

R / Bioconductor for Everyone — Exercises

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24 July 2012

Abstract

This lab provides an introduction to *R / Bioconductor* for high-throughput sequence analysis. It is designed for those who have some but not a lot of familiarity with *R* and *Bioconductor*. The first part of the lab will focus on *R* data types, functions, classes, methods, the package and help systems, and the *Bioconductor* web site. The second part of the lab takes a quick tour of essential packages, classes, and methods for sequence analysis. We will make brief stops at [GenomicRanges](#), [Biostrings](#), [GenomicFeatures](#), [ShortRead](#), [Rsamtools](#), [rtracklayer](#), [AnnotationDbi](#), and other packages of interest to participants.

Exercise 1

This exercise uses data describing 128 microarray samples as a basis for exploring R functions. Covariates such as age, sex, type, stage of the disease, etc., are in a data file pData.csv.

The following command creates a variable pdataFiles that is the location of a comma-separated value ('csv') file to be used in the exercise. A csv file can be created using, e.g., 'Save as...' in spreadsheet software.

```
> pdataFile <- "/home/martin/RBiocForEveryone/pData.csv"
```

Input the csv file using read.table, assigning the input to a variable pdata. Use dim to find out the dimensions (number of rows, number of columns) in the object. Are there 128 rows? Use names or colnames to list the names of the columns of pdata. Use summary to summarize each column of the data. What are the data types of each column in the data frame?

A data frame is a list of equal length vectors. Select the 'sex' column of the data frame using [[or \$. Pause to explain to your neighbor why this subsetting works. Since a data frame is a list, use sapply to ask about the class of each column in the data frame. Explain to your neighbor what this produces, and why.

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Use `table` to summarize the number of males and females in the sample. Consult the help page `?table` to figure out additional arguments required to include NA values in the tabulation.

The `mol.biol` column summarizes molecular biological attributes of each sample. Use `table` to summarize the different molecular biology levels in the sample. Use `%in%` to create a logical vector of the samples that are either BCR/ABL or NEG. Subset the original phenotypic data to contain those samples that are BCR/ABL or NEG.

After subsetting, what are the levels of the `mol.biol` column? Set the levels to be BCR/ABL and NEG, i.e., the levels in the subset.

One would like covariates to be similar across groups of interest. Use `t.test` to assess whether BCR/ABL and NEG have individuals with similar age. To do this, use a formula that describes the response age in terms of the predictor `mol.biol`. If age is not independent of molecular biology, what complications might this introduce into subsequent analysis? Use

Solution: Here we input the data and explore basic properties.

```
> pdata <- read.table(pdataFile)
> dim(pdata)

[1] 128 21

> names(pdata)

[1] "cod"           "diagnosis"     "sex"           "age"
[5] "BT"           "remission"     "CR"           "date.cr"
[9] "t.4.11."      "t.9.22."       "cyto.normal"  "citog"
[13] "mol.biol"     "fusion.protein" "mdr"          "kinet"
[17] "ccr"          "relapse"       "transplant"   "f.u"
[21] "date.last.seen"
```

A data frame can be subset as if it were a matrix, or a list of column vectors.

```
> head(pdata$sex) # same as pdata[, "sex"], pdata[["sex"]]

[1] M M F M M M
Levels: F M

> sapply(pdata, class)

      cod      diagnosis      sex      age      BT
"factor" "factor"      "factor" "integer" "factor"
remission      CR      date.cr      t.4.11.      t.9.22.
"factor"      "factor"      "factor" "logical" "logical"
cyto.normal      citog      mol.biol fusion.protein      mdr
"logical"      "factor"      "factor" "factor" "factor"
kinet      ccr      relapse      transplant      f.u
"factor"      "logical"      "logical" "logical" "factor"
date.last.seen
"factor"
```

The number of males and females, including NA, is

```
> table(pdata$sex, useNA="ifany")  
  
  F    M <NA>  
42  83    3
```

An alternative version of this uses the with function: with(pdata, table(sex, useNA="ifany")).

The mol.biol column contains the following samples:

```
> with(pdata, table(mol.biol, useNA="ifany"))  
  
mol.biol  
ALL1/AF4 BCR/ABL E2A/PBX1      NEG  NUP-98  p15/p16  
        10      37      5      74      1      1
```

A logical vector indicating that the corresponding row is either BCR/ABL or NEG is constructed as

```
> ridx <- pdata$mol.biol %in% c("BCR/ABL", "NEG")
```

We can get a sense of the number of rows selected via table or sum (discuss with your neighbor what sum does, and why the answer is the same as the number of TRUE values in the result of the table function).

```
> table(ridx)
```

```
ridx  
FALSE  TRUE  
   17   111
```

```
> sum(ridx)
```

```
[1] 111
```

The original data frame can be subset to contain only BCR/ABL or NEG samples using the logical vector ridx that we created.

```
> pdata1 <- pdata[ridx,]
```

The levels of each factor reflect the levels in the original object, rather than the levels in the subset object, e.g.,

```
> levels(pdata$mol.biol)
```

```
[1] "ALL1/AF4" "BCR/ABL" "E2A/PBX1" "NEG"      "NUP-98"  "p15/p16"
```

These can be re-coded by updating the new data frame to contain a factor with the desired levels.

```
> pdata1$mol.biol <- factor(pdata1$mol.biol)  
> table(pdata1$mol.biol)
```

BCR/ABL	NEG
37	74

To ask whether age differs between molecular biology types, we use a formula `age ~ mol.biol` to describe the relationship ('age as a function of molecular biology') that we wish to test

```
> with(pdata1, t.test(age ~ mol.biol))
```

Welch Two Sample t-test

```
data: age by mol.biol
t = 4.8172, df = 68.529, p-value = 8.401e-06
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
 7.13507 17.22408
sample estimates:
mean in group BCR/ABL      mean in group NEG
          40.25000          28.07042
```

This summary can be visualize with, e.g., the `boxplot` function

```
> ## not evaluated
> boxplot(age ~ mol.biol, pdata1)
```

Molecular biology seems to be strongly associated with age; individuals in the NEG group are considerably younger than those in the BCR/ABL group. We might wish to include age as a covariate in any subsequent analysis seeking to relate molecular biology to gene expression.

Exercise 2

This purpose of this exercise is to explore BAM files and the data they contain. We'll extract the sequences and qualities of aligned reads, and draw simple plots to illustrate their properties.

- Attach the [pasillaBamSubset](#) experiment data package (containing a subset of reads from [1], as described in the package help page). Attache the [Rsamtools](#) (for accessing BAM files) and [ShortRead](#) (for manipulating short reads) packages.
- Use `ScanBamParam` to create an object that restricts the information extracted from the BAM file. Restrict the information using `scanBamFlag` to select reads that align to the plus strand (`isMinusStrand=FALSE`). Further, extract only the sequence and quality information (using `what=c('seq', 'qual')`).
- Input the sequences and qualities using `scanBam`, providing the file path returned by `untreated1_chr4` (a 'convenience' function from the [pasillaBamSubset](#) package) and the `ScanBamParam` object created in the previous question.

- d. Use `alphabetByCycle` to summarize nucleotide use by cycle; plot the relationship between cycle and use. Where you expecting this? What is going on?
- e. Coerce the quality scores to a matrix, and summarize the distribution of qualities across cycle. Is this expected?

Solution: Here we attach the data and other packages.

```
> library(pasillaBamSubset)
> library(Rsamtools) # scanBam
> library(ShortRead) # alphabetByCycle
```

Now establish the `ScanBamParam` object to restrict information extracted from the BAM file.

```
> flag <- scanBamFlag(isMinusStrand=FALSE)
> param <- ScanBamParam(what=c("seq", "qual"), flag=flag)
```

Input the sequences and qualities using `scanBam`.

```
> fl <- untreated1_chr4()
> res <- scanBam(fl, param=param)[[1]]
```

Summarize nucleotide use by cycle and plot the result (Figure 1, left).

```
> abc <- alphabetByCycle(res$seq)
> matplot(t(abc[1:4,]), type="l", lty=1, lwd=2,
+         xlab="Cycle", ylab="Count")
> legend("topright", legend=rownames(abc)[1:4],
+         lty=1, lwd=2, col=1:4)
```

Convert encoding quality scores to a numeric matrix, and summarize these a boxplots, one for each cycle (column) of the matrix (Figure 1, right).

```
> qual <- as(res$qual, "matrix")
> boxplot(qual ~ col(qual), outline=FALSE,
+         xlab="Cycle", ylab="Quality")
```

Exercise 3

The purpose of this exercise is to count the number of reads overlapping genes. This is a typical first step in an RNA-seq differential expression analysis.

- a. Annotations (gene models) can come from different sources. We'll use a pre-packaged source here; a later exercise retrieves similar information from a GFF file. Attach the [TxDb.Dmelandogaster.UCSC.dm3.ensGene](#) annotation package, and use `exonsBy` to retrieve all exons, grouped by gene. Subset the result to contain just those annotations relevant to chromosome 4.

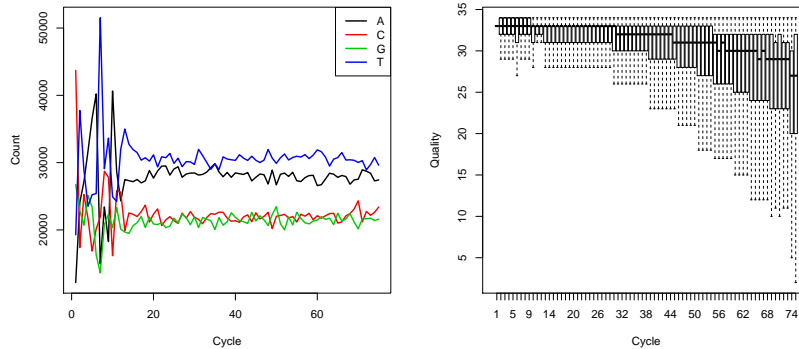


Figure 1: Nucleotide use (left) and base quality (right) in aligned reads.

- b. Create a character vector of file paths, using the helper functions `untreated1_chr4` and `untreated3_chr4`. Name the vector with an abbreviated name derived from the file name, and use this to invoke `BamFileList`, creating a list of paths to files that are known to be BAM files.
- c. Use `summarizeOverlaps` to count the number of reads in each region of interest. The RNA-seq protocol was not strand aware, so choose `ignore.strand=TRUE`. The object returned by `summarizeOverlaps` is an instance of class `SummarizedExperiment`. Explore it, and the counts that have been generated.
- d. Plot the counts in the two samples, perhaps transforming the counts with `asinh` (a log-like transform that deals with 0's better).
- e. As an additional exercise, use the `rtracklayer` package to retrieve and parse a GFF file containing similar annotations. How would this source of annotation tie into the work flow we have just performed?

Solution: Load the annotation package, extract exons grouped by gene, and select just those genes on chromosome 4.

```
> library(TxDb.Dmelanogaster.UCSC.dm3.ensGene) # genome coordinates
> exByGn <- exonsBy(TxDb.Dmelanogaster.UCSC.dm3.ensGene, "gene")
> chr4 <- exByGn[ all(seqnames(exByGn) == "chr4") ]
```

Create a `BamFileList` pointing to the BAM files we want to count.

```
> fls <- c(untreated1_chr4(), untreated3_chr4())
> names(fls) <- sub("_chr4.bam", "", basename(fls))
> bfl <- BamFileList(fls)
```

Count the number of reads overlapping each gene using `summarizeOverlaps`, remembering to the ignore strand to which the read aligns (why?).

```
> counts <- summarizeOverlaps(chr4, bfl, ignore.strand=TRUE)
> head(assay(counts))
```

	untreated1	untreated3
FBgn0002521	669	409
FBgn0004607	13	12
FBgn0004624	1168	816
FBgn0004859	388	199
FBgn0005558	8	22
FBgn0005561	7	11

Plot the results (Figure 2). Why are the points generally below the diagonal? Why do they have a funnel shape, with lots of variability between samples at low counts but very predictable numbers at high counts?

```
> plot(asinh(assay(counts)), asp=1, main="asinh(counts), chr4")
> abline(0, 1, lty=2)
```

The following illustrates an alternative source of annotation information. The annotations are in a GTF file on the ENSEMBL web site. We download them to our local disk, then use *rtracklayer*'s `import` to parse the file to a *GRanges* instance. We subset this to contain relevant information, then split the *GRanges* into a *GRangesList*, based on the `gene_id` column extracted from the GFF file. This could then be used in `summarizeOverlaps`, as above and without further change.

```
> library(rtracklayer) # import gff
> fl <- paste0("ftp://ftp.ensembl.org/pub/release-62/",
+            "gtf/drosophila_melanogaster/",
+            "Drosophila_melanogaster.BDGP5.25.62.gtf.gz")
> gffFile <- file.path(tempdir(), basename(fl))
> download.file(fl, gffFile)
> gff0 <- import(gffFile, asRangedData=FALSE)
> idx <- values(gff0)$source == "protein_coding" &
+       values(gff0)$type == "exon" &
+       seqnames(gff0) == "4"
> gff1 <- gff0[idx]
> chr4 <- split(gff1, values(gff1)$gene_id)
```

References

- [1] 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.

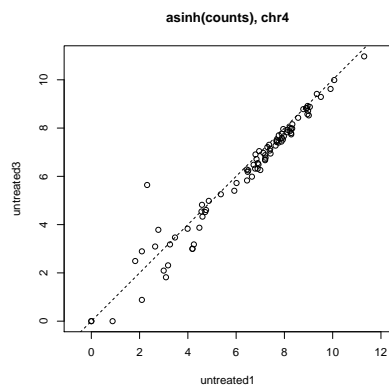


Figure 2: Count of reads overlapping genes on chr4.