

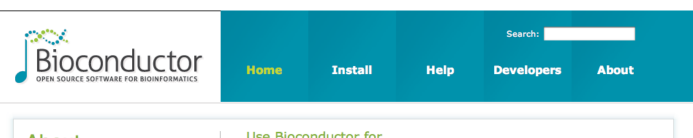
8. Bioconductor Intro and Annotation

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Auckland, November 2013

What is Bioconductor?



About Bioconductor

Bioconductor provides tools for the analysis and comprehension of highthroughput genomic data. Bioconductor uses the R statistical programming language, and is open source and open development. It has two releases each year, more than 460 packages, and an active user community.

Use Bioconductor for...

Microarrays

Import Affymetrix, Illumina, Nimblegen, Agilent, and other platforms. Perform quality assessment, normalization, differential expression, clustering, classification, gene set enrichment, genetical genomics and other workflows for expression, exon, copy number, SNP, methylation and other assays. Access GEO, ArrayExpress, Biomart, UCSC, and other community resources.

High Throughput Assays

Import, transform, edit, analyze and visualize flow cytometric, mass spec, HTqPCR, cell-based, and other assays.

Sequence Data

Import fasta, fastq, ELAND, MAQ, BWA, Bowtie, BAM, gff, bed, wig, and other sequence formats. Trim, transform, align, and manipulate sequences. Perform quality assessment, ChIP-seq, differential expression, RNA-seq, and other workflows. Access the Sequence Read Archive.

Annotation

Use microarray probe, gene, pathway, gene ontology, homology and other annotations. Access GO, KEGG, NCBI, Biomart, UCSC, vendor, and other



Mailing Lists

about an hour ago

about 2 hours ago

about 7 hours ago

Re: views on Rle using GRanges object

How to output Normalised count data f...

Re: EBS volumes with the Bioconductor...



Events

16 - 18 August 2011 — University of Warwick, Coventry, UK

Statistical Analyses for Next Generation

26 - 27 September 2011 — Birmingham, AL, USA

See all events »



News

BioC 2011 conference material

BioC 2011 conference material is now available.

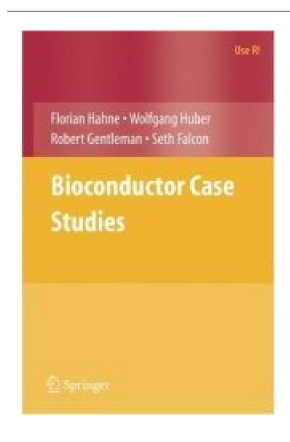
Bioconductor 2.8 released

Following the usual 6-month cycle, the Bioconductor community released Bioconductor 2.8 on April 14th, 2011. This release comprises 466 software packages and more than 500 up-todate annotation packages. It has been expressly designed to work with R 2.13.

What is Bioconductor?

- www.bioconductor.org
- Software project for analysis of genomic data and related tools, resources/datasets
- Open source and Open development
- Free

You could use commercial software; but experts typically write R code first. Also, the help manuals are not a sales pitch and encourage appropriate use.



- Begun in 2001, based at Harvard and now FHCRC (Seattle)
- A large collection of R packages (they also convert good software to R)
- Far too much for our little course!

We'll give examples of what Bioconductor can do, and how to learn more. Hahne et al (above) is a helpful reference text

Getting started...

Home » Install

• Install Packages • Find Packages • Update Packages • Install R

Install Bioconductor Packages

Use the biocLite.R script to install Bioconductor packages. To install a particular package, e.g., limma, type the following in an R command window:

```
source("http://bioconductor.org/biocLite.R")
biocLite("limma")
```

After downloading and installing this package, the script prints "Installation complete" and "TRUE". Install several packages, e.g., "GenomicFeatures" and "AnnotationDbi", with

```
biocLite(c("GenomicFeatures", "AnnotationDbi"))
```

To install a selection of core Bioconductor packages, use

```
biocLite()
```

Packages and their dependencies installed by this usage are: affy, affydata, affyPLM, affyQCReport, annaffy, annotate, Biobase, biomart, Biostrings, DynDoc, gcrma, genefilter, geneplotter, GenomicRanges, hgu95av2.db, limma, marray, multtest, vsn, and xtable. After downloading and installing these packages, the script prints "Installation complete" and "TRUE".

The biocLite.R script has arguments that change its default behavior:

```
pkgs
Character vector of Bioconductor packages to install.
destdir
File system directory for downloaded packages.
lib
R library where packages are installed.
```

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Bioconductor Release »

Packages in the stable, semi-annual release:

- BiocViews package discovery
- Software
- Metadata (Annotation, CDF and Probe)
- Experiment Data

Bioconductor is also available as an Amazon Machine Image (AMI).

Workflows »

Common Bioconductor workflows include:

- Oligonucleotide Arrays
- High-throughput Sequencing
- Annotation
- Flow Cytometry and other assays

Previous Versions »

For use with Bioconductor (R):

- 2.7 (2.12) 2.6 (2.11) 2.5 (2.10) • 2.4 (2.9) • 2.3 (2.8) • 2.2 (2.7) • 2.1 (2.6) • 2.0 (2.5) • 1.9 (2.4) • 1.8 (2.3)
- 1.7 (2.2) 1.6 (2.1)

```
> source("http://bioconductor.org/biocLite.R")
> biocLite()
installs the following general-purpose libraries;
Biobase, IRanges, AnnotationDbi
... then you use e.g. library("Biobase") as before. (NB older
versions used to download much more than this)
vignette(package="Biobase") tells you to look at vignette("esApply")
for a worked example – a very helpful introduction. (Or use e.g.
openVignette(), which is in the Biobase package itself)
```

To get other packages, use the source() command as before, then use e.g.

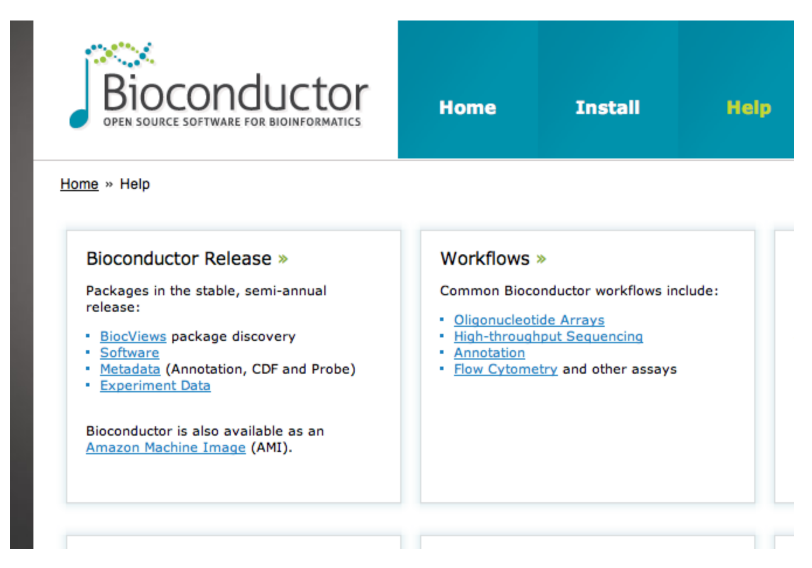
```
biocLite("SNPchip")
biocLite(c("limma", "siggenes"))
```

You do not need to type biocLite() again (even in a new R session). This would install the general-purpose packages again — which is harmless, but a waste of time.

Note; if, due to access privileges, you need to write to non-default directories, follow the onscreen commands and then start again. On Windows, 'Run as Administrator' may cut out this step.

What to install?

Back to the front page - click 'Help'



What to install?

- **Software** probably what you want
- Metadata e.g. annotation data, probe sequence data for microarrays of different types
- **Experiment data** e.g. datasets from hapmap.org, some expression datasets

Simple QC graphics

The "splots" package plots values from 96 or 384-well plates, for QC purposes

First, install it

biocLite("splots")

Then load into R

library("splots")

There is a single function: plotScreen() for displaying the results

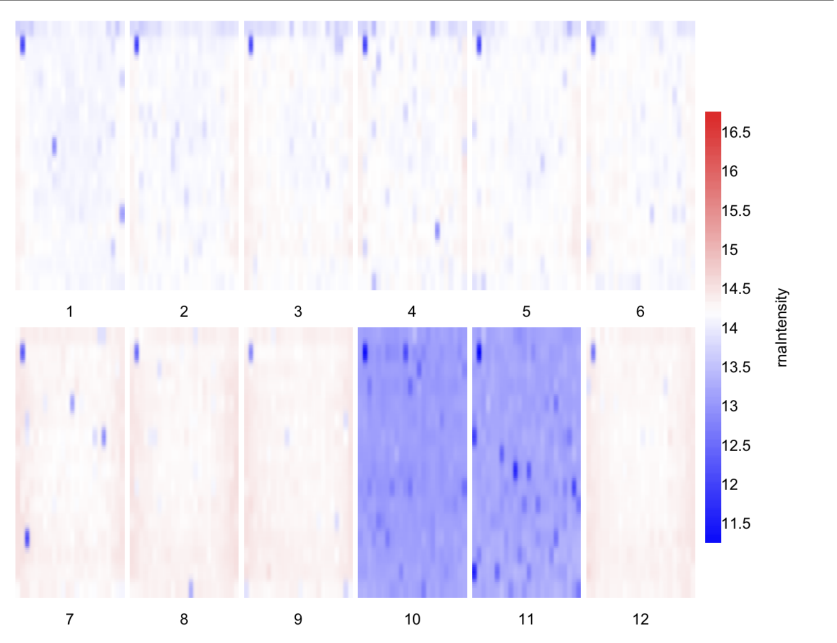
Example

The file "drosophila.rda" contains 12 of 114 plates from a RNAi gene-knockout study in fruit flies. Each spot represents a gene, and the intensity is low if knockout of that gene is lethal (data from the "RNAither" package)

```
load("drosophila.rda")
plotScreen(rnai)
```

The positive controls in the same position each plate are clear, and there are obvious plate effects that you might need to correct by normalization.

Example



Outline

One goal of Bioconductor is to provide efficient access inside R to the genome databases that are vital to interpreting associations.

We will look at a few of these

- annotate
- biomaRt.
- genomeGraphs

The reason to have an R interface to these databases is to be able to analyze annotation data for many SNPs or RNA transcripts.

Online or stored data

Annotation data can be downloaded in a single file or retrieved for each query from an online database.

Local storage is faster, but may require too much space (eg Ensembl) or become obsolete too quickly.

Local storage is ideal for fixed annotation data such as gene names for a microarray or SNP chip.

Types of database

Translations of names: Affy probe 32972_at is the gene NADPH oxidase 1 with symbol NOX1 and Ensembl gene id ENSG00000007952

Location: NOX1 is on Xq22.1, from 99984969 to 100015990, coded on the negative strand. There are 120 known polymorphisms (SNPs or indels) in this range.

Homology: The mouse version of NOX1 is also on the X chromosome, starting at 130621066 (and called Nox1)

Structure and function: NOX1 is a membrane protein (location), involved in voltage-gated ion channel activity (molecular function), and involved in signal transduction (biological process).

Annotaate

Bioconductor distributes annotation packages for a wide range of gene expression microarrays. The annotate package is one way to use this annotation information.

```
> library("annotate")
> library("hgu95av2.db")
> library("GO.db")
```

loads the annotate package and the databases for the Gene Ontology and one of the Affymetrix human microarray chips.

Lookups

The databases are queried with get() or mget() for multiple queries

```
> mget(c("738_at", "40840_at", "32972_at"), envir=hgu95av2GENENAME)
$'738 at'
[1] "5'-nucleotidase, cytosolic II"
$'40840_at'
[1] "peptidylprolyl isomerase F (cyclophilin F)"
$'32972_at'
[1] "NADPH oxidase 1"
> go<-get("738_at", envir=hgu95av2G0)</pre>
> names(go)
[1] "GD:0009117" "GD:0005829" "GD:0005737" "GD:0000166" "GD:0000287"
[6] "GD:0008253" "GD:0008253" "GD:0016787"
```

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Lookups

```
> get("GO:0009117", envir=GOTERM)
GOID: GO:0009117
Term: nucleotide metabolic process
Ontology: BP
Definition: The chemical reactions and pathways involving a
   nucleotide, a nucleoside that is esterified with (ortho)phosphate
   or an oligophosphate at any hydroxyl group on the glycose
```

moiety; may be mono-, di- or triphosphate; this definition

includes cyclic nucleotides (nucleoside cyclic phosphates).

Synonym: nucleotide metabolism

What lookups are there?

```
> library(help="hgu95av2.db")
hgu95av2ALIAS2PROBE Map between Common Gene Symbol Identifiers and
                    Manufacturer Identifiers
> get("NOX1", envir=hgu95av2ALIAS2PR0BE)
[1] "32972_at" "32973_s_at"
You can also reverse a lookup table with revmap()
> get("NOX1", envir=revmap(hgu95av2SYMBOL))
[1] "32972_at" "32973_s_at"
> get("X",revmap(hgu95av2CHR))
 [1] "1016_s_at" "107_at" "1100_at" "112_g_at" "1155_at"
 .... and lots more
```

BioMart (www.biomart.org) is a query-oriented data management system developed jointly by the European Bioinformatics Institute (EBI) and Cold Spring Harbor Laboratory (CSHL).

biomaRt is an R interface to BioMart systems, in particular to Ensembl (www.ensembl.org). Ensembl is a joint project between EMBL - European Bioinformatics Institute (EBI) and the Wellcome Trust Sanger Institute (WTSI) to develop a software system which produces and maintains automatic annotation on selected eukaryotic genomes.

We begin by choosing which BioMart to use

```
> library(biomaRt)
Loading required package: RCurl
> listMarts()
                               biomart
                          ensembl
                                                          ENSEMBL GENES 63 (SANGER U
1
2
                                                              VARIATION 63 (SANGER U
                                                    ENSEMBL
                              snp
3
                                                    ENSEMBL REGULATION 63 (SANGER U
             functional_genomics
4
                                                                           (SANGER U
                                                                  VEGA 43
                             vega
5
                                                          ENSEMBL BACTERIA 10 (EBI U
                bacteria_mart_10
6
                   fungi_mart_10
                                                             ENSEMBL FUNGI 10 (EBI U
7
             fungi_variations_10
                                                  ENSEMBL FUNGI VARIATION 10 (EBI U
8
                 metazoa mart 10
                                                           ENSEMBL METAZOA 10 (EBI U
9
                                                ENSEMBL METAZOA VARIATION 10 (EBI U
           metazoa_variations_10
60
              ENSEMBL_MART_PLANT
                                           GRAMENE 30 ENSEMBL GENES (CSHL/CORNELL U
61
          ENSEMBL_MART_PLANT_SNP
                                               GRAMENE 30 VARIATION (CSHL/CORNELL U
62
               GRAMENE MARKER 30
                                                 GRAMENE 30 MARKERS (CSHL/CORNELL U
                                                GRAMENE 30 MAPPINGS (CSHL/CORNELL U
63
                  GRAMENE MAP 30
64
                        QTL_MART
                                                  GRAMENE 32 QTL DB (CSHL/CORNELL U
                                             UNIGENE SALMO SALAR DATABASE (CMM CHIL
65
                salmosalar2 mart
66
                                 UNIGENE ONCORHYNCHUS MYKISS DATABASE (CMM CHIL
                     trucha_mart
> ens <- useMart("ensembl")</pre>
```

We then choose a database to use

```
> listDatasets(ens)
                           dataset
                                                                      description
                                                                    description
                           dataset
                                        Ornithorhynchus anatinus genes (OANA5)
           oanatinus_gene_ensembl
1
                                       Taeniopygia guttata genes (taeGut3.2.4)
            tguttata_gene_ensembl
3
          cporcellus_gene_ensembl
                                                Cavia porcellus genes (cavPor3)
          gaculeatus_gene_ensembl
                                        Gasterosteus aculeatus genes (BROADS1)
4
                                            Loxodonta africana genes (loxAfr3)
           lafricana_gene_ensembl
5
30
                                             Pteropus vampyrus genes (pteVam1)
           pvampyrus_gene_ensembl
58
                                                      Bos taurus genes (UMD3.1)
             btaurus_gene_ensembl
59
            meugenii_gene_ensembl
                                             Macropus eugenii genes (Meug_1.0)
           sharrisii_gene_ensembl
                                         Sarcophilus harrisii genes (DEVIL7.0)
60
         cfamiliaris_gene_ensembl
                                            Canis familiaris genes (CanFam3.1)
61
> hsap <- useDataset("hsapiens_gene_ensembl",mart=ens)</pre>
```

The getGene function queries the database for gene information. It accepts many forms of gene identifier, eg Entrez, HUGO, Affy transcript

```
> getGene(id=1440, type="entrezgene", mart=hsap)
  entrezgene hgnc_symbol
        1440
                    CSF3
                                                               description
1 colony stimulating factor 3 (granulocyte) [Source: HGNC Symbol; Acc: 2438]
  chromosome_name band strand start_position end_position ensembl_gene_id
                                     38171614
                                                  38174066 ENSG00000108342
1
               17 q21.1
                             1
> getGene(id=c("AGT","AGTR1"), type="hgnc_symbol", mart=hsap)
 hgnc_symbol hgnc_symbol
          AGT
                      AGT
1
        AGTR1
                    AGTR1
1 angiotensinogen (serpin peptidase inhibitor, clade A, member 8) [Source: HGNC Sym
2
                                  angiotensin II receptor, type 1 [Source: HGNC Sym
  chromosome_name band strand start_position end_position ensembl_gene_id
                1 q42.2
                                    230838269 230850043 ENSG00000135744
                            -1
2
                    q24
                             1
                                   148415571 148460795 ENSG00000144891
```

getBM is more general than getGene. It specifies a list of **filters** for selecting genes or SNPs and **attributes** to return from the database.

```
> affyids <- c("202763_at", "209310_s_at", "207500_at")</pre>
> getBM(attributes = c("affy_hg_u133_plus_2", "hgnc_symbol", "chromosome_name",
   "start_position", "end_position", "band"), filters = "affy_hg_u133_plus_2",
    values = affyids, mart = hsap)
    affy_hg_u133
                  hgnc chromosome_name start_position end_position band
                          CASP3
                                        4
                                              185785844
                                                           185807623 q35.1
           202763_at
1
                                   11
                                                           104384957 q22.3
           207500 at
                          CASP5
                                              104370180
         209310_s_at
                         CASP4 11
                                              104318804
                                                           104344535 q22.3
```

listAttributes(hsap) and listFilters(hsap) list the avilable attributes and filters (hundreds)

```
> getBM(mart=hsap, attributes=c("band", "hgnc_symbol"),
        filters=c("band_start","band_end","chromosome_name"),
        values=list("p21.33","p21.33",6))
     band hgnc_symbol
   p21.33
   p21.33
             SNORD117
3 p21.33
              SNORA38
4 p21.33
              SNORD48
5
   p21.33
              SNORD52
6
  p21.33
               MIR877
   p21.33
              MIR1236
  p21.33
8
               GTF2H4
  p21.33
9
                VARS2
10 p21.33
                SFTA2
11 p21.33
                DPCR1
12 p21.33
                MUC21
               HSPA1A
121 p21.33
122 p21.33
                 TNXB
123 p21.33
                STK19
124 p21.33
                  C4A
125 p21.33
                  C4B
```

Homology

getLDS() combines two data marts, for example to homologous genes in other species. We can look up the mouse equivalents of a particular Affy transcript, or of the NOX1 gene.

```
> human = useMart("ensembl", dataset = "hsapiens_gene_ensembl")
> mouse = useMart("ensembl", dataset = "mmusculus_gene_ensembl")
> getLDS(attributes = c("hgnc_symbol", "chromosome_name", "start_position"),
+ filters = "hgnc_symbol", values = "NOX1", mart = human,
+ attributesL = c("chromosome_name", "start_position", "external_gene_id"),
+ martL = mouse)
    V1 V2    V3 V4    V5    V6
1 NOX1    X 100098313    X 134086421 Nox1
```

The mouse gene name is the same as the human one apart from capitalisation.

Homology

The getSequence function looks up DNA or protein sequences by chromosome position or gene identifiers

```
> agt<-getSequence(id="AGT",type="hgnc_symbol", seqType="peptide",mart=hsap)
> agt
```

1 MRKRAPQSEMAPAGVSLRATILCLLAWAGLAAGDRVYIHPFHLVIHNESTCEQLAKANAGKPKDPTFIPAPIQAKTS PVDEKALQDQLVLVAAKLDTEDKLRAAMVGMLANFLGFRIYGMHSELWGVVHGATVLSPTAVFGTLASLYLGALDHTAD RLQAILGVPWKDKNCTSRLDAHKVLSALQAVQGLLVAQGRADSQAQLLLSTVVGVFTAPGLHLKQPFVQGLALYTPVVL PRSLDFTELDVAAEKIDRFMQAVTGWKTGCSLMGASVDSTLAFNTYVHFQGKMKGFSLLAEPQEFWVDNSTSVSVPMLS GMGTFQHWSDIQDNFSVTQVPFTESACLLLIQPHYASDLDKVEGLTFQQNSLNWMKKLSPRTIHLTMPQLVLQGSYDLQ DLLAQAELPAILHTELNLQKLSNDRIRVGEVLNSIFFELEADEREPTESTQQLNKPEVLEVTLNRPFLFAVYDQSATAL HFLGRVANPLSTA*

Example: finding chromosomes

We had a set 1524 SNPs, of which 409 did not have their chromosome listed.

I needed to know which SNPs were on the X chromosome, to estimate sex from DNA intensity and heterozygous X-chromosome loci, for QC.

> head(unknown)

- [1] "UGT1A3-001449-0_B_R_1538822" "LIPC-002761-0_B_R_1538453"
- [3] "CETP-001265-0_B_R_1538254" "F8-165293-0_T_F_1538626"
- [5] "CPB2-051208-0_B_F_1539402" "VDRDIL-1355-0_T_F_1539404"

A hand-search would be easy but tedious, so we want an automated approach

Example: finding chromosomes

First extract the gene names

```
genes <- sapply(unknown, function(snp) strsplit(snp,"-")[[1]][1])
ugenes <- unique(genes)</pre>
```

Now call out to Ensembl

```
getBM(attributes="chromosome_name", filters="hgnc_symbol",values=ugenes,
    mart=hsap)
```

works for all except VRDIL, which isn't recognized.

Finding SNPs

Human SNPs (and short indels) are in a separate database from gene information. We can look up known SNPs and other polymorphisms for the NOX1 gene

```
> snpmart = useMart("snp", dataset = "hsapiens_snp")
Checking attributes ... ok
Checking filters ... ok
> getBM(c("refsnp_id", "allele", "chrom_start", "chrom_strand"),
      filters = c("chr_name", "chrom_start", "chrom_end"),
      values = list("X",99984969,100015990), mart = snpmart)
                              allele chrom_start chrom_strand
        refsnp_id
     rs7054049
                            T/A
                                   99985184
1
                            G/T 99985304
2
    rs60975472
                            A/G 99985571
    rs58902780
                            G/A 99985618
4
   rs182188185
5
   rs186748080
                            A/G 99985798
```

More metadata

The citation() function prints out information about how to cite a package

> citation("biomaRt")

To cite the biomaRt package in publications use:

Mapping identifiers for the integration of genomic datasets with the R/Bioconductor package biomaRt. Steffen Durinck, Paul T. Spellman, Ewan Birney and Wolfgang Huber, Nature Protocols 4, 1184-1191 (2009).

BioMart and Bioconductor: a powerful link between biological databases and microarray data analysis. Steffen Durinck, Yves Moreau, Arek Kasprzyk, Sean Davis, Bart De Moor, Alvis Brazma and Wolfgang Huber, Bioinformatics 21, 3439-3440 (2005).

Citations are one way academic software authors can prove to funders and promotion committees that software is worthwhile.

GenomeGraphs

This package makes pretty pictures from the annotation data.

For example, a pictures showing the standard and alternative splices for the NOX1 gene and the location of the gene on the X chromosome

GenomeGraphs

