https://cran.rstudio.com/")
pac_isin("dplyr", repos = biocran_repos()[grep("Bio", names(biocran_repos()))])

## End(Not run)
```

---

pac_islast	<i>Checking if a package version is the most recent one</i>
------------	---

---

## Description

checking if a package version is the most recent one, by default the installed version is compared.

## Usage

```
pac_islast(pac, version = NULL, lib.loc = .libPaths(), repos = biocran_repos())
```

## Arguments

pac	character a package name.
version	character version of a package. Default: NULL
lib.loc	character vector of search paths with local packages. Default: .libPaths()
repos	character vector of repositories URLs to use. By default checking CRAN and newest Bioconductor per R version. Default pacs::biocran_repos()

## Value

logical if a package is inside repositories.

## Note

Results are cached for 30 minutes with memoise package.

## Examples

```
## Not run:
pac_islast("memoise")
pac_islast("dplyr", version = "1.0.0")
pac_islast("S4Vectors")
pac_islast("S4Vectors", version = pac_last("S4Vectors"))

## End(Not run)
```

---

pac\_last                      *Getting the most recent package version*

---

**Description**

using `utils::available.packages` to get the newest package version.

**Usage**

```
pac_last(pac, repos = biocran_repos())
```

**Arguments**

`pac`                      character a package name.  
`repos`                    character vector of repositories URLs to use. By default checking CRAN and newest Bioconductor per R version. Default `pac::biocran_repos()`

**Value**

character most recent package version.

**Note**

Results are cached for 30 minutes with `memoise` package. For Bioconductor the newest one per R version.

**Examples**

```
## Not run:  
pac_last("dplyr")  
pac_last("S4Vectors")  
  
## End(Not run)
```

---

pac\_lifeduration              *Package version life duration at specific Date or for a specific version*

---

**Description**

a package life duration for a certain version or at a specific Date. By default works for the newest package version.

**Usage**

```
pac_lifeduration(
  pac,
  version = NULL,
  at = NULL,
  lib.loc = .libPaths(),
  repos = biocran_repos(),
  source = c("crandb", "cran")
)
```

**Arguments**

pac	character a package name.
version	character version of a package. Default: NULL
at	Date from which to take the version. Default: NULL
lib.loc	character vector of search paths with local packages. Default: .libPaths()
repos	character vector of repositories URLs to use. By default checking CRAN and newest Bioconductor per R version. Default pacs::biocran_repos()
source	character one of c("crandb", "cran"). Using the MEATCRAN CRANDB or the direct web page download from CRAN. Default: "crandb"

**Value**

difftime, number of days package version was the newest one.

**Note**

Results are cached for 30 minutes with memoise package. The crandb R packages database is a part of METACRAN project, source: Csárdi G, Salmon M (2022). pkgsearch: Search and Query CRAN R Packages. <https://github.com/r-hub/pkgsearch>, <https://r-hub.github.io/pkgsearch/>. For source = "cran" the function will scrap two CRAN URLs. Works only with CRAN packages. Please as a courtesy to the R CRAN, don't overload their servers by constantly using this function.

**Examples**

```
## Not run:
pacs::pac_lifeduration("memoise")
pacs::pac_lifeduration("memoise", source = "cran")
pacs::pac_lifeduration("dplyr", version = "0.8.0")
pacs::pac_lifeduration("dplyr", at = as.Date("2019-02-14"))
# For Bioconductor packages it will work only for the newest per R version and installed ones.
pacs::pac_lifeduration("S4Vectors")

## End(Not run)
```

---

pac_namespace	<i>package NAMESPACE file</i>
---------------	-------------------------------

---

### Description

CRAN package NAMESPACE file taken locally or remotely from GITHUB CRAN mirror or CRAN website. By default works for the newest package version.

### Usage

```
pac_namespace(  
  pac,  
  version = NULL,  
  at = NULL,  
  local = FALSE,  
  lib.loc = .libPaths(),  
  repos = "https://cran.rstudio.com/"  
)
```

### Arguments

pac	character a package name.
version	character version of a package. Default: NULL
at	Date from which to take the version. Default: NULL
local	logical if to use local repository (or newest remote packages). Default: FALSE
lib.loc	character vector of search paths with local packages. Default: .libPaths()
repos	character vector repositories URLs to use. Used only for the validation. Default https://cran.rstudio.com/

### Value

list with names proper for NAMESPACE file, the same as format as returned by `base::parseNamespaceFile`.

### Note

Results are cached for 30 minutes with `memoise` package. This function is mainly built under source code from `base::parseNamespaceFile`.

### Examples

```
## Not run:  
pacs::pac_namespace("dplyr", version = "0.8.0")  
pacs::pac_namespace("dplyr", at = as.Date("2019-02-01"))  
pacs::pac_namespace("memoise", local = TRUE)  
  
## End(Not run)
```

---

pac_size	<i>Size of the package</i>
----------	----------------------------

---

**Description**

Size of package.

**Usage**

```
pac_size(pac, lib.loc = .libPaths())
```

**Arguments**

pac	character a package name.
lib.loc	character vector of search paths with local packages. Default: .libPaths()

**Value**

numeric size in bytes, to get MB ten divide by  $10^{**6}$ .

**Examples**

```
## Not run:
cat(pacs::pac_size("stats") / 10**6, "MB")

## End(Not run)
```

---

pac_timemachine	<i>Package metadata for all releases</i>
-----------------	--

---

**Description**

Using CRAN website to get a package metadata used at a specific Date or a Date interval or for specific version.

**Usage**

```
pac_timemachine(
  pac,
  at = NULL,
  from = NULL,
  to = NULL,
  version = NULL,
  source = c("crandb", "cran")
)
```



**Arguments**

pac	character a package name.
at	Date from which to take the version. Default: NULL
from	Date the lower limit. Default: NULL
to	Date the upper limit. Default: NULL
version	character version of a package. Default: NULL
source	character one of c("crandb", "cran"). Using the MEATCRAN CRANDB or the direct web page download from CRAN. Default: "crandb"

**Value**

data.frame with 7 columns

**Package** character package name.

**Version** character package version.

**Released** character release Date

**Archived** character archived Date.

**LifeDuration** difftime number of days the version was the newest one.

**URL** character the suffix of the base URL to tar.gz file. The base part of URL in the result is <https://cran.r-project.org/src/contrib/>.

**Size** character size of the tar.gz file.

**Note**

Results are cached for 30 minutes with memoise package. The crandb R packages database is a part of METACRAN project, source: Csárdi G, Salmon M (2022). pkgsearch: Search and Query CRAN R Packages. <https://github.com/r-hub/pkgsearch>, <https://r-hub.github.io/pkgsearch/>. For source = "cran" the function will scrap two CRAN URLs. Works only with CRAN packages. Please as a courtesy to the R CRAN, don't overload their servers by constantly using this function.

**Examples**

```
## Not run:
pacs::pac_timemachine("dplyr")
pacs::pac_timemachine("dplyr", at = as.Date("2017-02-02"))
pacs::pac_timemachine("dplyr", from = as.Date("2017-02-02"), to = as.Date("2018-04-02"))
pacs::pac_timemachine("dplyr", at = Sys.Date())
pacs::pac_timemachine("tidyr", from = as.Date("2020-06-01"), to = Sys.Date())

## End(Not run)
```

---

pac_true_size	<i>True size of the package</i>
---------------	---------------------------------

---

**Description**

True size of the package as it takes into account its all dependencies, recursively.

**Usage**

```
pac_true_size(
  pac,
  fields = c("Depends", "Imports", "LinkingTo"),
  lib.loc = .libPaths(),
  exclude_joint = 0L
)
```

**Arguments**

pac	character a package name.
fields	character vector listing the types of dependencies, a subset of c("Depends", "Imports", "LinkingTo", "Suggests", "Enhances"). Character string "all" is shorthand for that vector, character string "most" for the same vector without "Enhances", character string "strong" (default) for the first three elements of that vector. Default: c("Depends", "Imports", "LinkingTo")
lib.loc	character vector of search paths with local packages. Default: .libPaths()
exclude_joint	integer exclude packages which are dependencies of at least N other packages, not count main package dependencies. Default: 0

**Value**

numeric size in bytes, to get MB then divide by 10\*\*6.

**Note**

R base packages are not counted. The default value of fields should be suited for almost all scenarios.

**Examples**

```
## Not run:
# size in MB, with all its dependencies
pacs::pac_true_size("memoise") / 10**6

## End(Not run)
```

---

pac_validate	<i>Validate a specific local package</i>
--------------	--

---

### Description

Checking if installed package dependencies have correct versions taking into account their DESCRIPTION files requirements. Moreover identifying which packages are newest releases. Optionally we could add life duration and CRAN check page status for each dependency.

### Usage

```
pac_validate(
  pac,
  lib.loc = .libPaths(),
  fields = c("Depends", "Imports", "LinkingTo"),
  lifeduration = FALSE,
  checkedred = list(scope = character(0), flavors = NULL),
  repos = biocran_repos()
)
```

### Arguments

pac	character a package name.
lib.loc	character vector of search paths with local packages. Default: <code>.libPaths()</code>
fields	character vector listing the types of dependencies, a subset of <code>c("Depends", "Imports", "LinkingTo", "Suggests", "Enhances")</code> . Character string "all" is shorthand for that vector, character string "most" for the same vector without "Enhances", character string "strong" (default) for the first three elements of that vector. Default: <code>c("Depends", "Imports", "LinkingTo")</code>
lifeduration	logical if to assess life duration for each package in the library. For installed newest releases of packages, a local evaluation is used. MEATCRAN CRANDB is used for libraries with less than 500 packages. Otherwise the direct web page download from CRAN is used. Default: <code>'FALSE'</code>
checkedred	list with two named fields, scope and flavor. scope of R CRAN check pages statuses to consider, any of <code>c("ERROR", "FAIL", "WARN", "NOTE")</code> . flavor is a vector of CRAN machines to consider, which might be retrieved with <code>pac:::cran_flavors()\$Flavor</code> . By default an empty scope field deactivated assessment for checkedred column, and NULL flavor will results in checking all machines. Default: <code>list(scope = character(0), flavor = NULL)</code>
repos	character vector of repositories URLs to use. By default checking CRAN and newest Bioconductor per R version. Default <code>pac:::biocran_repos()</code>

### Value

data.frame with 5/7/8/9 columns.

**Package** character a package name.

**Version.expected.min** character expected by DESCRIPTION files minimal version. "" means not specified so the newest version.

**Version.have** character installed package version.

**version\_status** numeric -1/0/1 which comes from `utils::compareVersion` function. 0 means that we have the same version as required by DESCRIPTION files. -1 means we have too low version installed, this is an error. 1 means we have higher version.

**direct** logical if the package is in the first dependency layer, direct dependencies from DESCRIPTION file.

**newest** logical (Internet needed) if the installed version is the newest one.

**cran** logical (Internet needed) if the package is on CRAN, version is not taken into account here.

**checkred** (Optional) (Internet needed) logical if the NEWEST package contains any specified statuses on CRAN check page.

**lifeduration** (Optional) (Internet needed) integer number of days a package was released.

## Note

Version.expected.min column not count packages which are not a dependency for any package, so could not be find in DESCRIPTION files. When turn on the lifeduration option, calculations might be time consuming when there is more than 500 packages. Please as a courtesy to the R CRAN, don't overload their server by constantly using this function with lifeduration or checkred turned on. Results are cached with memoise package, memory cache. The cranR database is a part of METACRAN project, source: Csárdi G, Salmon M (2022). pkgsearch: Search and Query CRAN R Packages. <https://github.com/r-hub/pkgsearch>, <https://r-hub.github.io/pkgsearch/>

## Examples

```
## Not run:
pacs::pac_validate("memoise")
pacs::pac_validate(
  "memoise",
  lifeduration = TRUE,
  checkred = list(scope = c("ERROR", "FAIL"), flavors = NULL)
)
pacs::pac_validate(
  "memoise",
  lifeduration = TRUE,
  checkred = list(scope = c("ERROR", "FAIL"), flavors = pacs::match_flavors())
)

## End(Not run)
```

# Index

app\_deps, 2  
app\_size, 4

bio\_releases, 5  
biocran\_repos, 5

checked\_packages, 6  
compareVersionsMax, 7  
compareVersionsMin, 7  
cran\_flavors, 8

dir\_size, 8

lib\_validate, 9  
lock\_validate, 11

match\_flavors, 13

pac\_checkpage, 15  
pac\_checkred, 16  
pac\_compare\_namespace, 17  
pac\_compare\_versions, 18  
pac\_deps, 19  
pac\_deps\_dev, 20  
pac\_deps\_heavy, 21  
pac\_deps\_timemachine, 22  
pac\_deps\_user, 23  
pac\_description, 24  
pac\_health, 25  
pac\_isin, 27  
pac\_islast, 28  
pac\_last, 29  
pac\_lifeduration, 29  
pac\_namespace, 31  
pac\_size, 32  
pac\_timemachine, 32  
pac\_true\_size, 34  
pac\_validate, 35  
pacs\_base, 13  
pacs\_lifeduration, 14