Advanced BSgenomeForge usage

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1 Introduction

This vignette describes the process of forging a *BSgenome data package*. It is intended for Bioconductor users who want to make a new *BSgenome data package*, not for regular users of these packages.

IMPORTANT NOTE: If your genome assembly is an NCBI assembly or UCSC genome then please make sure to consult the introductory vignette (A quick introduction to the BSgenomeForge package) also provided in this package. This is the easy case and is covered by user-friendly functions forgeBSgenomeDataPkgFromNCBI and forgeBSgenomeDataPkgFromUCSC described in the introductory vignette. The vignette you are currently reading is for advanced users that need to forge a BSgenome data package for a genome assembly not covered by the introductory vignette.

Requirements:

- Some basic knowledge of the Unix/Linux command line is required. The commands that you will most likely need are: cd, mkdir, mv, rmdir, tar, gunzip, unzip, ftp and wget. Also you will need to create and edit some text files.
- You need access to a good Unix/Linux build machine with a decent amount of RAM (>= 4GB), especially if your genome is big. For smaller genomes, 2GB or even 1GB of RAM might be enough.
- You need the latest release versions of R plus the BSgenomeForge package installed on the build machine. To check your installation, start R and try to load the BSgenomeForge package.
- Finally, you need to obtain the source data files of the genome that you want to build a package for. There are 2 groups of source data files: (1) the files containing the sequence data (those files are required), and (2) the files containing the mask data (those files are optional). For most organisms, these files have been made publicly available on the internet by genome providers like UCSC, NCBI, FlyBase, TAIR, etc. The next sections of this vignette explain how to obtain and prepare these files.

Refer to the R Installation and Administration manual 1 if you need to install R or upgrade your R version, and to the Bioconductor - Install page 2 on the Bioconductor website if you need to install or update the Biostrings, BSgenome, or BSgenomeForge package.

 $^{^{1} \}verb|https://cran.r-project.org/doc/manuals/R-admin.html|$

²https://bioconductor.org/install/

Questions, comments or bug reports about this vignette or any of the functions described here are welcome. Please use the Bioconductor support site ³ if you have questions or need help to forge a *BSgenome data package*, or about anything related to the BSgenomeForge package. For bug reports or feature requests, please open an issue on GitHub at https://github.com/Bioconductor/BSgenomeForge/issues.

In the next section ("How to forge a BSgenome data package with bare sequences"), we describe how to forge a BSgenome data package with sequences that have no masks on them. If this is what you're after, then you'll be done at the end of the section. Only if you need to forge a package with masked sequences, you'll have to keep reading but you'll have to forge a BSgenome data package with bare sequences first.

In this vignette, we call *target packages* the *BSgenome data packages* that we're going to forge (one target package with bare sequences and possibly a 2nd target package with masked sequences).

2 How to forge a BSgenome data package with bare sequences

2.1 Obtain and prepare the sequence data

2.1.1 What do I need?

The sequence data must be in a single twoBit file (e.g. musFur1.2bit) or in a collection of FASTA files (possibly gzip-compressed).

If the latter, then you need 1 FASTA file per sequence that you want to put in the $target\ package$. In that case the name of each FASTA file must be of the form prefix><seqname><suffix> where <seqname> is the name of the sequence in it and prefix> and <suffix> are a prefix and a suffix (possibly empty) that are the same for all the FASTA files.

2.1.2 Examples

UCSC provides the genome sequences for Stickleback (gasAcu1) in various forms: (a) as a twoBit file (gasAcu1.2bit located in https://hgdownload.cse.ucsc.edu/goldenPath/gasAcu1/bigZips/, (b) as a big compressed tarball containing one file per chromosome (chromFa.tar.gz located in the same folder), and (c) as a collection of compressed FASTA files (chrI.fa.gz, chrII.fa.gz, chrIII.fa.gz, ..., chrXXI.fa.gz, chrM.fa.gz and chrUn.fa.gz located in https://hgdownload.cse.ucsc.edu/goldenPath/gasAcu1/chromosomes/). To forge a BSgenome data package for gasAcu1, the easiest would be to use (a) (the twoBit file). However it would also be possible to use (c) (the collection of compressed FASTA files). In that case the suffix would need to be set to .fa.gz. Note that the prefix here is empty and not chr, because chr is considered to be part of the sequence names (a commonly used chromosome naming convention at UCSC). Alternatively, the big compressed tarball (chromFa.tar.gz) could be used. After downloading and extracting it, we should end up with the same files as with (c).

You can use the fasta.seqlengths function from the Biostrings package to get the lengths of the sequences in a FASTA file:

```
> library(BSgenome)
> file <- system.file("extdata", "ce2chrM.fa.gz", package="BSgenome")
> fasta.seqlengths(file)
```

³https://support.bioconductor.org/

```
chrM
13794
```

For the BSgenome.Mfuro.UCSC.musFur1 package, the twoBit file was used and downloaded with:

```
wget https://hgdownload.soe.ucsc.edu/goldenPath/musFur1/bigZips/musFur1.2bit
```

For the BSgenome.Rnorvegicus.UCSC.rn4 package, the big compressed tarball was used. It was downloaded and extracted with:

```
wget https://hgdownload.cse.ucsc.edu/goldenPath/rn4/bigZips/chromFa.tar.gz
tar zxf chromFa.tar.gz
```

2.1.3 The $\langle seqs \; srcdir \rangle$ folder

So we assume that you've downloaded the *sequence data files* and that they are now located in a folder on your machine⁴. From now on, we'll refer to this folder as the *<seqs_srcdir>* folder.

Depending on what data files you downloaded (see previous sub-subsection), you might also need to extract and possibly rename the files. Note that all the sequence data files should be located directly in the <seqs_srcdir> folder, not in subfolders of this folder. For example, depending on the genome, UCSC provides either a big chromFa.tar.gz or chromFa.zip file that contains the sequence data for all the chromosomes. But it could be that, after extraction of this big file, the individual FASTA files for each chromosome end up being located one level down the <seqs_srcdir> folder (granted that you were in this folder when you extracted the file). If this is the case, then you will need to move them one level up (use mv -i */*.fa . for this, then remove all the empty subfolders with rmdir *).

2.2 Prepare the BSgenome data package seed file

2.2.1 Overview

The BSgenome data package seed file will contain all the information needed by the forgeBSgenomeDataPkg function to forge the target package.

The format of this file is DCF (Debian Control File), which is also the format used for the DESCRIPTION file of any R package. The valid fields of a *seed file* are divided in 3 categories:

- 1. Standard DESCRIPTION fields. These fields are actually the mandatory fields found in any DESCRIPTION file. They will be copied to the DESCRIPTION file of the target package.
- 2. Non-standard DESCRIPTION fields. These fields are specific to seed files and they will also be copied to the DESCRIPTION file of the target package. In addition, the values of these fields will be stored in the BSgenome object that will be contained in the target package. This means that the users of the target package will be able to retrieve these values via the accessor methods defined for BSgenome objects. See the man page for the BSgenome class (?'BSgenome-class') for a description of these methods.
- 3. Additional fields that don't fall in the first 2 categories.

⁴Note that checking the md5sums after download is always a good idea.

The 3 following sub-subsections give an extensive description of all the valid fields of a seed file.

Alternatively, the reader in a hurry can go directly to the last sub-subsection of this subsection for an example of a *seed file*.

2.2.2 Standard DESCRIPTION fields

- Package: Name to give to the target package. The convention used for the packages built by the Bioconductor project is to use a name made of 4 parts separated by a dot. Part 1 is always BSgenome. Part 2 is the abbreviated name of the organism (when the name of the organism is made of 2 words, we put together the first letter of the first word in upper case followed by the entire second word in lower case e.g. Rnorvegicus). Part 3 is the name of the organisation who provided the genome (e.g. UCSC). Part 4 is the release string or number used by this organisation to identify this version of the genome (e.g. rn4).
- Title: The title of the *target package*. E.g. Full genome sequences for Rattus norvegicus (UCSC version rn4).
- Description, Version, Author, Maintainer, License: Like the 2 previous fields, these are mandatory fields found in any DESCRIPTION file. Please refer to the *The DESCRIPTION file* subsection of the Writing R Extensions manual ⁵ for more information about these fields. If you plan to distribute the target package that you are about to forge, please pickup the license carefully and make sure that it is compatible with the license of the sequence data files if any.
- Suggests: [OPTIONAL] If you're going to add examples to the Examples section of the target package (see PkgExamples field below for how to do this), use this field to list the packages used in these examples.

2.2.3 Non-standard DESCRIPTION fields

- organism: The scientific name of the organism in the format Genus species (e.g. Rattus norvegicus, Homo sapiens) or Genus species subspecies (e.g. Homo sapiens neanderthalensis).
- common_name: The common name of the organism (e.g. Rat or Human). For organisms that have more than one commmon name (e.g. Bovine or Cow for Bos taurus), choose one. Note that for the packages built by the Bioconductor project from a UCSC genome, this field corresponds to the SPECIES column of the List of UCSC genome releases table ⁶.
- genome: The name of the genome. Typically the name of an NCBI assembly (e.g. GRCh38.p12, WBcel235, TAIR10.1, ARS-UCD1.2, etc...) or UCSC genome (e.g. hg38, bosTau9, galGal6, ce11, etc...). Should preferably match part 4 of the package name (field Package). For the packages built by the Bioconductor project from a UCSC genome, this field corresponds to the UCSC VERSION field of the List of UCSC genome releases table.
- provider: The provider of the sequence data files e.g. UCSC, NCBI, BDGP, FlyBase, etc... Should preferably match part 3 of the package name (field Package).
- release_date: When this assembly of the genome was released. For the packages built by the Bioconductor project from a UCSC genome, this field corresponds to the RELEASE DATE field of the *List of UCSC genome releases* table.

 $^{^5 \}mathrm{https://cran.r-project.org/doc/manuals/R-exts.html\#The-DESCRIPTION-file}$

⁶https://genome.ucsc.edu/FAQ/FAQreleases#release1

- source_url: The permanent URL where the sequence data files used to forge the target package can be found. If the target package is for an NCBI assembly, use the link to the NCBI landing page for the assembly e.g. https://www.ncbi.nlm.nih.gov/assembly/GCF_003254395.2/.
- organism_biocview: The official biocViews term for this organism. This is generally the same as the organism field except that spaces should be replaced with underscores. The value of this field matters only if the *target package* is going to be added to a Bioconductor repository, because it will determine where the package will show up in the biocViews tree ⁷. Note that this is the only field in this category that won't get stored in the *BSgenome* object that will get wrapped in the *target package*.

2.2.4 Other fields

- BSgenomeObjname: Should match part 2 of the package name (see Package field above).
- seqnames: [OPTIONAL] Needed only if you are using a collection of sequence data files. In that case seqnames must be an R expression returning the names of the single sequences to forge (in a character vector). E.g. paste("chr", c(1:20, "X", "M", "Un", paste(c(1:20, "X", "Un"), "_random", sep="")), sep="").
- circ_seqs: Not needed if your NCBI assembly or UCSC genome is registered in the GenomeInfoDb package. An R expression returning the names of the circular sequences (in a character vector). If the seqnames field is specified, then circ_seqs must be a subset of it. E.g. "chrM" for rn4 or c("chrM", "2micron") for the sacCer2 genome (Yeast) from UCSC. If the assembly or genome has no circular sequence, set circ_seqs to character(0).
- mseqnames: [OPTIONAL] An R expression returning the names of the multiple sequences to forge (in a character vector). E.g. c("random", "chrUn") for rn5. The default value for mseqnames is NULL (no multiple sequence).
- PkgDetails: [OPTIONAL] Some arbitrary text that will be copied to the Details section of the man page of the target package.
- SrcDataFiles: [OPTIONAL] Some arbitrary text that will be copied to the Note section of the man pages of the *target package*. SrcDataFiles should describe briefly where the *sequence data files* are coming from. Will typically contain URLs (permanent URLs are a must).
- PkgExamples: [OPTIONAL] Some R code (possibly with comments) that will be added to the Examples section of the man page of the *target package*. Don't forget to list the packages used in these examples in the Suggests field.
- seqs_srcdir: The absolute path to the folder containing the sequence data files.
- seqfile_name: Required if the sequence data files is a single twoBit file (e.g. musFur1.2bit). Ignored otherwise.
- seqfiles_prefix, seqfiles_suffix: [OPTIONAL] Needed only if you are using a collection of sequence data files. The common prefix and suffix that need to be added to all the sequence names (fields seqnames and mseqnames) to get the name of the corresponding FASTA file. Default values are the empty prefix for seqfiles_prefix and .fa for seqfiles_suffix.

 $^{^7 \}texttt{https://bioconductor.org/packages/release/BiocViews.html\#__Organism}$

• ondisk_seq_format: [OPTIONAL] Specifies how the single sequences should be stored in the target package. Can be 2bit, rda, fa.rz, or fa. If 2bit (the default), then all the single sequences are stored in a single twoBit file. If rda, then each single sequence is stored in a separated serialized XString object (one per single sequence). If fa.rz or fa, then all the single sequences are stored in a single FASTA file (compressed in the RAZip format if fa.rz).

2.2.5 An example

> library(BSgenomeForge)

The *seed files* used for the packages forged by the Bioconductor project are included in the BSgenomeForge package:

```
> seed_files <- system.file("extdata", "seeds", package="BSgenomeForge")
> tail(list.files(seed_files, pattern="-seed$"))
[1] "BSgenome.Sscrofa.UCSC.susScr3-seed"
[2] "BSgenome.Sscrofa.UCSC.susScr3.masked-seed"
[3] "BSgenome.Tguttata.UCSC.taeGut1-seed"
[4] "BSgenome.Tguttata.UCSC.taeGut1.masked-seed"
[5] "BSgenome.Tguttata.UCSC.taeGut2-seed"
[6] "BSgenome.influenza.NCBI.20100628-seed"
> ## Display seed file for musFur1:
> musFur1_seed <- list.files(seed_files, pattern="\\.musFur1-seed$", full.names=TRUE)
> cat(readLines(musFur1_seed), sep="\n")
Package: BSgenome.Mfuro.UCSC.musFur1
Title: Full genomic sequences for Mustela putorius furo (UCSC version musFur1)
Description: Full genomic sequences for Mustela putorius furo (Ferret) as provided by UCSC (musFur1, A
Version: 1.4.2
organism: Mustela putorius furo
common_name: Ferret
provider: UCSC
provider_version: musFur1
release_date: Apr. 2011
release_name: Ferret Genome Sequencing Consortium MusPutFur1.0
source_url: http://hgdownload.soe.ucsc.edu/goldenPath/musFur1/bigZips/
organism_biocview: Mustela_furo
BSgenomeObjname: Mfuro
SrcDataFiles: musFur1.2bit from http://hgdownload.soe.ucsc.edu/goldenPath/musFur1/bigZips/
PkgExamples: bsg$GL896898 # same as bsg[["GL896898"]]
seqs_srcdir: /fh/fast/morgan_m/BioC/BSgenomeForge/srcdata/BSgenome.Mfuro.UCSC.musFur1/seqs
seqfile_name: musFur1.2bit
> ## Display seed file for rn4:
> rn4_seed <- list.files(seed_files, pattern="\\.rn4-seed$", full.names=TRUE)
> cat(readLines(rn4_seed), sep="\n")
```

```
Package: BSgenome.Rnorvegicus.UCSC.rn4
Title: Full genomic sequences for Rattus norvegicus (UCSC version rn4)
Description: Full genomic sequences for Rattus norvegicus (Rat) as provided by UCSC (rn4, Nov. 2004) a
Version: 1.4.2
Suggests: TxDb.Rnorvegicus.UCSC.rn4.ensGene
organism: Rattus norvegicus
common_name: Rat
provider: UCSC
provider_version: rn4
release_date: Nov. 2004
release_name: Baylor College of Medicine HGSC v3.4
source_url: http://hgdownload.cse.ucsc.edu/goldenPath/rn4/bigZips/
organism_biocview: Rattus_norvegicus
BSgenomeObjname: Rnorvegicus
seqnames: paste("chr", c(1:20, "X", "M", "Un", paste(c(1:20, "X", "Un"), "_random", sep="")), sep="")
circ_seqs: "chrM"
SrcDataFiles: chromFa.tar.gz from http://hgdownload.cse.ucsc.edu/goldenPath/rn4/bigZips/
PkgExamples: bsg$chr1 # same as bsg[["chr1"]]
        ## -----
        ## Upstream sequences
        ## Starting with BioC 3.0, the upstream1000, upstream2000, and
        ## upstream5000 sequences for rn4 are not included in the BSgenome data
        ## package anymore. However they can easily be extracted from the full
        ## genomic sequences with something like:
        library(TxDb.Rnorvegicus.UCSC.rn4.ensGene)
        txdb <- TxDb.Rnorvegicus.UCSC.rn4.ensGene
        gn <- sort(genes(txdb))</pre>
        up1000 <- flank(gn, width=1000)
        up1000seqs <- getSeq(bsg, up1000)
        ## IMPORTANT: Make sure you use a TxDb package (or TxDb object),
        ## that contains a gene model based on the exact same reference genome
        ## as the BSgenome object you pass to getSeq(). Note that you can make
        ## your own custom TxDb object from various annotation resources.
        ## See the makeTxDbFromUCSC(), makeTxDbFromBiomart(), and
        ## makeTxDbFromGFF() functions in the txdbmaker package.
seqs_srcdir: /fh/fast/morgan_m/BioC/BSgenomeForge/srcdata/BSgenome.Rnorvegicus.UCSC.rn4/seqs
```

From now we assume that you have managed to prepare the seed file for your target package.

2.3 Forge the target package

To forge the *target package*, start R, load the BSgenomeForge package, and call the forgeBSgenomeDataPkg function on your *seed file*. For example, if the path to your *seed file* is "path/to/my/seed", do:

- > library(BSgenomeForge)
- > forgeBSgenomeDataPkg("path/to/my/seed")

Depending on the size of the genome and your hardware, this can take between 2 minutes and 1 or 2 hours. By default forgeBSgenomeDataPkg will create the source tree of the *target package* in the current directory.

Once forgeBSgenomeDataPkg is done, ignore the warnings (if any), quit R, and build the source package (tarball) with

```
R CMD build <pkgdir>
```

where <pkgdir> is the path to the source tree of the package.

Then check the package with

```
R CMD check <tarball>
```

where <tarball> is the path to the tarball produced by R CMD build.

Finally install the package with

```
R CMD INSTALL <tarball>
```

and use it!

3 How to forge a BSgenome data package with masked sequences

3.1 Obtain and prepare the mask data

3.1.1 What do I need?

The mask data are not available for all organisms. What you download exactly depends of course on what's available and also on what built-in masks you want to have in the 2nd target package. 4 kinds of built-in masks are currently supported by BSgenomeForge:

- the masks of assembly gaps, aka "the AGAPS masks";
- the masks of intra-contig ambiguities, aka "the AMB masks";
- the masks of repeat regions that were determined by the RepeatMasker software, aka "the RM masks";
- the masks of repeat regions that were determined by the Tandem Repeats Finder software (where only repeats with period less than or equal to 12 were kept), aka "the TRF masks".

For the AGAPS masks, you need UCSC "gap" or NCBI "agp" files. It can be one file per chromosome or a single big file containing the assembly gap information for all the chromosomes together. In the former

case, the name of each file must be of the form prefix><seqname><suffix>. Like for the FASTA files in group 1, <seqname> must be the name of the sequence (sequence names for FASTA files and AGAPS masks must match) and prefix> and <suffix> must be a prefix and a suffix (possibly empty) that are the same for all the files (this prefix/suffix doesn't need to be, and typically is not, the same as for the FASTA files in group 1).

You don't need any file for the AMB masks.

For the RM masks, you need RepeatMasker .out files. Like for the AGAPS masks, it can be one file per chromosome or a single big file containing the RepeatMasker information for all the chromosomes together. In the former case, the name of each file must also be of the form <prefix><seqname><suffix>. Same comments apply as for the AGAPS masks above.

For the TRF masks, you need Tandem Repeats Finder .bed files. Again, it can be one file per chromosome or a single big file. In the former case, the name of each file must also be of the form *pre-fix><seqname><suffix>)*. Same comments apply as for the AGAPS masks above.

Again, for some organisms none of the masks above are available or only some of them are.

3.1.2 An example

Here is how the *mask data files* for the BSgenome.Rnorvegicus.UCSC.rn4 package were obtained and prepared:

• AGAPS masks: all the chr*_gap.txt.gz files (UCSC "gap" files) were downloaded from the UCSC database folder ⁸ for rn4. This was done with the standard Unix/Linux ftp command:

```
ftp hgdownload.cse.ucsc.edu # login as "anonymous"
cd goldenPath/rn4/database
prompt
mget chr*_gap.txt.gz
```

Then all the downloaded files were uncompressed with:

```
for file in chr*_gap.txt.gz; do gunzip $file ; done
```

• RM masks: file chromOut.tar.gz was downloaded from the UCSC bigZips folder and extracted with:

```
tar zxf chromOut.tar.gz
```

• TRF masks: file chromTrf.tar.gz was downloaded from the UCSC bigZips folder and extracted with:

```
tar zxf chromTrf.tar.gz
```

 $^{^8}$ https://hgdownload.cse.ucsc.edu/goldenPath/rn4/database/

3.1.3 The $< masks \ srcdir >$ folder

From now we assume that you've downloaded (checking the md5sums is always a good idea) and possibly extracted all the mask data files, and that they are located in the <masks srcdir> folder.

Note that all the $mask\ data\ files$ should be located directly in the $< masks_srcdir>$ folder, not in subfolders of this folder.

3.2 Prepare the seed file for the masked BSgenome data package

3.2.1 Overview

The seed file for the BSgenome data package with masked sequences (i.e. 2nd target package) is similar to the seed file for the BSgenome data package containing the corresponding bare sequences (a.k.a. the reference target package i.e. the package you forged in the previous section). It will contain all the information needed by the forgeMaskedBSgenomeDataPkg function to forge the 2nd target package.

The fields of this *seed file* are described in the 3 following sub-subsections.

Alternatively, the reader in a hurry can go directly to the last sub-subsection of this subsection for an example of *seed file*.

3.2.2 Standard DESCRIPTION fields

- Package: Name to give to the 2nd target package. The convention used for the packages built by the Bioconductor project is to use the same name as the reference target package with the .masked suffix added to it.
- Title: The title of the package. E.g. Full masked genome sequences for Rattus norvegicus (UCSC version rn4).
- Description, Version, Author, Maintainer, License: See previous section.

3.2.3 Non-standard DESCRIPTION fields

- organism_biocview: Same as for the reference target package.
- source_url: The permanent URL where the mask data files used to forge this package can be found.

3.2.4 Other fields

- RefPkgname: The name of the reference target package.
- nmask_per_seq: The number of masks per sequence (1 to 4).
- PkgDetails, PkgExamples: See previous section.
- SrcDataFiles: [OPTIONAL] Some arbitrary text that will be copied to the Note section of the man pages of the *target package*. SrcDataFiles should describe briefly where the *mask data files* are coming from. Will typically contain URLs (permanent URLs are a must).

- masks_srcdir: The path to the <masks srcdir> folder.
- AGAPSfiles_type: [OPTIONAL] Must be gap (the default) if the source data files containing the AGAPS masks information are UCSC "gap" files, or agp if they are NCBI "agp" files.
- AGAPSfiles_name: [OPTIONAL] Omit this field if you have one *source data file* per single sequence for the AGAPS masks and use the AGAPSfiles_prefix and AGAPSfiles_suffix fields below instead. Otherwise, use this field to specify the name of the single big file.
- AGAPSfiles_prefix, AGAPSfiles_suffix: [OPTIONAL] Omit these fields if you have one single big source data file for all the AGAPS masks and use the AGAPSfiles_name field above instead. Otherwise, use these fields to specify the common prefix and suffix that need to be added to all the single sequence names (field seqnames) to get the name of the file that contains the corresponding AGAPS mask information. Default values are the empty prefix for AGAPSfiles_prefix and _gap.txt for AGAPSfiles_suffix.
- RMfiles_name, RMfiles_prefix, RMfiles_suffix: [OPTIONAL] Those fields work like the AGAPSfiles* fields above but for the RM masks. Default values are the empty prefix for RMfiles_prefix and .fa.out for RMfiles_suffix.
- TRFfiles_name, TRFfiles_prefix, TRFfiles_suffix: [OPTIONAL] Those fields work like the AGAPSfiles* fields above but for the TRF masks. Default values are the empty prefix for TRFfiles_prefix and .bed for TRFfiles_suffix.

3.2.5 An example

organism_biocview: Rattus_norvegicus

The *seed files* used for the packages forged by the Bioconductor project are included in the BSgenomeForge package:

```
> library(BSgenomeForge)
> seed_files <- system.file("extdata", "seeds", package="BSgenomeForge")
> tail(list.files(seed_files, pattern="\\.masked-seed$"))
[1] "BSgenome.Ptroglodytes.UCSC.panTro2.masked-seed"
[2] "BSgenome.Ptroglodytes.UCSC.panTro3.masked-seed"
[3] "BSgenome.Rnorvegicus.UCSC.rn4.masked-seed"
[4] "BSgenome.Rnorvegicus.UCSC.rn5.masked-seed"
[5] "BSgenome.Sscrofa.UCSC.susScr3.masked-seed"
[6] "BSgenome.Tguttata.UCSC.taeGut1.masked-seed"
> rn4_masked_seed <- list.files(seed_files, pattern="\\.rn4\\.masked-seed$", full.names=TRUE)
> cat(readLines(rn4_masked_seed), sep="\n")
Package: BSgenome.Rnorvegicus.UCSC.rn4.masked
Title: Full masked genomic sequences for Rattus norvegicus (UCSC version rn4)
Description: Full genomic sequences for Rattus norvegicus (Rat) as provided by UCSC (rn4, Nov. 2004) a
Version: 1.3.99
RefPkgname: BSgenome.Rnorvegicus.UCSC.rn4
source_url: http://hgdownload.cse.ucsc.edu/goldenPath/rn4/bigZips/
```

From now we assume that you have managed to prepare the seed file for the 2nd target package.

3.3 Forge the 2nd target package

To forge the package, start R, make sure the reference target package is installed (try to load it), load the BSgenomeForge package, and call the forgeMaskedBSgenomeDataPkg function on the seed file you made for the 2nd target package. For example, if the path to your seed file is "path/to/my/seed", do:

- > library(BSgenomeForge)
- > forgeMaskedBSgenomeDataPkg("path/to/my/seed")

Depending on the size of the genome and your hardware, this can take between 2 minutes and 1 or 2 hours. By default forgeMaskedBSgenomeDataPkg will create the source tree of the 2nd target package in the current directory.

Once forgeMaskedBSgenomeDataPkg is done, ignore the warnings (if any), quit R, and build the source package (tarball) with

```
R CMD build <pkgdir>
```

where <pkgdir> is the path to the source tree of the package.

Then check the package with

```
R CMD check <tarball>
```

where <tarball> is the path to the tarball produced by R CMD build.

Finally install the package with

```
R CMD INSTALL <tarball>
```

and use it!

If you want to distribute this package, you need to distribute it with the reference target package since it depends on it.

4 Session information

The output in this vignette was produced under the following conditions:

```
> sessionInfo()
R Under development (unstable) (2025-10-20 r88955)
Platform: x86_64-pc-linux-gnu
Running under: Ubuntu 24.04.3 LTS
Matrix products: default
BLAS:
        /home/biocbuild/bbs-3.23-bioc/R/lib/libRblas.so
LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.12.0 LAPACK version 3.12.0
locale:
 [1] LC_CTYPE=en_US.UTF-8
                                LC_NUMERIC=C
 [3] LC_TIME=en_GB
                                LC_COLLATE=C
 [5] LC_MONETARY=en_US.UTF-8
                                LC_MESSAGES=en_US.UTF-8
 [7] LC_PAPER=en_US.UTF-8
                                LC_NAME=C
 [9] LC_ADDRESS=C
                                LC_TELEPHONE=C
[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
time zone: America/New_York
tzcode source: system (glibc)
attached base packages:
[1] stats4
                       graphics grDevices utils
             stats
                                                      datasets methods
[8] base
other attached packages:
 [1] BSgenomeForge_1.11.0 BSgenome_1.79.1
                                               rtracklayer_1.71.0
 [4] BiocIO_1.21.0
                          GenomicRanges_1.63.0 Biostrings_2.79.1
 [7] XVector_0.51.0
                          GenomeInfoDb_1.47.0 Seqinfo_1.1.0
[10] IRanges_2.45.0
                          S4Vectors_0.49.0
                                                BiocGenerics_0.57.0
[13] generics_0.1.4
                          BiocStyle_2.39.0
loaded via a namespace (and not attached):
 [1] SummarizedExperiment_1.41.0 rjson_0.2.23
 [3] xfun_0.54
                                 bslib_0.9.0
 [5] devtools_2.4.6
                                 remotes_2.5.0
 [7] processx_3.8.6
                                 Biobase_2.71.0
 [9] lattice_0.22-7
                                 callr_3.7.6
[11] ps_1.9.1
                                 vctrs_0.6.5
[13] tools_4.6.0
                                 bitops_1.0-9
[15] curl_7.0.0
                                 parallel_4.6.0
[17] Matrix_1.7-4
                                 desc_1.4.3
[19] cigarillo_1.1.0
                                 lifecycle_1.0.4
[21] compiler_4.6.0
                                 Rsamtools_2.27.0
[23] codetools_0.2-20
                                 htmltools_0.5.8.1
[25] usethis_3.2.1
                                 sass_0.4.10
```

[27]	RCurl_1.98-1.17
[29]	crayon_1.5.3
[31]	ellipsis_0.3.2
[33]	DelayedArray_0.37.0
[35]	sessioninfo_1.2.3
[37]	digest_0.6.37
[39]	restfulr_0.0.16
[41]	fastmap_1.2.0
[43]	cli_3.6.5
[45]	magrittr_2.0.4
[47]	XML_3.99-0.19
[49]	UCSC.utils_1.7.0
[51]	httr_1.4.7
[53]	memoise_2.0.1
[55]	knitr_1.50
[57]	glue_1.8.0
[59]	pkgload_1.4.1
[61]	jsonlite_2.0.0
[63]	MatrixGenerics_1.23.0

[65] fs_1.6.6

yaml_2.3.10 jquerylib_0.1.4 BiocParallel_1.45.0 cachem_1.1.0 abind_1.4-8 purrr_1.2.0 bookdown_0.45 grid_4.6.0 SparseArray_1.11.1 S4Arrays_1.11.0 pkgbuild_1.4.8 rmarkdown_2.30 matrixStats_1.5.0 evaluate_1.0.5 rlang_1.1.6 BiocManager_1.30.26 rstudioapi_0.17.1 R6_2.6.1

 ${\tt GenomicAlignments_1.47.0}$