Template file for the documentation of the seqinR package

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1 Introduction

This is a template file to explain how to re-use data from or to contribute to the documentation of the seqinR package.

2 Where are the files

All the files used to compile the documentation are located in the src folder. We are currently in the template folder within the src folder, we move one step back in the hierarchy to show the content of the src folder:

The documentation is in fact a collection of independent LATEX articles generated by Sweaving a *.rnw file. They are merged into a single multi-author document by Sweaving the file book.rnw in the book folder. The articles to be incorporated in the final documentation are located in the frontmatter, mainmatter and appendix folders.

2.1 Bibliography

The references are in the book.bib file located in the config folder:

```
bib <- readLines("../config/book.bib")
cat(head(bib), sep = "\n")

@incollection{seqinr,
    author = {Charif, D. and Lobry, J.R.},
    title = {Seqin{R} 1.0-2: a contributed package to the {R} project for statistical computing devoted to biolog
    booktitle = {Structural approaches to sequence evolution: Molecules, networks, populations},
    year = {2007},
    editor = {U. Bastolla, M. Porto, H.E. Roman and M. Vendruscolo},</pre>
```

New references should be included in this file. There is a line at the end of all *.rnw files saying \bibliography{../config/book} so that this file can be used by bibTeX. References are called in the text with a \cite{} command, for instance \cite{seqinr} to cite the seqinR package [1].

2.2 Tables

The LATEX tables are all in the tables folder.

2.2.1 Regular LATEX tables

For example, the file templatetable.tex looks like:

```
ttab <- readLines("../tables/templatetable.tex")</pre>
 cat(ttab, sep = "\n")
% Example of latex table
\begin{table}[ht]
\begin{center}
\begin{tabular}{rll}
\hline
& Type & Description \\
\hline
1 &
                                   .PE protein coding region \
           CDS &
        LOCUS &
                                       sequenced DNA fragment \\
3 & MISC\_RNA & .RN other structural RNA coding region \\
                                   .RR mature ribosomal RNA \\
.SC small cytoplasmic RNA \\
.SN small nuclear RNA \\
.TR mature transfer RNA \\
4 &
5 &
         RRNA &
         SCRNA &
         SNRNA &
6 &
7 &
\hline
\caption{Example of latex table (available subsequences in genbank)}
\label{templatetable}
\end{center} \end{table}
```

This LATEX table can be included with a \input{../tables/templatetable.tex} command and called in the text with a \ref{templatetable} command (see table 1).

	Type	Description
1	CDS	.PE protein coding region
2	LOCUS	sequenced DNA fragment
3	$\mathrm{MISC}_{-}\mathrm{RNA}$.RN other structural RNA coding region
4	RRNA	.RR mature ribosomal RNA
5	SCRNA	.SC small cytoplasmic RNA
6	SNRNA	.SN small nuclear RNA
7	TRNA	.TR mature transfer RNA

Table 1: Example of latex table (available subsequences in genbank)

2.2.2 Tables generated with xtable()

Tables can also be generated on the fly at the Revel with xtable(). Here is an example:

```
data(toyaa)
print(xtable(toyaa,
   digits = rep(0,4),
   caption = "Data to be loaded with \\texttt{data(toyaa).}",
   label = "toyaa"),
   file = "../tables/toyaa.tex")
```

	Ala	Val	Cys
1	130	70	0
2	60	40	0
3	60	35	5

Table 2: Data to be loaded with data(toyaa).

This LATEX table is included as previously with a \input{../tables/toyaa.tex} command and called in the text with a \ref{toyaa} command (see table 2). Note the use of the double escape sequence \\ in the caption.

2.3 Figures

All figures are in the figs folder in PDF format.

2.3.1 Regular LATEX figures

The file Rlogo.pdf contains for instance the Qlogo. The following code includes figure 1:

```
\begin{figure}
\centering
\fbox{
\begin{minipage}{\textwidth}
\centering
\includegraphics[width=0.5\textwidth]{../figs/Rlogo}
\caption{The \Rlogo{} logo}
\label{Rlogofig}
\end{minipage}
}
\end{figure}
```

The following code include the figure in the margin:

```
\marginpar{
\includegraphics[width=\marginparwidth]{../figs/Rlogo}\\
\tiny{The \Rlogo{} logo}
}
```



2.3.2 Figures generated at the \mathbb{Q} level

Add a fig=T in the header to insert a figure. The following code chunk:

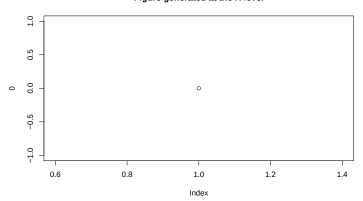
```
<<tryfig, width = 8, height = 5, fig=T>>= plot(0, main = "Figure generated at the R level")
```



yields:

plot(0, main = "Figure generated at the R level")

Figure generated at the R level



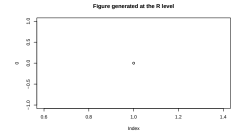
All the figures are generated in PDF format in the figs folder. To avoid names collisions they are prefixed with the name of the the current *.rnw file, that is with template- in this case:

```
list.files(path = "../figs/", pattern = "tryfig")
```

[1] "template-tryfig.pdf"

The default figure size in the final document can be changed. For instance after \setkeys{Gin}{width=0.5\textwidth} the same code chunk yields:

```
plot(0, main = "Figure generated at the R level")
```



The automatic inclusion of a figure after a code chunk can be neutralized with include=F in the header. For example after the following code chunk the figure is generated in the figs folder but not included:

```
<<showinclude, width = 7, height = 7, fig=T, include=F>>=
plot(0, main = "Figure generated at the R level")
list.files(path="../figs/", pattern = "showinclude")
@
plot(0, main = "Figure generated at the R level\n (width = 7, height = 7)")
```

You can then include it with its name, here template-showinclude:

```
list.files(path = "../figs/", pattern = "showinclude")
```

[1] "template-showinclude.pdf"

or better use .PrefixName so that if you change the file name the figure will still be found. The code source of this document (template.rnw) shows how figure 2 was inserted.

3 Configuration files

3.1 LaTeXpackages

The second line of the *.rnw files contains the following LATEX command:

```
\input{../config/commontex}

ctex <- readLines("../config/commontex.tex")
cat(ctex, sep = "\n")

\usepackage{graphicx} % Extended graphics inclusions
\usepackage{float}
\usepackage{float}
\usepackage{url} % For \url{}
\usepackage{amsfonts} % Needed for some fonts
\usepackage[usenames]{color} % Needed for colored R input/output
\usepackage{pdfcolmk} % Correct some problems with the color stack</pre>
```

Extra LATEX packages should go there so that the book could be compiled.

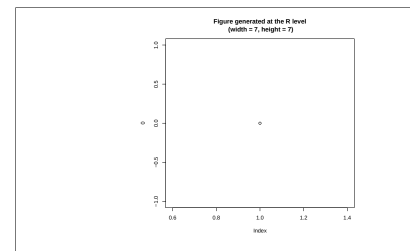


Figure 2: This figure was generated at the \mathbb{Q} level and inserted at the $\mathbb{E}^{T}_{E}X$ level.

3.2 Authors

The authors should be given with \and as separator as in:

```
\author{Lobry, J.R. \and Someone, E.L.S.E.}
```

The authors of the whole document are all contributors listed in alphabetical order. This is done with the script in the file book/book.rnw.

3.3 • packages

Required packages should be loaded in the config/commonrnw.rnw file which is included at the beginning of the *.rnw files with:

SweaveInput{../config/commonrnw.rnw}

The current list of \mathbb{Q} packages is as follows:

```
crnw <- readLines("../config/commonrnw.rnw")
cat(crnw[grep("library", crnw)], sep = "\n")

library(seqinr, lib = "../../../../seqinr.Rcheck")
library(xtable)
library(ade4)
library(ape)
library(MASS)</pre>
```

Important note: if you are not working with development version of seqinR, then the line:

```
library(seqinr, lib = "../../../seqinr.Rcheck")
```

should be changed to:

library(seqinr)

3.4 Don't remove me

There is a comment line at the beginning of the *.rnw file stating:

```
% BEGIN - DO NOT REMOVE THIS LINE
```

As suggested, do not removed it as it is used by the book/book.rnw script to merge the relevant parts of the individual articles into a book.

3.5 Extras in the commonrnw.rnw file

To be documented.

4 Compiling this document

There are two successive compilation steps.

$4.1 *.rnw \rightarrow *.tex$

Under \mathbb{Q} with template as the current working directory run Sweave() with template.rnw as argument:

```
getwd()
```

[1] "/Users/lobry/seqinr/inst/doc/src/template"

```
Sweave("template.rnw")
```

The output on your © console should look like:

```
Writing to file template.tex
Processing code chunks ...
1 : term hide (label=options)
2 : echo term verbatim (label=wherearethefiles)
3 : echo term verbatim (label=showbookbig)
4 : echo term verbatim (label=showtemplatetable)
[...snip...]
16 : term tex (label=sessionInfo)
```

You can now run LaTeX on 'template.tex'

4.1.1 Troubleshooting

The list of packages that must be installed is listed in the next section (Session Informations). If one package is missing you'll get an error message. It's advisable to install packages with all dependencies, for instance:

```
install.packages("ape", dependencies = TRUE)

If the seqinR package is installed but you get an error like:

Error in library(seqinr, lib = "../../../../seqinr.Rcheck")
   open the file config/commonrnw.rnw and change the line:

library(seqinr, lib = "../../../../seqinr.Rcheck")
   into:

library(seqinr)

4.2 *.tex → *.pdf

Use pdfIATeX to compile template.tex into a PDF file with something like:
unix$ pdflatex template.tex
unix$ bibtex template
unix$ pdflatex template.tex
```

You should obtain the PDF of the present document (template.pdf).

5 Session Informations

This part was compiled under the following \mathbb{Q} environment:

- R version 2.5.1 (2007-06-27), i386-apple-darwin8.9.1
- Locale: C
- Base packages: base, datasets, grDevices, graphics, methods, stats, utils
- Other packages: MASS 7.2-34, ade4 1.4-3, ape 1.9-4, gee 4.13-12, lattice 0.15-11, nlme 3.1-83, seqinr 1.1-2, xtable 1.4-3

References

[1] D. Charif and J.R. Lobry. SeqinR 1.0-2: a contributed package to the R project for statistical computing devoted to biological sequences retrieval and analysis. In H.E. Roman U. Bastolla, M. Porto and M. Vendruscolo, editors, *Structural approaches to sequence evolution: Molecules, networks, populations*, Biological and Medical Physics, Biomedical Engineering, pages 207–232. Springer Verlag, New York, USA, 2007. ISBN 978-3-540-35305-8.