

Authoring R Packages

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Source of information

The complete documentation for managing and authoring of R packages is the **Writing R Extensions** manual.

http:

`//cran.r-project.org/doc/manuals/R-exts.html`

Guidelines for authoring Bioconductor packages can be found here:

`http://wiki.fhcrc.org/bioc/Package_Guidelines`

What is a Package?

- collection of functions and data structures addressing a **particular problem**
- structure to **organize** code and its documentation
- vehicle to **distribute** and share software

Purpose of a Package

- keep all pieces of the software together
- common installation process
- allow for shared development (*versions*)
- unified way to access documentation
- allow for quality control and testing

R Functions for Package Management

- `search()`

```
[1] ".GlobalEnv"          "package:stats"
[3] "package:graphics"   "package:grDevices"
[5] "package:utils"      "package:datasets"
[7] "package:methods"    "Autoloads"
[9] "package:base"
```

- `.packages()`

```
[1] "stats"      "graphics"  "grDevices" "utils"
[5] "datasets"  "methods"   "base"
```

- `system.file(package="tools")`

```
[1] "/home/fhahne/R/bin/R-2-7-0/lib/R/library/tools"
```

R Functions for Package Management

- `packageDescription("base")`

```
Package: base
```

```
Version: 2.7.0
```

```
Priority: base
```

```
Title: The R Base Package
```

```
Author: R Development Core Team and contributors  
worldwide
```

```
Maintainer: R Core Team <R-core@r-project.org>
```

```
Description: Base R functions
```

```
License: GPL (>= 2)
```

```
Built: R 2.7.0; ; Mi 6. Feb 18:04:55 PST 2008; unix
```

```
-- File: /home/fhahne/R/bin/R-2-7-0/lib/R/library/ba
```

R Functions for Package Management

- `download.packages()`
- `install.packages()`
- `update.packages()`
- `remove.packages()`
- `available.packages()`
- `old.packages()`
- `new.packages()`
- `installed.packages()`

Libraries

A library stores a **collection of packages**

- multiple libraries (`R_LIBS`)

```
> .Library
```

```
[1] "/home/fhahne/R/bin/R-2-7-0/lib/R/library"
```

```
> .libPaths()
```

```
[1] "/home/fhahne/R/source/R-2-7-0/localPackages"
```

```
[2] "/home/fhahne/R/bin/R-2-7-0/lib/R/library"
```

- version specific libraries (`R_LIBS_USER`)
- site specific libraries (`R_LIBS_SITE`)

Folder Structure

On the system level a package is a **collection of folders**

- `./R`: all native R code for the package
- `./man`: the documentation for all (exported) code items
- `./src`: foreign code
- `./data`: data sets that can be loaded using `data()`
- `./inst/doc`: additional documentation (*Vignettes*)
- (distribution specific subdirectories `windows` and `unix` for R and `man`)

DESCRIPTION File

The `DESCRIPTION` file contains the basic information about a package and settings for package installation and loading

- **mandatory fields** `Package`, `Version`, `License`, `Description`, `Title`, `Author`, `Maintainer`
- **fields stating dependencies:** `Depends`, `Suggests`, `Imports`
- **installation and loading settings:** `Collate`, `LazyLoad`, `LazyData`
- `biocViews`: terms in a controlled vocabulary to characterize the package

```
Package: Biobase
Title: Biobase: Base functions for Bioconductor
Version: 1.17.8
Author: R. Gentleman, V. Carey, M. Morgan, S. Falcon
Description: Functions that are needed by many other packages or which replace R functions.
Suggests: tkWidgets, ALL
Depends: R (>= 2.6.0), tools, methods, utils
Maintainer: Biocore Team c/o BioC user list <bioconductor@stat.math.ethz.ch>
License: The Artistic License, Version 2.0
Collate: tools.R strings.R environment.R vignettes.R packages.R
         AllGeneric.R
         VersionsClass.R
         VersionedClasses.R methods-VersionedClass.R methods-VersionedNull.R
         DataClasses.R methods-aggregator.R methods-ANY.R
         methods-container.R methods-data.frame.R methods-phenoData.R
         methods-MIAME.R methods-annotatedDataset.R
         methods-AssayData.R methods-AnnotatedDataFrame.R
         methods-eSet.R methods-ExpressionSet.R methods-MultiSet.R
         methods-SnpSet.R methods-NChannelSet.R
         anyMissing.R
         methods-exprSet.R rowOp-methods.R updateObject.R updateObjectTo.R
         methods-ScalarObject.R
         zzz.R
LazyLoad: yes
biocViews: Infrastructure, Statistics
```

Tools to Create Package Structure

- `package.skeleton()` helps to build a basic package structure and templates for the documentation
- all objects in the working directory are added to the package
- this is just a **template**; there remains a lot for the user to do...

Code Organisation

Organisation of files in the `R` directory is up to the user, there are some **guidelines**:

- all class definitions in one file
- all generic function definitions in one file
- functions including all necessary helper functions into separate files or organize according to conceptual entities

Building Packages

There are a number of command line tools to build R packages

- R CMD `build`: **create .tar.gz** from package source
- R CMD `check`: **check documentation, code and run all examples and Vignettes**
- R CMD `INSTALL`: **install a package .tar.gz**

Help Pages

documentation of functions in the **help pages**, of concepts and work flows in **Vignettes**.

help pages:

- show capabilities, inputs and outputs
- quality control: examples are run during `R CMD check`

prompt Family

functions of the `prompt` family can help create template documentation files:

- `prompt()`: functions, objects
- `promptPackage()`: package-related documentation
- `promptClass()`: S4 class documentation
- `promptMethods()`: S4 method documentation

.Rd format

syntax similar to that of \LaTeX used for the documentation files

- special markup: `\kbd`, `\code`, ...
- links:

```
\code{\link{foo}}
```

```
\code{\link[pkg]{foo}}
```

```
\code{\link[pkg:bar]{foo}}
```

```
\name{channel}
\alias{channel}

\title{Create a new ExpressionSet instance by selecting a specific channel}
\description{
  This generic function extracts a specific element from an object,
  returning a instance of ExpressionSet.
}
\usage{
channel(object, name, ...)
}
\arguments{
  \item{object}{An S4 object, typically derived from class
    \code{\link{eSet}}}
  \item{name}{A (length one) character vector channel names.}
  \item{...}{Additional arguments.}
}
\value{
  Instance of class \code{\link{ExpressionSet}}.
}
\author{Biocore}

\examples{
obj <- new("NChannelSet",
          R=matrix(runif(100), 20, 5),
          G=matrix(runif(100), 20, 5))
## G channel as ExpressionSet
channel(obj, "G")
}
\keyword{manip}
```

Vignettes

document integrating code and text that describes how to perform a **specific task**.

- created using `Sweave()`
- \LaTeX document
- special markup for code chunks

```
<<chunkName, some options...>>=
```

```
  R code...
```

```
@
```

- and for inline notation `\Sexpr{R expression}`

Concept of Name Spaces

- control which symbols are visible and which values are used during evaluation
- control which functionality is exported
- prevent undeliberate shadowing of symbols in the search path
- reduce overhead of symbol lookup by distinguishing between package attachment and package loading

NAMESPACE file (genefilter)

```
import("methods")
import("AnnotationDbi")
importFrom("annotate", "getAnnMap")
import("Biobase")
import("survival")
importFrom("graphics", "plot")

useDynLib("genefilter")

export("Anova", "allNA", "anyNA", "coxfilter",
       "cv", "eSetFilter", "varFilter", "featureFilter",
       "fastT", "ttest", "shorth", "half.range.mode",
       "rowttests", "colttests", "rowFtests", "colFtests",
       "rowSds", "rowVars", "dist2",
       "filterfun", "findLargest", "gapFilter",
       "genefilter", "genescale", "getFilterNames",
       "getFuncDesc", "getRdAsText", "isESet", "kOverA", "maxA", "pOverA",
       "parseArgs", "parseDesc", "setESetArgs", "showESet")

exportClasses("rowROC")
exportMethods("genefinder", "show", "plot", "[", "sens", "spec",
              "area", "pAUC", "AUC", "rowpAUCs", "nsFilter")
```

My first Package

Let's try this out and build our own first package...