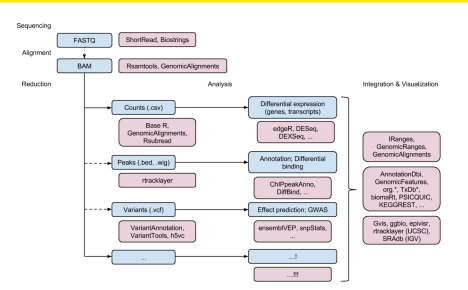
### **Bioconductor**

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Spring 2018

# High-throughput sequence workflow



### **Bioconductor**

Analysis and comprehension of high-throughput genomic data

- Statistical analysis designed for large genomic data
- Interpretation: biological context, visualization, reproducibility
- Support for all high-throughput technologies
  - Sequencing: RNASeq, ChIPSeq, variants, copy number, . . .
  - Microarrays: expression, SNP, ...
  - Flow cytometry, proteomics, images, ...

Bioconductor cheat sheet https://github.com/mikelove/bioc-refcard

# **Bioconductor packages**

- https://www.bioconductor.org/
- Over 1,400 packages
- Discover and navigate via biocView
- Informative package 'landing page'
  - Title, author / maintainer, short description, citation, installation instructions, ..., download statistics
- 'Release' (every six months) and 'devel' branches

 $http://bioconductor.org/packages/release/BiocViews.html\#\_\_\_Software$ 

## Reference manuals, vignettes

- All user-visible functions have help pages, most with runnable examples
- 'Vignettes' an important feature in Bioconductor narrative documents illustrating how to use the package, with integrated code
  - Example: AnnotationHub landing page, "AnnotationHub HOW TO's" vignette illustrating some fun use cases.

http://bioconductor.org/packages/devel/AnnotationHub

# **Objects**

- Bioconductor makes extensive use of classes to represent complicated data types
- Classes foster interoperability many different packages can work on the same data – but can be a bit intimidating
- Formal 'S4' object system
  - Often a class is described on a particular home page, e.g., ?GRanges, and in vignettes, e.g., vignette(package="GenomicRanges"), vignette("GenomicRangesIntroduction")
  - Many methods and classes can be discovered interactively, e.g., methods(class="GRanges") to find out what one can do with a GRanges instance, and methods(findOverlaps) for classes that the findOverlaps() function operates on.
  - In more advanced cases, one can look at the actual definition of a class or method using getClass(), getMethod()
- Interactive help
  - ?findOverlaps, <tab> to select help on a specific method, ?GRanges-class for help on a class.

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## High-throughput sequence data

**FASTA** 

**Biostrings** 

**FASTQ** 

ShortRead

BAM

GenomicAlignments

**VCF** 

VariantAnnotation

BED, GTF, WIG

rtracklayer

# DNA/amino acid sequences: FASTA files

 The Biostrings package, is used to represent DNA and other sequences, with many convenient sequence-related functions, e.g., ?consensusMatrix.

Input & manipulation, FASTA file example:

```
\label{eq:control_gamma} $\operatorname{gttggtggcccaccagtgccaaaatacacaagaagaagaaacagcatctt}$ gacactaaaatgcaaaaattgctttgcgtcaatgactcaaaacgaaaatg ...$ atgggtatcaagttgccccgtataaaaggcaagtttaccggttgcacggt $>\operatorname{NM}_001201794\_up\_2000\_chr2L\_8382455\_f chr2L:8382455-8384454 ttatttatgtaggcgcccgttcccgcagccaaagcactcagaattccggg cgtgtagcgcaacgaccatctacaaggcaatattttgatcgcttgttagg
```

>NM\_078863\_up\_2000\_chr2L\_16764737\_f chr2L:16764737-16766736

## Reads: FASTQ files

 The ShortRead package can be used for lower-level access to FASTQ files. readFastq(), FastqStreamer(), FastqSampler()

Input & manipulation, FASTQ file example:

http://bioconductor.org/packages/ShortRead

 $Quality\ scores:\ 'phred-like',\ encoded.\ See\ http://en.wikipedia.org/wiki/FASTQ\_format\#Encoding$ 

@ERR127302.1703 HWI-EAS350 0441:1:1:1460:19184#0/1

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## Biostrings, DNA or amino acid sequences

#### Classes

 XString, XStringSet, e.g., DNAString (genomes), DNAStringSet (reads)

#### Methods

- Manipulation, e.g., reverseComplement()
- Summary, e.g., letterFrequency()
- Matching, e.g., matchPDict(), matchPWM()

Related packages: BSgenome for working with whole genome sequences, e.g., ?"getSeq,BSgenome-method"

 $http://bioconductor.org/packages/release/bioc/vignettes/Biostrings/inst/doc/BiostringsQuickOverview.pdf \\ http://bioconductor.org/packages/BSgenome$ 

## Aligned reads: SAM/BAM files

#### Input & manipulation:

- 'low-level' Rsamtools scanBam(), BamFile()
- 'high-level' GenomicAlignments readGAlignments()

#### SAM Header example

```
@HD      VN:1.0      SO:coordinate
@SQ      SN:chr1      LN:249250621
@SQ      SN:chr10      LN:135534747
@SQ      SN:chr11      LN:135006516
...
@SQ      SN:chrY      LN:59373566
@PG      ID:TopHat      VN:2.0.8b
```

http://bioconductor.org/packages/Rsamtools

http://bioconductor.org/packages/GenomicAlignments

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CL:/home/hpages/tophat

## GenomicAlignments, Aligned reads

The GenomicAlignments package is used to input reads aligned to a reference genome. See for instance the ?readGAlignments help page and vignette(package="GenomicAlignments", "summarizeOverlaps")

Classes - GenomicRanges-like behaivor

GAlignments, GAlignmentPairs, GAlignmentsList

#### Methods

- readGAlignments(), readGAlignmentsList()
  - Easy to restrict input, iterate in chunks
- summarizeOverlaps()

### Called variants: VCF files

#### Input and manipulation:

 VariantAnnotation - readVcf(), readInfo(), readGeno() selectively with ScanVcfParam().

http://bioconductor.org/packages/VariantAnnotation

```
##fileformat=VCFv4.2
##fileDate=20090805
##source=myImputationProgramV3.1
##reference=file:///seq/references/1000GenomesPilot-NCBI36.fas
##contig=<ID=20,length=62435964,assembly=B36,md5=f126cdf8a6e06
##phasing=partial
##INFO=<ID=DP, Number=1, Type=Integer, Description="Total Depth"
##INFO=<ID=AF, Number=A, Type=Float, Description="Allele Frequence
##FILTER=<ID=q10, Description="Quality below 10">
##FILTER=<ID=s50, Description="Less than 50% of samples have days
##FORMAT=<ID=GT, Number=1, Type=String, Description="Genotype">
```

##FORMAT=<ID=GQ, Number=1, Type=Integer, Description="Genotype Qu

### **VCF** Location info

#CHROM	POS	ID	REF	ALT	QUAL	FILTER	
20	14370	rs6054257	G	A	29	PASS	
20	17330	•	T	Α	3	q10	
20	1110696	rs6040355	Α	G,T	67	PASS	
20	1230237	•	T	•	47	PASS	
20	1234567	microsat1	GTC	G,GTCT	50	PASS	

### **VCF Variant INFO**

```
#CHROM POS
                    INFO
20
       14370
                    NS=3;DP=14;AF=0.5;DB;H2
               . . .
20
       17330
               . . .
                    NS=3; DP=11; AF=0.017
20
       1110696 ...
                    NS=2;DP=10;AF=0.333,0.667;AA=T;DB
20
       1230237 ...
                    NS=3;DP=13;AA=T
20
       1234567 ... NS=3; DP=9; AA=G
```

## **Genotype FORMAT and samples**

```
... POS
                                                           NA
                FORMAT
                            NA00001
                                           NA00002
                GT:GQ:DP:HQ 0|0:48:1:51,51 1|0:48:8:51,51
                                                           1/:
... 14370
... 17330
                GT:GQ:DP:HQ 0|0:49:3:58,50 0|1:3:5:65,3
                                                           0/0
... 1110696 ...
                GT:GQ:DP:HQ 1|2:21:6:23,27 2|1:2:0:18,2
                                                           2/2
                                                           0/0
... 1230237 ...
                GT:GQ:DP:HQ 0|0:54:7:56,60 0|0:48:4:51,51
                GT:GQ:DP 0/1:35:4 0/2:17:2
                                                           1/:
... 1234567 ...
```

### VariantAnnotation, Called variants

#### Classes - GenomicRanges-like behavior

- VCF 'wide'
- VRanges 'tall'

#### Methods

- I/O and filtering: readVcf(), readGeno(), readInfo(), readGT(), writeVcf(), filterVcf()
- Annotation: locateVariants() (variants overlapping ranges), predictCoding(), summarizeVariants()
- SNPs: genotypeToSnpMatrix(), snpSummary()

# **VCF-Related packages**

- ensemblVEP- query the Ensembl Variant Effect Predictor
- VariantTools Explore, diagnose, and compare variant calls.
- VariantFiltering Filtering of coding and non-coding genetic variants.
- h5vc has variant calling functionality.
- snpStats Classes and statistical methods for large SNP association studies.

```
http://bioconductor.org/packages/ensembIVEP
```

http://bioconductor.org/packages/VariantTools

http://bioconductor.org/packages/VariantFiltering

http://bioconductor.org/packages/h5vc

https://bioconductor.org/packages/release/bioc/html/snpStats.html

Obenchain, V, Lawrence, M, Carey, V, Gogarten, S, Shannon, P, and Morgan, M. VariantAnnotation: a Bioconductor package for exploration and annotation of genetic variants. Bioinformatics, first published online March 28, 2014 doi:10.1093/bioinformatics/btu168, http://bioinformatics.oxfordjournals.org/content/early/2014/04/21/bioinformatics.btu168

 $Introduction\ to\ Variant Annotation,\ http://bioconductor.org/packages/release/bioc/vignettes/ShortRead/inst/doc/Overview.pdf$ 

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## Genome annotations: BED, WIG, GTF, etc. files

• The rtracklayer's import and export functions can read in many common file types, e.g., BED, WIG, GTF, ..., in addition to querying and navigating the UCSC genome browser. Check out the ?import page for basic usage.

Input: rtracklayer::import()

- BED: range-based annotation (see http://genome.ucsc.edu/FAQ/FAQformat.html for definition of this and related formats)
- WIG/bigWig: dense, continuous-valued data
- GTF: gene model

http://bioconductor.org/packages/rtracklayer

## **GTF** Component coordinates

7	protein_coding	gene	27221129	27224842	-
• • •					
7	protein_coding	transcript	27221134	27224835	-
7	protein_coding	exon	27224055	27224835	-
7	protein_coding	CDS	27224055	27224763	-
7	protein_coding	start_codon	27224761	27224763	-
7	protein_coding	exon	27221134	27222647	-
7	protein_coding	CDS	27222418	27222647	-
7	protein_coding	stop_codon	27222415	27222417	-
7	protein_coding	UTR	27224764	27224835	-
7	protein_coding	UTR	27221134	27222414	-

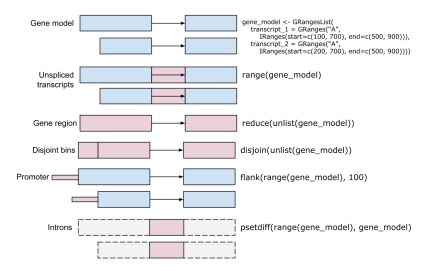
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```
gene_id "ENSG00000005073"; gene_name "HOXA11"; gene_source "en
...
... transcript_id "ENST00000006015"; transcript_name "HOXA11-(
... exon_number "1"; exon_id "ENSE00001147062";
... exon_number "1"; protein_id "ENSP00000006015";
... exon_number "1";
... exon_number "2"; exon_id "ENSE00002099557";
... exon_number "2"; protein_id "ENSP00000006015";
... exon_number "2"; protein_id "ENSP00000006015";
... exon_number "2";
```

Read GTF file into R, https://davetang.org/muse/2017/08/04/read-gtf-file-r/

Data representation in R / Bioconductor

## Ranges overview



## Ranges in Bioconductor

- IRanges
  - start() / end() / width()
  - List-like length(), subset, etc.
  - 'metadata', mcols()
- GRanges
  - 'seqnames' (chromosome), 'strand'
  - Seqinfo, including seqlevels and seqlengths

## Range methods

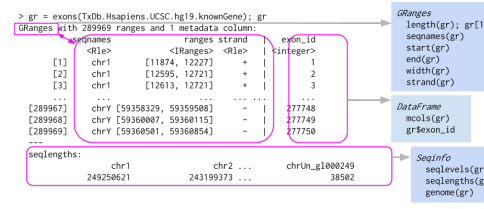
- Intra-range methods
  - Independent of other ranges in the same object
  - shift(), narrow(), flank(), promoters(), resize(), restrict(), trim()
  - See ?"intra-range-methods"
- Inter-range methods
  - Depends on other ranges in the same object
  - range(), reduce(), gaps(), disjoin(), coverage()
  - See ?"inter-range-methods"
- Between-range methods
  - Functions of two (or more) range objects
  - findOverlaps(), countOverlaps(), summarizeOverlaps(), ...,
    %over%, %within%, %outside%; union(), intersect(), setdiff(),
    punion(), pintersect(), psetdiff()

## **IRanges**

- The IRanges package defines an important class for specifying integer ranges
- There are many interesting operations to be performed on ranges, e.g, flank() identifies adjacent ranges
- IRanges extends the Ranges class

# **Genomic Ranges**

The GenomicRanges package extends the notion of ranges to include features relevant to application of ranges in sequence analysis, particularly the ability to associate a range with a sequence name (e.g., chromosome) and a strand.



## **GenomicRanges**

- Data (e.g., aligned reads, called peaks, copy number)
- Annotations (e.g., genes, exons, transcripts)
- Close relation to BED files (see rtracklayer::import.bed() and HelloRanges)
- Also vector interface length(), [, etc.

## **Lists of Genomic Ranges**

- List definition all elements of the same type
- E.g., lists of exons-within-transcripts, alignments-within-reads
- Many \*List-aware methods, but a common 'trick': apply a vectorized function to the unlisted representation, then re-list

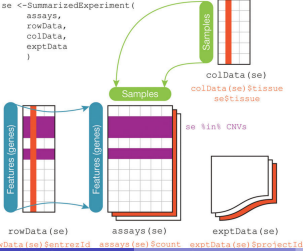
Lawrence M, Huber W, Pagès H, Aboyoun P, Carlson M, et al. (2013) Software for Computing and Annotating Genomic Ranges. PLoS Comput Biol 9(8): e1003118. doi:10.1371/journal.pcbi.1003118

# Lists of Genomic Ranges

```
> grl = exonsBy(TxDb.Hsapiens.UCSC.hg19.knownGene, "tx", use.names=TRUE); grl
GRangesList of length 82960:
                                                                                GRangesList
$uc001aaa.3
                                                                                  (list of GRanges)
GRanges with 3 ranges and 3 metadata columns:
                                                                                  length(grl)
      seanames
                      ranges strand |
                                      exon id exon name exon rank
                                                                                  gr1[1:3]
                   <IRanges> <Rle> | <integer> <character> <integer>
        <R1e>
                                                                                  shift(grl. 1)
        chr1 [11874, 12227]
                                                      <NA>
                                                                                  range(grl)
      chr1 [12613, 12721] + |
                                                      <NA>
  Г37
         chr1 [13221, 14409]
                                  + |
                                                      <N4>
Suc010nxa.1
                                                                               GRanges
GRanges with 3 ranges and 3 metadata columns:
                                                                                  gr1[[2]]
      segnames
                      ranges strand | exon id exon name exon rank
                                                                                  grl[["uc010nxq.1"]]
         chr1 [11874, 12227]
  Γ17
                                                  <NA>
 [2]
        chr1 [12595, 12721]
                                                  <NA>
  [3]
         chr1 [13403, 14409] + |
                                                  <NA>
                                                               3
$uc010nxr 1
GRanges with 3 ranges and 3 metadata columns:
                                                                         Two kinds of fun!
                      ranges strand | exon_id exon_name exon_rank
                                                                          introns =
      segnames
  Γ17
         chr1 [11874, 12227]
                                                  <NA>
                                                                            psetdiff(range(grl), grl)
  Γ27
       chr1 [12646, 12697]
                                                  <NA>
         chr1 [13221, 14409] + I
 [3]
                                                 <NA>
                                                                          grr = unlist(grl)
                                                                          ## transform grr. then...
                                                                          grl = relist(grr, grl)
<82957 more elements>
seginfo: 93 sequences (1 circular) from hg19 genome
                                                                                  'flesh'
                                                                                             'skeleton'
```

## **Summarized Experiments**

SummarizedExperiment - Rows are indexed by a dataframe of features. Accessible with rowData()



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## RangedSummarizedExperiment

 RangedSummarizedExperiment - Rows are indexed by genomic ranges. Accessble with rowRanges()

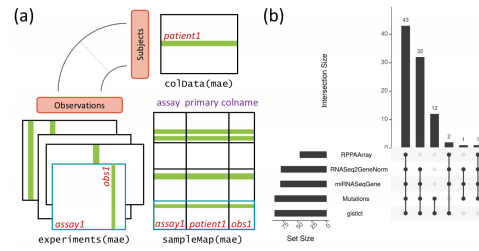
## SingleCellExperiment

- SingleCellExperiment an extension of RangedSummarizedExperiment with several internal slots
  - Has a slot for spike-in measures
  - Can store reduced dimensionality representation of the data

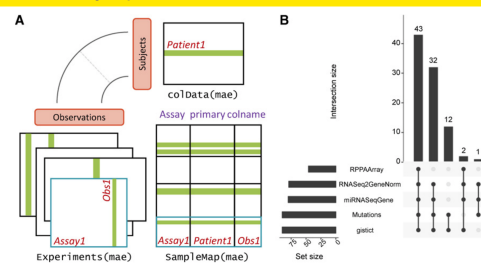
https://bioconductor.org/packages/3.7/bioc/vignettes/SingleCellExperiment/inst/doc/intro.html

## MultiAssayExperiment

Bioconductor package for management of multi-assay data. Especially useful for integrating TCGA datasets.



## MultiAssayExperiment



Ramos, Marcel, Lucas Schiffer, Angela Re, Rimsha Azhar, Azfar Basunia, Carmen Rodriguez Cabrera, Tiffany Chan, et al. "Software For The Integration Of Multi-Omics Experiments In Bioconductor," June 1, 2017. doi:10.1101/144774. http://biorxiv.org/content/early/2017/06/01/144774

## **Annotation packages**

- *Bioconductor* provides extensive access to 'annotation' resources, see the "AnnotationData" biocViews hierarchy.
- AnnotationDBI is a cornerstone of "AnnotationData" packages, provides user interface and database connection code for annotation data packages using SQLite data storage.

 $https://bioconductor.org/packages/release/BiocViews.html \#\_\_AnnotationData \\ http://bioconductor.org/packages/AnnotationDbi$ 

## **Annotation packages**

- org packages (e.g., org.Hs.eg.db) contain maps between different gene identifiers, e.g., ENTREZ and SYMBOL. The basic interface to these packages is described on the help page ?select
- TxDb packages (e.g., TxDb.Hsapiens.UCSC.hg38.knownGene) contain gene models (exon coordinates, exon / transcript relationships, etc) derived from common sources such as the hg38 knownGene track of the UCSC genome browser. These packages can be queried, e.g., as described on the ?exonsBy page to retrieve all exons grouped by gene or transcript.

https://bioconductor.org/packages/org.Hs.eg.db

https://bioconductor.org/packages/TxDb.Hsapiens.UCSC.hg38.knownGene

## **Annotation packages**

- EnsDb packages and databases (e.g. EnsDb.Hsapiens.v86) provide, similar to TxDb packages, gene models, but also protein annotations (protein sequences and protein domains within these) and additional annotation columns such as "gene\_biotype" or "tx\_biotype" defining the biotype of the features (e.g. lincRNA, protein\_coding, miRNA etc). EnsDb databases are designed for Ensembl annotations and contain annotations for all genes (protein coding and non-coding) for a specific Ensembl release.
- **BSgenome** packages (e.g., BSgenome.Hsapiens.UCSC.hg19) contain whole genomes of model organisms. See available.genomes() for pre-packaged genomes.

annotation work flow, http://bioconductor.org/help/workflows/annotation/annotation/

https://bioconductor.org/packages/release/data/annotation/html/EnsDb. Hsapiens. v86. html

https://bioconductor.org/packages/release/data/annotation/html/BSgenome. Hsapiens. UCSC.hg19.html/psgenome. UCSC.hg19.html/psgenome. Hsapiens. Hsapiens. Hsapiens. UCSC.hg19.html/psgenome. UCSC.hg19.html/psgenome. Hsapiens. UCSC.hg19.html/psgenome. Hsapiens. UCSC.hg19.html/psgenome. UCSC.hg19.html/psgenome. UCSC.hg19.html/psgenome. UCSC.hg19.html/psgenome. UCSC.hg19.html/psgenome. UCSC.hg19.html/psgenome. UCSC.hg19.html/psgenome. UCSC.hg19.html/psgenome. UCSC.hg19.html/psgenome. UCSC.hg19.html/psgeno

- Annotation packages usually 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().

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 keytyp
	select()	Retrieve annotations matching keys, keytype and

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Category	Function	Description
Manipulate	<pre>setdiff(), union(), intersect() duplicated(), unique() %in%, match() any(), all() merge()</pre>	Operations on sets Mark or remove duplicate Find matches Are any TRUE? Are all? Combine two different d

Category Function	Description
GRanges* transcripts(),         exons(), cds()         transcriptsBy(),         exonsBy()         cdsBy()	Features (transcripts, exons, coding sequence) as GRanges. Features group by gene, transcript, etc., as GRangesList.

#### **Biomart**

- Biomart R package, biomaRt, workflow:
  - Discover and select a mart and dataset,
  - Select filters, which IDs to convert from
  - Select attributes, which IDs to convert to
  - Run the query
- Biomart has a web interface, operating on the same principles

https://bioconductor.org/packages/biomaRt

http://bioconductor.org/packages/release/bioc/vignettes/biomaRt/inst/doc/biomaRt.html#selecting-a-biomart-database-and-dataset

http://bioconductor.org/packages/release/bioc/vignettes/biomaRt/inst/doc/biomaRt.html#annotate-a-set-of-entrezgene-identifiers-with-go-annotation

http://www.ensembl.org/biomart

### **KEGG**

- KEGG: Kyoto Encyclopedia of Genes and Genomes
- KEGG API R package, KEGGREST
  - Essential operations outlined in the vignette

http://www.genome.jp/kegg/pathway.html

https://bioconductor.org/packages/KEGGREST

http://bioconductor.org/packages/release/bioc/vignettes/KEGGREST/inst/doc/KEGGREST-vignette.html

### **PSICQUIC**

- PSICQUIC (the Proteomics Standard Initiative Common QUery InterfaCe) - standardized access to molecular interaction databases.
- Protein-protein interaction databases like "BioGrid", "Intact", "Reactome", "STRING", "BIND"
- Supports the molecular interaction query language (MIQL)

https://bioconductor.org/packages/release/bioc/html/PSICQUIC.html

### **AnnotationHub**

- AnnotationHub package curated database of large-scale whole-genome resources, e.g., regulatory elements from the Roadmap Epigenomics project, Ensembl GTF and FASTA files for model and other organisms, and the NHLBI grasp2db data base of GWAS results. Examples of use include:
  - Easily access and import Roadmap Epigenomics files.
  - liftOver genomic range-based annotations from one coordinate system (e.g, hg19) to another (e.g., GRCh38).
  - Create TranscriptDb and BSgenome-style annotation resources 'on the fly' for a diverse set of organisms.
  - Programmatically access the genomic coordinates of clinically relevant variants cataloged in dbSNP.
- Related packages: ExperimentHub curated data sets

https://bioconductor.org/packages/AnnotationHub

AnnotationHub HOW-TOs.

http://bioconductor.org/packages/devel/bioc/vignettes/AnnotationHub/inst/doc/AnnotationHub-HOWTO.html

https://bioconductor.org/packages/ExperimentHub

# **Domain-specific packages**

 Important packages for analysis of differential expression include edgeR and DESeq2; both have excellent vignettes for exploration.

```
\label{linear_http://bioconductor.org/packages/edgeR} $$ $$ $$ http://bioconductor.org/packages/DESeq2 $$
```

 Popular ChIP-seq packages include DiffBind and csaw for comparison of peaks across samples, ChIPQC for quality assessment, and ChIPpeakAnno and ChIPseeker for annotating results (e.g., discovering nearby genes).

```
http://bioconductor.org/packages/DiffBind
http://bioconductor.org/packages/csaw
http://bioconductor.org/packages/ChIPQC
http://bioconductor.org/packages/ChIPpeakAnno
http://bioconductor.org/packages/ChIPseeker
```

# **Domain-specific packages**

- Working with called variants (VCF files) is facilitated by packages such as VariantAnnotation, VariantFiltering and ensemblVEP.
- Packages for calling variants include, e.g., h5vc and VariantTools.

https://bioconductor.org/packages/release/bioc/html/VariantAnnotation.html

http://bioconductor.org/packages/VariantFiltering.html

https://bioconductor.org/packages/release/bioc/html/ensemblVEP.html

https://bioconductor.org/packages/release/bioc/html/h5vc.html

https://bioconductor.org/packages/release/bioc/html/VariantTools.html

# **Domain-specific packages**

- **Single-cell 'omics'** are increasingly important. From the biocView page, enter 'single cell' in the 'search table' field.
- Several packages identify copy number variants from sequence data, including cn.mops. The CNTools package provides some useful facilities for comparison of segments across samples.
- Microbiome and metagenomic analysis is facilitated by packages such as phyloseq and metagenomeSeq.
- Metabolomics, chemoinformatics, image analysis, and many other high-throughput analysis domains are also represented in *Bioconductor*, explore these via biocViews and title searches.

```
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### **Visiualization**

A number of *Bioconductor* packages help with visualization and reporting, in addition to functions provided by individual packages.

- Gviz provides a track-like visualization of genomic regions.
- ComplexHeatmap does an amazing job of all sorts of heatmaps, including OncoPrint-style summaries.
- ReportingTools provides a flexible way to generate static and dynamic HTML-based reports.

http://bioconductor.org/packages/Gviz

http://bioconductor.org/packages/ComplexHeatmap

http://bioconductor.org/packages/ReportingTools

# Working with 'big data'

- Much Bioinformatic data is very large and often cannot fit into memory
- Several general strategies for working with large data

#### Restriction to specific genomic regions

 e.g., ScanBamParam() limits input to desired data at specific genomic ranges

#### Iteration over pieces of genomic data

• e.g., yieldSize argument of BamFile(), or FastqStreamer() allows iteration through large files.

# Working with 'big data'

#### Compression

- Genomic vectors represented as Rle (run-length encoding) class
- Lists e.g., GRangesList are efficiently maintain the illusion that vector elements are grouped.

#### Parallel processing

e.g., via BiocParallel package

https://bioconductor.org/packages/release/bioc/html/BiocParallel.html

Lawrence, M and Morgan, M. Scalable Genomic Computing and Visualization with R and Bioconductor. Statistical Science 29 (2) (2014), 214-226.

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## **Code optimization**

- aprof Amdahl's Profiler, Directed Optimization Made Easy.
- profvis Visualize R profiling data.
- microbenchmark Accurate Timing Functions.

http://cran.r-project.org/web/packages/aprof/index.html

https://rstudio.github.io/profvis. Examples: https://rpubs.com/wch/123888

http://cran.r-project.org/web/packages/microbenchmark/index.html

### **Summary**

- *Bioconductor* is a large collection of R packages for the analysis and comprehension of high-throughput genomic data.
- Bioconductor relies on formal classes to represent genomic data, so it is important to develop a rudimentary comfort with classes, including seeking help for classes and methods.
- *Bioconductor* uses vignettes to augment traditional help pages; these can be very valuable in illustrating overall package use.