Sequences, Genomes, and Genes in ${\it R}~/~{\it Bioconductor}$

Martin Morgan

October 21, 2013

Contents

1	Intro	oduction 3
	1.1	High-throughput workflows
		1.1.1 Technologies
		1.1.2 Research questions
		1.1.3 Analysis
	1.2	R and Bioconductor
		1.2.1 Essential <i>R</i>
		1.2.2 Bioconductor for high-throughput analysis
		1.2.3 Strategies for working with large data
2	Seq	uences 19
	2.1	Biostrings and GenomicRanges
	2.2	From whole genome to short read
		2.2.1 Large and whole-genome sequences
		2.2.2 Short reads
	2.3	Exercises
3	Gen	es and Genomes 32
	3.1	Gene annotation
		3.1.1 Bioconductor data annotation packages
		3.1.2 Internet resources
		3.1.3 Exercises
	3.2	Genome annotation
		3.2.1 Bioconductor transcript annotation packages
		3.2.2 AnnotationHub
		3.2.3 <i>rtracklayer</i>
		3.2.4 VariantAnnotation
		3.2.5 Exercises
	3.3	Visualization
	2.0	······································

Bibliography

44

Chapter 1

Introduction

The first part of today's activities provide an introduction to high-throughput sequence analysis, including key 'infrastructure' in R and *Bioconductor*. The main objectives are to arrive at a common language for discussing sequence analysis, and to become familiar with concepts in R and *Bioconductor* that are necessary for effective analysis and comprehension of high-throughput sequence data. An approximate schedule is in Table 1.1.

1.1 High-throughput workflows

Recent technological developments introduce high-throughput sequencing approaches. A variety of experimental protocols and analysis work flows address gene expression, regulation, encoding of genetic variants, and microbial community structure. Experimental protocols produce a large number (tens of millions per sample) of short (e.g., 35-150, single or paired-end) nucleotide sequences. These are aligned to a reference or other genome. Analysis work flows use the alignments to infer levels of gene expression (RNA-seq), binding of regulatory elements to genomic locations (ChIP-seq), or prevalence of structural variants (e.g., SNPs, short indels, large-scale genomic rearrangements). Sample sizes range from minimal replication (e.g., 2 samples per treatment group) to thousands of individuals.

1.1.1 Technologies

The most common 'second generation' technologies readily available to labs are

- Illumina single- and paired-end reads. Short (100 150 per end) and very numerous. Flow cell, lane, bar-code.
- Roche 454. 100's of nucleotides, 100,000's of reads.
- Life Technologies SOLiD. Unique 'color space' model.
- Complete Genomics. Whole genome sequence / variants / etc as a service; end user gets derived results.

Figure 1.1 illustrates Illumina and 454 sequencing. *Bioconductor* has good support for Illumina and derived data such as aligned reads or called variants, and some support for Roche 454 sequencing; use of SOLiD color space reads typically requires conversion to FASTQ files that undermine the benefit of the color space model.

Table 1.1:	Partial	agenda,	Day 1.
------------	---------	---------	--------

Time	Торіс
8:30	
09:15	Introduction to high-throughput workflows, with R examples
10:15	Tea / coffee break
10:30	Sequences Long and Short
11:30	Genomes and Genes
12:30	Lunch
13:30	Genomes and Genes (continued)
14:00	

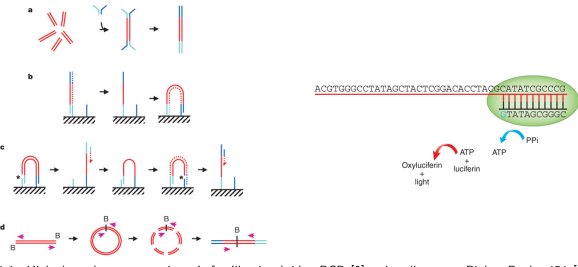


Figure 1.1: High-throughput sequencing. Left: Illumina bridge PCR [2]; mis-call errors. Right: Roche 454 [15]; ho-mopolymer errors.

All second-generation technologies rely on PCR and other techniques to generate reads from samples that represent aggregations of many DNA molecules. 'Third-generation' technologies shift to single-molecule sequencing, with relevant players including Pacific Biosciences and IonTorent. This very exciting data (e.g., [16]) will not be discussed further.

1.1.2 Research questions

Sequence data can be derived from a tremendous diversity of research questions. Some of the most common include:

- Variation DNA-Seq. Sequencing of whole or targeted (e.g., exome) genomic DNA. Common goals include SNP detection, indel and other large-scale structural polymorphisms, and CNV (copy number variation). DNA-seq is also used for de novo assembly, but de novo assembly is not an area where Bioconductor contributes. Key reference: [7, 11].
- **Expression** RNA-seq. Sequencing of reverse-complemented mRNA from the entire expressed transcriptome, typically. Used for *differential expression* studies like micro-arrays, or for *novel transcript discovery*.
- **Regulation** ChIP-seq. ChIP (chromatin immuno-precipitation) is used to enrich genomic DNA for regulatory elements, followed by sequencing and mapping of the enriched DNA to a reference genome. The initial statistical challenge is to identify regions where the mapped reads are enriched relative to a sample that did not undergo ChIP[12]; a subsequent task is to identify differential binding across a designed experiment, e.g., [14]. Survey of diversity of methods: [6].
- **Metagenomics** Sequencing generates sequences from samples containing multiple species, typically microbial communities sampled from niches such as the human oral cavity. Goals include inference of *species composition* (when sequencing typically targets phylogenetically informative genes such as 16S) or *metabolic contribution*. [10, 5]

Special challenges Non-model organisms. Small budgets.

1.1.3 Analysis

Work flows RNA-seq to measure gene expression through assessment of mRNA abundance represents major steps in a typical high-throughput sequence work flow. Typical steps include;

- 1. Experimental design.
- 2. Wet-lab protocols for mRNA extraction and reverse transcription to cDNA.
- 3. Sequencing; QA.
- 4. Alignment of sequenced reads to a reference genome; QA.
- 5. Summarizing of the number of reads aligning to a region; QA.
- 6. Normalization of samples to accommodate purely technical differences in preparation.

nates.

File FASTQ

BAM

VCF

BED

2bit

GFF, GTF

WIG, bigWig

Table 1.2: Common file types and <i>Bioconductor</i> packages used for input	ıt.	
Description	Package	
Unaligned sequences: identifier, sequence, and encoded quality score	ShortRead	
tuples		
Aligned sequences: identifier, sequence, reference sequence name, strand Rsamtools		
position, cigar and additional tags		
Called single nucleotide, indel, copy number, and structural variants,	VariantAnnotation	
often compressed and indexed (with <i>Rsamtools</i> bgzip, indexTabix)		
Gene annotations: reference sequence name, data source, feature type, rtracklayer		

7. Statistical assessment, including specification of an appropriate error model.

'Continuous' single-nucleotide annotation.

Compressed FASTA files with 'masks'

start and end positions, strand, etc.

8. Interpretation of results in the context of original biological questions; QA.

The central inference is that higher levels of gene expression translate to more abundant cDNA, and greater numbers of reads aligned to the reference genome.

Range-based annotation: reference sequence name, start, end coordi-

Common file formats The 'big data' component of high-throughput sequence analyses seems to be a tangle of transformations between file types; common files are summarized in Table 1.2. FASTQ and BAM (sometimes CRAM) files are the primary formats for representing raw sequences and their alignments. VCF are used to summarize called variants in DNA-seq; BED and sometimes WIG files are used to represent ChIP and other regulatory peaks and 'coverage'. GTF / GFF files are important for providing feature annotations, e.g., of exons organization into transcripts and genes.

Third-party (non-R) tools Common analyses often use well-established third-party tools for initial stages of the analysis; some of these have *Bioconductor* counterparts that are particularly useful when the question under investigation does not meet the assumptions of other facilities. Some common work flows (a more comprehensive list is available on the SeqAnswers wiki¹) include:

DNA-seq especially variant calling can be facilitated by software such as the $GATK^2$ toolkit.

RNA-seq In addition to the aligners mentioned above, RNA-seq for differential expression might use the *HTSeq*³ python tools for counting reads within regions of interest (e.g., known genes) or a pipeline such as the *bowtie* (basic alignment) / *tophat* (splice junction mapper) / *cufflinks* (esimated isoform abundance) (e.g., ⁴) or *RSEM*⁵ suite of tools for estimating transcript abundance.

ChIP-seq ChIP-seq experiments typically use DNA sequencing to identify regions of genomic DNA enriched in prepared samples relative to controls. A central task is thus to identify peaks, with common tools including *MACS* and *PeakRanger*.

Programs such as those outlined the previous paragraph often rely on information about gene or other structure as input, or produce information about chromosomal locations of interesting features. The GTF and BED file formats are common representations of this information. Representing these files as R data structures is often facilitated by the *rtracklayer* package. We explore these files in Chapter 3.2.3. Variants are very commonly represented in VCF (Variant Call Format) files; these are explored in Chapter 3.2.4.

Common statistical issues Important statistical issues are summarized in Table 1.3. These will be discussed further in later parts of the course, but similar types of issues are relevant in all high-throughput sequence work flows. An important general point is that wet-lab protocols, sequencing reactions, and alignment or other technological processing

rtracklaver

rtracklayer

¹http://seqanswers.com/wiki/RNA-Seq

²http://www.broadinstitute.org/gatk/

³http://www-huber.embl.de/users/anders/HTSeq/doc/overview.html

⁴http://bowtie-bio.sourceforge.net/index.shtml

⁵http://deweylab.biostat.wisc.edu/rsem/

Analysis stage	Issues
Experimental design	Technical versus biological replication, sample size, complexity of design, feasibility
	of intended analysis
Batch effects	Known and unknown factors; technical artifacts.
Summary	Data reduction without loss of information, e.g., counts versus RPKM.
Normalization	Robust estimates of library size.
Differential expression	Appropriate error model (Negative Binomial, Poisson,); 'shrinkage' to balance
	accuracy of per-gene estimates with precision of experiment-wide estimates.
Testing	Filtering to reduce multiple comparisons & false discovery rate.

Table 1.3: Common statistical issues in RNA-seq differential expression and other high-throughput experiments.

steps introduce artifacts that need to be acknowledged and, if possible, accommodated in down-stream analysis, e.g., through modeling or remediation of *batch effects*.

1.2 R and Bioconductor

R is an open-source statistical programming language. It is used to manipulate data, to perform statistical analysis, and to present graphical and other results. R consists of a core language, additional 'packages' distributed with the R language, and a very large number of packages contributed by the broader community. Packages add specific functionality to an R installation. R has become the primary language of academic statistical analysis, and is widely used in diverse areas of research, government, and industry.

R has several unique features. It has a surprisingly 'old school' interface: users type commands into a console; scripts in plain text represent work flows; tools other than R are used for editing and other tasks. R is a flexible programming language, so while one person might use functions provided by R to accomplish advanced analytic tasks, another might implement their own functions for novel data types. As a programming language, R adopts syntax and grammar that differ from many other languages: objects in R are 'vectors', and functions are 'vectorized' to operate on all elements of the object; R objects have 'copy on change' and 'pass by value' semantics, reducing unexpected consequences for users at the expense of less efficient memory use; common paradigms in other languages, such as the 'for' loop, are encountered much less commonly in R. Many authors contribute to R, so there can be a frustrating inconsistency of documentation and interface. R grew up in the academic community, so authors have not shied away from trying new approaches. Common statistical analysis functions are very well-developed.

A first session Opening an R session results in a prompt. The user types instructions at the prompt. Here is an example:

```
## assign values 5, 4, 3, 2, 1 to variable 'x'
x <- c(5, 4, 3, 2, 1)
x
## [1] 5 4 3 2 1</pre>
```

The first line starts with a # to represent a comment; the line is ignored by R. The next line creates a variable x. The variable is assigned (using <-, we could have used = almost interchangeably) a value. The value assigned is the result of a call to the c function. That it is a function call is indicated by the symbol named followed by parentheses, c(). The c function takes zero or more arguments, and returns a vector. The vector is the value assigned to x. R responds to this line with a new prompt, ready for the next input. The next line asks R to display the value of the variable x. R responds by printing [1] to indicate that the subsequent number is the first element of the vector. It then prints the value of x.

R has many features to aid common operations. Entering sequences is a very common operation, and expressions of the form 2:4 create a sequence from 2 to 4. Sub-setting one vector by another is enabled with [. Here we create an integer sequence from 2 to 4, and use the sequence as an index to select the second, third, and fourth elements of x

Category	Function	Description
Vectors	integer, numeric	Vectors of length $>= 0$ holding a single data type
	complex, character	
	raw, factor	
Statistical	NA, factor	Essential statistical concepts, integral to the language.
List-like	list	Arbitrary collections of elements
	data.frame	List of equal-length vectors
	environment	Pass-by-reference data storage; hash
Array-like	data.frame	Homogeneous columns; row- and column indexing
	array	0 or more dimensions
	matrix	Two-dimensional, homogeneous types
Classes	'S3'	List-like structured data; simple inheritance & dispatch
	'S4'	Formal classes, multiple inheritance & dispatch
Functions	'function'	A simple function with arguments, body, and return value
	'generic'	A (S3 or S4) function with associated methods
	'method'	A function implementing a generic for an S3 or S4 class

Table 1.4:	Essential	aspects	of the	R	language.

x[2:4]

[1] 4 3 2

Index values can be repeated, and if outside the domain of x return the special value NA. Negative index values remove elements from the vector. Logical and character vectors (described below) can also be used for sub-setting.

R functions operate on variables. Functions are usually vectorized, acting on all elements of their argument and obviating the need for explicit iteration. Functions can generate warnings when performing suspect operations, or errors if evaluation cannot proceed; try log(-1).

log(x)

[1] 1.6094 1.3863 1.0986 0.6931 0.0000

1.2.1 Essential R

Built-in (atomic) data types *R* has a number of built-in data types, summarized in Table 1.4. These represent integer, numeric (floating point), complex, character, logical (Boolean), and raw (byte) data. It is possible to convert between data types, and to discover the type or mode of a variable.

```
c(1.1, 1.2, 1.3)  # numeric
## [1] 1.1 1.2 1.3
c(FALSE, TRUE, FALSE)  # logical
## [1] FALSE TRUE FALSE
c("foo", "bar", "baz")  # character, single or double quote ok
## [1] "foo" "bar" "baz"
as.character(x)  # convert 'x' to character
## [1] "5" "4" "3" "2" "1"
class(x)  # the number 5 is numeric
## [1] "numeric"
```

R includes data types particularly useful for statistical analysis, including factor to represent categories and NA (used in any vector) to represent missing values.

```
sex <- factor(c("Male", "Female", NA), levels=c("Female", "Male"))
sex
## [1] Male Female <NA>
## Levels: Female Male
```

Lists, data frames, and matrices All of the vectors mentioned so far are homogeneous, consisting of a single type of element. A list can contain a collection of different types of elements and, like all vectors, these elements can be named to create a key-value association.

```
lst <- list(a=1:3, b=c("foo", "bar"), c=sex)
lst
## $a
## [1] 1 2 3
##
## $b
## [1] "foo" "bar"
##
## $c
## [1] Male Female <NA>
## Levels: Female Male
```

Lists can be subset like other vectors to get another list, or subset with [[to retrieve the actual list element; as with other vectors, sub-setting can use names

```
lst[c(3, 1)] # another list -- class isomorphism
## $c
## [1] Male Female <NA>
## Levels: Female Male
##
## $a
## [1] 1 2 3
lst[["a"]] # the element itself, selected by name
## [1] 1 2 3
```

A data.frame is a list of equal-length vectors, representing a rectangular data structure not unlike a spread sheet. Each column of the data frame is a vector, so data types must be homogeneous within a column. A data.frame can be subset by row or column, and columns can be accessed with \$ or [[.

df[c(1, 3),]
age sex
1 27 Male
3 19 Male
df[df\$age > 20,]
age sex
1 27 Male
2 32 Female

A matrix is also a rectangular data structure, but subject to the constraint that all elements are the same type. A matrix is created by taking a vector, and specifying the number of rows or columns the vector is to represent.

```
m <- matrix(1:12, nrow=3)</pre>
m
##
         [,1] [,2] [,3] [,4]
## [1,]
            1
                 4
                       7
                           10
## [2,]
            2
                 5
                       8
                           11
## [3,]
            3
                 6
                       9
                           12
m[c(1, 3), c(2, 4)]
         [,1] [,2]
##
## [1,]
           4
                10
## [2,]
            6
                12
```

On sub-setting, *R* coerces a single column data.frame or single row or column matrix to a vector if possible; use drop=FALSE to stop this behavior.

```
m[, 3]
## [1] 7 8 9
m[, 3, drop=FALSE]
## [,1]
## [1,] 7
## [2,] 8
## [3,] 9
```

An array is a data structure for representing homogeneous, rectangular data in higher dimensions.

S3 (and S4) classes More complicated data structures are represented using the 'S3' or 'S4' object system. Objects are often created by functions (for example, lm, below), with parts of the object extracted or assigned using *accessor* functions. The following generates 1000 random normal deviates as x, and uses these to create another 1000 deviates y that are linearly related to x but with some error. We fit a linear regression using a 'formula' to describe the relationship between variables, summarize the results in a familiar ANOVA table, and access fit (an S3 object) for the residuals of the regression, using these as input first to the var (variance) and then sqrt (square-root) functions. Objects can be interrogated for their class.

x <- rnorm(1000, sd=1)
y <- x + rnorm(1000, sd=.5)</pre>

```
fit <- lm(y ~ x)
                       # formula describes linear regression
                       # an 'S3' object
fit
##
## Call:
## lm(formula = y ~ x)
##
## Coefficients:
## (Intercept)
                          x
        0.0104
                     0.9847
##
anova(fit)
## Analysis of Variance Table
##
## Response: y
##
              Df Sum Sq Mean Sq F value Pr(>F)
             1 1023
                           1023
                                  4141 <2e-16 ***
## x
## Residuals 998
                  247
                              0
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
sqrt(var(resid(fit))) # residuals accessor and subsequent transforms
## [1] 0.4969
class(fit)
## [1] "lm"
```

Many *Bioconductor* packages implement S4 objects to represent data. S3 and S4 systems are quite different from a programmer's perspective, but conceptually similar from a user's perspective: both systems encapsulate complicated data structures, and allow for methods specialized to different data types; accessors are used to extract information from the objects. A quick guide to using S4 methods is in Table 1.5.

Functions R has a very large number of functions; Table 1.6 provides a brief list of those that might be commonly used and particularly useful. See the help pages (e.g., ?lm) and examples (example(match)) for more details on each of these functions.

R functions accept arguments, and return values. Arguments can be required or optional. Some functions may take variable numbers of arguments, e.g., the columns in a data.frame

```
y <- 5:1
log(y)
## [1] 1.6094 1.3863 1.0986 0.6931 0.0000
args(log)  # arguments 'x' and 'base'; see ?log
## function (x, base = exp(1))
## NULL
log(y, base=2)  # 'base' is optional, with default value
## [1] 2.322 2.000 1.585 1.000 0.000
try(log())  # 'x' required; 'try' continues even on error
args(data.frame) # ... represents variable number of arguments
```

Table 1.5:	Using S4	classes	and	methods.

Best practices	
gr <- GRanges()	'Constructor'; create an instance of the GRanges class
seqnames(gr)	'Accessor', extract information from an instance
<pre>countOverlaps(gr1, gr2)</pre>	A method implementing a generic with useful functionality
Older packages	
s <- new("MutliSet)	A constructor
s@annotation	A 'slot' accessor
Help	
class(gr)	Discover class of instance
getClass(gr)	Display class structure, e.g., inheritance
<pre>showMethods(findOverlaps)</pre>	Classes for which methods of findOverlaps are implemented
showMethods(class="GRanges"	', where=search())
	Generics with methods implemented for the GRanges class, limited to
	currently loaded packages.
class?GRanges	Documentation for the GRanges class.
method?"findOverlaps,GRange	es,GRanges"
	Documentation for the findOverlaps method when the two arguments
	are both <i>GRanges</i> instances.
<pre>selectMethod(findOverlaps,</pre>	c("GRanges", "GRanges"))
	View source code for the method, including method 'dispatch'

Table 1.6: A selection of R function.

- dir, read.table (and friends), scan List files in a directory, read spreadsheet-like data into R, efficiently read homogeneous data (e.g., a file of numeric values) to be represented as a matrix.
- c, factor, data.frame, matrix Create a vector, factor, data frame or matrix.
- summary, table, xtabs Summarize, create a table of the number of times elements occur in a vector, cross-tabulate two or more variables.
- t.test, aov, lm, anova, chisq.test Basic comparison of two (t.test) groups, or several groups via analysis of variance / linear models (aov output is probably more familiar to biologists), or compare simpler with more complicated models (anova); χ^2 tests.
- dist, hclust Cluster data.

plot Plot data.

- ls, str, library, search List objects in the current (or specified) workspace, or peak at the structure of an object; add a library to or describe the search path of attached packages.
- lapply, sapply, mapply, aggregate Apply a function to each element of a list (lapply, sapply), to elements of several lists (mapply), or to elements of a list partitioned by one or more factors (aggregate).
- with Conveniently access columns of a data frame or other element without having to repeat the name of the data frame.
- match, %in% Report the index or existence of elements from one vector that match another.
- split, cut, unlist Split one vector by an equal length factor, cut a single vector into intervals encoded as levels of a factor, unlist (concatenate) list elements.
- strsplit, grep, sub Operate on character vectors, splitting it into distinct fields, searching for the occurrence of a patterns using regular expressions (see ?regex, or substituting a string for a regular expression.

biocLite, install.packages Install a package from an on-line repository into your R.

traceback, debug, browser Report the sequence of functions under evaluation at the time of the error; enter a debugger when a particular function or statement is invoked.

Package	Description
base	Data input and manipulation; scripting and programming.
stats	Essential statistical and plotting functions.
lattice, ggplot2	Approaches to advanced graphics.
methods	'S4' classes and methods.
parallel	Facilities for parallel evaluation.
Matrix	Diverse matrix representations
data.table	Efficient management of large data tables

Table 1.7: Selected base and contributed packages.

```
## function (..., row.names = NULL, check.rows = FALSE, check.names = TRUE,
## stringsAsFactors = default.stringsAsFactors())
## NULL
```

Arguments can be matched by name or position. If an argument appears after ..., it must be named.

```
log(base=2, y) # match argument 'base' by name, 'x' by position
```

[1] 2.322 2.000 1.585 1.000 0.000

A function such as anova is a *generic* that provides an overall signature but dispatches the actual work to the *method* corresponding to the class(es) of the arguments used to invoke the generic. A generic may have fewer arguments than a method, as with the S3 function anova and its method anova.glm.

```
args(anova)
## function (object, ...)
## NULL
args(anova.glm)
## function (object, ..., dispersion = NULL, test = NULL)
## NULL
```

The ... argument in the anova generic means that additional arguments are possible; the anova generic passes these arguments to the method it dispatches to.

Packages Packages provide functionality beyond that available in base *R*. There are over 4000 packages in CRAN (Comprehensive *R* Archive Network) and 749 *Bioconductor* packages. Packages are contributed by diverse members of the community; they vary in quality (many are excellent) and sometimes contain idiosyncratic aspects to their implementation. Table 1.7 outlines key base packages and selected contributed packages; see a local CRAN mirror (including the task views summarizing packages in different domains) and *Bioconductor* for additional contributed packages. New packages (from *Bioconductor* or CRAN) can be added to an *R* installation using biocLite():

```
source("http://bioconductor.org/biocLite.R")
biocLite(c("GenomicRanges", "ShortRead"))
```

A package is installed only once per *R* installation, but needs to be loaded (with library) in each session in which it is used. Loading a package also loads any package that it depends on. Packages loaded in the current session are displayed with search. The ordering of packages returned by search represents the order in which the global environment (where commands entered at the prompt are evaluated) and attached packages are searched for symbols.

length(search())
[1] 37
head(search(), 3)
[1] ".GlobalEnv" "package:EMB02013" "package:rtracklayer"

It is possible for a package earlier in the search path to mask symbols later in the search path; these can be disambiguated using ::.

pi <- 3.2 ## http://en.wikipedia.org/wiki/Indiana_Pi_Bill
base::pi
[1] 3.142
rm(pi) ## remove from the .GlobalEnv</pre>

Help! Find help using the *R* help system. Start a web browser with help.start(). The 'Search Engine and Keywords' link is helpful in day-to-day use.

Manual pages provided detailed descriptions of the arguments and return values of functions, and the structure and methods of classes. Find help within an R session as

?data.frame ?lm ?anova ?anova.lm

S3 and S4 methods can be queried interactively. For S3,

```
methods(anova)
## [1] anova.glm anova.glmlist anova.lm anova.loess* anova.mlm anova.nls*
##
## Non-visible functions are asterisked
```

methods(class="glm")

##	[1]	add1.glm*	anova.glm	confint.glm*	cooks.distance.glm*
##	[5]	deviance.glm*	drop1.glm*	effects.glm*	extractAIC.glm*
##	[9]	family.glm*	formula.glm*	influence.glm*	logLik.glm*
##	[13]	model.frame.glm	nobs.glm*	predict.glm	print.glm
##	[17]	residuals.glm	rstandard.glm	rstudent.glm	summary.glm
##	[21]	vcov.glm*	weights.glm*		
##					
##	N	on-visible functions	are asterisked		

It is often useful to view a method definition, either by typing the method name at the command line or, for 'non-visible' methods, using getAnywhere:

anova.lm
getAnywhere("anova.loess")

For instance, the source code of a function is printed if the function is invoked without parentheses. Here we discover that the function head (which returns the first 6 elements of anything) defined in the *utils* package, is an S3 generic

(indicated by UseMethod) and has several methods. We use head to look at the first six lines of the head method specialized for matrix objects.

```
utils::head
## function (x, ...)
## UseMethod("head")
## <environment: namespace:utils>
methods(head)
## [1] head.data.frame* head.default*
                                          head.ftable*
                                                           head.function*
                                                                             head.matrix
## [6] head.table*
                        head.Vector
##
      Non-visible functions are asterisked
##
head(head.matrix)
##
## 1 function (x, n = 6L, ...)
## 2 {
## 3
         stopifnot(length(n) == 1L)
## 4
         n <- if (n < OL)
             max(nrow(x) + n, OL)
## 5
         else min(n, nrow(x))
## 6
```

Vignettes, especially in *Bioconductor* packages, provide an extensive narrative describing overall package functionality. Use

vignette(package="GenomicRanges")

to see a list of vignettes available in the *GenomicRanges* package; add the short name of the vignette to view in your web browser. Vignettes usually consist of text with embedded *R* code, a form of literate programming. The vignette can be read as a PDF document, while the *R* source code is present as a script file ending with extension .R. The script file can be sourced or copied into an *R* session to evaluate exactly the commands used in the vignette. For *Bioconductor* packages, vignettes are available on the package 'landing page', e.g., for *GenomicRanges*.

Scripts Many users implement analyses as scripts that load packages and input data (including massaging raw data into formats that are conducive to down-stream analysis), and then perform one or several statistical analyses to generate output in the form of summary tables or figures. R scripts are plain text files, so easily shared.

Experienced users rapidly migrate to several 'best practices' for managing their scripts. (1) R has the notion of a *vignette* that integrates a textual description with the actual analysis code, a form of literate programming. Simple and elegant vignettes can be constructed using markdown and the *knitr* package (*RStudio* has great integration of these technologies); more elaborate vignettes are based on LATEX Sweave documents. (2) *Version control*, including *git* running on your own computer or in *github* is really amazing, with a relatively moderate speed-bump to get going. Version control allows one to 'check in' a current working version of a script and data, and proceed with modifications without worrying about arbitrary file naming conventions or corrupting a previously working version. (3) It is surprisingly easy to *create an R package* that coordinates scripts, specialized and potentially re-usable functions, data, and vignettes into an easy-to-share (with lab mates or more broadly) package.

1.2.2 Bioconductor for high-throughput analysis

Bioconductor is a collection of *R* packages for the analysis and comprehension of high-throughput genomic data. *Bioconductor* started more than 10 years ago, and is widely used (Figure 1.2). It gained credibility for its statistically rigorous approach to microarray pre-processing and analysis of designed experiments, and integrative and reproducible approaches

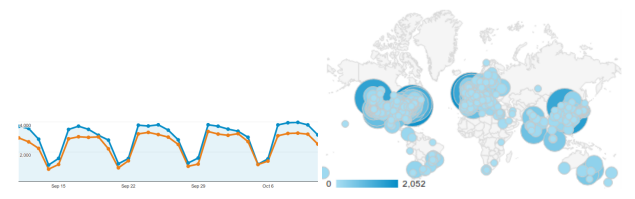


Figure 1.2: Bioconductor use, September-October 2012 (orange) and 2013 (blue).

to bioinformatic tasks. There are now 749 *Bioconductor* packages for expression and other microarrays, sequence analysis, flow cytometry, imaging, and other domains. The *Bioconductor* web site provides installation, package repository, help, and other documentation.

- The *Bioconductor* web site is at bioconductor.org. Features include:
- Introductory work flows.
- A manifest of *Bioconductor* packages arranged in BiocViews.
- Annotation (data bases of relevant genomic information, e.g., Entrez gene ids in model organisms, KEGG pathways) and experiment data (containing relatively comprehensive data sets and their analysis) packages.
- Mailing lists, including searchable archives, as the primary source of help.
- Course and conference information, including extensive reference material.
- General information about the project.
- Package developer resources, including guidelines for creating and submitting new packages.

Table 1.8 enumerates some of the packages available for sequence analysis. The table includes packages for representing sequence-related data (e.g., *GenomicRanges*, *Biostrings*), as well as domain-specific analysis such as RNA-seq (e.g., *edgeR*, *DEXSeq*), ChIP-seq (e.g., *ChIPpeakAnno*, *DiffBind*), variants (e.g., *VariantAnnotation*, *VariantTools*), and SNPs and copy number variation (e.g., *genoset*, *ggtools*). [1] illustrate integration of *Bioconductor* packages into a typical high-throughput work flow, in this case for RNA-seq analysis.

1.2.3 Strategies for working with large data

Bioinformatics data is now very large; it is not reasonable to expect all of a FASTQ or BAM file, for instance, to fit into memory. How is this data to be processed? This challenge confronts us in whatever language or tool we are using. In R and *Bioconductor*, the main approaches are to: (1) write *efficient* R code; (2) *restrict* data input to an interesting subset of the larger data set; (3) *sample* from the large data, knowing that an appropriately sized sample will accurately estimate statistics we are interested in; (4) *iterate* through large data in chunks; and (5) use *parallel* evaluation on one or several computers.

There are often many ways to accomplish a result in R, but these different ways often have very different speed or memory requirements. For small data sets these performance differences are not that important, but for large data sets (e.g., high-throughput sequencing; genome-wide association studies, GWAS) or complicated calculations (e.g., bootstrapping) performance can be important. Several approaches to achieving efficient R programming are summarized in Table 1.9; common tools used to help with assessing performance (including comparison of results from different implementations!) are in Table 1.10. Several common performance bottlenecks often have easy or moderate solutions; the most common of these are highlighted here.

R is vectorized, so traditional programming for loops are often not necessary. Rather than calculating 100000 random numbers one at a time, or squaring each element of a vector, or iterating over rows and columns in a matrix to calculate row sums, invoke the single function that performs each of these operations.

x <- runif(100000); x2 <- x^2
m <- matrix(x2, nrow=1000); y <- rowSums(m)</pre>

Concept	Packages
Data representation	IRanges, GenomicRanges, Biostrings, BSgenome, VariantAnnotation.
Input / output	ShortRead (FASTQ), Rsamtools (BAM), rtracklayer (GFF, WIG, BED), Vari-
	antAnnotation (VCF).
Annotation	AnnotationHub, biomaRt, GenomicFeatures, TxDb.*, org.*, ChIPpeakAnno,
	VariantAnnotation.
Alignment	gmapR, Rsubread, Biostrings.
Visualization	ggbio, Gviz.
Quality assessment	qrqc, seqbias, ReQON, htSeqTools, TEQC, ShortRead.
RNA-seq	BitSeq, cqn, cummeRbund, DESeq2, DEXSeq, EDASeq, edgeR, gage, goseq,
	iASeq, tweeDEseq.
ChIP-seq, etc.	BayesPeak, baySeq, ChIPpeakAnno, chipseq, ChIPseqR, ChIPsim, CSAR,
	DiffBind, MEDIPS, mosaics, NarrowPeaks, nucleR, PICS, PING, REDseq,
	Repitools, TSSi.
Variants	VariantAnnotation, VariantTools, gmapR
SNPs	snpStats, GWASTools, SeqVarTools, hapFabia, GGtools
Copy number	cn.mops, genoset, fastseq, CNAnorm, exomeCopy, seqmentSeq.
Motifs	MotifDb, BCRANK, cosmo, MotIV, seqLogo, rGADEM.
3C, etc.	HiTC, r3Cseq.
Microbiome	phyloseq, DirichletMultinomial, clstutils, manta, mcaGUI.
Work flows	QuasR, ReportingTools, easyRNASeq, ArrayExpressHTS, oneChannelGUI.
Database	SRAdb, GEOquery.

Table 1.8: Selected Bioconductor packages for high-throughput sequence analysis.

Table 1.9: Common ways to improve efficiency of R code.

Easy	Moderate
1. Selective input	1. Know relevant packages
2. Vectorize	2. Understand algorithm complexity
3. Pre-allocate and fill	3. Use parallel evaluation
4. Avoid expensive conveniences	4. Exploit libraries and C++ code

This often requires a change of thinking, turning the sequence of operations 'inside-out'. For instance, calculate the log of the square of each element of a vector by calculating the square of all elements, followed by the log of all elements $x^2 < x^2$; $x^3 < \log(x^2)$, or simply $\log x^2 < \log(x^2)$.

It may sometimes be natural to formulate a problem as a for loop, or the formulation of the problem may require that a for loop be used. In these circumstances the appropriate strategy is to pre-allocate the result object, and to fill the result during loop iteration.

```
result <- numeric(nrow(df))  ## pre-allocate
for (i in seq_len(nrow(df)))
result[[i]] <- some_calc(df[i,]) ## fill</pre>
```

T I I I I A	T 1 C		c
Table 1.10:	lools for	measuring	performance.

Function	Description
identical, all.equal	Compare content of objects.
system.time	Time required to evaluate an expression
Rprof	Time spent in each function; also summaryRprof.
tracemem	Indicate when memory copies occur (R must be configured to support this).
microbenchmark	Packages for standardizing speed measurment

Failure to pre-allocate and fill is the second cirle of R hell [4].

Using appropriate functions can greatly influence performance; it takes experience to know when an appropriate function exists. For instance, the 1m function could be used to assess differential expression of each gene on a microarray, but the *limma* package implements this operation in a way that takes advantage of the experimental design that is common to each probe on the microarray, and does so in a very efficient manner.

```
## not evaluated
library(limma) # microarray linear models
fit <- lmFit(eSet, design)</pre>
```

Using appropriate algorithms can have significant performance benefits, especially as data becomes larger. This solution requires moderate skills, because one has to be able to think about the complexity (e.g., expected number of operations) of an algorithm, and to identify algorithms that accomplish the same goal in fewer steps. For example, a naive way of identifying which of 100 numbers are in a set of size 10 might look at all 100×10 combinations of numbers (i.e., polynomial time), but a faster way is to create a 'hash' table of one of the set of elements and probe that for each of the other elements (i.e., linear time). The latter strategy is illustrated with

```
x <- 1:100; s <- sample(x, 10)
inS <- x %in% s
```

Restriction Just because a data file contains a lot of data does not mean that we are interested in all of it. In base *R*, one might use the colClasses argument to read.delim or similar function (e.g., setting some elements to NULL) to read only some columns of a large comma-separated value file.

```
## not evaluated
colClasses <-
    c("NULL", "integer", "numeric", "NULL")
df <- read.table("myfile", colClasses=colClasses)</pre>
```

In addition to the obvious benefit of using less memory than if all of the file had been read in, input will be substantially faster because less computation needs to be done to coerce values from their representation in the file to their representation in R's memory.

A variation on the idea of restricting data input is to organize the data on disk into a representation that facilitates restriction. In *R*, large data might be stored in a relational data base like the *sqlite* data bases made available by the *RSQLite* package and used in the *AnnotationDbi Bioconductor* packages. In addition to facilitating restriction, these approaches are typically faster than parsing a plain text file, because the data base software has stored data in a way that efficiently transforms from on-disk to in-memory representation.

Restriction is such a useful concept that many *Bioconductor* high throughput sequence analysis functions enable doing the right thing. Functions such as coverage, BamFile-method or readGAlignments use restrictions to read in the specific data required for them to compute the statistic of interest. Most "higher level" functions have a param=ScanBamParam() argument to allow the user to specify additional fields if desired.

Sampling R is after all a statistical language, and it sometimes makes sense to draw inferences from a sample of large data. For instance, many quality assessment statistics summarize overall properties (e.g., GC content or per-nucleotide base quality of FASTQ reads) that don't require processing of the entire data. For these statistics to be valid, the sample from the file needs to be a random sample, rather than a sample of convenience.

There are two advantages to sampling from a FASTQ (or BAM) file. The sample uses less memory than the full data. And, because less data needs to be parsed from the on-disk to in-memory representation, the input is faster.

Iteration Restriction may not be enough to wrestle large data down to size, and sampling may be inappropriate for the task at hand. A solution is then to iterate through the file. An example in base R is to open a file connection, and then read and process successive chunks of the file, e.g., reading chunks of 10000 lines

```
## NOT RUN
con <- file("<hypothetical-file>.txt")
open(f)
repeat {
    x <- readLines(f, n=10000)  # or other input function
    if (length(x) == 0)
        break
    ## work on character vector 'x'
}
close(f)</pre>
```

This paradigm extends to parsing FASTQ, BAM, VCF, and other files.

Parallel evaluation The preceding paragraphs emphasize that the starting point for analysis of large data is efficient, vectorized code. Performance differences between poorly written versus well written R code can easily span two orders of magnitude, whereas parallel processing can only increase throughput by an amount inversely proportional to the number of processing units (e.g., CPUs) available. The memory management techniques outlined earlier in this chapter are important in a parallel evaluation context. This is because we will typically be trying to exploit multiple processing cores on a single computer, and the cores will be competing for the same pool of shared memory. We thus want to arrange for the collection of processors to cooperate in dividing available memory between them, i.e., each processor needs to use only a fraction of total memory.

There are a number of ways in which R code can be made to run in parallel. The least painful and most effective will use 'multicore' functionality provided by the *parallel* package; the *parallel* package is installed by default with base R, and has a useful vignette [13].

Parallel evaluation on several cores of a single Linux or MacOS computer is particularly easy to achieve when the code is already vectorized. The solution on these operating systems is to use the functions mclapply or pvec. These functions allow the 'master' process to 'fork' processes for parallel evaluation on each of the cores of a single machine. The forked processes initially share memory with the master process, and only make copies when the forked process modifies a memory location ('copy on change' semantics). On Linux and MacOS, the mclapply function is meant to be a 'drop-in' replacement for lapply, but with iterations being evaluated on different cores.

The *parallel* package does not support fork-like behavior on Windows, where users need to more explicitly create a cluster of *R* workers and arrange for each to have the same data loaded into memory; similarly, parallel evaluation across computers (e.g., in a cluster) require more elaborate efforts to coordinate workers; this is typically done using lapply-like functions provided by the *parallel* package but specialized for simple ('snow') or more robust ('MPI') communication protocols between workers.

Data movement and random numbers are two important additional considerations in parallel evaluation. Moving data to and from cores to the manager can be expensive, so strategies that minimize explicit movement (e.g., passing file names data base queries rather than R objects read from files; reducing data on the worker before transmitting results to the manager) can be important. Random numbers need to be synchronized across cores to avoid generating the same sequences on each 'independent' computation.

Chapter 2

Sequences

Biostrings and GenomicRanges 2.1

Biostrings The Biostrings package provides tools for working with sequences. The essential data structures are DNAString and DNAStringSet, for working with one or multiple DNA sequences. The Biostrings package contains additional classes for representing amino acid and general biological strings. The BSgenome and related packages (e.g., BSgenome.Dmelanogaster.UCSC.dm3) are used to represent whole-genome sequences. Table 2.2 summarizes common operations.

```
library(Biostrings)
DNAString(c("ACACTTG"))
##
   7-letter "DNAString" instance
## seq: ACACTTG
dna <- DNAStringSet(c("AACCAA", "GCCGTCGNM"))</pre>
dna
##
   A DNAStringSet instance of length 2
##
      width seq
## [1]
       6 AACCAA
## [2]
          9 GCCGTCGNM
alphabetFrequency(dna, baseOnly=TRUE)
##
       A C G T other
## [1,] 4 2 0 0
                    0
## [2,] 0 3 3 1
```

2

Package	Description
Biostrings	Classes (e.g., DNAStringSet) and methods (e.g., alphabetFrequency,
	pairwiseAlignment) for representing and manipulating DNA and other bio-logical sequences.
BSgenome	Representation and manipulation of large (e.g., whole-genome) sequences.
ShortRead	I/O and manipulation of FASTQ files.

Table 2.1: Selected *Bioconductor* packages for representing strings and reads.

	Function	Description
Access	length, names	Number and names of sequences
	[, head, tail, rev	Subset, first, last, or reverse sequences
	С	Concatenate two or more objects
	width, nchar	Number of letters of each sequence
	Views	Light-weight sub-sequences of a sequence
Compare	==, !=, match, %in%	Element-wise comparison
•	duplicated, unique	Analog to duplicated and unique on character vectors
	sort, order	Locale-independent sort, order
	split, relist	Split or relist objects to, e.g., DNAStringSetList
Edit	subseq, subseq<-	Extract or replace sub-sequences in a set of sequences
	reverse, complement	Reverse, complement, or reverse-complement DNA
	reverseComplement	
	translate	Translate DNA to Amino Acid sequences
	chartr	Translate between letters
	replaceLetterAt	Replace letters at a set of positions by new letters
	trimLRPatterns	Trim or find flanking patterns
Count	alphabetFrequency	Tabulate letter occurrence
	letterFrequency	
	letterFrequencyInSlidingVi	ew
	consensusMatrix	Nucleotide $ imes$ position summary of letter counts
	dinucleotideFrequency	2-mer, 3-mer, and k-mer counting
	trinucleotideFrequency	,,
	oligonucleotideFrequency	
	nucleotideFrequencyAt	Nucleotide counts at fixed sequence positions
Match	matchPattern, countPattern	Short patterns in one or many $(v*)$ sequences
	vmatchPattern, vcountPatter	
	matchPDict, countPDict	Short patterns in one or many (v*) sequences (mismatch only)
	whichPDict, vcountPDict	
	vwhichPDict	
	pairwiseAlignment	Needleman-Wunsch, Smith-Waterman, etc. pairwise alignment
	matchPWM, countPWM	Occurrences of a position weight matrix
	matchProbePair	Find left or right flanking patterns
	findPalindromes	Palindromic regions in a sequence. Also
		findComplementedPalindromes
	stringDist	Levenshtein, Hamming, or pairwise alignment scores
I/0	readDNAStringSet	FASTA (or sequence only from FASTQ). Also
1/0	1 caubinabel ingbet	readBStringSet, readRNAStringSet, readAAStringSet
	writeXStringSet	Teaubouringbet, readimentingbet, reademotringbet
	writePairwiseAlignments	Write pairwiseAlignment as "pair" format
	readDNAMultipleAlignment	Multiple alignments (FASTA, "stockholm", or "clustal"). Also
	readuratettettettettette	readRNAMultipleAlignment, readAAMultipleAlignment
	write.phylip	readaminiterpresententione, readaminiterpresententient
-	#1100.bultth	

Table 2.2: Operations on strings in the *Biostrings* package.

GenomicRanges Ranges describe both features of interest (e.g., genes, exons, promoters) and reads aligned to the genome. *Bioconductor* has very powerful facilities for working with ranges, some of which are summarized in Table 2.3. These are implemented in the *GenomicRanges* package; see [9] for a more comprehensive conceptual orientation.

The GRanges class Instances of *GRanges* are used to specify genomic coordinates. Suppose we wish to represent two *D. melanogaster* genes. The first is located on the positive strand of chromosome 3R, from position 19967117 to 19973212. The second is on the minus strand of the X chromosome, with 'left-most' base at 18962306, and right-most

Package	Description
IRanges	Defines important classes (e.g., <i>IRanges</i> , <i>Rle</i>) and methods (e.g., findOverlaps,
	countOverlaps) for representing and manipulating ranges of consecutive values.
	Also introduces DataFrame, SimpleList and other classes tailored to representing
	very large data.
GenomicRanges	Range-based classes tailored to sequence representation (e.g., GRanges, GRanges-
	List), with information about strand and sequence name.
GenomicFeatures	Foundation for manipulating data bases of genomic ranges, e.g., representing coor-
	dinates and organization of exons and transcripts of known genes.

Table 2.3: Selected Bioconductor packages for representing and manipulating ranges, strings, and other data structures.

base at 18962925. The coordinates are 1-based (i.e., the first nucleotide on a chromosome is numbered 1, rather than 0), *left-most* (i.e., reads on the minus strand are defined to 'start' at the left-most coordinate, rather than the 5' coordinate), and *closed* (the start and end coordinates are included in the range; a range with identical start and end coordinates has width 1, a 0-width range is represented by the special construct where the end coordinate is one less than the start coordinate). A complete definition of these genes as *GRanges* is:

The components of a *GRanges* object are defined as vectors, e.g., of seqnames, much as one would define a *data.frame*. The start and end coordinates are grouped into an *IRanges* instance. The optional seqlengths argument specifies the maximum size of each sequence, in this case the lengths of chromosomes 3R and X in the 'dm2' build of the *D*. *melanogaster* genome. This data is displayed as

GRanges with 2 ranges and 0 metadata columns: seqnames ## ranges strand ## <Rle> <IRanges> <Rle> ## [1] chr3R [19967117, 19973212] + ## [2] chrX [18962306, 18962925] ## ## seqlengths: ## chr3R chrX 27905053 22422827

The GRanges class has many useful methods defined on it. Consult the help page

?GRanges

genes

and package vignettes

vignette(package="GenomicRanges")

for a comprehensive introduction. A GRanges instance can be subset, with accessors for getting and updating information.

```
genes<sup>[2]</sup>
## GRanges with 1 range and 0 metadata columns:
        seqnames ranges strand
<Rle> <IRanges> <Rle>
##
##
            chrX [18962306, 18962925]
##
    [1]
##
    ____
##
    seqlengths:
##
     chr3R
                   chrX
     27905053 22422827
##
strand(genes)
## factor-Rle of length 2 with 2 runs
##
   Lengths: 1 1
   Values : + -
##
## Levels(3): + - *
width(genes)
## [1] 6096 620
length(genes)
## [1] 2
names(genes) <- c("FBgn0039155", "FBgn0085359")</pre>
genes # now with names
## GRanges with 2 ranges and 0 metadata columns:
##
                seqnames ranges strand
                  <Rle> <IRanges> <Rle>
##
    FBgn0039155 chr3R [19967117, 19973212]
##
                 chrX [18962306, 18962925]
##
    FBgn0085359
##
     ____
##
     seqlengths:
##
        chr3R
                   chrX
      27905053 22422827
##
```

strand returns the strand information in a compact representation called a *run-length encoding*. The 'names' could have been specified when the instance was constructed; once named, the *GRanges* instance can be subset by name like a regular vector.

As the GRanges function suggests, the *GRanges* class extends the *IRanges* class by adding information about seqnames, strand, and other information particularly relevant to representing ranges that are on genomes. The *IRanges* class and related data structures (e.g., *RangedData*) are meant as a more general description of ranges defined in an arbitrary space. Many methods implemented on the *GRanges* class are 'aware' of the consequences of genomic location, for instance treating ranges on the minus strand differently (reflecting the 5' orientation imposed by DNA) from ranges on the plus strand.

Operations on ranges The *GRanges* class has many useful methods. We use *IRanges* to illustrate these operations to avoid complexities associated with strand and seqnames, but the operations are comparable on *GRanges*. We begin with a simple set of ranges:

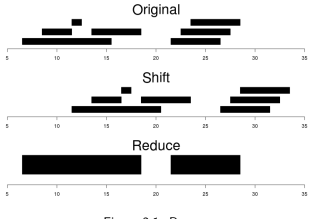


Figure 2.1: Ranges

These and some common operations are illustrated in the upper panel of Figure 2.1 and summarized in Table 2.4. Common operations on ranges are summarized in Table 2.4. Methods on ranges can be grouped as follows:

Intra-range methods act on each range independently. These include flank, narrow, reflect, resize, restrict, and shift, among others. An illustration is shift, which translates each range by the amount specified by the shift argument. Positive values shift to the right, negative to the left; shift can be a vector, with each element of the vector shifting the corresponding element of the *IRanges* instance. Here we shift all ranges to the right by 5, with the result illustrated in the middle panel of Figure 2.1.

shift(ir, 5)

##	IRaı	iges	of	leı	ngth	7
##		star	ct	end	widt	th
##	[1]	1	12	20		9
##	[2]	1	14	16		3
##	[3]	1	17	17		1
##	[4]	1	19	23		5
##	[5]	2	27	31		5
##	[6]	2	28	32		5
##	[7]	2	29	33		5

Inter-range methods act on the collection of ranges as a whole. These include disjoin, reduce, gaps, and range. An illustration is reduce, which reduces overlapping ranges into a single range, as illustrated in the lower panel of Figure 2.1.

```
reduce(ir)
```

IRanges of length 2
start end width
[1] 7 18 12
[2] 22 28 7

coverage is an inter-range operation that calculates how many ranges overlap individual positions. Rather than returning ranges, coverage returns a compressed representation (run-length encoding)

coverage(ir)

integer-Rle of length 28 with 12 runs
Lengths: 6 2 4 1 2 3 3 1 1 3 1 1
Values : 0 1 2 1 2 1 0 1 2 3 2 1

The run-length encoding can be interpreted as 'a run of length 6 of nucleotides covered by 0 ranges, followed by a run of length 2 of nucleotides covered by 1 range...'.

Between methods act on two (or sometimes more) IRanges instances. These include intersect, setdiff, union,

Category	Function	Description
Accessors	start, end, width	Get or set the starts, ends and widths
	names	Get or set the names
	mcols, metadata	Get or set metadata on elements or object
	length	Number of ranges in the vector
	range	Range formed from min start and max end
Ordering	<, <=, >, >=, ==, !=	Compare ranges, ordering by start then width
	sort, order, rank	Sort by the ordering
	duplicated	Find ranges with multiple instances
	unique	Find unique instances, removing duplicates
Arithmetic	r + x, r - x, r * x	Shrink or expand ranges r by number x
	shift	Move the ranges by specified amount
	resize	Change width, anchoring on start, end or mid
	distance	Separation between ranges (closest endpoints)
	restrict	Clamp ranges to within some start and end
	flank	Generate adjacent regions on start or end
Set operations	reduce	Merge overlapping and adjacent ranges
	intersect, union, setdiff	Set operations on reduced ranges
	pintersect, punion, psetdiff	Parallel set operations, on each x[i], y[i]
	gaps, pgap	Find regions not covered by reduced ranges
	disjoin	Ranges formed from union of endpoints
Overlaps	findOverlaps	Find all overlaps for each x in y
	countOverlaps	Count overlaps of each x range in y
	nearest	Find nearest neighbors (closest endpoints)
	precede, follow	Find nearest y that \mathbf{x} precedes or follows
	x %in% y	Find ranges in ${f x}$ that overlap range in ${f y}$
Coverage	coverage	Count ranges covering each position
Extraction	r[i]	Get or set by logical or numeric index
	r[[i]]	Get integer sequence from start[i] to end[i]
	subsetByOverlaps	Subset \mathbf{x} for those that overlap in \mathbf{y}
	head, tail, rev, rep	Conventional R semantics
Split, combine	split	Split ranges by a factor into a RangesList
	с	Concatenate two or more range objects

Table 2.4: Common operations on IRanges, GRanges and GRangesList.

pintersect, psetdiff, and punion.

The countOverlaps and findOverlaps functions operate on two sets of ranges. countOverlaps takes its first argument (the query) and determines how many of the ranges in the second argument (the subject) each overlaps. The result is an integer vector with one element for each member of query. findOverlaps performs a similar operation but returns a more general matrix-like structure that identifies each pair of query / subject overlaps. Both arguments allow some flexibility in the definition of 'overlap'.

mcols and metadata The *GRanges* class (actually, most of the data structures defined or extending those in the *IRanges* package) has two additional very useful data components. The mcols function allows information on each range to be stored and manipulated (e.g., subset) along with the *GRanges* instance. The element metadata is represented as a *DataFrame*, defined in *IRanges* and acting like a standard *R data.frame* but with the ability to hold more complicated data structures as columns (and with element metadata of its own, providing an enhanced alternative to the *Biobase* class *AnnotatedDataFrame*).

metadata allows addition of information to the entire object. The information is in the form of a list; any data can be provided.

```
metadata(genes) <- list(CreatedBy="A. User", Date=date())</pre>
```

The GRangesList class The GRanges class is extremely useful for representing simple ranges. Some next-generation sequence data and genomic features are more hierarchically structured. A gene may be represented by several exons within it. An aligned read may be represented by discontinuous ranges of alignment to a reference. The *GRangesList* class represents this type of information. It is a list-like data structure, which each element of the list itself a *GRanges* instance. The gene FBgn0039155 contains several exons, and can be represented as a list of length 1, where the element of the list contains a *GRanges* object with 7 elements:

```
## GRangesList of length 1:
  $FBgn0039155
##
##
  GRanges with 7 ranges and 2 metadata columns:
##
         seqnames
                                  ranges strand |
                                                      exon_id
                                                                 exon_name
##
             <Rle>
                               <IRanges> <Rle> |
                                                   <integer> <character>
##
     [1]
             chr3R [19967117, 19967382]
                                               + |
                                                        50486
                                                                       <NA>
     [2]
             chr3R [19970915, 19971592]
##
                                               + |
                                                        50487
                                                                       <NA>
##
     [3]
             chr3R [19971652, 19971770]
                                               + 1
                                                        50488
                                                                       <NA>
##
     [4]
             chr3R [19971831, 19972024]
                                               + |
                                                        50489
                                                                       <NA>
##
     [5]
             chr3R [19972088, 19972461]
                                               + |
                                                                       <NA>
                                                        50490
             chr3R [19972523, 19972589]
##
     [6]
                                               + |
                                                        50491
                                                                       <NA>
##
     [7]
             chr3R [19972918, 19973212]
                                               + |
                                                        50492
                                                                       <NA>
##
##
  ___
## seqlengths:
##
       chr3R
##
   27905053
```

The *GRangesList* object has methods one would expect for lists (e.g., length, sub-setting). Many of the methods introduced for working with *IRanges* are also available, with the method applied element-wise.

2.2 From whole genome to short read

2.2.1 Large and whole-genome sequences

There are three ways in which whole-genome sequences are represented in *Bioconductor*. For model organisms, the *BSgenome* package and suite of annotations (e.g., *BSgenome.Hsapiens.UCSC.hg19*) can be used to query genome coordinates and to load whole chromosomes into memory. The annotation packages contain optional 'masks', e.g., of repeat regions, and are explored in an exercise at the end of this section.

The *Rsamtools* package provides an interface to indexed FASTA files via the FaFile function; this can be used to input whole genomes or, more usefully, along with *GRanges* instances to input selected sequences. This is explored in an exercise related to the *AnnotationHub* package later today.

Finally, the rtracklayer package enables import of '2bit' FASTA format files.

2.2.2 Short reads

Short read formats The Illumina GAII and HiSeq technologies generate sequences by measuring incorporation of florescent nucleotides over successive PCR cycles. These sequencers produce output in a variety of formats, but *FASTQ* is ubiquitous. Each read is represented by a record of four components:

The first and third lines (beginning with @ and + respectively) are unique identifiers. The identifier produced by the sequencer typically includes a machine id followed by colon-separated information on the lane, tile, x, and y coordinate of the read. The example illustrated here also includes the SRA accession number, added when the data was submitted to the archive. The machine identifier could potentially be used to extract information about batch effects. The spatial coordinates (lane, tile, x, y) are often used to identify optical duplicates; spatial coordinates can also be used during quality assessment to identify artifacts of sequencing, e.g., uneven amplification across the flow cell, though these spatial effects are rarely pursued.

The second and fourth lines of the FASTQ record are the nucleotides and qualities of each cycle in the read. This information is given in 5' to 3' orientation as seen by the sequencer. A letter N in the sequence is used to signify bases that the sequencer was not able to call. The fourth line of the FASTQ record encodes the quality (confidence) of the corresponding base call. The quality score is encoded following one of several conventions, with the general notion being that letters later in the visible ASCII alphabet

```
## !"#$%&'()*+,-./0123456789:;<=>?@ABCDEFGHIJKLMN0
## PQRSTUVWXYZ[\]^_`abcdefghijklmnopqrstuvwxyz{|}~
```

are of higher quality; this is developed further below. Both the sequence and quality scores may span multiple lines.

Technologies other than Illumina use different formats to represent sequences. Roche 454 sequence data is generated by 'flowing' labeled nucleotides over samples, with greater intensity corresponding to longer runs of A, C, G, or T. This data is represented as a series of 'flow grams' (a kind of run-length encoding of the read) in Standard Flowgram Format (SFF). The *Bioconductor* package *R453Plus1Toolbox* has facilities for parsing SFF files, but after quality control steps the data are frequently represented (with some loss of information) as FASTQ. SOLiD technologies produce sequence data using a 'color space' model. This data is not easily read in to *R*, and much of the error-correcting benefit of the color space model is lost when converted to FASTQ; SOLiD sequences are not well-handled by *Bioconductor* packages.

Short reads in R FASTQ files can be read in to *R* using the readFastq function from the *ShortRead* package. Use this function by providing the path to a FASTQ file. There are sample data files available in the bigdata folder

```
bigdata <- file.choose()</pre>
```

Each file consists of 1 million reads from a lane of the Pasilla data set.

```
library(ShortRead)
fastqDir <- file.path(bigdata, "fastq")
fastqFiles <- dir(fastqDir, full=TRUE)
fq <- readFastq(fastqFiles[1])
fq
## class: ShortReadQ
## length: 1000000 reads; width: 37 cycles</pre>
```

The data are represented as an object of class *ShortReadQ*.

head(sread(fq), 3)

```
## A DNAStringSet instance of length 3
## width seq
## [1] 37 GTTTTGTCCAAGTTCTGGTAGCTGAATCCTGGGGCGCC
## [2] 37 GTTGTCGCATTCCTTACTCTCATTCGGGAATTCTGTT
## [3] 37 GAATTTTTTGAGAGCGAAATGATAGCCGATGCCCTGA
```

head(quality(fq), 3)

```
## class: FastqQuality
## quality:
## A BStringSet instance of length 3
## width seq
## [2]
      ## [3]
      head(id(fq), 3)
   A BStringSet instance of length 3
##
##
    width seq
## [1]
      58 SRR031724.1 HWI-EAS299_4_30M2BAAXX:5:1:1513:1024 length=37
      57 SRR031724.2 HWI-EAS299_4_30M2BAAXX:5:1:937:1157 length=37
## [2]
```

[3] 58 SRR031724.4 HWI-EAS299_4_30M2BAAXX:5:1:1443:1122 length=37

The ShortReadQ class illustrates class inheritance. It extends the ShortRead class

```
getClass("ShortReadQ")
## Class "ShortReadQ" [package "ShortRead"]
##
## Slots:
##
               quality
                                              id
## Name:
                              sread
## Class: QualityScore DNAStringSet BStringSet
##
## Extends:
## Class "ShortRead", directly
## Class ".ShortReadBase", by class "ShortRead", distance 2
##
## Known Subclasses: "AlignedRead"
```

Methods defined on ShortRead are available for ShortReadQ.

showMethods(class="ShortRead", where="package:ShortRead")

For instance, the width can be used to demonstrate that all reads are of the same width:

```
table(width(fq))
##
##
27
```

37 ## 1000000

The alphabetByCycle function summarizes use of nucleotides at each cycle in a (equal width) ShortReadQ or DNAStringSet instance.

```
abc <- alphabetByCycle(sread(fq))
abc[1:4, 1:8]
## cycle
## alphabet [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8]</pre>
```

 ##
 A
 78194
 153156
 200468
 230120
 283083
 322913
 162766
 220205

 ##
 C
 439302
 265338
 362839
 251434
 203787
 220855
 253245
 287010

 ##
 G
 397671
 270342
 258739
 356003
 301640
 247090
 227811
 246684

 ##
 T
 84833
 311164
 177954
 162443
 211490
 209142
 356178
 246101

We mentioned sampling and iteration as strategies for dealing with large data. A very common reason for looking at FASTQ data is to explore sequence quality. In these circumstances it is often not necessary to parse the entire FASTQ file. Instead create a representative sample

```
sampler <- FastqSampler(fastqFiles[1], 1000000)
yield(sampler) # sample of 1000000 reads</pre>
```

```
## class: ShortReadQ
## length: 1000000 reads; width: 37 cycles
```

A second common scenario is to pre-process reads, e.g., trimming low-quality tails, adapter sequences, or artifacts of sample preparation. The *FastqStreamer* class can be used to 'stream' over the fastq files in chunks, processing each chunk independently.

Quality assessment ShortRead contains facilities for quality assessment of FASTQ files. Here we generate a report from a sample of 1 million reads from each of our files and display it in a web browser

```
qa <- qa(dirname(fastqFiles), "*.fastq", type="fastq")
rpt <- report(qa, dest=tempfile())
browseURL(rpt)</pre>
```

A report from a larger subset of the experiment is available

2.3 Exercises

Exercise 1 Develop a function gcFunction to calculate GC content; likely your function will use alphabetFrequency. Demonstrate its use on a DNAStringSet instance.

Solution: Here is the gcFunction helper function to calculate GC content:

```
gcFunction <-
   function(x)
{
    alf <- alphabetFrequency(x, as.prob=TRUE)
    rowSums(alf[,c("G", "C")])
}</pre>
```

The gcFunction is really straight-forward: it invokes the function alphabetFrequency from the *Biostrings* package. This returns a simple matrix of exon \times nucleotide probabilities. The row sums of the G and C columns of this matrix are the GC contents of each exon. Here's a simple illustration:

gcFunction(dna)
[1] 0.3333 0.6667

28

Exercise 2 This exercise calculates the GC content of two genes. We make use of a BSgenome package, and DNAString and GRanges classes.

Load the BSgenome.Dmelanogaster.UCSC.dm3 data package, containing the UCSC representation of D. melanogaster genome assembly dm3. Discover the content of the package by evaluating Dmelanogaster.

Create the genes object (a GRanges instance), from a page or so ago. Use getSeq to retrieve a DNAStringSet representing the sequence of each gene. Use gcFunction to calculate the GC content of each gene. Compare this to the density of short reads, calcuated later.

Solution: Here we load the *D. melanogaster* genome and some genome coordinates. We then select a single chromosome, and create Views that reflect the ranges of the FBgn0002183.

```
library(BSgenome.Dmelanogaster.UCSC.dm3)
Dmelanogaster
## Fly genome
## |
## | organism: Drosophila melanogaster (Fly)
## | provider: UCSC
## | provider version: dm3
## | release date: Apr. 2006
## | release name: BDGP Release 5
## |
## | single sequences (see '?seqnames'):
## | chr2L chr2R
                            chr3L
                                      chr3R
                                                 chr4
                                                            chrX
                                                                       chrU
                                                                                 chrM
## | chr2LHet chr2RHet
                           chr3LHet
                                      chr3RHet
                                                 chrXHet
                                                            chrYHet
                                                                       chrUextra
## |
## | multiple sequences (see '?mseqnames'):
## | upstream1000 upstream2000 upstream5000
## |
## | (use the '$' or '[[' operator to access a given sequence)
data(genes)
genes
## GRanges with 15682 ranges and 1 metadata column:
##
                                                     seqnames
                                     ranges strand
                                                               gene_id
##
                   <Rle>
                                    <IRanges> <Rle>
                                                      | <CharacterList>
##
    FBgn0000003 chr3R [ 2648220, 2648518]
                                                 +
                                                      FBgn000003
##
    FBgn0000008 chr2R [18024494, 18060346]
                                                     FBgn000008
                                                  +
    FBgn0000014 chr3R [12632936, 12655767]
##
                                                     FBgn0000014
    FBgn0000015
                  chr3R [12752932, 12797958]
                                                      FBgn0000015
##
                  chr3L [16615470, 16640982]
##
    FBgn0000017
                                                      FBgn0000017
                                                  _
##
            . . .
                   . . .
                                          . . .
                                                 . . .
                                                    . . .
    FBgn0264723 chr3L [12238610, 12239148]
##
                                                            FBgn0264723
    FBgn0264724 chr3L [15327882, 15329271]
##
                                                  +
                                                      FBgn0264724
##
    FBgn0264725 chr3L [12025657, 12026099]
                                                  +
                                                      FBgn0264725
##
    FBgn0264726 chr3L [12020901, 12021253]
                                                      FBgn0264726
                   chr3L [22065469, 22065720]
##
    FBgn0264727
                                                      FBgn0264727
##
##
    seqlengths:
##
                   chr2R
                             chr3L
         chr2L
                                      chr3R ... chr3RHet
                                                            chrXHet
                                                                      chrYHet chrUextra
##
      23011544 21146708 24543557 27905053 ...
                                                  2517507
                                                             204112
                                                                      347038 29004656
```

seq <- getSeq(Dmelanogaster, genes)</pre>

The subject GC content is

gc <- gcFunction(seq)</pre>

Exercise 3 Use the file path bigdata and the file.path and dir functions to locate the fastq file from [3] (the file was obtained as described in the pasilla experiment data package).

Input the fastq files using readFastq from the ShortRead package.

Use alphabetFrequency to summarize the GC content of all reads (hint: use the sread accessor to extract the reads, and the collapse=TRUE argument to the alphabetFrequency function). Using the helper function gcFunction defined elsewhere in this document, draw a histogram of the distribution of GC frequencies across reads.

Use alphabetByCycle to summarize the frequency of each nucleotide, at each cycle. Plot the results using matplot, from the graphics package.

As an advanced exercise, and if on Mac or Linux, use the parallel package and mclapply to read and summarize the GC content of reads in two files in parallel.

Use gcFunction to calculate the GC content in each gene.

Solution: Discovery:

```
dir(bigdata)
```

```
## [1] "bam" "fastq"
```

```
fls <- dir(file.path(bigdata, "fastq"), full=TRUE)</pre>
```

Input:

```
fq <- readFastq(fls[1])</pre>
```

A histogram of the GC content of individual reads is obtained with

```
gc <- gcFunction(sread(fq))
hist(gc)</pre>
```

Alphabet by cycle:

```
abc <- alphabetByCycle(sread(fq))
matplot(t(abc[c("A", "C", "G", "T"),]), type="l")</pre>
```

(Mac, Linux only): processing on multiple cores.

```
library(parallel)
gc0 <- mclapply(fls, function(fl) {
  fq <- readFastq(fl)
  gc <- gcFunction(sread(fq))
  table(cut(gc, seq(0, 1, .05)))
})
## simplify list of length 2 to 2-D array
gc <- simplify2array(gc0)
matplot(gc, type="s")</pre>
```

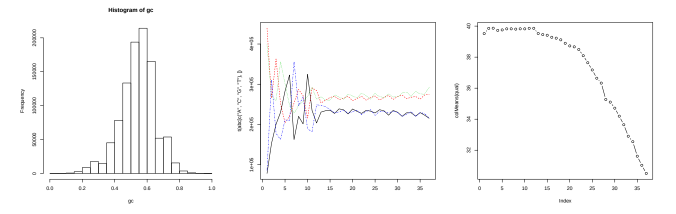


Figure 2.2: Short read GC content (left), alphabet-by-cycle (middle), and quality scores (right)

Exercise 4 Use quality to extract the quality scores of the short reads. Interpret the encoding qualitatively. Convert the quality scores to a numeric matrix, using as. Inspect the numeric matrix (e.g., using dim) and understand what it represents.

Use colMeans to summarize the average quality score by cycle. Use plot to visualize this.

Solution:

```
head(quality(fq))
## class: FastqQuality
##
 quality:
##
   A BStringSet instance of length 6
##
     width seq
       ##
  [1]
  [2]
       ##
##
  [3]
       [4]
##
       37 IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII. $
##
 [5]
## [6]
       37 III.IIIIIIIIIIIIIIIII//IIE(-EIH<IIII
qual <- as(quality(fq), "matrix")</pre>
dim(qual)
## [1] 1000000
              37
plot(colMeans(qual), type="b")
```

Exercise 5 As an independent exercise, visit the qrqc landing page and explore the package vignette. Use the qrqc package (you may need to install this) to generate base and average quality plots for the data, like those in the report generated by ShortRead.

Chapter 3

Genes and Genomes

Bioconductor provides extensive annotation resources. These can be *gene-*, or *genome-*centric. Annotations can be provided in packages curated by *Bioconductor*, or obtained from web-based resources. Gene-centric *AnnotationDbi* packages include:

- Organism level: e.g. org.Mm.eg.db, Homo.sapiens.
- Platform level: e.g. hgu133plus2.db, hgu133plus2.probes, hgu133plus2.cdf.
- Homology level: e.g. hom.Dm.inp.db.
- System biology level: GO.db, KEGG.db, Reactome.db.

Examples of genome-centric packages include:

- *GenomicFeatures*, to represent genomic features, including constructing reproducible feature or transcript data bases from file or web resources.
- Pre-built transcriptome packages, e.g. *TxDb.Hsapiens.UCSC.hg19.knownGene* based on the *H. sapiens* UCSC hg19 knownGenes track.
- BSgenome for whole genome sequence representation and manipulation.
- Pre-built genomes, e.g., BSgenome. Hsapiens. UCSC. hg19 based on the H. sapiens UCSC hg19 build.

Web-based resources include

- *biomaRt* to query biomart resource for genes, sequence, SNPs, and etc.
- rtracklayer for interfacing with browser tracks, especially the UCSC genome browser.

3.1 Gene annotation

3.1.1 Bioconductor data annotation packages

Organism-level ('org') packages contain mappings between a central identifier (e.g., Entrez gene ids) and other identifiers (e.g. GenBank or Uniprot accession number, RefSeq id, etc.). The name of an org package is always of the form org.<Sp>.<id>.db (e.g. org.Sc.sgd.db) where <Sp> is a 2-letter abbreviation of the organism (e.g. Sc for Saccharomyces cerevisiae) and <id> is an abbreviation (in lower-case) describing the type of central identifier (e.g. sgd for gene identifiers assigned by the Saccharomyces Genome Database, or eg for Entrez gene ids). The "How to use the '.db' annotation packages' vignette in the AnnotationDbi package (org packages are only one type of ".db" annotation packages) is a key reference. The '.db' and most other Bioconductor annotation packages are updated every 6 months.

Annotation packages contain an object named after the package itself. These objects are collectively called *AnnotationDb* objects, with more specific classes named *OrgDb*, *ChipDb* or *TranscriptDb* objects. Methods that can be applied to these objects include cols, keys, keytypes and select. Common operations for retrieving annotations are summarized in Table 3.1.

3.1.2 Internet resources

A short summary of select *Bioconductor* packages enabling web-based queries is in Table 3.2.

Category	Function	Description
Discover	columns	List the kinds of columns that can be returned
	keytypes	List columns that can be used as keys
	keys	List values that can be expected for a given keytype
	select	Retrieve annotations matching keys, keytype and columns
Manipulate	setdiff, union, intersect	Operations on sets
	duplicated, unique	Mark or remove duplicates
	%in%, match	Find matches
	any, all	Are any TRUE? Are all?
	merge	Combine two different data.frames based on shared keys
GRanges*	transcripts, exons, cds	Features (transcripts, exons, coding sequence) as <i>GRanges</i> .
	transcriptsBy , exonsBy	Features group by gene, transcript, etc., as GRangesList.
	cdsBy	

Table 3.1: Common operations for retrieving and manipulating annotations.

Table 3.2: Selected packages querying web-based annotation services.

Package	Description
AnnotationHub	Ensembl, Encode, dbSNP, UCSC data objects
biomaRt	http://biomart.org, Ensembl and other annotations
PSICQUIC	https://code.google.com/p/psicquic.org, protein interactions
uniprot.ws	http://uniprot.org, protein annotations
KEGGREST	http://www.genome.jp/kegg, KEGG pathways
SRAdb	http://www.ncbi.nlm.nih.gov/sra, sequencing experiments.
rtracklayer	http://genome.ucsc.edu, genome tracks.
GEOquery	http://www.ncbi.nlm.nih.gov/geo/, array and other data
ArrayExpress	http://www.ebi.ac.uk/arrayexpress/, array and other data

Using biomaRt The *biomaRt* package offers access to the online biomart resource. this consists of several data base resources, referred to as 'marts'. Each mart allows access to multiple data sets; the *biomaRt* package provides methods for mart and data set discovery, and a standard method getBM to retrieve data.

3.1.3 Exercises

Exercise 6 What is the name of the org package for Drosophila? Load it. Display the OrgDb object for the org.Dm.eg.db package. Use the columns method to discover which sorts of annotations can be extracted from it.

Use the keys method to extract UNIPROT identifiers and then pass those keys in to the select method in such a way that you extract the SYMBOL (gene symbol) and KEGG pathway information for each.

Use select to retrieve the ENTREZ and SYMBOL identifiers of all genes in the KEGG pathway 00310.

Solution: The *OrgDb* object is named org.Dm.eg.db.

```
library(org.Dm.eg.db)
columns(org.Dm.eg.db)
##
    [1] "ENTREZID"
                         "ACCNUM"
                                         "ALIAS"
                                                         "CHR"
                                                                         "CHRLOC"
    [6] "CHRLOCEND"
                         "ENZYME"
                                         "MAP"
                                                                         "PMID"
##
                                                         "PATH"
   [11] "REFSEQ"
##
                         "SYMBOL"
                                         "UNIGENE"
                                                         "ENSEMBL"
                                                                         "ENSEMBLPROT"
   [16] "ENSEMBLTRANS" "GENENAME"
                                        "UNIPROT"
                                                        "GO"
                                                                         "EVIDENCE"
##
## [21] "ONTOLOGY"
                         "GOALL"
                                         "EVIDENCEALL"
                                                        "ONTOLOGYALL"
                                                                        "FLYBASE"
## [26] "FLYBASECG"
                        "FLYBASEPROT"
```

keytypes(org.Dm.eg.db)

```
##
    [1] "ENTREZID"
                        "ACCNUM"
                                       "ALIAS"
                                                       "CHR"
                                                                      "CHRLOC"
   [6] "CHRLOCEND"
                        "ENZYME"
                                       "MAP"
                                                       "PATH"
                                                                      "PMID"
##
## [11] "REFSEQ"
                        "SYMBOL"
                                       "UNIGENE"
                                                       "ENSEMBL"
                                                                      "ENSEMBLPROT"
## [16] "ENSEMBLTRANS" "GENENAME"
                                       "UNIPROT"
                                                      "GO"
                                                                      "EVIDENCE"
## [21] "ONTOLOGY"
                       "GOALL"
                                       "EVIDENCEALL"
                                                      "ONTOLOGYALL" "FLYBASE"
## [26] "FLYBASECG"
                       "FLYBASEPROT"
uniprotKeys <- head(keys(org.Dm.eg.db, keytype="UNIPROT"))</pre>
cols <- c("SYMBOL", "PATH")</pre>
select(org.Dm.eg.db, keys=uniprotKeys, columns=cols, keytype="UNIPROT")
##
     UNIPROT SYMBOL PATH
## 1 Q8IRZO CG3038 <NA>
## 2 Q95RP8 CG3038 <NA>
## 3 M9PGH7
                 G9a 00310
## 4 Q95RU8
                 G9a 00310
## 5 Q9W5H1 CG13377 <NA>
## 6 P39205
              cin <NA>
```

Selecting UNIPROT and SYMBOL ids of KEGG pathway 00310 is very similar:

kegg <- select(org.Dm.eg.db, "00310", c("UNIPROT", "SYMBOL"), "PATH")
Warning: 'select' resulted in 1:many mapping between keys and return rows
nrow(kegg)
[1] 35
head(kegg, 3)
PATH UNIPROT SYMBOL
1 00310 M9PGH7 G9a
2 00310 Q95RU8 G9a
3 00310 M9NE25 Hmt4-20</pre>

Exercise 7 Load the biomaRt package and list the available marts. Choose the ensembl mart and list the datasets for that mart. Set up a mart to use the ensembl mart and the hsapiens_gene_ensembl dataset.

A biomaRt dataset can be accessed via getBM. In addition to the mart to be accessed, this function takes filters and attributes as arguments. Use filterOptions and listAttributes to discover values for these arguments. Call getBM using filters and attributes of your choosing.

Solution:

Use head(res) to see the results.

Exercise 8 As an exercise for later in this course, annotate the genes that are differentially expressed in Dr. Huber's lab, e.g., find the GENENAME associated with the five most differentially expressed genes. Do these make biological sense? Can you merge the annotation results with the 'top table' results to provide a statistically and biologically informative summary?

3.2 Genome annotation

There are a diversity of packages and classes available for representing large genomes. Several include:

TxDb.* For transcript and other genome / coordinate annotation.

BSgenome For whole-genome representation. See available.packages for pre-packaged genomes, and the vignette 'How to forge a BSgenome data package' in the

Homo.sapiens For integrating *TxDb** and *org.** packages.

SNPlocs.* For model organism SNP locations derived from dbSNP.

FaFile (Rsamtools) for accessing indexed FASTA files.

SIFT.*, PolyPhen, ensemblVEP Variant effect scores.

3.2.1 Bioconductor transcript annotation packages

Genome-centric packages are very useful for annotations involving genomic coordinates. It is straight-forward, for instance, to discover the coordinates of coding sequences in regions of interest, and from these retrieve corresponding DNA or protein coding sequences. Other examples of the types of operations that are easy to perform with genome-centric annotations include defining regions of interest for counting aligned reads in RNA-seq experiments and retrieving DNA sequences underlying regions of interest in ChIP-seq analysis, e.g., for motif characterization.

3.2.2 AnnotationHub

The AnnotationHub package makes it easier to access genome-scale resources. It consists of an R 'client' that queries an AnnotationHub server. There server contains lightly-curated versions of large genome resources such as UCSC or ENCODE tracks and Ensembl gtf or fasta files. There are a large number of resources available.

```
library(AnnotationHub)
hub <- AnnotationHub()
hub
## class: AnnotationHub
## length: 8674
## filters: none
## hubUrl: http://annotationhub.bioconductor.org/ah
## snapshotVersion: 2.13; snapshotDate: 2013-06-29
## hubCache: /home/mtmorgan/.AnnotationHub</pre>
```

The hub can be queried for metadata metadata(hub) or explored using tab completion hub\$ensembl<tab>.

'Light curation' means that they have been transformed to data structures that are particularly easy to use from *Bioconductor*. Here we retrieve the complete sequence of *Pan troglodytes* (this is about 865M of data to download).

pan <-

hub\$ensembl.release.72.fasta.pan_troglodytes.dna.Pan_troglodytes.CHIMP2.1..4.72.dna.toplevel.fa.rz

pan is a razip-compressed file FaFile from the Rsamtools package. We can discover sequences the file contains

```
seqinfo(pan)
```

and retrieve arbitrary regions

```
roi <- GRanges("22", IRanges(c(30000000, 40000000), width=100000))
getSeq(pan, roi)</pre>
```

Retrievals from the AnnotationHub server are cached, so the next time access is fast It's cached, so the next time it's fast

```
system.time(
   hub$ensembl.release.72.fasta.pan_troglodytes.dna.Pan_troglodytes.CHIMP2.1.4.72.dna.toplevel.fa.rz
)
### user system elapsed
### 0.024 0.000 0.421
```

3.2.3 rtracklayer

rtracklayer

```
library(rtracklayer)
```

3.2.4 VariantAnnotation

A major product of DNASeq experiments are catalogs of called variants (e.g., SNPs, indels); recent scenarios use public consortium called variants to develop novel predictive filters of regulatory function [8]. We will use the *VariantAnnotation* package to explore this type of data. Sample data included in the package are a subset of chromosome 22 from the 1000 Genomes project. Variant Call Format (VCF; full description) text files contain meta-information lines, a header line with column names, data lines with information about a position in the genome, and optional genotype information on samples for each position.

Important operations on VCF files available with the VariantAnnotation package are summarized in Table 3.3.

Data input VariantAnnotation input of whole VCF files (readVcf), or more conveniently restriction to specific fields (e.g., readGeno, readInfo), samples or ranges (using ScanVcfParam). It is also straight-forward to iterate through large VCF files by using a TabixFile with a yieldSize specification, or to use filterVcf to transform a large VCF to a subset of variants relevant in a particular study.

SNP Annotation Variants can be easily identified according to region such as coding, intron, intergenic, spliceSite etc. Amino acid coding changes are computed for the non-synonymous variants. SIFT and PolyPhen databases provide predictions of how severely the coding changes affect protein function. Additional annotations are easily crafted using the *GenomicRanges* and *GenomicFeatures* software in conjunction with *Bioconductor* and broader community annotation resources.

Locating variants in and around genes Variant location with respect to genes can be identified with the locateVariants function. Regions are specified in the region argument and can be one of the following constructors: CodingVariants(), IntronVariants(), FiveUTRVariants(), ThreeUTRVariants(), IntergenicVariants(), SpliceSiteVariants(), or AllVariants(). Location definitions are shown in Table 3.4.

Category	Function	Description
Read	scanVcfHeader	Retrieve file header information
	scanVcfParam	Select fields to read in
	readVcf	Read VCF file into a VCF class
	scanVcf	Read VCF file into a list
Filter	filterVcf	Filter a VCF from one file to another
Write	writeVcf	Write a VCF file to disk
Annotate	locateVariants	Identify where variant overlaps a gene annotation
	predictCoding	Amino acid changes for variants in coding regions
	summarizeVariants	Summarize variant counts by sample
SNPs	genotypeToSnpMatrix	Convert genotypes to a SnpMatrix
	GLtoGP	Convert genotype likelihoods to genotypes
	snpSummary	Counts and distribution statistics for SNPs
Manipulate	expand	Convert CompressedVCF to ExpandedVCF
	cbind, rbind	Combine by column or row

Table 3.3:	Working	with	VCF	files	and	data.

```
Table 3.4: Variant locations
```

Location	Details
coding	Within a coding region
fiveUTR	Within a 5' untranslated region
threeUTR	Within a 3' untranslated region
intron	Within an intron region
intergenic	Not within a transcript associated with a gene
spliceSite	Overlaps any of the first or last 2 nucleotides of an intron

Amino acid coding changes predictCoding computes amino acid coding changes for non-synonymous variants. Only ranges in query that overlap with a coding region in subject are considered. Reference sequences are retrieved from either a BSgenome or fasta file specified in seqSource. Variant sequences are constructed by substituting, inserting or deleting values in the varAllele column into the reference sequence. Amino acid codes are computed for the variant codon sequence when the length is a multiple of 3.

The query argument to predictCoding can be a GRanges or VCF. When a GRanges is supplied the varAllele argument must be specified. In the case of a VCF object, the alternate alleles are taken from alt(<VCF>) and the varAllele argument is not specified.

The result is a modified query containing only variants that fall within coding regions. Each row represents a variant-transcript match so more than one row per original variant is possible.

```
library(VariantAnnotation)
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
txdb <- TxDb.Hsapiens.UCSC.hg19.knownGene
library(BSgenome.Hsapiens.UCSC.hg19)
fl <- system.file("extdata", "chr22.vcf.gz", package="VariantAnnotation")
vcf <- readVcf(fl, "hg19")
seqlevels(vcf, force=TRUE) <- c("22"="chr22")
coding <- predictCoding(vcf, txdb, seqSource=Hsapiens)
coding[5:9]</pre>
```

##	GRanges with 5 range	es and 1	17 metadata	columns:			
##	Se	eqnames		ranges	strand	paramRangeID	REF
##		<rle></rle>		<iranges></iranges>	<rle></rle>	<factor></factor>	<dnastringset></dnastringset>
##	22:50301584_C/T	chr22	[50301584,	50301584]	-	<na></na>	С
##	rs114264124	chr22	[50302962,	50302962]	-	<na></na>	С

##	rs149209714	chr22 [5030	200E E02	1000F]		-	<na></na>	С
## ##			•			1	<na></na>	T
	22:50303554_T/C	chr22 [5030	•			-		
##	rs12167668	chr22 [5030	•			-	<na></na>	C
##			ALT	QUAL		ILTER	varAllele	CDSLOC
##		<dnastringsetl< th=""><th></th><th></th><th><charao< th=""><th></th><th></th><th><iranges></iranges></th></charao<></th></dnastringsetl<>			<charao< th=""><th></th><th></th><th><iranges></iranges></th></charao<>			<iranges></iranges>
##	22:50301584_C/T		T	100		PASS		777, 777]
##	rs114264124		Т	100		PASS	-	698, 698]
##	rs149209714		G	100		PASS		665, 665]
##	22:50303554_T/C		С	100		PASS		652, 652]
##	rs12167668		Т	100		PASS	A [645, 645]
##		PROTEINLOC	QUERYID		TXID	CDSID	GENEID	CONSEQUENCE
##		<integerlist></integerlist>	<integer></integer>	<chara< th=""><th>acter></th><th><integer></integer></th><th><character></character></th><th><factor></factor></th></chara<>	acter>	<integer></integer>	<character></character>	<factor></factor>
##	22:50301584_C/T	259	28		75253	218562	79087	synonymous
##	rs114264124	233	57		75253	218563	79087	nonsynonymous
##	rs149209714	222	58		75253	218563	79087	nonsynonymous
##	22:50303554_T/C	218	73		75253	218564	79087	nonsynonymous
##	rs12167668	215	74		75253	218564	79087	synonymous
##		REFCODON	VAI	RCODON		REFAA	VARAA	
##		<dnastringset></dnastringset>	<dnastri< th=""><th>ngSet></th><th><aasti< th=""><th>ringSet> <</th><th>AAStringSet></th><th></th></aasti<></th></dnastri<>	ngSet>	<aasti< th=""><th>ringSet> <</th><th>AAStringSet></th><th></th></aasti<>	ringSet> <	AAStringSet>	
##	22:50301584_C/T	CCG		CCA		Р	P	
##	rs114264124	CGG		CAG		R	Q	
##	rs149209714	GGA		GCA		G	A	
##	22:50303554_T/C	ATC		GTC		I	V	
##	rs12167668	CCG		CCA		Р	Р	
##								
##	seqlengths:							
##	chr22							
##	NA							

Using variant rs114264124 as an example, we see varAllele A has been substituted into the refCodon CGG to produce varCodon CAG. The refCodon is the sequence of codons necessary to make the variant allele substitution and therefore often includes more nucleotides than indicated in the range (i.e. the range is 50302962, 50302962, width of 1). Notice it is the second position in the refCodon that has been substituted. This position in the codon, the position of substitution, corresponds to genomic position 50302962. This genomic position maps to position 698 in coding region-based coordinates and to triplet 233 in the protein. This is a non-synonymous coding variant where the amino acid has changed from R (Arg) to Q (Gln).

When the resulting varCodon is not a multiple of 3 it cannot be translated. The consequence is considered a frameshift and varAA will be missing.

coding[coding\$CONSEQUENCE == "frameshift"]

```
## GRanges with 2 ranges and 17 metadata columns:
```

	0								
##		seqnames		ranges	strand	paramRa	angeID	REF	
##		<rle></rle>		<iranges></iranges>	<rle></rle>	<fa< th=""><th>actor> <dna< th=""><th>AStringSet></th><th></th></dna<></th></fa<>	actor> <dna< th=""><th>AStringSet></th><th></th></dna<>	AStringSet>	
##	22:50317001_G/GCACT	chr22 [503	17001,	50317001]	+		<na></na>	G	ł
##	22:50317001_G/GCACT	chr22 [503	17001,	50317001]	+		<na></na>	G	ł
##			ALT	QUAL	FIL	TER	varAllele	CDSLOC	
##		<dnastringset< th=""><th>List> <</th><th>numeric></th><th><charact< th=""><th>er> <dnas< th=""><th>StringSet></th><th><iranges></iranges></th><th></th></dnas<></th></charact<></th></dnastringset<>	List> <	numeric>	<charact< th=""><th>er> <dnas< th=""><th>StringSet></th><th><iranges></iranges></th><th></th></dnas<></th></charact<>	er> <dnas< th=""><th>StringSet></th><th><iranges></iranges></th><th></th></dnas<>	StringSet>	<iranges></iranges>	
##	22:50317001_G/GCACT		GCACT	233	Р	ASS	GCACT	[808, 808]	
##	22:50317001_G/GCACT		GCACT	233	Р	ASS	GCACT	[628, 628]	
##		PROTEINLOC	QUER	YID	TXID	CDSID	GENE	ID	
##		<integerlist></integerlist>	<integ< th=""><th>er> <char< th=""><th>acter> <</th><th>integer></th><th><character< th=""><th>r></th><th></th></character<></th></char<></th></integ<>	er> <char< th=""><th>acter> <</th><th>integer></th><th><character< th=""><th>r></th><th></th></character<></th></char<>	acter> <	integer>	<character< th=""><th>r></th><th></th></character<>	r>	
##	22:50317001_G/GCACT	270		359	74357	216303	791	74	
##	22:50317001_G/GCACT	210		359	74358	216303	791	74	

##		CONSEQUENCE	REFCODON	VARCODON	REFAA
##		<factor></factor>	<dnastringset></dnastringset>	<dnastringset></dnastringset>	<aastringset></aastringset>
##	22:50317001_G/GCACT	frameshift	GCC	GCC	А
##	22:50317001_G/GCACT	frameshift	GCC	GCC	A
##		VARA	AA		
##		<aastringset< td=""><td>t></td><td></td><td></td></aastringset<>	t>		
##	22:50317001_G/GCACT				
##	22:50317001_G/GCACT				
##					
##	seqlengths:				
##	chr22				
##	NA				

Annotation with ensemblVEP fixme: Material for ensemblVEP

3.2.5 Exercises

Exercise 9 Load the 'transcript.db' package relevant to the dm3 build of D. melanogaster. Use select and friends to select the Flybase gene ids of the top table tt and the Flybase transcript names (TXNAME) and Entrez gene identifiers (GENEID).

Use cdsBy to extract all coding sequences, grouped by transcript. Subset the coding sequences to contain just the transcripts relevant to the top table. How many transcripts are there? What is the structure of the first transcript's coding sequence?

Load the 'BSgenome' package for the dm3 build of D. melanogaster. Use the coding sequences ranges of the previous part of this exercise to extract the underlying DNA sequence, using the extractTranscriptsFromGenome function. Use Biostrings' translate to convert DNA to amino acid sequences.

Solution: The following loads the relevant Transcript.db package, and creates a more convenient alias to the *TranscriptDb* instance defined in the package.

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

We also need the data - flybase IDs from our differential expression analysis.

```
library(EMB02013)
data(topTable)
fbids <- rownames(topTable)</pre>
```

We can discover available keys (using keys) and columns (columns) in txdb, and then use select to retrieve the transcripts associated with each differentially expressed gene. The mapping between gene and transcript is not one-to-one – some genes have more than one transcript.

```
txnm <- select(txdb, fbids, "TXNAME", "GENEID")</pre>
```

Warning: 'select' resulted in 1:many mapping between keys and return rows

nrow(txnm)

[1] 207

head(txnm, 3)

 ##
 GENEID
 TXNAME

 ##
 1
 FBgn0039155
 FBtr0084549

 ##
 2
 FBgn0039827
 FBtr0085755

 ##
 3
 FBgn0039827
 FBtr0085756

The *TranscriptDb* instances can be queried for data that is more structured than simple data frames, and in particular return *GRanges* or *GRangesList* instances to represent genomic coordinates. These queries are performed using cdsBy (coding sequence), transcriptsBy (transcripts), etc., where a function argument by specifies how coding sequences or transcripts are grouped. Here we extract the coding sequences grouped by transcript, returning the transcript names, and subset the resulting *GRangesList* to contain just the transcripts of interest to us. The first transcript is composed of 6 distinct coding sequence regions.

```
cds <- cdsBy(txdb, "tx", use.names=TRUE)</pre>
txnm <- txnm[txnm$TXNAME %in% names(cds),]</pre>
cds <- cds[txnm$TXNAME]
length(cds)
## [1] 202
cds[1]
## GRangesList of length 1:
  $FBtr0084549
##
## GRanges with 6 ranges and 3 metadata columns:
##
         seqnames
                                 ranges strand |
                                                               cds_name exon_rank
                                                     cds_id
##
            <Rle>
                              <IRanges> <Rle> | <integer> <character> <integer>
            chr3R [19970946, 19971592]
##
     [1]
                                             + |
                                                      41058
                                                                   <NA>
                                                                                 2
     [2]
            chr3R [19971652, 19971770]
                                              + |
                                                                                 3
##
                                                      41059
                                                                   <NA>
                                                                                 4
##
     [3]
            chr3R [19971831, 19972024]
                                             + |
                                                      41060
                                                                   <NA>
##
     [4]
            chr3R [19972088, 19972461]
                                             + |
                                                      41061
                                                                    <NA>
                                                                                 5
##
     [5]
            chr3R [19972523, 19972589]
                                             + |
                                                      41062
                                                                    <NA>
                                                                                 6
                                                                                 7
            chr3R [19972918, 19973094]
##
     [6]
                                             + |
                                                      41063
                                                                    <NA>
##
##
  ___
## seqlengths:
##
        chr2L
                  chr2R
                             chr3L
                                       chr3R ... chr3RHet
                                                               chrXHet
                                                                         chrYHet chrUextra
    23011544 21146708 24543557 27905053 ... 2517507 204112
##
                                                                          347038 29004656
```

The following code loads the appropriate BSgenome package; the Dmelanogaster object refers to the whole genome sequence represented in this package. The remaining steps extract the DNA sequence of each transcript, and translates these to amino acid sequences. Issues of strand are handled correctly.

```
library(BSgenome.Dmelanogaster.UCSC.dm3)
txx <- extractTranscriptsFromGenome(Dmelanogaster, cds)</pre>
length(txx)
## [1] 202
head(txx, 3)
##
     A DNAStringSet instance of length 3
##
       width seq
                                                                           names
## [1]
      1578 ATGGGCAGCATGCAAGTGGCGCTGCTGG...CAAGCTGCAGATCAAGTGCAGCGACTAG FBtr0084549
##
  [2]
        2760 ATGCTGCGTTATCTGGCGCGTTTCGGAGG...GATTGTTGCTGCCCCCATTCGAACTTTAG FBtr0085755
## [3] 2217 ATGGCACTCAAGTTTCCCACAGTCAAGC...GATTGTTGCTGCCCCATTCGAACTTTAG FBtr0085756
head(translate(txx), 3)
     A AAStringSet instance of length 3
##
##
       width seq
## [1]
         526 MGSMQVALLALLVLGQLFPSAVANGSSSYSSTSTSASNQ...SSPNSVLDDSRNVFTFTTPKCENFRKRFPKLQIKCSD*
## [2]
         920 MLRYLALSEAGIAKLPRPQSRCYHSEKGVWGYKPIAQRE...GQQLHYCGRCEAPTPATGIGKVHKREVDEIVAAPFEL*
## [3] 739 MALKFPTVKRYGGEGAESMLAFFWQLLRDSVQANIEHVV...GQQLHYCGRCEAPTPATGIGKVHKREVDEIVAAPFEL*
```

Exercise 10 The objective of this exercise is to compare the quality of called SNPs that are located in dbSNP, versus those that are novel.

Locate the sample data in the file system. Explore the metadata (information about the content of the file) using scanVcfHeader. Discover the 'info' fields VT (variant type), and RSQ (genotype imputation quality).

Input the sample data using readVcf. You'll need to specify the genome build (genome="hg19") on which the variants are annotated. Take a peak at the rowData to see the genomic locations of each variant.

Data resource often adopt different naming conventions for sequences. For instance, dbSNP uses abbreviations such as ch22 to represent chromosome 22, whereas our VCF file uses 22. Use rowData and seqlevels<- to extract the row data of the variants, and rename the chromosomes.

Solution: Explore the header:

```
library(VariantAnnotation)
fl <- system.file("extdata", "chr22.vcf.gz", package="VariantAnnotation")
(hdr <- scanVcfHeader(fl))
## class: VCFHeader
## samples(5): HG00096 HG00097 HG00099 HG00100 HG00101
## meta(1): fileformat
## fixed(1): ALT
## info(22): LDAF AVGPOST ... VT SNPSOURCE
## geno(3): GT DS GL
info(hdr)[c("VT", "RSQ"),]
## DataFrame with 2 rows and 3 columns</pre>
```

##		Number	Туре	Description
##		<character></character>	<character></character>	<character></character>
## \	VΤ	1	String	indicates what type of variant the line represents
## F	RSQ	1	Float	Genotype imputation quality from MaCH/Thunder
	``			

Input the data and peak at their locations:

```
vcf <- readVcf(fl, "hg19")
head(rowData(vcf), 3)</pre>
```

##	GRanges with 3	3 ranges and	5 metada	ata column	s:			
##		seqnames		ranges	strand	pa	ramRangeID	REF
##		<rle></rle>		<iranges></iranges>	<rle></rle>		<factor></factor>	<dnastringset></dnastringset>
##	rs7410291	22 [50	300078,	50300078]	*		<na></na>	A
##	rs147922003	22 [50	300086,	50300086]	*		<na></na>	С
##	rs114143073	22 [50	300101,	50300101]	*		<na></na>	G
##			ALT	QUAL	FIL	TER		
##		<dnastringse< th=""><th>tList> ·</th><th><numeric></numeric></th><th><charact< th=""><th>er></th><th></th><th></th></charact<></th></dnastringse<>	tList> ·	<numeric></numeric>	<charact< th=""><th>er></th><th></th><th></th></charact<>	er>		
##	rs7410291		G	100	F	PASS		
##	rs147922003		Т	100	F	PASS		
##	rs114143073		А	100	F	PASS		
##								
##	seqlengths:							
##	22							
##	NA							

Rename chromosome levels:

```
seqlevels(vcf, force=TRUE) <- c("22"="ch22")</pre>
```

Exercise 11 Load the TxDb.Hsapiens.UCSC.hg19.knownGene annotation package, and read in the chr22.vcf.gz example file from the VariantAnnotation package.

Remembering to re-name sequence levels, use the locateVariants function to identify coding variants.

Summarize aspects of your data, e.g., did any coding variants match more than one gene? How many coding variants are there per gene ID?

Solution: Here we open the known genes data base, and read in the VCF file.

```
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
txdb <- TxDb.Hsapiens.UCSC.hg19.knownGene
fl <- system.file("extdata", "chr22.vcf.gz", package="VariantAnnotation")
vcf <- readVcf(fl, "hg19")
seqlevels(vcf, force=TRUE) <- c("22"="chr22")</pre>
```

The next lines locate coding variants.

```
rd <- rowData(vcf)
loc <- locateVariants(rd, txdb, CodingVariants())
head(loc, 3)</pre>
```

```
## GRanges with 3 ranges and 7 metadata columns:
##
                                ranges strand | LOCATION
                                                            QUERYID
                                                                         TXID
                                                                                  CDSID
        seqnames
##
            <Rle>
                             <IRanges> <Rle> | <factor> <integer> <integer> <integer>
     [1] chr22 [50301422, 50301422]
                                           * |
##
                                                  coding
                                                                 24
                                                                        75253
                                                                                 218562
##
     [2]
           chr22 [50301476, 50301476]
                                             * |
                                                  coding
                                                                 25
                                                                        75253
                                                                                 218562
##
     [3]
            chr22 [50301488, 50301488]
                                            * |
                                                   coding
                                                                 26
                                                                        75253
                                                                                 218562
              GENEID
                           PRECEDEID
                                            FOLLOWID
##
         <character> <CharacterList> <CharacterList>
##
     [1]
              79087
##
##
     [2]
              79087
##
     [3]
               79087
##
     ___
     seqlengths:
##
      chr22
##
##
        NA
```

To answer gene-centric questions data can be summarized by gene regardless of transcript.

```
## Did any coding variants match more than one gene?
splt <- split(loc$GENEID, loc$QUERYID)</pre>
table(sapply(splt, function(x) length(unique(x)) > 1))
##
## FALSE TRUE
##
    965
            15
## Summarize the number of coding variants by gene ID
splt <- split(loc$QUERYID, loc$GENEID)</pre>
head(sapply(splt, function(x) length(unique(x))), 3)
## 113730
            1890 23209
   22
##
          15 30
```

3.3 Visualization

The *Gviz* package produces very elegant data organized in a more-or-less familiar 'track' format. The following exercises walk through the *Gviz* User guide Section 2.

Load the *Gviz* package and sample *GRanges* containing genomic coordinates of CpG islands. Create a couple of variables with information on the chromosome and genome of the data.

```
library(Gviz)
data(cpgIslands)
chr <- "chr7"
genome <- "hg19"</pre>
```

The basic idea is to create a track, perhaps with additional attributes, and to plot it. There are different types of track, and we create these one at a time. We start with a simple annotation track

atrack <- AnnotationTrack(cpgIslands, name="CpG")</pre>

(This track could be plotted with plotTrack(atrack)). Then add a track that represents genomic coordinates. Tracks are combined when plotted, as a simple list. The vertical ordering of tracks is determined by their position in the list.

```
gtrack <- GenomeAxisTrack()</pre>
```

(Plot this with plotTracks(list(gtrack, atrack))). We can add an ideogram to provide overall orientation...

```
itrack <- IdeogramTrack(genome=genome, chromosome=chr)</pre>
```

and a more elaborate gene model, as an data.frame or GRanges object with specific columns of metadata.

Plot this as

plotTracks(tracks)

Zoom out to change the location box on the ideogram

```
plotTracks(tracks, from=2.5e7, to=2.8e7)
```

When zoomed in we can add sequence data

```
library(BSgenome.Hsapiens.UCSC.hg19)
strack <- SequenceTrack(Hsapiens, chromosome=chr)</pre>
```

plotTracks(c(tracks, strack), from=26450430, to=26450490, cex=.8)

As the *Gviz* vignette humbly says, 'so far we have replicated the features of a whole bunch of other genome browser tools out there'. We'd like to be able integrate our data into these plots, with a rich range of plotting options. The key is the DataTrack function, which we demonstrate with some simulated data; this final result is shown in Figure 3.1.

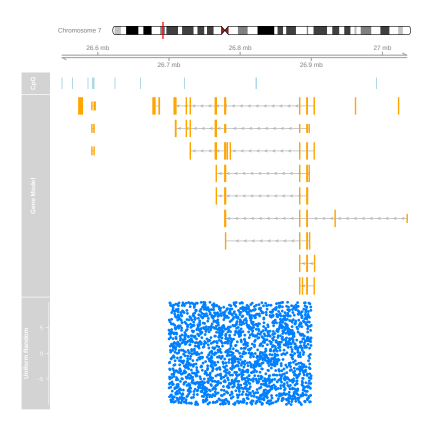


Figure 3.1: Gviz ideogram, genome coordinate, annotation, and data tracks.

```
plotTracks(c(tracks, dtrack))
```

Section 4.3 of the Gviz vignette illustrates flexibility of the data track.

Bibliography

- S. Anders, D. J. McCarthy, Y. Chen, M. Okoniewski, G. K. Smyth, W. Huber, and M. D. Robinson. Count-based differential expression analysis of RNA sequencing data using R and Bioconductor. *Nat Protoc*, 8(9):1765–1786, Sep 2013.
- [2] D. Bentley, S. Balasubramanian, H. Swerdlow, G. Smith, J. Milton, C. Brown, K. Hall, D. Evers, C. Barnes, H. Bignell, et al. Accurate whole human genome sequencing using reversible terminator chemistry. *Nature*, 456(7218):53–59, 2008.
- [3] A. N. Brooks, L. Yang, M. O. Duff, K. D. Hansen, J. W. Park, S. Dudoit, S. E. Brenner, and B. R. Graveley. Conservation of an RNA regulatory map between Drosophila and mammals. *Genome Research*, pages 193–202, 2011.
- [4] P. Burns. The R inferno. Technical report, 2011.
- [5] J. G. Caporaso, C. L. Lauber, W. A. Walters, D. Berg-Lyons, J. Huntley, N. Fierer, S. M. Owens, J. Betley, L. Fraser, M. Bauer, et al. Ultra-high-throughput microbial community analysis on the illumina hiseq and miseq platforms. *The ISME journal*, 6(8):1621–1624, 2012.
- [6] T. S. Furey. Chip-seq and beyond: new and improved methodologies to detect and characterize protein-dna interactions. *Nature Reviews Genetics*, 13(12):840–852, 2012.
- [7] D. B. Goldstein, A. Allen, J. Keebler, E. H. Margulies, S. Petrou, S. Petrovski, and S. Sunyaev. Sequencing studies in human genetics: design and interpretation. *Nat. Rev. Genet.*, 14(7):460–470, Jul 2013.
- [8] E. Khurana, Y. Fu, V. Colonna, X. J. Mu, H. M. Kang, T. Lappalainen, A. Sboner, L. Lochovsky, J. Chen, A. Harmanci, J. Das, A. Abyzov, S. Balasubramanian, K. Beal, D. Chakravarty, D. Challis, Y. Chen, D. Clarke, L. Clarke, F. Cunningham, U. S. Evani, P. Flicek, R. Fragoza, E. Garrison, R. Gibbs, Z. H. Gümüş, J. Herrero, N. Kitabayashi, Y. Kong, K. Lage, V. Liluashvili, S. M. Lipkin, D. G. MacArthur, G. Marth, D. Muzny, T. H. Pers, G. R. S. Ritchie, J. A. Rosenfeld, C. Sisu, X. Wei, M. Wilson, Y. Xue, F. Yu, . G. P. Consortium, E. T. Dermitzakis, H. Yu, M. A. Rubin, C. Tyler-Smith, and M. Gerstein. Integrative annotation of variants from 1092 humans: Application to cancer genomics. *Science*, 342(6154), 2013.
- [9] M. Lawrence, W. Huber, H. Pagès, P. Aboyoun, M. Carlson, R. Gentleman, M. T. Morgan, and V. J. Carey. Software for computing and annotating genomic ranges. *PLoS Comput Biol*, 9(8):e1003118, 08 2013.
- [10] B. A. Methe et al. A framework for human microbiome research. Nature, 486(7402):215–221, Jun 2012.
- [11] J. C. Mwenifumbo and M. A. Marra. Cancer genome-sequencing study design. Nat. Rev. Genet., 14(5):321–332, May 2013.
- [12] P. J. Park. ChIP-seq: advantages and challenges of a maturing technology. Nat. Rev. Genet., 10:669–680, Oct 2009. [PubMed Central:PMC3191340] [DOI:10.1038/nrg2641] [PubMed:19736561].
- [13] R Core Team. R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria, 2013. ISBN 3-900051-07-0.
- [14] C. Ross-Innes, R. Stark, A. Teschendorff, K. Holmes, H. Ali, M. Dunning, G. Brown, O. Gojis, I. Ellis, A. Green, et al. Differential oestrogen receptor binding is associated with clinical outcome in breast cancer. *Nature*, 481(7381):389– 393, 2012.

- [15] J. Rothberg and J. Leamon. The development and impact of 454 sequencing. Nature biotechnology, 26(10):1117– 1124, 2008.
- [16] A. Sottoriva, I. Spiteri, D. Shibata, C. Curtis, and S. Tavaré. Single-molecule genomic data delineate patient-specific tumor profiles and cancer stem cell organization. *Cancer research*, 73(1):41–49, 2013.