Annotating genes, genomes, and variants

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What is 'Annotation'?

- Genes classification schemes (e.g., Entrez, Ensembl), pathway membership, . . .
- Genomes reference genomes; exons, transcripts, coding sequence; coding consequences
- System / network biology pathways, biochemical reactions,...
- 'Consortium' resources, TCGA, ENCODE, dbSNP, GTEx, ...

Other defintions (not covered here)

- ► SNP (and similar) consequences (*VariantAnnotation*, *VariantFiltering*, *ensemblVEP*)
- Assign function to novel sequences

Bioconductor Annotation Resources - Packages

Model organism annotation packages

- ▶ org.* gene names and pathways
- ► *TxDb.** gene models
- ► *BSgenome.** whole-genome sequences

org.* packages

The 'select' interface:

Discovery: keytypes, columns, keys

Retrieval: select, mapIds

```
library(org.Hs.eg.db)
keytypes(org.Hs.eg.db)
columns(org.Hs.eg.db)
egid <-
    select(org.Hs.eg.db, "BRCA1", "ENTREZID", "SYMBOL")</pre>
```

org.* (and other annotation) packages – Under the hood...

SQL (sqlite) data bases

- org.Hs.eg_dbconn() to query using RSQLite package
- ▶ org.Hs.eg_dbfile() to discover location and query outside R.

TxDb.* packages

- Gene models for common model organsisms / genome builds / known gene schemes
- Supports the 'select' interface (keytypes, columns, keys, select)
- 'Easy' to build custom packages when gene model exist

Retrieving genomic ranges

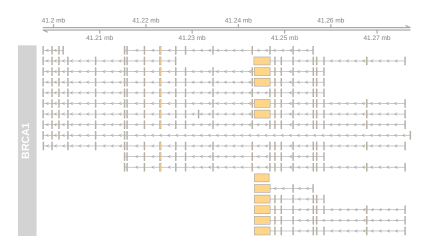
- transcripts, exons, cds,
- transcriptsBy , exonsBy, cdsBy group by gene, transcirpt, etc.

```
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
txdb <- TxDb.Hsapiens.UCSC.hg19.knownGene
cdsByTx <- cdsBy(txdb, "tx")</pre>
```

Example: Visualize BRCA1 Transcripts

```
library(org.Hs.eg.db)
eid <- mapIds(org.Hs.eg.db, "BRCA1", "ENTREZID",
  "SYMBOL")
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
txdb <- TxDb.Hsapiens.UCSC.hg19.knownGene
txid <- select(txdb, eid, "TXNAME", "GENEID")[["TXNAME"]]</pre>
cds <- cdsBy(txdb, by="tx", use.names=TRUE)</pre>
brca1cds <- cds[names(cds) %in% txid]
library(Gviz)
tx <- rep(names(brca1cds), elementLengths(brca1cds))</pre>
id <- unlist(brca1cds)$cds id
grt <- GeneRegionTrack(brca1cds, name="BRCA1", id=tx,</pre>
  gene="BRCA1", feature=tx, transcript=tx, exon=id)
plotTracks(list(GenomeAxisTrack(), grt))
```

Example: Visualize BRCA1 Transcripts



BSgenome.* Packages: Whole-Genome Sequences

- 'Masks' when available, e.g., repeat regions
- Load chromosomes, range-based queries: getSeq, extactTranscriptSeqs

```
library(BSgenome.Hsapiens.UCSC.hg19)
extractTranscriptSeqs(Hsapiens, brca1cds)
##
     A DNAStringSet instance of length 20
##
        width seq
                                           names
         2280 ATGGATTTATCTG...AGCCACTACTGA uc010whl.2
##
    [1]
##
   [2] 5379 ATGAGCCTACAAG...AGCCACTACTGA uc002icp.4
   [3] 522 ATGGATGCTGAGT...AGCCACTACTGA uc010whm.2
##
##
   [18]
##
         3954 ATGCTGAAACTTC...GATTCAAACTTA uc010cyz.2
         4017 ATGGATTTATCTG...GATTCAAACTTA uc010cza.2
##
   Г197
   [20]
         3207 ATGAATGTAGAAA...GATTCAAACTTA uc010wht.1
##
```

Web-based resources

AnnotationHub Ensembl, Encode, dbSNP, UCSC data objects, ...

biomaRt Ensembl and other annotations, url

PSICQUIC Protein interactions, url uniprot.ws Protein annotations, url KEGGREST KEGG pathways, url

SRAdb Sequencing experiments, url

rtracklayer genome tracks, url

GEOquery Array and other data, url
ArrayExpress Array and other data, url

Web-based resources

Demo

Summary

Genes

org.* packages, columns(), keys(), mapIds(), select().

Genomes

- ► TxDb.* packages. select(), exons(), exonsBy() & friends.
- ▶ BSgenome.* packages. FaFile, TwoBitFile files.

Variants

► VariantAnnotation, VariantFiltering, ensemblVEP.

Web-based resources

▶ biomaRt, AnnotationHub, and others.

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