Software Manual





msa

An R Package for Multiple Sequence Alignment

Enrico Bonatesta, Christoph Kainrath, and Ulrich Bodenhofer

Institute of Bioinformatics, Johannes Kepler University Linz Altenberger Str. 69, 4040 Linz, Austria msa@bioinf.jku.at

Version 1.22.0, August 12, 2020



Scope and Purpose of this Document

This document provides a gentle introduction into the R package msa. Not all features of the R package are described in full detail. Such details can be obtained from the documentation enclosed in the R package. Further note the following: (1) this is not an introduction to multiple sequence alignment or algorithms for multiple sequence alignment; (2) this is not an introduction to R or any of the Bioconductor packages used in this document. If you lack the background for understanding this manual, you first have to read introductory literature on the subjects mentioned above.

Co	4	L	4
ιn	m	\mathbf{A}	nte
\sim			

1	Introduction	4									
2	Installation										
3	msa for the Impatient	5									
4	Functions for Multiple Sequence Alignment in More Detail 4.1 ClustalW-Specific Parameters	11 13 13 14									
5	Printing Multiple Sequence Alignments										
6	Processing Multiple Alignments 6.1 Methods Inherited From Biostrings	21 21 24 30									
7	Pretty-Printing Multiple Sequence Alignments	34									
	7.1 Consensus Sequence and Sequence Logo 7.2 Color Shading Modes 7.3 Subsetting 7.4 Additional Customizations 7.5 Sweave or knitr Integration 7.6 Sequence Names 7.7 Pretty-Printing Wide Alignments 7.8 Further Caveats	34 35 37 37 37 38 38 39									
8	8 Known Issues										
9	Future Extensions	40									
10	How to Cite This Package	41									

4 2 Installation

1 Introduction

Multiple sequence alignment is one of the most fundamental tasks in bioinformatics. Algorithms like ClustalW [13], ClustalOmega [12], and MUSCLE [3, 4] are well known and widely used. However, all these algorithms are implemented as stand-alone commmand line programs without any integration into the R/Bioconductor ecosystem. Before the msa package, only the muscle package has been available in R, but no other multiple sequence alignment algorithm, although the Biostrings package has provided data types for representing multiple sequence alignments for quite some time. The msa package aims to close that gap by providing a unified R interface to the multiple sequence alignment algorithms ClustalW, ClustalOmega, and MUSCLE. The package requires no additional software packages and runs on all major platforms. Moreover, the msa package provides an R interface to the powerful LATEX package TEX shade [1] which allows for a highly customizable plots of multiple sequence alignments. Unless some very special features of TEX shade are required, users can pretty-print multiple sequence alignments without the need to know the details of LATEX or TEX shade.

2 Installation

The msa R package (current version: 1.22.0) is available via Bioconductor. The simplest way to install the package is the following:

```
if (!requireNamespace("BiocManager", quietly=TRUE))
   install.packages("BiocManager")
BiocManager::install("msa")
```

To test the installation of the msa package, enter

```
library(msa)
```

in your R session. If this command terminates without any error message or warning, you can be sure that the msa package has been installed successfully. If so, the msa package is ready for use now and you can start performing multiple sequence alignments.

To make use of all functionalities of msaPrettyPrint(), a TEX/LATEX system [5] must be installed. To make use of LATEX code created by msaPrettyPrint() or to use the output of msaPrettyPrint() in Sweave [6] or knitr [15] documents, the LATEX package TEXSTATE (file texshade.sty) [1] must be accessible to the LATEX system too. The file texshade.sty is shipped with the msa package. To determine where the file is located, enter the following command in your R session:

```
system.file("tex", "texshade.sty", package="msa")
```

Alternatively, TeXshade can be installed directly from the Comprehensive TeX Archive Network (CTAN).¹

¹https://www.ctan.org/pkg/texshade

3 msa for the Impatient

In order to illustrate the basic workflow, this section presents a simple example with default settings and without going into the details of each step. Let us first load amino acid sequences from one of the example files that are supplied with the msa package:

```
mySequenceFile <- system.file("examples", "exampleAA.fasta", package="msa")</pre>
mySequences <- readAAStringSet(mySequenceFile)</pre>
mySequences
## AAStringSet object of length 9:
##
       width seq
                                                  names
         452 MSTAVLENPGLGRKLS...NSEIGILCSALQKIK PH4H_Homo_sapiens
## [1]
## [2]
         453 MAAVVLENGVLSRKLS...SEVGILCNALQKIKS PH4H_Rattus_norve...
## [3]
         453 MAAVVLENGVLSRKLS...SEVGILCHALQKIKS PH4H_Mus_musculus
## [4]
         297 MNDRADFVVPDITTRK...LNAGDRQGWADTEDV PH4H_Chromobacter...
## [5]
         262 MKTTQYVARQPDDNGF...RLGLHAPLFPPKQAA PH4H_Pseudomonas_...
## [6]
         451 MSALVLESRALGRKLS...SSEVEILCSALQKLK PH4H_Bos_taurus
##
  [7]
         313 MAIATPTSAAPTPAPA...LNAGTREGWADTADI PH4H_Ralstonia_so...
## [8]
         294 MSGDGLSNGPPPGARP...AYATAGGRLAGAAAG PH4H_Caulobacter_...
         275 MSVAEYARDCAAQGLR...VARRKDQKALDPATV PH4H_Rhizobium_loti
## [9]
```

Now that we have loaded the sequences, we can run the msa() function which, by default, runs ClustalW with default parameters:

```
myFirstAlignment <- msa(mySequences)</pre>
## use default substitution matrix
myFirstAlignment
## CLUSTAL 2.1
##
## Call:
     msa(mySequences)
##
##
## MsaAAMultipleAlignment with 9 rows and 456 columns
##
      aln
                                         names
## [1] MAAVVLENGVLSRKLSDF...SINSEVGILCNALQKIKS PH4H_Rattus_norve...
## [2] MAAVVLENGVLSRKLSDF...SINSEVGILCHALQKIKS PH4H_Mus_musculus
## [3] MSTAVLENPGLGRKLSDF...SINSEIGILCSALQKIK- PH4H_Homo_sapiens
  [4] MSALVLESRALGRKLSDF...SISSEVEILCSALQKLK- PH4H_Bos_taurus
## [5] ----- PH4H_Chromobacter...
         ----- PH4H_Ralstonia_so...
## [7] -----PH4H_Caulobacter_...
```

```
## [8] ------ PH4H_Pseudomonas_...
## [9] ------ PH4H_Rhizobium_loti
## Con ----- Consensus
```

Obviously, the default printing function shortens the alignment for the sake of compact output. The print() function provided by the msa package provides some ways for customizing the output, such as, showing the entire alignment split over multiple blocks of sub-sequences:

```
print(myFirstAlignment, show="complete")
##
## MsaAAMultipleAlignment with 9 rows and 456 columns
  [1] MAAVVLENGVLSRKLSDFGQETSYIEDNSNQNGAISLIF PH4H_Rattus_norve...
  [2] MAAVVLENGVLSRKLSDFGQETSYIEDNSNQNGAVSLIF PH4H_Mus_musculus
  [3] MSTAVLENPGLGRKLSDFGQETSYIEDNCNQNGAISLIF PH4H_Homo_sapiens
##
  [4] MSALVLESRALGRKLSDFGQETSYIEGNSDQN-AVSLIF PH4H_Bos_taurus
  [5] ----- PH4H_Chromobacter...
  [6] ----- PH4H_Ralstonia_so...
##
  [7] ----- PH4H_Caulobacter_...
  [8] ----- PH4H_Pseudomonas_...
##
  [9] ----- PH4H_Rhizobium_loti
  Con ----- Consensus
##
##
##
     aln (40..78)
                                  names
  [1] SLKEEVGALAKVLRLFEENDINLTHIESRPSRLNKDEYE PH4H_Rattus_norve...
##
  [2] SLKEEVGALAKVLRLFEENEINLTHIESRPSRLNKDEYE PH4H_Mus_musculus
  [3] SLKEEVGALAKVLRLFEENDVNLTHIESRPSRLKKDEYE PH4H_Homo_sapiens
##
##
  [4] SLKEEVGALARVLRLFEENDINLTHIESRPSRLRKDEYE PH4H_Bos_taurus
  [5] ----- PH4H_Chromobacter...
  [6] ----- PH4H_Ralstonia_so...
##
  [7] ----- PH4H_Caulobacter_...
##
  [8] ----- PH4H_Pseudomonas_...
##
##
  [9] ----- PH4H_Rhizobium_loti
##
  Con ----- Consensus
##
##
     aln (79..117)
                                  names
  [1] FFTYLDKRTKPVLGSIIKSLRNDIGATVHELSRDKEKNT PH4H_Rattus_norve...
##
  [2] FFTYLDKRSKPVLGSIIKSLRNDIGATVHELSRDKEKNT PH4H_Mus_musculus
  [3] FFTHLDKRSLPALTNIIKILRHDIGATVHELSRDKKKDT PH4H_Homo_sapiens
##
##
  [4] FFTNLDQRSVPALANIIKILRHDIGATVHELSRDKKKDT PH4H_Bos_taurus
  [5] ----- PH4H_Chromobacter...
  [6] ----- PH4H_Ralstonia_so...
##
  [7] ----- PH4H_Caulobacter_...
  [8] ----- PH4H_Pseudomonas_...
##
 [9] ----- PH4H_Rhizobium_loti
```

```
## Con ----- Consensus
##
##
      aln (118..156)
                                          names
## [1] VPWFPRTIQELDRFANQILSYGAELDADHPGFKDPVYRA PH4H_Rattus_norve...
## [2] VPWFPRTIQELDRFANQILSYGAELDADHPGFKDPVYRA PH4H_Mus_musculus
  [3] VPWFPRTIQELDRFANQILSYGAELDADHPGFKDPVYRA PH4H_Homo_sapiens
## [4] VPWFPRTIQELDNFANQVLSYGAELDADHPGFKDPVYRA PH4H_Bos_taurus
## [5] -----MNDRADFVVPD----ITTRKNVG PH4H_Chromobacter...
## [7] -----MSG-----DGLSNG PH4H_Caulobacter_...
## [8] -----PH4H_Pseudomonas_...
## Con -----?????????????D?????D????? Consensus
##
##
      aln (157..195)
                                          names
## [1] RRKQFADIAYNYRHGQPIPRVEYTEEEKQTWGTVFRTLK PH4H_Rattus_norve...
## [2] RRKQFADIAYNYRHGQPIPRVEYTEEERKTWGTVFRTLK PH4H_Mus_musculus
## [3] RRKQFADIAYNYRHGQPIPRVEYMEEEKKTWGTVFKTLK PH4H_Homo_sapiens
## [4] RRKQFADIAYNYRHGQPIPRVEYTEEEKKTWGTVFRTLK PH4H_Bos_taurus
## [5] LSHDAN-----DFTLPQPLDRYSAEDHATWATLYQRQC PH4H_Chromobacter...
## [6] FAEGLDGQTLRPDFTMEQPVHRYTAADHATWRTLYDRQE PH4H_Ralstonia_so...
## [7] PPPGAR----PDWTIDQGWETYTQAEHDVWITLYERQT PH4H_Caulobacter_...
## [8] VARQPD-----DNGFIHYPETEHQVWNTLITRQL PH4H_Pseudomonas_...
## [9] LRGDYS--VCRADFTVAQDYD-YSDEEQAVWRTLCDRQT PH4H_Rhizobium_loti
## Con ?R?Q??????????P?P???YTEEE??TW?TL??RQ? Consensus
##
##
      aln (196..234)
                                          names
## [1] ALYKTHACYEHNHIFPLLEKYCGFREDNIPQLEDVSQFL PH4H_Rattus_norve...
## [2] ALYKTHACYEHNHIFPLLEKYCGFREDNIPQLEDVSQFL PH4H_Mus_musculus
## [3] SLYKTHACYEYNHIFPLLEKYCGFHEDNIPQLEDVSQFL PH4H_Homo_sapiens
## [4] SLYKTHACYEHNHIFPLLEKYCGFREDNIPQLEEVSQFL PH4H_Bos_taurus
## [5] KLLPGRACDEFMEGL----ERLEVDADRVPDFNKLNQKL PH4H_Chromobacter...
## [6] ALLPGRACDEFLQGL----STLGMSREGVPSFDRLNETL PH4H_Ralstonia_so...
## [7] DMLHGRACDEFMRGL----DALDLHRSGIPDFARINEEL PH4H_Caulobacter_...
## [8] KVIEGRACQEYLDGI----EQLGLPHERIPQLDEINRVL PH4H_Pseudomonas_...
## [9] KLTRKLAHHSYLDGV----EKLGL-LDRIPDFEDVSTKL PH4H_Rhizobium_loti
## Con ?L????AC?E???G?----??LG???D?IPQLE?VSQ?L Consensus
##
##
      aln (235..273)
                                          names
## [1] QTCTGFRLRPVAGLLSSRDFLGGLAFRVFHCTQYIRHGS PH4H_Rattus_norve...
## [2] QTCTGFRLRPVAGLLSSRDFLGGLAFRVFHCTQYIRHGS PH4H_Mus_musculus
## [3] QTCTGFRLRPVAGLLSSRDFLGGLAFRVFHCTQYIRHGS PH4H_Homo_sapiens
## [4] QSCTGFRLRPVAGLLSSRDFLGGLAFRVFHCTQYIRHGS PH4H_Bos_taurus
## [5] MAATGWKIVAVPGLIPDDVFFEHLANRRFPVTWWLREPH PH4H_Chromobacter...
## [6] MRATGWQIVAVPGLVPDEVFFEHLANRRFPASWWMRRPD PH4H_Ralstonia_so...
## [7] KRLTGWTVVAVPGLVPDDVFFDHLANRRFPAGQFIRKPH PH4H_Caulobacter_...
```

```
[8] QATTGWRVARVPALIPFQTFFELLASQQFPVATFIRTPE PH4H_Pseudomonas_...
  [9] RKLTGWEIIAVPGLIPAAPFFDHLANRRFPVTNWLRTRQ PH4H_Rhizobium_loti
## Con Q??TGWR???VPGL?P???FF??LA?R?FP?TQ?IR??? Consensus
##
##
       aln (274..312)
                                                names
## [1] KPMYTPEPDICHELLGHVPLFSDRSFAOFSQEIG-LASL PH4H Rattus norve...
## [2] KPMYTPEPDICHELLGHVPLFSDRSFAQFSQEIG-LASL PH4H_Mus_musculus
## [3] KPMYTPEPDICHELLGHVPLFSDRSFAQFSQEIG-LASL                               PH4H_Homo_sapiens
## [4] KPMYTPEPDICHELLGHVPLFSDRSFAQFSQEIG-LASL PH4H_Bos_taurus
## [5] QLDYLQEPDVFHDLFGHVPLLINPVFADYLEAYGKGGVK PH4H_Chromobacter...
## [6] QLDYLQEPDGFHDIFGHVPLLINPVFADYMQAYGQGGLK PH4H_Ralstonia_so...
## [7] ELDYLQEPDIFHDVFGHVPMLTDPVFADYMQAYGEGGRR PH4H_Caulobacter_...
## [8] ELDYLQEPDIFHEIFGHCPLLTNPWFAEFTHTYGKLGLK PH4H_Pseudomonas_...
## [9] ELDYIVEPDMFHDFFGHVPVLSQPVFADFMQMYGKKAGD PH4H_Rhizobium_loti
## Con ?LDY??EPDIFHELFGHVPLLSDP?FA?F?Q?YG?LA?? Consensus
##
##
       aln (313..351)
                                                names
## [1] GAPDEYIEKLATIYWFTVEFGLCKEG-DSIKAYGAGLLS PH4H_Rattus_norve...
## [2] GAPDEYIEKLATIYWFTVEFGLCKEG-DSIKAYGAGLLS PH4H_Mus_musculus
## [3] GAPDEYIEKLATIYWFTVEFGLCKQG-DSIKAYGAGLLS PH4H_Homo_sapiens
## [4] GAPDEYIEKLATIYWFTVEFGLCKQG-DSIKAYGAGLLS PH4H_Bos_taurus
## [5] AKALGALPMLARLYWYTVEFGLINTP-AGMRIYGAGILS PH4H_Chromobacter...
## [6] AARLGALDMLARLYWYTVEFGLIRTP-AGLRIYGAGIVS PH4H_Ralstonia_so...
## [7] ALGLGRLANLARLYWYTVEFGLMNTP-AGLRIYGAGIVS PH4H_Caulobacter_...
## [8] ASKE-ERVFLARLYWMTIEFGLVETD-QGKRIYGGGILS PH4H_Pseudomonas_...
## [9] IIALGGDEMITRLYWYTAEYGLVQEAGQPLKAFGAGLMS PH4H_Rhizobium_loti
## Con ?A?????E?LARLYW?TVEFGL????-???KAYGAGLLS Consensus
##
##
       aln (352..390)
                                                names
## [1] SFGELQYCLSD-KPKLLPLELEKTACQEYSVTEFQPLYY PH4H_Rattus_norve...
## [2] SFGELQYCLSD-KPKLLPLELEKTACQEYTVTEFQPLYY PH4H_Mus_musculus
## [3] SFGELQYCLSE-KPKLLPLELEKTAIQNYTVTEFQPLYY PH4H_Homo_sapiens
## [4] SFGELQYCLSD-KPKLLPLELEKTAVQEYTITEFQPLYY PH4H_Bos_taurus
## [5] SKSESIYCLDSASPNRVGFDLMRIMNTRYRIDTFQKTYF PH4H_Chromobacter...
## [6] SKSESVYALDSASPNRIGFDVHRIMRTRYRIDTFQKTYF PH4H_Ralstonia_so...
## [7] SRTESIFALDDPSPNRIGFDLERVMRTLYRIDDFQQVYF PH4H_Caulobacter_...
## [8] SPKETVYSLSD-EPLHQAFNPLEAMRTPYRIDILQPLYF PH4H_Pseudomonas_...
## [9] SFTELQFAVEGKDAHHVPFDLETVMRTGYEIDKFQRAYF PH4H_Rhizobium_loti
## Con SF?ELQYCLSD-?P???PF?LE??M?T?Y?ID?FQPLYF Consensus
##
       aln (391..429)
##
                                                names
## [1] VAESFSDAKEKVRTFAATIPRPFSVRYDPYTQRVEVLDN PH4H_Rattus_norve...
## [2] VAESFNDAKEKVRTFAATIPRPFSVRYDPYTQRVEVLDN PH4H_Mus_musculus
## [3] VAESFNDAKEKVRNFAATIPRPFSVRYDPYTQRIEVLDN PH4H_Homo_sapiens
## [4] VAESFNDAKEKVRNFAATIPRPFSVHYDPYTQRIEVLDN PH4H_Bos_taurus
## [5] VIDSFKQLFDATA-PDFAPLYLQLADAQPWGAGDVAPDD PH4H_Chromobacter...
```

```
[6] VIDSFEQLFDATR-PDFTPLYEALGTLPTFGAGDVVDGD PH4H_Ralstonia_so...
  [7] VIDSIQTLQEVTL-RDFGAIYERLASVSDIGVAEIVPGD PH4H_Caulobacter_...
  [8] VLPDLKRLFQLAQ-EDIMALVHEAMRLG-LHAPLFPPKQ PH4H_Pseudomonas_...
## [9] VLPSFDALRDAFQTADFEAIVARRKDQKALDPATV---- PH4H_Rhizobium_loti
  Con V??SF??L?E??R??D?T???????P?????V?D? Consensus
##
##
##
      aln (430..456)
                                 names
  [1] TQQLKILADSINSEVGILCNALQKIKS PH4H_Rattus_norve...
  [2] TQQLKILADSINSEVGILCHALQKIKS PH4H_Mus_musculus
## [3] TQQLKILADSINSEIGILCSALQKIK- PH4H_Homo_sapiens
## [4] TQQLKILADSISSEVEILCSALQKLK- PH4H_Bos_taurus
## [5] LVLNAGDRQGWADTEDV----- PH4H_Chromobacter...
## [6] AVLNAGTREGWADTADI----- PH4H_Ralstonia_so...
## [7] AVLTRGT-QAYATAGGRLAGAAAG--- PH4H_Caulobacter_...
## [8] AA-----PH4H_Pseudomonas_...
## [9] ----- PH4H_Rhizobium_loti
## Con ?????????????IL??A???--- Consensus
```

The msa package additionally offers the function msaPrettyPrint() which allows for pretty-printing multiple alignments using the LATEX package TEXshade. As an example, the following R code creates a PDF file myfirstAlignment.pdf which is shown in Figure 1:

In the above call to msaPrettyPrint(), the printing of sequence names has been suppressed by showNames="none". The settings askForOverwrite=FALSE and verbose=FALSE are necessary for building this vignette, but, in an interactive R session, they are not necessary.

Almost needless to say, the file names created by msaPrettyPrint() are customizable. By default, the name of the argument is taken as file name. More importantly, the actual output of msaPrettyPrint() is highly customizable, too. For more details, see the Section 7 and the help page of the function (?msaPrettyPrint).

The msaPrettyPrint() function is particularly useful for pretty-printing multiple sequence alignments in Sweave [6] or knitr [15] documents. More details are provided in Section 7. Here, we restrict to a teasing example:

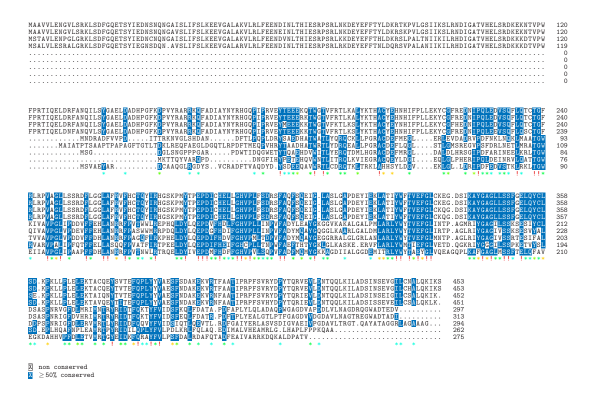


Figure 1: The PDF file myfirstAlignment.pdf created with msaPrettyPrint().

```
IAYNYRHGQPIPRVEYTEEEKQTWGTVFRTLKALYKTHACYEHNHIFPLL 213
IAYNYRHGQPIPRVEYTEEERKTWGTVFRTLKALYKTHACYEHNHIFPLL 213
IAYNYRHGQPIPRVEYMEEEKKTWGTVFKTLKSLYKTHACYEYNHIFPLL 213
IAYNYRHGQPIPRVEYTEEEKKTWGTVFRTLKSLYKTHACYEYNHIFPLL 212
....DFTLPQPLDRYSAEDHATWATLYQRQCKLLPGRACDEFMEGL... 67
QTLRPDFTMEQPVHRYTAADHATWRTLYDRQEALLPGRACDEFLQGL... 83
....PDWTIDQGWETYTQAEHDVWITLYERQTDMLHGRACDEFMRGL... 58
.....DNGFIHYPETEHQVWNTLITRQLKVIEGRACQEYLDGI... 50
.VCRADFTVAQDYD.YSDEEQAVWRTLCDRQTKLTRKLAHHSYLDGV... 65

* * !**** *! !* ** !* **

X non conserved

X > 50% conserved
```

4 Functions for Multiple Sequence Alignment in More Detail

The example in Section 3 above simply called the function msa() without any additional arguments. We mentioned already that, in this case, ClustalW is called with default parameters. We can also explicitly request ClustalW or one of the two other algorithms ClustalOmega or MUSCLE:

```
myClustalWAlignment <- msa(mySequences, "ClustalW")</pre>
## use default substitution matrix
myClustalWAlignment
## CLUSTAL 2.1
##
## Call:
     msa(mySequences, "ClustalW")
##
##
## MsaAAMultipleAlignment with 9 rows and 456 columns
##
      aln
                                        names
##
  [1] MAAVVLENGVLSRKLSDF...SINSEVGILCNALQKIKS PH4H_Rattus_norve...
  [2] MAAVVLENGVLSRKLSDF...SINSEVGILCHALQKIKS PH4H_Mus_musculus
  [3] MSTAVLENPGLGRKLSDF...SINSEIGILCSALQKIK- PH4H_Homo_sapiens
##
  [4] MSALVLESRALGRKLSDF...SISSEVEILCSALQKLK- PH4H_Bos_taurus
  [5] ----- PH4H_Chromobacter...
##
  [6]
         ----- PH4H_Ralstonia_so...
##
                   ---...AYATAGGRLAGAAAG--- PH4H_Caulobacter_...
##
         ----- PH4H_Pseudomonas_...
##
         ----- PH4H_Rhizobium_loti
           ----- Consensus
myClustalOmegaAlignment <- msa(mySequences, "ClustalOmega")</pre>
```

```
## using Gonnet
myClustalOmegaAlignment
## ClustalOmega 1.2.0
##
## Call:
    msa(mySequences, "ClustalOmega")
##
##
## MsaAAMultipleAlignment with 9 rows and 467 columns
##
## [1] MSALVLESRALGRKLSDF...SISSEVEILCSALQKLK- PH4H_Bos_taurus
## [2] MSTAVLENPGLGRKLSDF...SINSEIGILCSALQKIK- PH4H_Homo_sapiens
## [3] MAAVVLENGVLSRKLSDF...SINSEVGILCNALQKIKS PH4H_Rattus_norve...
## [4] MAAVVLENGVLSRKLSDF...SINSEVGILCHALQKIKS PH4H_Mus_musculus
## [5] ----- PH4H_Pseudomonas_...
## [6] ----- PH4H_Rhizobium_loti
## [7] ----- PH4H_Caulobacter_...
## [8] ----- PH4H_Chromobacter...
## [9] ----- PH4H_Ralstonia_so...
## Con ----- Consensus
myMuscleAlignment <- msa(mySequences, "Muscle")</pre>
myMuscleAlignment
## MUSCLE 3.8.31
##
## Call:
    msa(mySequences, "Muscle")
##
##
## MsaAAMultipleAlignment with 9 rows and 460 columns
##
## [1] MAAVVLENGVLSRKLSDF...SINSEVGILCNALQKIKS PH4H_Rattus_norve...
## [2] MAAVVLENGVLSRKLSDF...SINSEVGILCHALQKIKS PH4H_Mus_musculus
## [3] MSTAVLENPGLGRKLSDF...SINSEIGILCSALQKIK- PH4H_Homo_sapiens
## [4] MSALVLESRALGRKLSDF...SISSEVEILCSALQKLK- PH4H_Bos_taurus
## [5] ----- PH4H_Pseudomonas_...
## [6] ----- PH4H_Rhizobium_loti
## [7] -----PH4H_Caulobacter_...
## [8] ----- PH4H_Chromobacter...
## [9] MAIATPTSAAPTPAPAGF...EGWADTADI----- PH4H_Ralstonia_so...
## Con M?????????????DF...???????L??A???--- Consensus
```

Please note that the call msa(mySequences, "ClustalW", ...) is just a shortcut for the call msaClustalW(mySequences, ...), analogously for msaClustalOmega() and msaMuscle().

In other words, msa() is nothing else but a wrapper function that provides a unified interface to the three functions msaClustalW(), msaClustalOmega(), and msaMuscle().

All three functions msaClustalW(), msaClustalOmega(), and msaMuscle() have the same parameters: The input sequences are passed as argument inputSeqs, and all functions have the following arguments: cluster, gapOpening, gapExtension, maxiters, substitutionMatrix, order, type, and verbose. The ways these parameters are interpreted, are largely analogous, although there are some differences, also in terms of default values. See the subsections below and the man page of the three functions for more details. All of the three functions msaClustalW(), msaClustalOmega(), and msaMuscle(), however, are not restricted to the parameters mentioned above. All three have a '...' argument through which several other algorithm-specific parameters can be passed on to the underlying library. The following subsections provide an overview of which parameters are supported by each of the three algorithms.

4.1 ClustalW-Specific Parameters

The original implementation of ClustalW offers a lot of parameters for customizing the way a multiple sequence alignment is computed. Through the '...' argument, msaClustalW() provides an interface to make use of most these parameters (see the documentation of ClustalW² for a comprehensive overview). Currently, the following restrictions and caveats apply:

- The parameters infile, clustering, gapOpen, gapExt, numiters, matrix, and outorder have been renamed to the standardized argument names inputSeqs, cluster, gapOpening, gapExtension, maxiters, substitutionMatrix, and order in order to provide a consistent interface for all three multiple sequence alignment algorithms.
- Boolean flags must be passed as logical values, e.g. verbose=TRUE.
- The parameter quiet has been replaced by verbose (with the exact opposite meaning).
- The following parameters are (currently) not supported: bootstrap, check, fullhelp, interactive, maxseqlen, options, and tree.
- For the parameter output, only the choice "clustal" is available.

4.2 ClustalOmega-Specific Parameters

In the same way as ClustalW, the original implementation of ClustalOmega also offers a lot of parameters for customizing the way a multiple sequence alignment is computed. Through the '...' argument, msaClustalOmega() provides an interface to make use of most these parameters (see the documentation of ClustalOmega³ for a comprehensive overview). Currently, the following restrictions and caveats apply:

■ The parameters infile, cluster-size, iterations, and output-order have been renamed to the argument names inputSeqs, cluster, maxiters, and order in order to provide a consistent interface for all three multiple sequence alignment algorithms.

²http://www.clustal.org/download/clustalw_help.txt

³http://www.clustal.org/omega/README

- ClustalOmega does not allow for setting custom gap penalties. Therefore, setting the parameters gapOpening and gapExtension currently has no effect and will lead to a warning. These arguments are only defined for future extensions and consistency with the other algorithms available in msa.
- ClustalOmega only allows for choosing substitution matrices from a pre-defined set of names, namely "BLOSUM30", "BLOSUM40", "BLOSUM50", "BLOSUM65", "BLOSUM80", and "Gonnet". This is a new feature the original ClustalOmega implementation does not allow for using any custom substitution matrix. However, since these are all amino acid substitution matrices, ClustalOmega is still hardly useful for multiple alignments of nucleotide sequences.
- Boolean flags must be passed as logical values, e.g. verbose=TRUE.
- The following parameters are (currently) not supported: maxSeqLength and help.
- For the parameter outFmt, only the choice "clustal" is available.

4.3 MUSCLE-Specific Parameters

Finally, also MUSCLE offers a lot of parameters for customizing the way a multiple sequence alignment is computed. Through the '...' argument, msaMuscle() provides an interface to make use of most these parameters (see the documentation of MUSCLE⁴ for a comprehensive overview). Currently, the following restrictions and caveats apply:

- The parameters in, gapOpen, gapExtend, matrix, and seqtype have been renamed to inputSeqs, gapOpening, gapExtension, substitutionMatrix and type in order to provide a consistent interface for all three multiple sequence alignment algorithms.
- Boolean flags must be passed as logical values, e.g. verbose=TRUE.
- The parameter quiet has been replaced by verbose (with the exact opposite meaning).
- The following parameters are currently not supported: clw, clwstrict, fastaout, group, html, in1, in2, log, loga, msaout, msf, out, phyi, phyiout, phys, physout, refine, refinew, scorefile, spscore, stable, termgaps4, termgapsfull, termgapshalf, termgapshalflonger, tree1, tree2, usetree, weight1, and weight2.

5 Printing Multiple Sequence Alignments

As already shown above, multiple sequence alignments can be shown in plain text format on the R console using the print() function (which is implicitly called if just the object name is entered on the R console). This function allows for multiple customizations, such as, enabling/disabling to display a consensus sequence, printing the entire alignment or only a subset, enabling/disabling to display sequence names, and adjusting the width allocated for sequence names. For more information, the reader is referred to the help page of the print function:

⁴http://www.drive5.com/muscle/muscle.html

```
help("print, MsaDNAMultipleAlignment-method")
```

We only provide some examples here:

```
print(myFirstAlignment)
## CLUSTAL 2.1
##
## Call:
##
    msa(mySequences)
##
## MsaAAMultipleAlignment with 9 rows and 456 columns
##
     aln
                                     names
## [1] MAAVVLENGVLSRKLSDF...SINSEVGILCNALQKIKS PH4H_Rattus_norve...
## [2] MAAVVLENGVLSRKLSDF...SINSEVGILCHALQKIKS PH4H_Mus_musculus
## [3] MSTAVLENPGLGRKLSDF...SINSEIGILCSALQKIK- PH4H_Homo_sapiens
## [4] MSALVLESRALGRKLSDF...SISSEVEILCSALQKLK- PH4H_Bos_taurus
## [5] ----- PH4H_Chromobacter...
## [6] ----- PH4H_Ralstonia_so...
## [7] -----PH4H_Caulobacter_...
## [8] ----- PH4H_Pseudomonas_...
## [9] ----- PH4H_Rhizobium_loti
## Con ----- Consensus
print(myFirstAlignment, show="complete")
##
## MsaAAMultipleAlignment with 9 rows and 456 columns
     aln (1..39)
## [1] MAAVVLENGVLSRKLSDFGQETSYIEDNSNQNGAISLIF PH4H_Rattus_norve...
## [2] MAAVVLENGVLSRKLSDFGQETSYIEDNSNQNGAVSLIF PH4H_Mus_musculus
## [3] MSTAVLENPGLGRKLSDFGQETSYIEDNCNQNGAISLIF PH4H_Homo_sapiens
## [4] MSALVLESRALGRKLSDFGQETSYIEGNSDQN-AVSLIF PH4H_Bos_taurus
        ----- PH4H_Chromobacter...
## [6] ----- PH4H_Ralstonia_so...
## [7] ----- PH4H_Caulobacter_...
## [8] ----- PH4H_Pseudomonas_...
## [9] ----- PH4H_Rhizobium_loti
## Con ----- Consensus
##
     aln (40..78)
                                     names
## [1] SLKEEVGALAKVLRLFEENDINLTHIESRPSRLNKDEYE PH4H_Rattus_norve...
## [2] SLKEEVGALAKVLRLFEENEINLTHIESRPSRLNKDEYE PH4H_Mus_musculus
## [3] SLKEEVGALAKVLRLFEENDVNLTHIESRPSRLKKDEYE PH4H_Homo_sapiens
## [4] SLKEEVGALARVLRLFEENDINLTHIESRPSRLRKDEYE PH4H_Bos_taurus
```

```
[5] ----- PH4H_Chromobacter...
  [6] ----- PH4H_Ralstonia_so...
  [7] ----- PH4H_Caulobacter_...
 [8] ----- PH4H_Pseudomonas_...
 [9] ----- PH4H_Rhizobium_loti
## Con ----- Consensus
##
##
     aln (79..117)
## [1] FFTYLDKRTKPVLGSIIKSLRNDIGATVHELSRDKEKNT PH4H_Rattus_norve...
## [2] FFTYLDKRSKPVLGSIIKSLRNDIGATVHELSRDKEKNT PH4H_Mus_musculus
## [3] FFTHLDKRSLPALTNIIKILRHDIGATVHELSRDKKKDT                                PH4H_Homo_sapiens
## [4] FFTNLDQRSVPALANIIKILRHDIGATVHELSRDKKKDT PH4H_Bos_taurus
 [5] ----- PH4H_Chromobacter...
 [6] ----- PH4H_Ralstonia_so...
 [7] ----- PH4H_Caulobacter_...
## [8] ----- PH4H_Pseudomonas_...
 [9] ----- PH4H_Rhizobium_loti
## Con ----- Consensus
##
##
     aln (118...156)
                                     names
## [1] VPWFPRTIQELDRFANQILSYGAELDADHPGFKDPVYRA PH4H_Rattus_norve...
 [2] VPWFPRTIQELDRFANQILSYGAELDADHPGFKDPVYRA PH4H_Mus_musculus
## [3] VPWFPRTIQELDRFANQILSYGAELDADHPGFKDPVYRA PH4H_Homo_sapiens
## [4] VPWFPRTIQELDNFANQVLSYGAELDADHPGFKDPVYRA PH4H_Bos_taurus
## [5] ------MNDRADFVVPD----ITTRKNVG PH4H_Chromobacter...
## [6] -----MAIATPTSAAPTPAPAGFTGTLTDKLREQ PH4H_Ralstonia_so...
 [7] -----MSG------DGLSNG PH4H_Caulobacter_...
## [8] -----MKTTQY PH4H_Pseudomonas_...
## Con -----?????????????D?????D????? Consensus
##
##
     aln (157..195)
                                     names
## [1] RRKQFADIAYNYRHGQPIPRVEYTEEEKQTWGTVFRTLK PH4H_Rattus_norve...
 [2] RRKQFADIAYNYRHGQPIPRVEYTEEERKTWGTVFRTLK PH4H_Mus_musculus
 [3] RRKQFADIAYNYRHGQPIPRVEYMEEEKKTWGTVFKTLK PH4H_Homo_sapiens
## [4] RRKQFADIAYNYRHGQPIPRVEYTEEEKKTWGTVFRTLK PH4H_Bos_taurus
 [5] LSHDAN-----DFTLPQPLDRYSAEDHATWATLYQRQC PH4H_Chromobacter...
## [6] FAEGLDGQTLRPDFTMEQPVHRYTAADHATWRTLYDRQE PH4H_Ralstonia_so...
## [7] PPPGAR----PDWTIDQGWETYTQAEHDVWITLYERQT PH4H_Caulobacter_...
## [8] VARQPD-----DNGFIHYPETEHQVWNTLITRQL PH4H_Pseudomonas_...
 [9] LRGDYS--VCRADFTVAQDYD-YSDEEQAVWRTLCDRQT PH4H_Rhizobium_loti
## Con ?R?Q???????????P?P???YTEEE??TW?TL??RQ? Consensus
##
##
     aln (196..234)
                                     names
## [1] ALYKTHACYEHNHIFPLLEKYCGFREDNIPQLEDVSQFL PH4H_Rattus_norve...
## [2] ALYKTHACYEHNHIFPLLEKYCGFREDNIPQLEDVSQFL PH4H_Mus_musculus
```

```
## [3] SLYKTHACYEYNHIFPLLEKYCGFHEDNIPQLEDVSQFL PH4H_Homo_sapiens
## [4] SLYKTHACYEHNHIFPLLEKYCGFREDNIPQLEEVSQFL PH4H_Bos_taurus
## [5] KLLPGRACDEFMEGL----ERLEVDADRVPDFNKLNQKL PH4H_Chromobacter...
## [6] ALLPGRACDEFLQGL----STLGMSREGVPSFDRLNETL PH4H_Ralstonia_so...
## [7] DMLHGRACDEFMRGL----DALDLHRSGIPDFARINEEL PH4H_Caulobacter_...
## [8] KVIEGRACQEYLDGI----EQLGLPHERIPQLDEINRVL PH4H_Pseudomonas_...
## [9] KLTRKLAHHSYLDGV----EKLGL-LDRIPDFEDVSTKL PH4H_Rhizobium_loti
## Con ?L?????AC?E???G?----??LG???D?IPQLE?VSQ?L Consensus
##
##
       aln (235..273)
                                                names
## [1] QTCTGFRLRPVAGLLSSRDFLGGLAFRVFHCTQYIRHGS PH4H_Rattus_norve...
## [2] QTCTGFRLRPVAGLLSSRDFLGGLAFRVFHCTQYIRHGS PH4H_Mus_musculus
## [3] QTCTGFRLRPVAGLLSSRDFLGGLAFRVFHCTQYIRHGS PH4H_Homo_sapiens
## [4] QSCTGFRLRPVAGLLSSRDFLGGLAFRVFHCTQYIRHGS PH4H_Bos_taurus
## [5] MAATGWKIVAVPGLIPDDVFFEHLANRRFPVTWWLREPH PH4H_Chromobacter...
## [6] MRATGWQIVAVPGLVPDEVFFEHLANRRFPASWWMRRPD PH4H_Ralstonia_so...
## [7] KRLTGWTVVAVPGLVPDDVFFDHLANRRFPAGQFIRKPH PH4H_Caulobacter_...
## [8] QATTGWRVARVPALIPFQTFFELLASQQFPVATFIRTPE PH4H_Pseudomonas_...
## [9] RKLTGWEIIAVPGLIPAAPFFDHLANRRFPVTNWLRTRQ PH4H_Rhizobium_loti
## Con Q??TGWR???VPGL?P???FF??LA?R?FP?TQ?IR??? Consensus
##
##
       aln (274..312)
                                                names
## [1] KPMYTPEPDICHELLGHVPLFSDRSFAQFSQEIG-LASL PH4H_Rattus_norve...
## [2] KPMYTPEPDICHELLGHVPLFSDRSFAQFSQEIG-LASL PH4H_Mus_musculus
## [3] KPMYTPEPDICHELLGHVPLFSDRSFAQFSQEIG-LASL PH4H_Homo_sapiens
## [4] KPMYTPEPDICHELLGHVPLFSDRSFAQFSQEIG-LASL PH4H_Bos_taurus
## [5] QLDYLQEPDVFHDLFGHVPLLINPVFADYLEAYGKGGVK PH4H_Chromobacter...
## [6] QLDYLQEPDGFHDIFGHVPLLINPVFADYMQAYGQGGLK PH4H_Ralstonia_so...
## [7] ELDYLQEPDIFHDVFGHVPMLTDPVFADYMQAYGEGGRR PH4H_Caulobacter_...
## [8] ELDYLQEPDIFHEIFGHCPLLTNPWFAEFTHTYGKLGLK PH4H_Pseudomonas_...
## [9] ELDYIVEPDMFHDFFGHVPVLSQPVFADFMQMYGKKAGD PH4H_Rhizobium_loti
## Con ?LDY??EPDIFHELFGHVPLLSDP?FA?F?Q?YG?LA?? Consensus
##
##
       aln (313..351)
                                                names
## [1] GAPDEYIEKLATIYWFTVEFGLCKEG-DSIKAYGAGLLS PH4H_Rattus_norve...
## [2] GAPDEYIEKLATIYWFTVEFGLCKEG-DSIKAYGAGLLS PH4H_Mus_musculus
## [3] GAPDEYIEKLATIYWFTVEFGLCKQG-DSIKAYGAGLLS                                PH4H_Homo_sapiens
## [4] GAPDEYIEKLATIYWFTVEFGLCKQG-DSIKAYGAGLLS PH4H_Bos_taurus
## [5] AKALGALPMLARLYWYTVEFGLINTP-AGMRIYGAGILS PH4H_Chromobacter...
## [6] AARLGALDMLARLYWYTVEFGLIRTP-AGLRIYGAGIVS PH4H_Ralstonia_so...
## [7] ALGLGRLANLARLYWYTVEFGLMNTP-AGLRIYGAGIVS PH4H_Caulobacter_...
## [8] ASKE-ERVFLARLYWMTIEFGLVETD-QGKRIYGGGILS PH4H_Pseudomonas_...
## [9] IIALGGDEMITRLYWYTAEYGLVQEAGQPLKAFGAGLMS PH4H_Rhizobium_loti
## Con ?A?????E?LARLYW?TVEFGL????-???KAYGAGLLS Consensus
##
## aln (352..390)
                                                names
```

```
[1] SFGELQYCLSD-KPKLLPLELEKTACQEYSVTEFQPLYY PH4H_Rattus_norve...
## [2] SFGELQYCLSD-KPKLLPLELEKTACQEYTVTEFQPLYY PH4H_Mus_musculus
## [3] SFGELQYCLSE-KPKLLPLELEKTAIQNYTVTEFQPLYY PH4H_Homo_sapiens
## [4] SFGELQYCLSD-KPKLLPLELEKTAVQEYTITEFQPLYY PH4H_Bos_taurus
## [5] SKSESIYCLDSASPNRVGFDLMRIMNTRYRIDTFQKTYF PH4H_Chromobacter...
## [6] SKSESVYALDSASPNRIGFDVHRIMRTRYRIDTFQKTYF PH4H_Ralstonia_so...
## [7] SRTESIFALDDPSPNRIGFDLERVMRTLYRIDDFQQVYF PH4H_Caulobacter_...
## [8] SPKETVYSLSD-EPLHQAFNPLEAMRTPYRIDILQPLYF PH4H_Pseudomonas_...
## [9] SFTELQFAVEGKDAHHVPFDLETVMRTGYEIDKFQRAYF PH4H_Rhizobium_loti
## Con SF?ELQYCLSD-?P???PF?LE??M?T?Y?ID?FQPLYF Consensus
##
      aln (391..429)
##
                                              names
## [1] VAESFSDAKEKVRTFAATIPRPFSVRYDPYTQRVEVLDN PH4H_Rattus_norve...
## [2] VAESFNDAKEKVRTFAATIPRPFSVRYDPYTQRVEVLDN PH4H_Mus_musculus
## [3] VAESFNDAKEKVRNFAATIPRPFSVRYDPYTQRIEVLDN PH4H_Homo_sapiens
## [4] VAESFNDAKEKVRNFAATIPRPFSVHYDPYTQRIEVLDN PH4H_Bos_taurus
## [5] VIDSFKQLFDATA-PDFAPLYLQLADAQPWGAGDVAPDD PH4H_Chromobacter...
## [6] VIDSFEQLFDATR-PDFTPLYEALGTLPTFGAGDVVDGD PH4H_Ralstonia_so...
## [7] VIDSIQTLQEVTL-RDFGAIYERLASVSDIGVAEIVPGD PH4H_Caulobacter_...
## [8] VLPDLKRLFQLAQ-EDIMALVHEAMRLG-LHAPLFPPKQ PH4H_Pseudomonas_...
## [9] VLPSFDALRDAFQTADFEAIVARRKDQKALDPATV---- PH4H_Rhizobium_loti
## Con V??SF??L?E??R??D?T???????P?????V?D? Consensus
##
##
      aln (430..456)
                                  names
## [1] TQQLKILADSINSEVGILCNALQKIKS PH4H_Rattus_norve...
## [2] TQQLKILADSINSEVGILCHALQKIKS PH4H_Mus_musculus
## [3] TQQLKILADSINSEIGILCSALQKIK- PH4H_Homo_sapiens
## [4] TQQLKILADSISSEVEILCSALQKLK- PH4H_Bos_taurus
## [5] LVLNAGDRQGWADTEDV----- PH4H_Chromobacter...
## [6] AVLNAGTREGWADTADI----- PH4H_Ralstonia_so...
## [7] AVLTRGT-QAYATAGGRLAGAAAG--- PH4H_Caulobacter_...
## [8] AA----- PH4H_Pseudomonas_...
## [9] ----- PH4H_Rhizobium_loti
## Con ?????????????IL??A???--- Consensus
print(myFirstAlignment, showConsensus=FALSE, halfNrow=3)
## CLUSTAL 2.1
##
## Call:
##
     msa(mySequences)
##
## MsaAAMultipleAlignment with 9 rows and 456 columns
## [1] MAAVVLENGVLSRKLSDF...SINSEVGILCNALQKIKS PH4H_Rattus_norve...
## [2] MAAVVLENGVLSRKLSDF...SINSEVGILCHALQKIKS PH4H_Mus_musculus
```

```
## [3] MSTAVLENPGLGRKLSDF...SINSEIGILCSALQKIK- PH4H_Homo_sapiens
## [7] -----PH4H_Caulobacter_...
## [8] ----- PH4H_Pseudomonas_...
## [9] ----- PH4H_Rhizobium_loti
print(myFirstAlignment, showNames=FALSE, show="complete")
##
## MsaAAMultipleAlignment with 9 rows and 456 columns
     aln (1..60)
## [1] MAAVVLENGVLSRKLSDFGQETSYIEDNSNQNGAISLIFSLKEEVGALAKVLRLFEENDI
## [2] MAAVVLENGVLSRKLSDFGQETSYIEDNSNQNGAVSLIFSLKEEVGALAKVLRLFEENEI
## [3] MSTAVLENPGLGRKLSDFGQETSYIEDNCNQNGAISLIFSLKEEVGALAKVLRLFEENDV
## [4] MSALVLESRALGRKLSDFGQETSYIEGNSDQN-AVSLIFSLKEEVGALARVLRLFEENDI
## [6] -----
## [8] -----
## [9] -----
##
##
     aln (61..120)
## [1] NLTHIESRPSRLNKDEYEFFTYLDKRTKPVLGSIIKSLRNDIGATVHELSRDKEKNTVPW
 [2] NLTHIESRPSRLNKDEYEFFTYLDKRSKPVLGSIIKSLRNDIGATVHELSRDKEKNTVPW
## [3] NLTHIESRPSRLKKDEYEFFTHLDKRSLPALTNIIKILRHDIGATVHELSRDKKKDTVPW
## [4] NLTHIESRPSRLRKDEYEFFTNLDQRSVPALANIIKILRHDIGATVHELSRDKKKDTVPW
## [5] ----
## [6] -----
## [8] -----
## [9] -----
## Con -----
##
##
     aln (121..180)
## [1] FPRTIQELDRFANQILSYGAELDADHPGFKDPVYRARRKQFADIAYNYRHGQPIPRVEYT
## [2] FPRTIQELDRFANQILSYGAELDADHPGFKDPVYRARRKQFADIAYNYRHGQPIPRVEYT
  [3] FPRTIQELDRFANQILSYGAELDADHPGFKDPVYRARRKQFADIAYNYRHGQPIPRVEYM
## [4] FPRTIQELDNFANQVLSYGAELDADHPGFKDPVYRARRKQFADIAYNYRHGQPIPRVEYT
## [5] -----DFTLPQPLDRYS
## [6] -----MAIATPTSAAPTPAPAGFTGTLTDKLREQFAEGLDGQTLRPDFTMEQPVHRYT
## [7] -----MSG-----MSG-----DGLSNGPPPGAR----PDWTIDQGWETYT
## [8] -----DNGFIHYP
## [9] -----MSVAEYAR-----DCAAQGLRGDYS--VCRADFTVAQDYD-YS
##
```

```
aln (181..240)
   [1] EEEKQTWGTVFRTLKALYKTHACYEHNHIFPLLEKYCGFREDNIPQLEDVSQFLQTCTGF
  [2] EEERKTWGTVFRTLKALYKTHACYEHNHIFPLLEKYCGFREDNIPQLEDVSQFLQTCTGF
## [3] EEEKKTWGTVFKTLKSLYKTHACYEYNHIFPLLEKYCGFHEDNIPQLEDVSQFLQTCTGF
  [4] EEEKKTWGTVFRTLKSLYKTHACYEHNHIFPLLEKYCGFREDNIPQLEEVSQFLQSCTGF
##
  [5] AEDHATWATLYQRQCKLLPGRACDEFMEGL---ERLEVDADRVPDFNKLNQKLMAATGW
  [6] AADHATWRTLYDRQEALLPGRACDEFLQGL---STLGMSREGVPSFDRLNETLMRATGW
  [7] QAEHDVWITLYERQTDMLHGRACDEFMRGL----DALDLHRSGIPDFARINEELKRLTGW
  [8] ETEHQVWNTLITRQLKVIEGRACQEYLDGI----EQLGLPHERIPQLDEINRVLQATTGW
  [9] DEEQAVWRTLCDRQTKLTRKLAHHSYLDGV----EKLGL-LDRIPDFEDVSTKLRKLTGW
## Con EEE??TW?TL??RQ??L????AC?E???G?----??LG???D?IPQLE?VSQ?LQ??TGW
##
##
       aln (241..300)
## [1] RLRPVAGLLSSRDFLGGLAFRVFHCTQYIRHGSKPMYTPEPDICHELLGHVPLFSDRSFA
  [2] RLRPVAGLLSSRDFLGGLAFRVFHCTQYIRHGSKPMYTPEPDICHELLGHVPLFSDRSFA
## [3] RLRPVAGLLSSRDFLGGLAFRVFHCTQYIRHGSKPMYTPEPDICHELLGHVPLFSDRSFA
## [4] RLRPVAGLLSSRDFLGGLAFRVFHCTQYIRHGSKPMYTPEPDICHELLGHVPLFSDRSFA
## [5] KIVAVPGLIPDDVFFEHLANRRFPVTWWLREPHQLDYLQEPDVFHDLFGHVPLLINPVFA
  [6] QIVAVPGLVPDEVFFEHLANRRFPASWWMRRPDQLDYLQEPDGFHDIFGHVPLLINPVFA
## [7] TVVAVPGLVPDDVFFDHLANRRFPAGQFIRKPHELDYLQEPDIFHDVFGHVPMLTDPVFA
## [8] RVARVPALIPFQTFFELLASQQFPVATFIRTPEELDYLQEPDIFHEIFGHCPLLTNPWFA
  [9] EIIAVPGLIPAAPFFDHLANRRFPVTNWLRTRQELDYIVEPDMFHDFFGHVPVLSQPVFA
## Con R???VPGL?P???FF??LA?R?FP?TQ?IR????LDY??EPDIFHELFGHVPLLSDP?FA
##
##
       aln (301..360)
## [1] QFSQEIG-LASLGAPDEYIEKLATIYWFTVEFGLCKEG-DSIKAYGAGLLSSFGELQYCL
  [2] QFSQEIG-LASLGAPDEYIEKLATIYWFTVEFGLCKEG-DSIKAYGAGLLSSFGELQYCL
  [3] QFSQEIG-LASLGAPDEYIEKLATIYWFTVEFGLCKQG-DSIKAYGAGLLSSFGELQYCL
  [4] QFSQEIG-LASLGAPDEYIEKLATIYWFTVEFGLCKQG-DSIKAYGAGLLSSFGELQYCL
##
  [5] DYLEAYGKGGVKAKALGALPMLARLYWYTVEFGLINTP-AGMRIYGAGILSSKSESIYCL
   [6] DYMQAYGQGGLKAARLGALDMLARLYWYTVEFGLIRTP-AGLRIYGAGIVSSKSESVYAL
  [7] DYMQAYGEGGRRALGLGRLANLARLYWYTVEFGLMNTP-AGLRIYGAGIVSSRTESIFAL
  [8] EFTHTYGKLGLKASKE-ERVFLARLYWMTIEFGLVETD-QGKRIYGGGILSSPKETVYSL
  [9] DFMQMYGKKAGDIIALGGDEMITRLYWYTAEYGLVQEAGQPLKAFGAGLMSSFTELQFAV
## Con ?F?Q?YG?LA???A?????E?LARLYW?TVEFGL????-???KAYGAGLLSSF?ELQYCL
##
##
       aln (361..420)
## [1] SD-KPKLLPLELEKTACQEYSVTEFQPLYYVAESFSDAKEKVRTFAATIPRPFSVRYDPY
  [2] SD-KPKLLPLELEKTACQEYTVTEFQPLYYVAESFNDAKEKVRTFAATIPRPFSVRYDPY
## [3] SE-KPKLLPLELEKTAIQNYTVTEFQPLYYVAESFNDAKEKVRNFAATIPRPFSVRYDPY
## [4] SD-KPKLLPLELEKTAVQEYTITEFQPLYYVAESFNDAKEKVRNFAATIPRPFSVHYDPY
  [5] DSASPNRVGFDLMRIMNTRYRIDTFQKTYFVIDSFKQLFDATA-PDFAPLYLQLADAQPW
  [6] DSASPNRIGFDVHRIMRTRYRIDTFQKTYFVIDSFEQLFDATR-PDFTPLYEALGTLPTF
## [7] DDPSPNRIGFDLERVMRTLYRIDDFQQVYFVIDSIQTLQEVTL-RDFGAIYERLASVSDI
## [8] SD-EPLHQAFNPLEAMRTPYRIDILQPLYFVLPDLKRLFQLAQ-EDIMALVHEAMRLG-L
## [9] EGKDAHHVPFDLETVMRTGYEIDKFQRAYFVLPSFDALRDAFQTADFEAIVARRKDQKAL
```

```
Con SD-?P???PF?LE??M?T?Y?ID?FQPLYFV??SF??L?E??R??D?T????????P?
##
##
      aln (421..456)
  [1] TQRVEVLDNTQQLKILADSINSEVGILCNALQKIKS
##
##
  [2] TQRVEVLDNTQQLKILADSINSEVGILCHALQKIKS
  [3] TQRIEVLDNTQQLKILADSINSEIGILCSALQKIK-
  [4] TQRIEVLDNTQQLKILADSISSEVEILCSALQKLK-
  [5] GAGDVAPDDLVLNAGDRQGWADTEDV-----
  [6] GAGDVVDGDAVLNAGTREGWADTADI-----
  [7] GVAEIVPGDAVLTRGT-QAYATAGGRLAGAAAG---
  [8] HAPLFPPKQAA-----
## [9] DPATV-----
## Con ?????V?D????????????????IL??A???---
```

6 Processing Multiple Alignments

6.1 Methods Inherited From Biostrings

The classes defined by the msa package for storing multiple alignment results have been derived from the corresponding classes defined by the Biostrings package. Therefore, all methods for processing multiple alignments are available and work without any practical limitation. In this section, we highlight some of these.

The classes used for storing multiple alignments allow for defining masks on sequences and sequence positions via their row and column mask slots. They can be set by rowmask() and colmask() functions which serve both as setter and getter functions. To set row or column masks, an IRanges object must be supplied:

```
myMaskedAlignment <- myFirstAlignment</pre>
colM <- IRanges(start=1, end=100)</pre>
colmask(myMaskedAlignment) <- colM</pre>
myMaskedAlignment
## CLUSTAL 2.1
##
## Call:
##
      msa(mySequences)
##
## MsaAAMultipleAlignment with 9 rows and 456 columns
##
                                                names
##
  [1] #################...SINSEVGILCNALQKIKS PH4H_Rattus_norve...
  [2] ###############...SINSEVGILCHALQKIKS PH4H_Mus_musculus
  [3] ################...SINSEIGILCSALQKIK- PH4H_Homo_sapiens
## [4] ###############...SISSEVEILCSALQKLK- PH4H_Bos_taurus
## [5] ################...GWADTEDV----- PH4H_Chromobacter...
```

The unmasked() allows for removing these masks, thereby casting the multiple alignment to a set of aligned Biostrings sequences (class AAStringSet, DNAStringSet, or RNAStringSet):

```
unmasked(myMaskedAlignment)
## AAStringSet object of length 9:
     width seq
##
                                      names
       456 MAAVVLENGVLSRKLS...SEVGILCNALQKIKS PH4H_Rattus_norve...
## [1]
## [2]
       456 MAAVVLENGVLSRKLS...SEVGILCHALQKIKS PH4H_Mus_musculus
      456 MSTAVLENPGLGRKLS...SEIGILCSALQKIK- PH4H_Homo_sapiens
## [3]
      456 MSALVLESRALGRKLS...SEVEILCSALQKLK- PH4H_Bos_taurus
## [4]
      456 ----- PH4H_Chromobacter...
## [5]
       456 ----- PH4H_Ralstonia_so...
## [6]
## [7]
       456 ----- PH4H_Caulobacter_...
       456 ----- PH4H_Pseudomonas_...
## [8]
## [9]
       456 ----- PH4H_Rhizobium_loti
```

Consensus matrices can be computed conveniently as follows:

```
conMat <- consensusMatrix(myFirstAlignment)</pre>
dim(conMat)
## [1] 21 456
conMat[, 101:110]
##
     [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8] [,9] [,10]
## -
       5
            5
                 5
                     5
                          5
                               5
                                    5
                                             5
## A
       0
            0
                 0
                     4
                          0
                               0
                                    0
                                         0
                                             0
                                                   0
       0
            0
                0
                     0
                          0
                               0
                                    0
                                         0
                                                   0
## C
## D
       4
           0
              0
                     0
                          0
                               0
                                    0
                                         0
                                             0
                                                   0
       0
         0 0
                     0
                          0
                               0
                                    0
                                        4
                                             0
                                                   0
## E
       0 0 0
                     0
                          0
                               0
## F
                                    0
                                        0
                                             0
                                                   0
## G
       0
         0
              4
                     0
                          0
                               0
                                    0
                                        0
                                             0
                                                   0
## H
       0
          0
                0
                     0
                          0
                               0
                                    4
                                        0
                                             0
                                                   0
## I
       0
         4 0
                     0
                          0
                               0
                                    0
                                             0
                                                   0
            0 0
                     0
                          0
                                        0
## K
       0
                               0
                                    0
                                             0
                                                   0
     0 0 0 0
                          0 0
                                    0
## L
```

```
0
##
           0
                  0
                          0
                                 0
                                               0
                                                       0
                                                                     0
                                                                              0
   M
##
   N
           0
                  0
                          0
                                 0
                                        0
                                               0
                                                       0
                                                              0
                                                                     0
                                                                              0
                                                                     0
                                                                              0
##
   Ρ
           0
                  0
                          0
                                 0
                                        0
                                               0
                                                      0
                                                              0
                                 0
                                                                     0
                                                                              0
##
   Q
           0
                  0
                          0
                                        0
                                               0
                                                       0
                                                              0
##
   R
           0
                  0
                          0
                                 0
                                        0
                                               0
                                                       0
                                                              0
                                                                     0
                                                                              0
   S
           0
                          0
                                 0
                                        0
                                               0
                                                              0
                                                                     0
                                                                              4
##
                  0
                                                       0
           0
                          0
                                 0
                                        4
                                               0
                                                              0
                                                                     0
                                                                              0
##
   Τ
                  0
                                                       0
           0
                  0
                          0
                                 0
                                        0
                                               4
                                                       0
                                                              0
                                                                     0
                                                                              0
## V
                                               0
                                                              0
                                                                     0
## W
           0
                  0
                          0
                                 0
                                        0
                                                       0
                                                                              0
## Y
           0
                  0
                         0
                                 0
                                        0
                                               0
                                                       0
                                                              0
                                                                     0
                                                                              0
```

If called on a masked alignment, consensusMatrix() only uses those sequences/rows that are not masked. If there are masked columns, the matrix contains NA's in those columns:

```
conMat <- consensusMatrix(myMaskedAlignment)</pre>
conMat[, 95:104]
##
            [,2]
                   [,3]
                         [,4]
                               [,5]
                                      [,6]
                                            [,7]
                                                  [,8]
                                                         [,9]
                                                            5
##
        NA
               NA
                     NA
                           NA
                                  NA
                                        NA
                                               5
                                                      5
                                                                    5
                                                      0
                                                            0
                                                                    4
## A
        NA
               NA
                     NA
                                 NA
                                        NA
                                               0
                           NA
##
   C
        NA
               NA
                     NA
                           NA
                                        NA
                                               0
                                                      0
                                                            0
                                                                    0
                                 NA
##
   D
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                               4
                                                      0
                                                            0
                                                                    0
##
   Ε
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                               0
                                                      0
                                                            0
                                                                    0
## F
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                               0
                                                      0
                                                            0
                                                                    0
                                                            4
##
   G
        NA
               NA
                     NA
                           NA
                                  NA
                                        NA
                                               0
                                                      0
                                                                    0
                                                                    0
##
   Η
                                                      0
                                                            0
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                               0
##
   Ι
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                                      4
                                                            0
                                                                    0
                                               0
                                                      0
                                                            0
                                                                    0
## K
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                               0
## L
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                               0
                                                      0
                                                            0
                                                                    0
## M
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                               0
                                                      0
                                                            0
                                                                    0
                                                      0
                                                            0
                                                                    0
##
   N
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                               0
##
   Ρ
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                               0
                                                      0
                                                            0
                                                                    0
##
   Q
        NA
               NA
                           NA
                                 NA
                                        NA
                                               0
                                                      0
                                                            0
                                                                    0
                     NA
## R
               NA
                                               0
                                                      0
                                                            0
                                                                    0
        NA
                     NA
                           NA
                                 NA
                                        NA
## S
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                               0
                                                      0
                                                            0
                                                                    0
                                                                    0
##
   Τ
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                               0
                                                      0
                                                            0
                                                      0
                                                            0
                                                                    0
## V
        NA
               NA
                     NA
                                        NA
                                               0
                           NA
                                  NA
                                                      0
                                                            0
                                                                    0
## W
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                               0
## Y
        NA
               NA
                     NA
                           NA
                                 NA
                                        NA
                                               0
                                                      0
                                                                    0
```

Multiple alignments also inherit the consensusString() method from the Biostrings package. However, for more flexibility and consistency, we rather advise users to use the method msaConsensusSequence() method (see below).

6.2 Consensus Sequences and Conservation Scores

With version 1.7.1 of msa, new methods have been provided that allow for the computation of consensus sequences and conservation scores. By default, the msaConsensusSequence() method is a wrapper around the consensusString() method from the Biostrings:

However, there is also a second method for computing consensus sequence that has been implemented in line with a consensus sequence method implemented in Texshade that allows for specify an upper and a lower conservation threshold (see example below). This method can be accessed via the argument type="upperlower". Additional customizations are available, too:

Regardless of which method is used, masks are taken into account: masked rows/sequences are neglected and masked columns are shown as "#" in the consensus sequence:

The main purpose of consensus sequences is to get an impression of conservation at individual positions/columns of a multiple alignment. The msa package also provides another means of analyzing conservation: the method msaConservationScore() computes sums of pairwise scores for a given substitution/scoring matrix. Thereby, conservation can also be analyzed in a more sensible way than by only taking relative frequencies of letters into account as msaConsensusSequence() does.

```
data(BLOSUM62)
msaConservationScore(myFirstAlignment, BLOSUM62)
##
##
    -55
         -95
              -94 -109
                         -71
                              -71
                                   -55
                                        -71 -119 -121
                                                        -71
                                                             -95
                                                                   -55
##
                                    _
##
    -55
         -71
              -71
                   -39
                         -39
                              -39
                                   -55
                                        -55
                                             -55
                                                   -71
                                                        -23
                                                             -71
##
                              -39
                                             -79
                                                                  -39
##
    -81
         -39
              -96
                   -69
                         -55
                                   -54
                                        -71
                                                   -71
                                                        -71
                                                             -71
##
##
    -71
         -71
              -55
                   -55
                         -55
                              -71
                                   -39
                                        -71
                                              -71
                                                   -71
                                                        -73
                                                             -71
##
                          _
                              -39
                                        -77
##
    -55
         -71
              -39
                   -55
                         -55
                                   -64
                                             -39
                                                   -71
                                                        -55
                                                                  -71
##
                         -71
##
    -55
         -71
              -55
                   -23
                              -55
                                   -71
                                        -97
                                              -55
                                                   -39
                                                        -55
                                                             -23
##
##
    -39
         -39
              -55
                   -91
                         -71
                              -39
                                   -79
                                        -55
                                             -88 -121
                                                        -23 -103
                                                                  -71
##
                                   -71
##
   -110
         -87
              -71
                   -71
                         -55 -119
                                        -55
                                             -71
                                                   -39
                                                        -71
                                                             -39
##
##
    -55
         -71
                   -55
                         -71
                              -71
                                   -55
                                        -39
                                             -55
                                                   -87
                                                        -55
                                                             -79
                                                                  -55
##
                                                          ?
                                                               ?
                   -39
                        -23
                             -55
                                  -55
                                             -55
##
    -71
         -23
               41
                                        -71
                                                  -55
                                                        -59
                                                             -60 -113
     ?
               ?
                    ?
                         ?
                               ?
                                   ?
                                         Y
                                              ?
                                                   ?
##
                                             -45
##
    -60
         -75
                9
                    44
                        -47
                              -59
                                  -11
                                         18
                                                  -58
                                                        -71
                                                             -91
   ? ? ? ? ? D ? ?
```

-42 ## -74 -66 -27 -59 -27 -51 246 52 74 52 97 74 ? ? ? ? ? ? ? ## ? R ? Q ? ? 100 125 60 51 -50 -99 -62 -36 -19 -32 49 ## -5 77 ## ? ? ? ? Ρ ? ? ? Υ Τ Е Ē Р ## 96 49 -2 166 30 218 6 21 34 567 141 160 165 ? ? ? Τ ? ? ? ## Ē Τ W L R Q 325 179 71 216 891 38 405 204 208 79 165 109 42 ## ? ## ? L ? ? ? Α C ? Ε ? ? ? 536 249 113 106 172 324 109 324 261 72 ## 79 93 157 ## G ? ? ? L G ? ? ? -23 -71 -71 -55 163 60 204 -19 60 ## 54 156 267 153 ? Ρ E ? ? ## D Ι Q L V S Q L 196 189 324 ## 288 141 296 567 183 181 145 228 236 20 Q ? ? Τ G R ? ? ? V Ρ G ## W 171 145 ## 74 405 486 411 167 216 36 125 324 199 388 ? ? ## L Ρ ? ? ? F F L Α ? 42 486 214 324 324 92 ## 324 216 199 86 86 125 108 ? Ρ ? ? ? ? ## R F Τ Q ? Ι R 343 233 117 ## 341 45 486 223 172 184 106 405 117 100 ## ? L D Y ? ? Е Ρ D Ι F Η E ## 214 209 92 110 567 124 132 405 567 486 149 648 301 ? ## L F G Η V Ρ L L S D Ρ F 567 196 486 ## 186 214 486 648 249 249 108 239 175 27 ? ? ? ? Y G ? ? F Q L Α ## Α 90 199 486 -73 7 196 3 26 ## 324 213 387 286 70 ## ? Α ? ? ? ? ? Ε ? L L Α R ## 124 83 76 29 49 40 71 108 71 292 261 165 244 ## Υ W ? Τ V Ε F G L ? ? ? ? 129 301 405 244 405 439 486 324 173 157 97 ## 567 891 ? ? ? Υ ## K Α G Α G L L S 6 140 285 124 502 486 262 486 244 217 324 ## 92 143 ? ## S F ? Ε L Q Υ С L S D ## 324 52 93 405 77 81 451 261 276 171 183 -117 131 ? ? ? ? ## Ρ ? Ρ F L Ε ? ? 175 149 8 ## 436 104 45 137 163 214 254 131 175 120 ? ? ? Υ F ## Τ Y Ι D F Q Ρ L 190 108 ## 165 0 567 102 288 73 388 405 169 567 382 ? ? ## V? S F ? L ? Ε ? ? R 100 ## 324 182 262 306 118 82 124 62 264 64 95 129 ? ? ? ? ? ? ? ? ## ? D Τ ? ? ## -75 -10 134 92 87 44 68 51 56 18 41 44 61 ? ? ? Ρ ? ? ? ? ? V ? ? ## D ## 79 83 91 161 69 51 12 42 66 -40 78 116 56 ? ? ? ? ? ? ? ? ? ? ? ? ? ## ## 43 -40 -71 -11 -86 -70 -67 37 -8 -38 1 -41-11? ## Ι L Α ?

```
## -28 -87 -7 -5 -44 4 -110 -44 -84 -68 -74 -83 -55
## -
## -47
```

As the above example shows, a substitution matrix must be provided. The result is obviously a vector as long as the alignment has columns. The entries of the vector are labeled by the consensus sequence. The way the consensus sequence is computed can be customized:

```
msaConservationScore(myFirstAlignment, BLOSUM62, gapVsGap=0,
                          type="upperlower", thresh=c(40, 20))
##
                                                     -144
     -80
         -120
               -119
                      -134
                                    -96
                                                -96
                                                                       -120
                                                                               -80
##
                             -96
                                          -80
                                                           -146
                                                                   -96
##
##
     -80
           -96
                 -96
                       -64
                             -64
                                    -64
                                          -80
                                                -80
                                                      -80
                                                            -96
                                                                   -48
                                                                         -96
                                                                               -80
##
##
   -106
           -64
                -121
                       -94
                             -80
                                    -64
                                          -90
                                                -96
                                                     -104
                                                            -96
                                                                   -96
                                                                         -96
                                                                               -64
##
##
     -96
                 -80
                             -80
                                    -96
                                          -64
                                                -96
                                                                               -96
           -96
                       -80
                                                      -96
                                                            -96
                                                                   -98
                                                                         -96
##
##
     -80
           -96
                 -64
                       -80
                              -80
                                    -64
                                          -89
                                               -102
                                                       64
                                                             -96
                                                                   -80
                                                                         -32
                                                                               -96
##
                                              -122
##
     -80
           -96
                 -80
                       -48
                             -96
                                    -80
                                          -96
                                                      -80
                                                            -64
                                                                   -80
                                                                         -48
                                                                               -80
##
##
     -64
           -64
                 -80
                      -116
                             -96
                                    -64
                                         -104
                                                -80
                                                     -113
                                                           -146
                                                                   -48
                                                                       -128
                                                                               -96
##
##
    -135
          -112
                 -96
                       -96
                             -80
                                  -144
                                          -96
                                                -80
                                                      -96
                                                             64
                                                                   -96
                                                                         -64
                                                                               -96
##
##
     -80
           -96
                 -32
                       -80
                             -96
                                   -96
                                          -80
                                                -64
                                                      -80 -112
                                                                   -80
                                                                       -104
                                                                               -80
##
##
     -96
           -48
                  16
                       -64
                             -48
                                    -80
                                          -80
                                                -96
                                                      -80
                                                            -80
                                                                   -75
                                                                         -76
                                                                             -129
##
                                                                                 d
                                                  у
##
     -76
           -91
                   8
                        43
                              -48
                                    -63
                                          -15
                                                 14
                                                      -49
                                                            -62
                                                                   -80
                                                                       -100
                                                                               -10
##
                                                  d
     -90
           -82
                 -43
                             -75
                                                246
                                                       52
                                                                    52
##
                       -58
                                    -36
                                          -60
                                                              74
                                                                          97
                                                                                74
##
             r
                          q
##
           100
                       125
                               60
                                     51
                                          -66
                                              -115
                                                      -71
                                                             -45
                                                                   -28
                                                                         -36
                                                                                48
      -5
                  77
##
                                                               Υ
                                                                     t
                                                                                 е
                         p
                                      p
                                                                           е
            48
                                            6
                                                 21
##
      95
                  -3
                       166
                               30
                                    218
                                                       33
                                                            567
                                                                   141
                                                                         160
                                                                               165
                                            Τ
                                                  1
##
                          t
                                W
       е
                                                                     r
                                                                           q
##
     325
           179
                  71
                       216
                             891
                                     38
                                          405
                                                204
                                                      208
                                                              79
                                                                   165
                                                                         109
                                                                                42
##
             1
                                            Α
                                                  С
                                                               е
##
      79
           249
                 113
                       106
                               93
                                   172
                                          324
                                                536
                                                      109
                                                            324
                                                                  261
                                                                          72
                                                                               157
##
                                                         1
       g
                                                               g
##
      54
           156
                 -48
                       -96
                             -96
                                    -80
                                          163
                                                 60
                                                      204
                                                            267
                                                                   153
                                                                         -20
                                                                                60
##
                   i
                         Ρ
                                      1
                                                                                 L
                                q
                                            е
                                                         V
                                                               S
```

```
##
     288
           141
                  296
                        567
                                     196
                                           181
                                                               236
                                                                      189
                                                                                  324
                               183
                                                  145
                                                         228
                                                                             20
##
                          Τ
                                 G
                                                                        V
       q
                                        W
                                              r
                                                                              р
                                                                                     g
##
     171
            74
                  145
                        405
                               486
                                     411
                                           167
                                                  216
                                                          36
                                                               125
                                                                      324
                                                                            199
                                                                                  388
##
       L
                                              F
                                                    f
                                                                        L
                                                                              Α
                    p
##
     324
           216
                  199
                         86
                                86
                                      42
                                           486
                                                  214
                                                        125
                                                               108
                                                                     324
                                                                            324
                                                                                    92
##
                    F
                                        t
                                                           i
                                                                 R
       r
                          p
                                              q
                               172
                                                         233
                                                               405
##
     341
             45
                  486
                        223
                                     184
                                           106
                                                  343
                                                                      117
                                                                            117
                                                                                   100
              1
                          Υ
                                              Е
                                                    Ρ
                                                           D
                                                                 i
                                                                        f
                                                                              Η
##
                    d
                                                                                     е
                  110
                               124
                                                  567
##
     209
            92
                        567
                                     132
                                           405
                                                        486
                                                               149
                                                                     214
                                                                            648
                                                                                  301
##
       1
              f
                    G
                          Η
                                        Ρ
                                              1
                                                    1
                                                                 d
                                                                                     F
                                 V
                                                           S
                                                                        р
##
     186
           214
                  486
                        648
                               249
                                     567
                                            249
                                                  196
                                                        108
                                                               239
                                                                      175
                                                                             27
                                                                                  486
##
       Α
                    f
                                                    G
                                                                  1
                                 q
                                              У
                                                                        а
##
     324
           213
                  387
                         90
                               286
                                      70
                                           199
                                                  486
                                                         -89
                                                                 7
                                                                      196
                                                                              3
                                                                                    26
                                                                 1
##
                                                                                     1
              а
                                                                              r
                                                     е
                                                                        а
            83
##
     124
                   76
                         29
                                48
                                      40
                                             71
                                                  108
                                                          71
                                                               292
                                                                     261
                                                                            165
                                                                                  244
##
       Υ
              W
                          Τ
                                        Ε
                                              f
                                                    G
                                                           L
                                 V
##
     567
           891
                  301
                        405
                               244
                                     405
                                            439
                                                  486
                                                        324
                                                               173
                                                                      157
                                                                            129
                                                                                    97
##
                                 k
                                        а
                                              У
                                                     G
                                                           а
                                                                 G
                                                                        1
                                                                               1
                                                                                     S
     -58
            92
                  143
                                     124
                                           502
                                                  486
                                                        262
                                                               486
                                                                     244
                                                                            217
                                                                                  324
##
                        140
                               285
##
       S
              f
                          Ε
                                 1
                                                           1
                                                                        d
                                                    С
                                                                  S
                                        q
                                              У
                                77
##
     324
            52
                   93
                        405
                                      81
                                            451
                                                  261
                                                        276
                                                               171
                                                                      183
                                                                          -142
                                                                                   131
##
                                        f
                                                    1
                                                           е
                                 p
                                                                              m
       p
##
     436
           104
                   45
                        137
                               163
                                     214
                                            254
                                                  175
                                                         131
                                                               175
                                                                      120
                                                                            149
                                                                                     8
                                                                                     f
                    Υ
                                 i
                                                    f
                                                                               Υ
##
       t
                                        d
                                                           Q
                                                                 p
                                                                        1
                                                        405
     165
              0
                  567
                        102
                                     190
                                             73
                                                  388
                                                               169
                                                                      108
                                                                                  382
##
                               288
                                                                            567
       V
##
                           S
                                 f
                                                    1
                                                                  е
                                                                                     r
     324
                                                  124
##
           100
                  182
                        262
                               306
                                     118
                                             82
                                                          62
                                                               264
                                                                       64
                                                                             95
                                                                                   129
##
                    d
                                 t
                  134
                         92
                                                          56
##
     -91
           -10
                                87
                                      44
                                             68
                                                   51
                                                                18
                                                                       41
                                                                             44
                                                                                    61
##
                                                                               d
                                                                 V
                    p
##
      79
            83
                   90
                        161
                                56
                                      69
                                             51
                                                   12
                                                          42
                                                                65
                                                                      -41
                                                                             77
                                                                                   115
##
                  -44
                                     -90
                                                  -76
                                                          33
##
      42
              0
                        -75
                               -15
                                            -74
                                                               -12
                                                                      -45
                                                                            -42
                                                                                   -15
##
                           i
                                 1
                                                     а
##
     -32
           -91
                  -11
                         -9
                               -60
                                     -12 -126
                                                  -60 -100
                                                               -84
                                                                      -90 -108
                                                                                  -80
##
##
     -96
```

The additional argument gapVsGap allows for controlling how pairs of gap are taken into account when computing pairwise scores (see ?msaConservationScore for more details).

Conservation scores can also be computed from masked alignments. For masked columns, NA's are returned:

##	#	#	#	#	#	#	#	#	#	#	#	#	#
##	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
##	#	#	#	#	#	#	#	#	#	#	#	#	#
##	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
##	#	#	#	#	#	#	#	#	#	#	#	#	#
##	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
##	#	#	#	#	#	#	#	#	#	#	#	#	#
##	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
##	#	#	#	#	#	#	#	#	#	#	#	#	#
##	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
##	#	#	#	#	#	#	#	#	#	#	#	#	#
##	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
##	#	#	#	#	#	#	#	#	#	#	#	#	#
##	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
##	#	#	#	#	#	#	#	#	#	_	_	_	_
##	ΝA	NA	NA	NA	NA	NA	NA	NA	NA	-64	-96	-64	-96
##	_	_	_	_	_	_	_	_	_	_	_	_	_
##	-80	-96	-32	-80	-96	-96	-80	-64		-112			-80
##	_	_	_	_	_	_	_	_	_	_	•		
##	-96	-48	16	-64	-48	-80	-80	-96	-80		-75	-76	-129
##								у					
##	-76	-91	8	43	-48	-63	-15	14	-49	-62		-100	
##								d					
##	-90	-82	-43	-58	-75	-36	-60	246	52	74	52	97	74
##		r											
##	-5	100	77	q 125	60	51		-115	-71	-45	-28	-36	48
##				p		р				Y	t	е	e
##	95	48	-3	166	30	218	6		33	567	141	160	165
##	e	•		t	W	210	Т	1			r		
##	325	179	71	216	891	38	405	204	208	79	165	q 109	42
##		1	, ,				403 A	204 C	200	e			
##	79	249	113	106	93	172	324	536	109	324	261	72	157
##		249			93					324 g	201		
##	g 54		-48					60	204				
##	d		-40 i	-90 P			103 e	•					L
##	288	141	296	567	q 183	196	181	145	v 228	s 236	q 189		324
				307 T									
##	q 171	7/	1/5		G 186	W ⊿11	167			125	3.3.4 V	100	300 g
##	171 T	74	145	405	486	411	167	216	36	125	324	199	388
##	L		p 100	. 06	. 06	40	F	f	105		204 L	A 224	
##	324	216	199	86	86	42	486	214	125	108	324	324	92
##	r		F	р	170	t	q	242	i	R	117		100
##	341	45	486	223	172	184	106	343	233	405	117	117	100
##		1	d	Υ	104		E	P	D	i	f	H	e
##	209	92	110	567	124	132	405	567	486	149	214	648	301
##	1	f	G	H	V	P	1	1	S	d	p		
##	186	214	486	648	249	567	249	196	108	239	175	27	486

```
##
                     f
                                                      G
        Α
                                                У
                                  q
##
     324
            213
                  387
                          90
                                286
                                        70
                                             199
                                                    486
                                                          -89
                                                                    7
                                                                        196
                                                                                 3
                                                                                      26
                                                                    1
##
                                                                                        1
              а
                                                      е
                                                                          а
                                                                                 r
                                                                 292
##
     124
             83
                   76
                          29
                                 48
                                        40
                                              71
                                                    108
                                                            71
                                                                        261
                                                                              165
                                                                                     244
##
        Y
              W
                           Τ
                                  V
                                         Ε
                                               f
                                                      G
                                                             L
                                             439
                                                    486
                                                          324
                                                                                      97
##
     567
            891
                  301
                         405
                                244
                                      405
                                                                 173
                                                                        157
                                                                              129
                                                                    G
                                                                                 1
##
                                  k
                                                      G
                                                                          1
                                                                                        S
                                         а
                                               У
                                                             а
                         140
                                                          262
                                                                              217
     -58
                  143
                                285
                                      124
                                             502
                                                    486
                                                                 486
                                                                        244
                                                                                     324
##
             92
##
        S
              f
                           Ε
                                  1
                                                      С
                                                             1
                                                                    S
                                                                          d
                                               У
                                         q
##
     324
             52
                   93
                         405
                                 77
                                        81
                                             451
                                                    261
                                                          276
                                                                 171
                                                                        183
                                                                             -142
                                                                                     131
##
                                         f
                                                      1
       p
                                  p
                                                             е
                                                                                 m
                         137
                                      214
                                             254
                                                                                        8
##
     436
            104
                   45
                                163
                                                    175
                                                          131
                                                                 175
                                                                        120
                                                                              149
##
        t
                     Υ
                                  i
                                         d
                                                      f
                                                             Q
                                                                    p
                                                                          1
                                                                                 Y
                                                                                        f
     165
                  567
                         102
                                288
                                      190
                                              73
                                                    388
                                                          405
                                                                 169
                                                                        108
                                                                              567
                                                                                     382
##
              0
##
        V
                                  f
                                                      1
                            S
                                                                    6
                                                                                        r
##
     324
            100
                  182
                         262
                                306
                                      118
                                              82
                                                    124
                                                            62
                                                                 264
                                                                         64
                                                                                95
                                                                                     129
##
                     d
                                  t
##
     -91
            -10
                  134
                          92
                                 87
                                        44
                                              68
                                                     51
                                                            56
                                                                  18
                                                                         41
                                                                                44
                                                                                      61
##
                     р
                                                                    V
                                                                                 d
      79
             83
                   90
                         161
                                 56
                                        69
                                              51
                                                     12
                                                            42
                                                                  65
                                                                                77
                                                                                     115
##
                                                                        -41
##
##
      42
              0
                   -44
                         -75
                                -15
                                      -90
                                             -74
                                                    -76
                                                            33
                                                                 -12
                                                                        -45
                                                                               -42
                                                                                     -15
##
                            i
                                  1
                                                      а
     -32
##
                          -9
                                -60
                                      -12
                                           -126
                                                    -60
            -91
                  -11
                                                         -100
                                                                 -84
                                                                        -90
                                                                             -108
                                                                                     -80
##
     -96
##
```

6.3 Interfacing to Other Packages

There are also other sequence analysis packages that use or make use of multiple sequence alignments. The msa package does not directly interface to these packages in order to avoid dependencies and possible incompatibilities. However, msa provides a function msaConvert() that allows for converting multiple sequence alignment objects to other types/classes. Currently, five such conversions are available, namely to the classes alignment (seqinr package [2]), align (bios2mds package [14]), AAbin/DNAbin (ape package [10]), and phyDat (phangorn package [11]). Except for the conversion to the class phyDat, these conversion are performed without loading or depending on the respective packages.

In the following example, we perform a multiple alignment of Hemoglobin alpha example sequences and convert the result for later processing with the seqinr package:

```
hemoAln
## CLUSTAL 2.1
##
## Call:
     msa(hemoSeq)
##
##
## MsaAAMultipleAlignment with 17 rows and 143 columns
##
                                             names
   [1] -VLSPADKTNVKAAWGKV...LDKFLASVSTVLTSKYR HBA1_Homo_sapiens
##
##
   [2] MVLSPADKTNVKAAWGKV...LDKFLASVSTVLTSKYR HBA1_Pan_troglodytes
   [3] -VLSPADKSNVKAAWGKV...LDKFLASVSTVLTSKYR HBA1_Macaca_mulatta
    [4] -VLSAADKGNVKAAWGKV...LDKFLANVSTVLTSKYR HBA1_Bos_taurus
##
   [5] -VLSPADKTNVKGTWSKI...LDKFLASVSTVLTSKYR HBA1_Tursiops_tru...
##
   [6] -VLSGEDKSNIKAAWGKI...LDKFLASVSTVLTSKYR HBA1_Mus_musculus
##
##
   [7] MVLSADDKTNIKNCWGKI...LDKFLASVSTVLTSKYR HBA1_Rattus_norve...
##
   [8] -VLSATDKANVKTFWGKL...LDKFLATVATVLTSKYR HBA1_Erinaceus_eu...
##
   [9] -VLSAADKSNVKACWGKI...LDKFFSAVSTVLTSKYR HBA1_Felis_silves...
## [10] -VLSPADKTNIKSTWDKI...LDKFFTAVSTVLTSKYR HBA1_Chrysocyon_b...
## [11] -VLSDNDKTNVKATWSKV...LDKFLSNVSTVLTSKYR HBA1_Loxodonta_af...
## [12] -VLSAADKTNVKAAWSKV...LDKFLALLSTVLTSKYR HBA1_Monodelphis_...
## [13] -MLTDAEKKEVTALWGKA...MDKFLSKVATVLTSKYR HBA1_Ornithorhync...
## [14] -VLSAADKNNVKGIFTKI...LDKFLCAVGTVLTAKYR HBA1_Gallus_gallus
## [15] -HLTADDKKHIKAIWPSV...LDKFLVSVSNVLTSKYR HBA1_Xenopus_trop...
## [16] -VLTEEDKARVRVAWVPV...VDKFLGQISKVLASRYR HBA1_Microcephalo...
## Con -VLS?ADK?NVKA?WGK?...LDKFLA?VSTVLTSKYR Consensus
hemoAln2 <- msaConvert(hemoAln, type="seqinr::alignment")
```

Now we compute a distance matrix using the dist.alignment() function from the seqinr package:

```
library(seqinr)

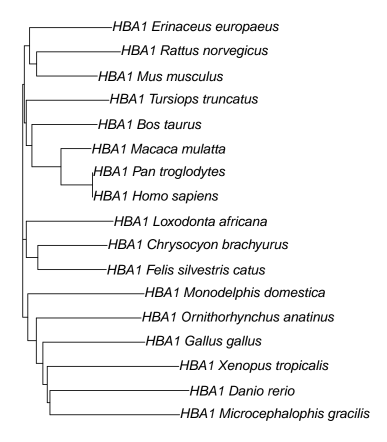
d <- dist.alignment(hemoAln2, "identity")
as.matrix(d)[2:5, "HBA1_Homo_sapiens", drop=FALSE]

## HBA1_Homo_sapiens
## HBA1_Pan_troglodytes 0.0000000
## HBA1_Macaca_mulatta 0.1684304
## HBA1_Bos_taurus 0.3472281
## HBA1_Tursiops_truncatus 0.4038819</pre>
```

Now we can construct a phylogenetic tree with the neighbor joining algorithm using the nj () function from the ape package:

```
library(ape)
hemoTree <- nj(d)
plot(hemoTree, main="Phylogenetic Tree of Hemoglobin Alpha Sequences")</pre>
```

Phylogenetic Tree of Hemoglobin Alpha Sequences



The following example shows how to convert a multiple alignment object in an object of class align as defined by the bios2mds package:

```
hemoAln3 <- msaConvert(hemoAln, type="bios2mds::align")
str(hemoAln3)

## List of 17
## $ HBA1_Homo_sapiens : chr [1:143] "-" "V" "L" "S" ...
## $ HBA1_Pan_troglodytes : chr [1:143] "M" "V" "L" "S" ...
```

```
: chr [1:143] "-" "V" "L" "S"
   $ HBA1_Macaca_mulatta
                                   : chr [1:143] "-" "V" "L" "S"
##
   $ HBA1_Bos_taurus
                                  : chr [1:143] "-" "V" "L" "S"
   $ HBA1_Tursiops_truncatus
##
                                  : chr [1:143] "-" "V" "L" "S"
   $ HBA1_Mus_musculus
##
##
   $ HBA1_Rattus_norvegicus
                                  : chr [1:143] "M" "V" "L" "S"
   $ HBA1_Erinaceus_europaeus : chr [1:143] "-" "V" "L" "S"
##
                                 : chr [1:143] "-" "V" "L" "S"
##
   $ HBA1_Felis_silvestris_catus
   $ HBA1_Chrysocyon_brachyurus : chr [1:143] "-" "V" "L" "S"
##
                                  : chr [1:143] "-" "V" "L" "S"
   $ HBA1_Loxodonta_africana
##
## $ HBA1_Monodelphis_domestica : chr [1:143] "-" "V" "L" "S"
## $ HBA1_Ornithorhynchus_anatinus : chr [1:143] "-" "M" "L" "T"
                                                 "-" "V" "L" "S"
   $ HBA1_Gallus_gallus
                                   : chr [1:143]
##
                                  : chr [1:143] "-" "H" "L" "T"
## $ HBA1_Xenopus_tropicalis
   $ HBA1_Microcephalophis_gracilis: chr [1:143] "-" "V" "L" "T"
                                   : chr [1:143] "-" "S" "L" "S"
## $ HBA1_Danio_rerio
## - attr(*, "class")= chr "align"
```

The conversions to the standard Biostrings classes are straightforward using standard as() methods and not provided by the msaConvert() function. The following example converts a multiple alignment object to class BStringSet (e.g. the msaplot() function from the ggtree package [16] accepts BStringSet objects):

```
hemoAln4 <- as(hemoAln, "BStringSet")
hemoAln4
## BStringSet object of length 17:
##
       width seq
                                                names
##
   [1]
        143 -VLSPADKTNVKAAW...KFLASVSTVLTSKYR HBA1_Homo_sapiens
##
   [2] 143 MVLSPADKTNVKAAW...KFLASVSTVLTSKYR HBA1_Pan_troglodytes
##
    [3]
         143 -VLSPADKSNVKAAW...KFLASVSTVLTSKYR HBA1 Macaca mulatta
##
  [4]
         143 -VLSAADKGNVKAAW...KFLANVSTVLTSKYR HBA1_Bos_taurus
## [5]
         143 -VLSPADKTNVKGTW...KFLASVSTVLTSKYR HBA1_Tursiops_tru...
##
## [13]
         143 -MLTDAEKKEVTALW...KFLSKVATVLTSKYR HBA1_Ornithorhync...
         143 -VLSAADKNNVKGIF...KFLCAVGTVLTAKYR HBA1_Gallus_gallus
## [14]
## [15]
         143 -HLTADDKKHIKAIW...KFLVSVSNVLTSKYR HBA1_Xenopus_trop...
          143 -VLTEEDKARVRVAW...KFLGQISKVLASRYR HBA1_Microcephalo...
## [16]
         143 -SLSDTDKAVVKAIW...KFFNNLALALSEKYR HBA1_Danio_rerio
## [17]
```

Note: The msaConvert() function has been introduced in version 1.3.3 of the msa package. So, to have this function available, at least Bioconductor 3.3 is required, which requires at least R 3.3.0.

7 Pretty-Printing Multiple Sequence Alignments

As already mentioned above, the msa package offers the function msaPrettyPrint() which allows for pretty-printing multiple sequence alignments using the LATEX package TeXshade [1]. Which prerequisites are necessary to take full advantage of the msaPrettyPrint() function is described in Section 2.

The msaPrettyPrint() function writes a multiple sequence alignment to an alignment (.aln) file and then creates LATEX code for pretty-printing the multiple sequence alignment on the basis of the LATEX package TEXSMace. Depending on the choice of the output argument, the function msaPrettyPrint() either prints a LATEX fragment to the R session (choice output="asis") or writes a LATEX source file (choice output="tex") that it processes to a DVI file (choice output="dvi") or PDF file (choice output="pdf"). Note that no extra software is needed for choices output="asis" and output="tex". For output="dvi" and output="pdf", however, a TEX/LATEX distribution must be installed in order to translate the LATEX source file into the desired target format (DVI or PDF).

The function msaPrettyPrint() allows for making the most common settings directly and conveniently via an R interface without the need to know the details of LATEX or TEX shade. In the following, we will describe some of these customizations. For all possibilities, the user is referred to the documentation of TEX shade.⁵

7.1 Consensus Sequence and Sequence Logo

The consensus sequence of the alignment is one of the most important results of a multiple sequence alignment. msaPrettyPrint() has a standard possibility to show this consensus sequence with the parameter showConsensus. The default value is "bottom", which results in the following:

```
IAYNYRHGQPIPRVEYTEEEKQTWGTVFRTLKALYKTHACYEHNHIFPLL 213
IAYNYRHGQPIPRVEYTEEERKTWGTVFRTLKALYKTHACYEHNHIFPLL 213
IAYNYRHGQPIPRVEYMEEEKKTWGTVFKTLKSLYKTHACYEYNHIFPLL 213
IAYNYRHGQPIPRVEYTEEEKKTWGTVFRTLKSLYKTHACYEYNHIFPLL 212
....DFTLPQPLDRYSAEDHATWATLYQRQCKLLPGRACDEFMEGL... 67
QTLRPDFTMEQPVHRYTAADHATWRTLYDRQEALLPGRACDEFLQGL... 83
```

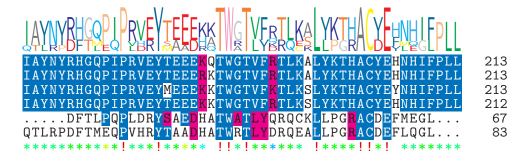
Consensus sequences can also be displayed on top of a multiple sequence alignment or omitted completely.

⁵https://www.ctan.org/pkg/texshade

In the above example, an exclamation mark '!' in the consensus sequence stands for a conserved letter, i.e. a sequence positions in which all sequences agree, whereas an asterisk '*' stands for positions in which there is a majority of sequences agreeing. Positions in which the sequences disagree are left blank in the consensus sequence. For a more advanced example how to customize the consensus sequence, see the example in Subsection 7.4 below.

The color scheme of the consensus sequence can be configured with the consensusColors parameter. Possible values are "ColdHot", "HotCold", "BlueRed", "RedBlue", "GreenRed", "RedGreen", or "Gray". The above example uses the color scheme "RedGreen".

Additionally, msaPrettyPrint() also offers a more sophisticated visual representation of the consensus sequence — sequence logos. Sequence logos can be displayed either on top of the multiple sequence alignment (showLogo="top"), below the multiple sequence alignment (showLogo="bottom"), or omitted at all (showLogo="none"):



The color scheme of the sequence logo can be configured with the logoColors parameter. Possible values are "chemical", "rasmol", "hydropathy", "structure", "standard area", and "accessible area". The above example uses the color scheme "rasmol".

Note that a consensus sequence and a sequence logo can be displayed together, but only on opposite sides.

Finally, a caveat: for computing consensus sequences, msaPrettyPrint() uses the functionality provided by TeXshade, therefore, the results need not match to the results of the methods described in Section 6 above.

7.2 Color Shading Modes

TeXshade offers different shading schemes for displaying the multiple sequence alignment itself. The following schemes are available: "similar", "identical", and "functional". Moreover, there are five different color schemes available for shading: "blues", "reds", "greens", "grays", or "black". The following example uses the shading mode "similar" along with the color scheme "blues":

83

58 50

65

```
msaPrettyPrint(myFirstAlignment, output="asis", y=c(164, 213),
                         showNames="none", shadingMode="similar",
                         shadingColors="blues", showLogo="none",
                         showLegend=FALSE, askForOverwrite=FALSE)
    IAYNYRHGQPIPRVEYTEEEKQTWGTVFRTLKALYKTHACYEHNHIFPLL
IAYNYRHGQPIPRVEYTEEERKTWGTVFRTLKALYKTHACYEHNHIFPLL
                                                                                                          213
                                                          RTLKALYKTHACYEHNHIFPLL
                                                                                                          213
    IAYNYRHGQPIPRVEYMEEEKKTWGTVF
                                                          KTLKS<mark>LYK</mark>TH<mark>AC</mark>YE<mark>Y</mark>NHIFPLL
                                                                                                          213
    IAYNYRHGQPIPRVEYTEEEKKTWGTVFRTLKSLYKTHACYEHNHIFPLL ....DFTLPQPLDRYSAEDHATWATLYQRQCKLLPGRACDEFMEGL ...
                                                                                                          212
                                                                                                           67
    QTLRPDFTMEQPVHRYTAADHATWRTLYDRQEALLPGRACDEFLQGL...

OTLRPDFTMEQPVHRYTAADHATWRTLYDRQEALLPGRACDEFLQGL...

PDWTIDQGWETYTQAEHDVWITLYERQTDMLHGRACDEFMRGL...

ONGFTHYPETEHQVWNTLITRQLKVIEGRACQEYLDGI...

VCRADFTVAQDYD.YSDEEQAVWRTLCDRQTKLTRKLAHHSYLDGV...
                                                                                                           83
                                                                                                           58
                                                                                                           50
```

If the shading modes "similar" or "identical" are used, the shadingModeArg argument allows for setting a similarity threshold (a numerical value between 0 and 100). For shading mode "functional", the following settings of the shadingModeArg argument are possible: "charge", "hydropathy", "structure", "hemical", "rasmol", "standard area", and "accessible area". The following example uses shading mode "functional" along with shadingModeArg set to "structure":

```
msaPrettyPrint(myFirstAlignment, output="asis", y=c(164, 213),
showNames="none", shadingMode="functional",
shadingModeArg="structure",
askForOverwrite=FALSE)

IAYNYRHGQPIPRVEYTEEEKQTWGTVFRTLKALYKTHACYEHNHIEPLL 213
IAYNYRHGQPIPRVEYTEEEKKTWGTVFRTLKALYKTHACYEHNHIEPLL 213
IAYNYRHGQPIPRVEYMEEEKKTWGTVFKTLKSLYKTHACYEYHNHIEPLL 213
IAYNYRHGQPIPRVEYMEEEKKTWGTVFKTLKSLYKTHACYEYHNHIEPLL 213
IAYNYRHGQPIPRVEYTEEEKKTWGTVFRTLKSLYKTHACYEHNHIEPLL 212
.....DFTLPQPLDRYSAEDHATWATLYQRQCKLLPGRACDEFMEGL... 67
```

QTLRPDFTMEQPVHRYTAADHATWRTLYDRQEALLPGRACDEFLQGL...

....PDWTIDQGWETYTQAEHDVWITLYERQTDMLHGRACDEFMRGL...
.....DNGFIHYPETEHQVWNTLITRQLKVIEGRACQEYLDGI...

. VCRADFTVAQDYD. YSDEEQAVWRTLCDRQTKLTRKLAHHSYLDGV...

X external
X ambivalent
X internal

In the above example, a legend is shown that specifies the meaning of the color codes with which the letters are shaded. In some of the other examples above, we have suppressed this legend with the option showLegend=FALSE. The default, however, is that a legend is printed underneath the multiple sequence alignment like in the previous example.

7.3 Subsetting

In case that not the complete multiple sequence alignment should be printed, msaPrettyPrint() offers two ways of sub-setting. On the one hand, the subset argument allows for selecting only a subset of sequences. Not surprisingly, subset must be a numeric vector with indices of sequences to be selected. On the other hand, it is also possible to slice out certain positions of the multiple sequence alignment using the y argument. In the simplest case, y can be a numeric vector with two elements in ascending order which correspond to the left and right bounds between which the multiple sequence alignment should be displayed. However, it is also possible to slice out multiple windows. For this purpose, the argument y must be an IRanges object containing the starts and ends of the windows to be selected.

7.4 Additional Customizations

The msaPrettyPrint() function provides an interface to the most common functionality of TeXshade in a way that the user does not need to know the specific commands of TeXshade. TeXshade, however, provides a host of additional customizations many of which are not covered by the interface of the msaPrettyPrint() function. In order to allow users to make use of all functionality of TeXshade, msaPrettyPrint() offers the furtherCode argument through which users can add LaTeX code to the texshade environment that is created by msaPrettyPrint(). Moreover, the code argument can be used to bypass all of msaPrettyPrint()'s generation of TeXshade code.

Here is an example how to use the furtherCode argument in order to customize the consensus sequence and to show a ruler on top:

```
IAYNYRHGQPIPRVEYTEEEKQTWGTVFRTLKALYKTHACYEHNHIFPLL 213
IAYNYRHGQPIPRVEYTEEERKTWGTVFRTLKALYKTHACYEHNHIFPLL 213
IAYNYRHGQPIPRVEYMEEEKKTWGTVFKTLKSLYKTHACYEYNHIFPLL 213
IAYNYRHGQPIPRVEYTEEEKKTWGTVFRTLKSLYKTHACYEHNHIFPLL 212
....DFTLPQPLDRYSAEDHATWATLYQRQCKLLPGRACDEFMEGL... 67
QTLRPDFTMEQPVHRYTAADHATWRTLYDRQEALLPGRACDEFLQGL... 83
iaynyrhgqpiPrveYteeek.TWgTvfrtlk.LykthACyE.nhifpll
```

7.5 Sweave or knitr Integration

The function msaPrettyPrint() is particularly well-suited for pretty-printing multiple alignments in Sweave [6] or knitr [15] documents. The key is to set output to "asis" when calling

msaPrettyPrint() and, at the same time, to let the R code chunk produce output that is directly included in the resulting LATEX document as it is. This can be accomplished with the code chunk option results="tex" in Sweave and with the code chunk option results="asis" in knitr. Here is an example of a Sweave code chunk that displays a pretty-printed multiple sequence alignment inline:

```
<<AnyChunkName,results="tex">>=
msaPrettyPrint(myFirstAlignment, output="asis")
@
```

The same example in knitr:

```
<<AnyChunkName,results="asis">>=
msaPrettyPrint(myFirstAlignment, output="asis")
@
```

Note that, for processing the resulting LATEX source document, the TEX shade package must be installed (see Section 2) and the TEX shade package must be loaded in the preamble:

```
\usepackage{texshade}
```

7.6 Sequence Names

The Biostrings package does not impose any restrictions on the names of sequences. Consequently, msa also allows all possible ASCII strings as sequence (row) names in multiple alignments. As soon as msaPrettyPrint() is used for pretty-printing multiple sequence alignments, however, the sequence names are interpreted as plain LATEX source code. Consequently, LATEX errors may arise because of characters or words in the sequence names that LATEX does not or cannot interpret as plain text correctly. This particularly includes appearances of special characters and backslash characters in the sequence names.

The msa package offers a function msaCheckNames() which allows for finding and replacing potentially problematic characters in the sequence names of multiple alignment objects (see ?msaCheckNames). However, the best solution is to check sequence names carefully and to avoid problematic sequence names from the beginning. Note, moreover, that too long sequence names will lead to less appealing outputs, so users are generally advised to consider sequence names carefully.

7.7 Pretty-Printing Wide Alignments

If the alignment to be printed with msaPrettyPrint() is wide (thousands of columns or wider), LATEX may terminate prematurely because of exceeded TEX capacity. Unfortunately, this problem remains opaque to the user, since texi2dvi() and texi2pdf() do not convey much details about LATEX problems when typesetting a document. We recommend the following if a user encounters problems with running msaPrettyPrint()'s output with texi2dvi() and texi2pdf():

8 Known Issues 39

1. Run pdflatex on the generated .tex file to see whether it is actually a problem with TeX capacity.

2. If so, split the alignment into multiple chunks and run msaPrettyPrint() on each chunk separately.

The following example demonstrates this approach for a multiple alignment object 'aln':

This creates multiple PDF files all of which show one part of the alignment. Please note, however, that the numbering of columns is restarted for each chunk.

7.8 Further Caveats

- Note that texi2dvi() and texi2pdf() always save the resulting DVI/PDF files to the current working directory, even if the LATEX source file is in a different directory. That is also the reason why the temporary file is created in the current working directory in the example below.
- TeXshade has a wide array of functionalities. Only the most common ones have been tested for interoperability with R. So the use of the arguments furtherCode and code is the user's own risk!

8 Known Issues

Memory Leaks

The original implementations of ClustalW, ClustalOmega, and MUSCLE are stand-alone command line programs which are only run once each time a multiple sequence alignment is performed. During the development of the msa package, we performed memory management checks using Valgrind [8] and discovered multiple memory leaks in ClustalW and MUSCLE. These memory leaks have no effect for the command line tools, since the program is closed each time the alignment is finished. In the implementation of the msa package, however, these memory leaks may have an effect if the same algorithm is run multiple times.

For MUSCLE, we managed to eliminate all memory leaks by deactivating the two parameters weight1 and weight2. ClustalOmega did not show any memory leaks. ClustalW indeed has several memory leaks which are benign if the algorithm is run only a few times, but which may have more severe effects if the algorithm is run many times. ClustalOmega also has a minor memory leak, but the loss of data is so small that no major problems are to be expected except for thousands of executions of ClustalOmega.

ClustalOmega vs. Older GCC Versions on Linux/Unix

We have encountered peculiar behavior of ClustalOmega if the package was built using an older GCC version: if we built the package on an x86_64 Linux system with GCC 4.4.7, ClustalOmega built smoothly and could be executed without any errors. However, the resulting multiple sequence alignment was more than sub-optimal. We could neither determine the source of this problem nor which GCC versions show this behavior. We therefore recommend Linux/Unix users to use an up-to-date GCC version (we used 4.8.2 during package development, which worked nicely) or, in case they encounter dubious results, to update to a newer GCC version and re-install the package.

ClustalOmega: OpenMP Support on Mac OS

ClustalOmega is implemented to make use of OpenMP (if available on the target platform). Due to issues on one of the Bioconductor build servers running Mac OS, we had to deactivate OpenMP generally for Mac OS platforms. If a Mac OS user wants to re-activate OpenMP, he/she should download the source package tarball, untar it, comment/uncomment the corresponding line in msa/src/ClustalOmega/msaMakefile (see first six lines), and build/install the package from source.

Build/installation issues

Some users have reported compiler and linker errors when building msa from source on Linux systems. In almost all cases, these could have been tracked down to issues with the R setup on those systems (e.g. a Rprofile.site file that makes changes to the R environment that are not compatible with msa's Makefiles). In most cases, these issues can be avoided by installing msa in a "vanilla R session", i.e. starting R with option --vanilla when installing msa.

9 Future Extensions

We envision the following changes/extensions in future versions of the package:

- Integration of more multiple sequence alignment algorithms, such as, T-Coffee [9] or DI-ALIGN [7]
- Support for retrieving guide trees from the multiple sequence alignment algorithms

⁶See, e.g., https://support.bioconductor.org/p/90735/

- Interface to methods computing phylogenetic trees (e.g. as contained in the original implementation of ClustalW)
- Elimination of memory leaks described in Section 8 and re-activation of parameters that have been deactivated in order to avoid memory leaks
- More tolerant handling of custom substitution matrices (MUSCLE interface)

10 How to Cite This Package

If you use this package for research that is published later, you are kindly asked to cite it as follows:

U. Bodenhofer, E. Bonatesta, C. Horejš-Kainrath, and S. Hochreiter (2015). msa: an R package for multiple sequence alignment. *Bioinformatics* **31**(24):3997–3999. DOI: bioinformatics/btv494.

To obtain a BibT_EX entries of the reference, enter the following into your R session:

```
toBibtex(citation("msa"))
```

Moreover, we insist that, any time you cite the package, you also cite the original paper in which the original algorithm has been introduced (see bibliography below).

References

- [1] E. Beitz. TeXshade: shading and labeling of multiple sequence alignments using LATeX2e. *Bioinformatics*, 16(2):135–139, 2000.
- [2] 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 U. Bastolla, M. Porto, H. E. Roman, and M. Vendruscolo, editors, *Structural approaches to sequence evolution: Molecules, networks, populations*, Biological and Medical Physics, Biomedical Engineering, pages 207–232. Springer, New York, 2007.
- [3] R. C. Edgar. MUSCLE: a multiple sequence alignment method with reduced time and space complexity. *BMC Bioinformatics*, 5(5):113, 2004.
- [4] R. C. Edgar. MUSCLE: multiple sequence alignment with high accuracy and high throughput. *Nucleic Acids Res.*, 32(5):1792–1797, 2004.
- [5] L. Lamport. ETeX A Document Preparation System. User's Guide and Reference Manual. Addison-Wesley Longman, Amsterdam, 1999.
- [6] F. Leisch. Sweave: dynamic generation of statistical reports using literate data analysis. In W. Härdle and B. Rönz, editors, *Compstat 2002 Proceedings in Computational Statistics*, pages 575–580, Heidelberg, 2002. Physica-Verlag.

42 References

[7] B. Morgenstern. DIALIGN 2: improvement of the segment-to-segment approach to multiple sequence alignment. *Bioinformatics*, 15(3):211–218, 1999.

- [8] N. Nethercote and J. Seward. Valgrind: A framework for heavyweight dynamic binary instrumentation. In *Proc. of the ACM SIGPLAN 2007 Conf. on Programming Language Design and Implementation*, San Diego, CA, 2007.
- [9] C. Notredame, D. G. Higgins, and J. Heringa. T-Coffee: A novel method for fast and accurate multiple sequence alignment. *J. Mol. Biol.*, 302(1):205–217, 2000.
- [10] E. Paradis, J. Claude, and K. Strimmer. APE: analyses of phylogenetics and evolution in R language. *Bioinformatics*, 20:289–290, 2004.
- [11] K. P. Schliep. phangorn: phylogenetic analysis in R. Bioinformatics, 27(4):592–593, 2011.
- [12] F. Sievers, A. Wilm, D. Dineen, T. J. Gibson, K. Karplus, W. Li, R. Lopez, H. McWilliam, M. Remmert, J. Söding, J. D. Thompson, and D. G. Higgins. Fast, scalable generation of high-quality protein multiple sequence alignments using Clustal Omega. *Mol. Syst. Biol.*, 7:539, 2011.
- [13] J. D. Thompson, D. G. Higgins, and T. J. Gibson. CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. *Nucleic Acids Res.*, 22(22):4673–4680, 2004.
- [14] J. Pele with J.-M. Becu, H. Abdi, and M. Chabbert. *bios2mds: From BIOlogical Sequences to MultiDimensional Scaling*, 2012. R package version 1.2.2.
- [15] Y. Xie. Dynamic Documents with R and knitr. Chapman & Hall/CRC, 2014.
- [16] G. Yu, D. Smith, H. Zhu, Y. Guan, and T. T. Y. Lam. ggtree: an R package for visualization and annotation of phylogenetic tree with different types of meta-data. submitted.