An Introduction to GenomeInfoDb

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1 Introduction

The *GenomeInfoDb* provides an interface to access seqlevelsStyles (such as UCSC, NCBI, Ensembl) and their supported mappings for organisms. For instance, for Homo sapiens, seqlevelsStyle "UCSC" maps to "chr1", "chr2", ..., "chrX", "chrY". The section below introduces these functions with examples.

2 Functionality for all existing organisms

2.1 genomeStyles

The genomeStyles lists out for each organism, the seglevelsStyles and their mappings.

```
seqmap <- genomeStyles()</pre>
head(seqmap, n=2)
## $Arabidopsis_thaliana
## circular auto sex NCBI TAIR9 Ensembl
## 1
       FALSE TRUE FALSE 1 Chr1
## 2
       FALSE TRUE FALSE 2 Chr2
## 3
       FALSE TRUE FALSE 3 Chr3
                                       3
## 4
       FALSE TRUE FALSE
                          4 Chr4
       FALSE TRUE FALSE 5 Chr5
                                       5
      TRUE FALSE FALSE MT ChrM
                                      Μt
## 7
       TRUE FALSE TRUE Pltd ChrC
                                      Pt
## $Caenorhabditis_elegans
    circular auto sex NCBI
       FALSE TRUE FALSE
## 1
                        I
                              chrI
                                        Т
## 2
       FALSE TRUE FALSE
                        II chrII
## 3
                                      III
       FALSE TRUE FALSE III chrIII
       FALSE TRUE FALSE IV chrIV
                                      ΙV
       FALSE TRUE FALSE
                              chrV
                                        ٧
## 5
                          V
## 6
       FALSE FALSE TRUE
                          Χ
                              chrX
                                        Χ
       TRUE TRUE FALSE MT
                              chrM
                                     MtDNA
```

Oragnism's supported by GenomeInfoDb can be found by :

If one knows the organism one is interested in, then we can directly access the information for the given organism along. Each function accepts an argument called species which as "genus species", the default is "Homo sapiens". In the following example we list out only the first five entries returned by the code snippet.

```
head(genomeStyles("Homo_sapiens"),5)
    circular auto sex NCBI UCSC dbSNP Ensembl
## 1
       FALSE TRUE FALSE
                          1 chr1 ch1
## 2
                                   ch2
                                             2
       FALSE TRUE FALSE
                          2 chr2
       FALSE TRUE FALSE
                          3 chr3 ch3
                                             3
## 4
       FALSE TRUE FALSE
                          4 chr4 ch4
                                             4
       FALSE TRUE FALSE
                          5 chr5
```

We can also check if a given style is supported by GenomeInfoDb for a given species. For example, if we want to know if "UCSC" mapping is supported for "Homo sapiens" we can ask:

```
"UCSC" %in% names(genomeStyles("Homo_sapiens"))
## [1] TRUE
```

2.2 extractSeqlevels

We can also extract the desired seqlevelsStyle from a given organism using the extractSe qlevels

```
extractSeqlevels(species="Arabidopsis_thaliana", style="NCBI")
## [1] "1" "2" "3" "4" "5" "MT" "Pltd"
```

2.3 extractSeqlevelsByGroup

We can also extract the desired seqlevelsStyle from a given organism based on a group (Group - 'auto' denotes autosomes, 'circular' denotes circular chromosomes and 'sex' denotes sex chromosomes; the default is all chromosomes are returned).

2.4 seglevelsStyle

We can find the seqname Style for a given character vector by using the seqlevelsStyle

```
seqlevelsStyle(paste0("chr",c(1:30)))
## [1] "UCSC"
seqlevelsStyle(c("2L","2R","X","Xhet"))
## [1] "NCBI"
```

2.5 seglevelsInGroup

We can also subset a given character vector containing seqnames using the seqlevelsInGroup. We currently support 3 groups: 'auto' for autosomes, 'sex' for allosomes/sex chromosomes and circular for 'circular' chromosomes. The user can also provide the style and species they are working with. In the following examples, we extract the sex, auto and circular chromosomes for Homo sapiens:

```
newchr <- paste0("chr",c(1:22,"X","Y","M","1_gl000192_random","4_ctg9_hap1"))
seqlevelsInGroup(newchr, group="sex")</pre>
```

```
## [1] "chrX" "chrY"

seqlevelsInGroup(newchr, group="auto")

## [1] "chr1" "chr2" "chr3" "chr4" "chr5" "chr6" "chr7" "chr8" "chr9"

## [10] "chr10" "chr11" "chr12" "chr13" "chr14" "chr15" "chr16" "chr17" "chr18"

## [19] "chr19" "chr20" "chr21" "chr22"

seqlevelsInGroup(newchr, group="circular")

## [1] "chrM"

seqlevelsInGroup(newchr, group="sex","Homo_sapiens","UCSC")

## [1] "chrX" "chrY"
```

if we have a vector containing seqnames and we want to verify the species and style for them , we can use:

```
seqnames <- c("chr1", "chr9", "chr2", "chr3", "chr10")
all(seqnames %in% extractSeqlevels("Homo_sapiens", "UCSC"))
## [1] TRUE</pre>
```

2.6 orderSeqlevels

The <u>orderSeqlevels</u> can return the order of a given character vector which contains seqnames. In the following example, we show how you can find the order for a given seqnames character vector.

```
seqnames <- c("chr1","chr9", "chr2", "chr3", "chr10")
orderSeqlevels(seqnames)

## [1] 1 3 4 2 5
seqnames[orderSeqlevels(seqnames)]

## [1] "chr1" "chr2" "chr3" "chr9" "chr10"</pre>
```

2.7 rankSeqlevels

The rankSeqlevels can return the rank of a given character vector which contains seqnames. In the following example, we show how you can find the rank for a given seqnames character vector.

```
seqnames <- c("chr1","chr9", "chr2", "chr3", "chr10")
rankSeqlevels(seqnames)
## [1] 1 4 2 3 5</pre>
```

2.8 mapSeglevels

Returns a matrix with 1 column per supplied sequence name and 1 row per sequence renaming map compatible with the specified style. If best.only is TRUE (the default), only the "best" renaming maps (i.e. the rows with less NAs) are returned.

```
mapSeqlevels(c("chrII", "chrIII", "chrM"), "NCBI")
## chrII chrIII chrM
## "II" "III" "MT"
```

We also have several seqlevel utility functions.Let us construct a basic GRanges and show how these functions can be used. .

```
gr <- GRanges(paste0("ch",1:35), IRanges(1:35, width=5))</pre>
gr
## GRanges object with 35 ranges and 0 metadata columns:
##
          segnames
                       ranges strand
##
             <Rle> <IRanges> <Rle>
##
      [1]
               ch1
                          1-5
##
      [2]
               ch2
                          2-6
                          3-7
##
      [3]
               ch3
      [4]
               ch4
                          4-8
##
      [5]
               ch5
                          5-9
##
      . . .
               . . .
                          . . .
##
     [31]
              ch31
                       31-35
     [32]
              ch32
                       32-36
                       33-37
##
     [33]
              ch33
##
     [34]
              ch34
                        34-38
                        35-39
##
              ch35
     [35]
##
     seqinfo: 35 sequences from an unspecified genome; no seqlengths
```

As you can see , we have "ch" instead of "chr" for chromosome names. We can use rename Seqlevels to change the "ch" to "chr"

2.9 renameSeqlevels

As the first argument - it takes the object whose seqlevels we need to change, and as the second argument it takes a named vector which has the changes.

```
newnames <- paste0("chr",1:35)</pre>
names(newnames) <- paste0("ch",1:35)</pre>
head(newnames)
      ch1
           ch2 ch3
                            ch4
                                 ch5
## "chr1" "chr2" "chr3" "chr4" "chr5" "chr6"
gr <- renameSeqlevels(gr,newnames)</pre>
## GRanges object with 35 ranges and 0 metadata columns:
                       ranges strand
##
          segnames
##
             <Rle> <IRanges> <Rle>
      [1]
              chr1
                          1-5
                          2-6
##
      [2]
              chr2
                          3-7
##
      [3]
              chr3
      [4]
              chr4
                          4-8
##
##
      [5]
              chr5
                          5-9
```

```
##
##
     [31]
             chr31
                        31-35
##
     [32]
                        32-36
             chr32
##
     [33]
             chr33
                        33-37
##
                        34-38
     [34]
             chr34
##
     [35]
             chr35
                        35-39
##
     seqinfo: 35 sequences from an unspecified genome; no seqlengths
```

Humans have just 22 primary chromosomes - but here we have some extra seqlevels which we want to remove - there are several ways we can achieve this:

2.10 dropSeglevels

Here the second argument is the seqlevels that you want to drop. Because these seqlevels are in use (i.e. have ranges on them), the ranges on these sequences need to be removed before the seqlevels can be dropped. We call this *pruning*. The pruning.mode argument controls how to prune gr. Unlike for list-like objects (e.g. GRangesList) for which pruning can be done in various ways, pruning a GRanges object is straightforward and achieved by specifying pruning.mode="coarse".

```
dropSeqlevels(gr, paste0("chr", 23:35), pruning.mode="coarse")
## GRanges object with 22 ranges and 0 metadata columns:
##
                       ranges strand
          segnames
##
             <Rle> <IRanges> <Rle>
##
      [1]
              chr1
                          1-5
##
      [2]
              chr2
                          2-6
##
      [3]
              chr3
                          3-7
      [4]
##
              chr4
                          4-8
##
      [5]
              chr5
                          5-9
##
               . . .
                          . . .
##
     [18]
             chr18
                        18-22
##
     [19]
             chr19
                        19-23
##
                        20-24
     [20]
             chr20
##
     [21]
             chr21
                        21-25
##
     [22]
             chr22
                        22-26
##
##
     seqinfo: 22 sequences from an unspecified genome; no seqlengths
```

2.11 keepSeqlevels

Here the second argument is the seglevels that you want to keep.

```
keepSeqlevels(gr, paste0("chr",1:22), pruning.mode="coarse")

## GRanges object with 22 ranges and 0 metadata columns:

## seqnames ranges strand

## <Rle> <IRanges> <Rle>

## [1] chr1 1-5 *

## [2] chr2 2-6 *
```

```
[3]
              chr3
                          3-7
##
      [4]
              chr4
                          4-8
##
      [5]
              chr5
                          5-9
##
      . . .
               . . .
##
     [18]
             chr18
                        18-22
##
     [19]
             chr19
                        19-23
##
     [20]
             chr20
                        20-24
                        21-25
##
     [21]
             chr21
     [22]
             chr22
                        22-26
##
     seqinfo: 22 sequences from an unspecified genome; no seqlengths
```

2.12 keepStandardChromosomes

This function internally uses the pre-defined tables inside GenomeInfoDb to find the correct seqlevels according to the sequence style of the object.

```
keepStandardChromosomes(gr, pruning.mode="coarse")
## GRanges object with 35 ranges and 0 metadata columns:
##
          seqnames
                       ranges strand
##
             <Rle> <IRanges> <Rle>
##
      [1]
              chr1
                          1-5
      [2]
              chr2
                          2-6
##
##
      [3]
              chr3
                          3-7
##
      [4]
              chr4
                          4-8
##
      [5]
              chr5
                          5-9
##
      . . .
              . . .
                          . . .
##
     [31]
                       31-35
             chr31
##
     [32]
             chr32
                       32-36
                       33-37
##
     [33]
             chr33
     [34]
             chr34
                       34-38
##
     [35]
                       35-39
             chr35
##
##
     seqinfo: 35 sequences from an unspecified genome; no seqlengths
```

One can also specify the optional species argument to bemore precise.

```
plantgr <- GRanges(c(1:5, "MT", "Pltd"), IRanges(1:7, width=5))</pre>
keepStandardChromosomes(plantgr, species="Arabidopsis thaliana",
                                   pruning.mode="coarse")
## GRanges object with 7 ranges and 0 metadata columns:
##
         seqnames
                      ranges strand
##
            <Rle> <IRanges> <Rle>
##
                1
                         1-5
     [1]
##
     [2]
                2
                         2-6
##
     [3]
                3
                         3-7
##
     [4]
                4
                         4-8
                5
                         5-9
##
     [5]
##
     [6]
               MT
                        6-10
                        7-11
##
     [7]
             Pltd
```

```
## -----
## seqinfo: 7 sequences from an unspecified genome; no seqlengths
```

3 Seqinfo objects

```
## Note that all the arguments (except 'genome') must have the
## same length. 'genome' can be of length 1, whatever the lengths
## of the other arguments are.
x <- Seqinfo(seqnames=c("chr1", "chr2", "chr3", "chrM"),</pre>
             seqlengths=c(100, 200, NA, 15),
             isCircular=c(NA, FALSE, FALSE, TRUE),
             genome="toy")
length(x)
## [1] 4
segnames(x)
## [1] "chr1" "chr2" "chr3" "chrM"
names(x)
## [1] "chr1" "chr2" "chr3" "chrM"
seglevels(x)
## [1] "chr1" "chr2" "chr3" "chrM"
seqlengths(x)
## chr1 chr2 chr3 chrM
## 100 200 NA 15
isCircular(x)
## chr1 chr2 chr3 chrM
     NA FALSE FALSE TRUE
genome(x)
## chr1 chr2 chr3 chrM
## "toy" "toy" "toy" "toy"
x[c("chrY", "chr3", "chr1")] # subset by names
## Seqinfo object with 3 sequences from 2 genomes (NA, toy):
## seqnames seqlengths isCircular genome
##
    chrY
                     NA
                                 NA <NA>
    chr3
                     NA
                              FALSE
                                       toy
                     100
    chr1
## Rename, drop, add and/or reorder the sequence levels:
seqlevels(xx) <- sub("chr", "ch", seqlevels(xx)) # rename</pre>
## Seqinfo object with 4 sequences (1 circular) from toy genome:
## seqnames seqlengths isCircular genome
```

```
100
##
     ch1
                                  NA
                                        toy
##
     ch2
                     200
                              FALSE
                                        toy
                              FALSE
##
     ch3
                      NA
                                        toy
##
     chM
                      15
                               TRUE
                                        toy
seqlevels(xx) <- rev(seqlevels(xx)) # reorder</pre>
XX
## Seqinfo object with 4 sequences (1 circular) from toy genome:
     segnames seglengths isCircular genome
                      15
                               TRUE
                                        toy
##
    ch3
                      NA
                              FALSE
                                        toy
                              FALSE
##
    ch2
                     200
                                        toy
##
    ch1
                     100
                                  NA
                                        toy
seqlevels(xx) <- c("ch1", "ch2", "chY") # drop/add/reorder</pre>
XX
## Seqinfo object with 3 sequences from 2 genomes (toy, NA):
    seqnames seqlengths isCircular genome
##
    ch1
                     100
                                  NA
                                        toy
##
     ch2
                     200
                              FALSE
                                        toy
    chY
                      NA
                                  NA
                                       <NA>
seqlevels(xx) <- c(chY="Y", ch1="1", "22") # rename/reorder/drop/add</pre>
## Seqinfo object with 3 sequences from 2 genomes (NA, toy):
    seqnames seqlengths isCircular genome
##
    Υ
                      NA
                                 NA
                                       <NA>
##
    1
                     100
                                  NA
                                        toy
                      NA
                                 NA
                                      <NA>
y <- Seginfo(segnames=c("chr3", "chr4", "chrM"),
             seqlengths=c(300, NA, 15))
## Seqinfo object with 3 sequences from an unspecified genome:
     segnames seglengths isCircular genome
     chr3
                     300
                                       <NA>
                                 NA
##
    chr4
                      NA
                                  NA
                                       <NA>
                      15
    chrM
                                  NA
                                       <NA>
merge(x, y) # rows for chr3 and chrM are merged
## Warning in .merge_two_Seqinfo_objects(x, y): Each of the 2 combined objects
has sequence levels not in the other:
    - in 'x': chr1, chr2
##
     - in 'y': chr4
    Make sure to always combine/compare objects based on the same reference
    genome (use suppressWarnings() to suppress this warning).
## Seqinfo object with 5 sequences (1 circular) from 2 genomes (toy, NA):
     seqnames seqlengths isCircular genome
##
##
     chr1
                     100
                                 NA
                                        toy
##
     chr2
                     200
                              FALSE
                                        toy
```

```
FALSE
##
     chr3
                     300
                                        toy
##
                      15
                                TRUE
     chrM
                                        toy
##
     chr4
                      NA
                                  NA
                                       <NA>
suppressWarnings(merge(x, y))
## Seqinfo object with 5 sequences (1 circular) from 2 genomes (toy, NA):
     segnames seglengths isCircular genome
##
     chr1
                     100
                                  NA
##
     chr2
                     200
                               FALSE
                                        toy
##
                     300
                               FALSE
     chr3
                                        toy
                      15
                                TRUE
     chrM
                                        toy
##
     chr4
                      NA
                                  NA
                                       <NA>
## Note that, strictly speaking, merging 2 Seqinfo objects is not
## a commutative operation, i.e., in general z1 < merge(x, y)
## is not identical to 'z2 <- merge(y, x)'. However 'z1' and 'z2'
## are quaranteed to contain the same information (i.e. the same
## rows, but typically not in the same order):
suppressWarnings(merge(y, x))
## Seqinfo object with 5 sequences (1 circular) from 2 genomes (toy, NA):
##
     seqnames seqlengths isCircular genome
##
     chr3
                     300
                               FALSE
##
     chr4
                      NA
                                  NA
                                       <NA>
##
     chrM
                      15
                                TRUE
                                        toy
##
     chr1
                     100
                                  NA
                                        toy
                     200
                               FALSE
     chr2
                                        toy
## This contradicts what 'x' says about circularity of chr3 and chrM:
isCircular(y)[c("chr3", "chrM")] <- c(TRUE, FALSE)</pre>
У
## Seqinfo object with 3 sequences (1 circular) from an unspecified genome:
     segnames seglengths isCircular genome
                                TRUE
##
     chr3
                     300
                                       <NA>
##
     chr4
                      NA
                                  NA
                                       <NA>
##
     chrM
                      15
                               FALSE
                                       <NA>
if (interactive()) {
  merge(x, y) # raises an error
}
```

4 Examples

4.1 converting seqlevel styles (eg:UCSC to NCBI)

A quick example using Drosophila Melanogaster. The txdb object contains seqlevels in UCSC style, we want to convert them to NCBI

```
txdb <- TxDb.Dmelanogaster.UCSC.dm3.ensGene
seqlevels(txdb)</pre>
```

```
"chr2R"
    [1] "chr2L"
                                "chr3L"
                                            "chr3R"
                                                         "chr4"
                                                                     "chrX"
    [7] "chrU"
                    "chrM"
                                "chr2LHet"
                                            "chr2RHet"
##
                                                         "chr3LHet"
                                                                     "chr3RHet"
## [13] "chrXHet"
                    "chrYHet"
                                "chrUextra"
genomeStyles("Drosophila melanogaster")
                 sex auto NCBI
                                                              Ensembl
      circular
                                      UCSC
## 1
         FALSE FALSE
                      TRUE
                              2L
                                     chr2L
                                                                   2L
## 2
         FALSE FALSE TRUE
                              2R
                                     chr2R
                                                                   2R
## 3
         FALSE FALSE
                      TRUE
                              3L
                                     chr3L
                                                                   3L
## 4
         FALSE FALSE
                     TRUE
                                                                   3R
                              3R
                                     chr3R
## 5
         FALSE FALSE
                      TRUE
                               4
                                      chr4
                                                                    4
## 6
         FALSE TRUE FALSE
                               Χ
                                      chrX
                                                                   Χ
## 7
         FALSE TRUE FALSE
                                      chrY
                                                                    Υ
## 8
         TRUE FALSE FALSE
                              MT
                                      chrM dmel_mitochondrion_genome
## 9
         FALSE FALSE 2LHet
                                  chr2LHet
                                                                2LHet
                                  chr2RHet
                                                                2RHet
## 10
         FALSE FALSE 2Rhet
## 11
         FALSE FALSE 3LHet chr3LHet
                                                                3LHet
## 12
         FALSE FALSE 3RHet
                                  chr3RHet
                                                                3RHet
## 13
         FALSE FALSE Xhet
                                   chrXHet
                                                                 XHet
## 14
         FALSE FALSE Yhet
                                   chrYHet
                                                                 YHet
## 15
         FALSE FALSE
                                      chrU
                                                                    U
                              Un
         FALSE FALSE FALSE <NA> chrUextra
## 16
                                                               Uextra
mapSeglevels(seglevels(txdb), "NCBI")
##
       chr2L
                 chr2R
                           chr3L
                                     chr3R
                                                 chr4
                                                           chrX
                                                                     chrU
##
        "2L"
                  "2R"
                            "3L"
                                      "3R"
                                                  "4"
                                                            "X"
                                                                     "Un"
##
        chrM chr2LHet chr2RHet chr3LHet
                                            chr3RHet
                                                        chrXHet
                                                                  chrYHet
        "MT"
               "2LHet"
                         "2Rhet"
                                                         "Xhet"
                                                                   "Yhet"
##
                                   "3LHet"
                                             "3RHet"
## chrUextra
          NA
```

4.2 converting styles and removing unwanted seqlevels

Suppose we read in a Bam file or a BED file and the resulting GRanges have a lot of seqlevels which are not required by your analysis or you want to rename the seqlevels from the current style to your own style (eg:USCS to NCBI), we can use the functionality provided by GenomeInfoDb to do that.

Let us say that we have extracted the seqlevels of the Seqinfo object(say GRanges from a BED file) in a variable called "sequence".

```
## sequence is in UCSC format and we want NCBI style
newStyle <- mapSeqlevels(sequence, "NCBI")
newStyle <- newStyle[complete.cases(newStyle)] # removing NA cases.

## rename the seqlevels
x <- renameSeqlevels(x,newStyle)

## keep only the seqlevels you want (say autosomes)</pre>
```

5 Session Information

Here is the output of sessionInfo on the system on which this document was compiled:

toLatex(sessionInfo())

- R version 4.3.2 Patched (2023-11-13 r85521), x86_64-pc-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_GB, LC_COLLATE=C, LC_MONETARY=en_US.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=en_US.UTF-8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.UTF-8, LC_IDENTIFICATION=C
- Time zone: America/New_York
- TZcode source: system (glibc)
- Running under: Ubuntu 22.04.3 LTS
- Matrix products: default
- BLAS: /home/biocbuild/bbs-3.18-bioc/R/lib/libRblas.so
- LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.10.0
- Base packages: base, datasets, grDevices, graphics, methods, stats, stats4, utils
- Other packages: AnnotationDbi 1.64.1, Biobase 2.62.0, BiocGenerics 0.48.1, BiocStyle 2.30.0, GenomeInfoDb 1.38.6, GenomicFeatures 1.54.3, GenomicRanges 1.54.1, IRanges 2.36.0, S4Vectors 0.40.2, TxDb.Dmelanogaster.UCSC.dm3.ensGene 3.2.2
- Loaded via a namespace (and not attached): BiocFileCache 2.10.1, BiocIO 1.12.0, BiocManager 1.30.22, BiocParallel 1.36.0, Biostrings 2.70.2, DBI 1.2.1, DelayedArray 0.28.0, GenomeInfoDbData 1.2.11, GenomicAlignments 1.38.2, KEGGREST 1.42.0, Matrix 1.6-5, MatrixGenerics 1.14.0, R6 2.5.1, RCurl 1.98-1.14, RSQLite 2.3.5, Rsamtools 2.18.0, S4Arrays 1.2.0, SparseArray 1.2.3, SummarizedExperiment 1.32.0, XML 3.99-0.16.1, XVector 0.42.0, abind 1.4-5, biomaRt 2.58.2, bit 4.0.5, bit64 4.0.5, bitops 1.0-7, blob 1.2.4, bookdown 0.37, bslib 0.6.1, cachem 1.0.8, cli 3.6.2, codetools 0.2-19, compiler 4.3.2, crayon 1.5.2, curl 5.2.0, dbplyr 2.4.0, digest 0.6.34, dplyr 1.1.4, evaluate 0.23, fansi 1.0.6, fastmap 1.1.1, filelock 1.0.3, generics 0.1.3, glue 1.7.0, grid 4.3.2, highr 0.10, hms 1.1.3, htmltools 0.5.7, httr 1.4.7, jquerylib 0.1.4, jsonlite 1.8.8, knitr 1.45, lattice 0.22-5, lifecycle 1.0.4, magrittr 2.0.3, matrixStats 1.2.0, memoise 2.0.1, parallel 4.3.2, pillar 1.9.0, pkgconfig 2.0.3, png 0.1-8, prettyunits 1.2.0, progress 1.2.3, rappdirs 0.3.3, restfulr 0.0.15, rjson 0.2.21, rlang 1.1.3, rmarkdown 2.25, rtracklayer 1.62.0, sass 0.4.8, stringi 1.8.3, stringr 1.5.1, tibble 3.2.1, tidyselect 1.2.0, tools 4.3.2, utf8 1.2.4, vctrs 0.6.5, xfun 0.42, xml2 1.3.6, yaml 2.3.8, zlibbioc 1.48.0