Extending ensemb1db: MySQL backend and protein annotations

Johannes Rainer (EURAC research, Italy), Sebastian Gibb, Laurent Gatto (CPU Cambridge, UK)

December 7, 2016

Introduction

- ensembldb: retrieve gene & transcript annotations.
- ensembldb package defines the EnsDb class:
 - Wrapper to access annotations from an SQLite database.
 - Same functionality than the GenomicFeatures package (TxDb object).
 - Filter framework to enable specific and fast queries.
 - NEW: MySQL backend support.
 - NEW: provide protein annotations.

Basic usage

- Available methods to extract data: genes, transcripts, transcriptsBy, exons, exonsBy, cdsBy, fiveUTRsByTranscripts, threeUTRsByTranscripts, proteins (NEW).
- Example: get all lincRNA genes encoded on chromosome Y.

```
## Load an EnsDb package matching Ensembl version 86
library(EnsDb.Hsapiens.v86)
edb <- EnsDb.Hsapiens.v86

## Retrieve all lincRNAs encoded on chromosome Y.
## Create the filter objects
sf <- SeqnameFilter("Y")
gbf <- GenebiotypeFilter("lincRNA")</pre>
```

Basic usage

Example: (continued)

```
## Retrieve the data.
genes(edb, filter = list(sf, gbf))
GRanges object with 52 ranges and 6 metadata columns:
                      ranges strand |
 seanames
                                     gene id
    <Rle>
                   <IRanges> <Rle> | <character>
 ENSG00000278847
                      Y [2934406, 2934771] - | ENSG00000278847
                      Y [3002912, 3102272] + | ENSG00000231535
 FNSG00000231535
                      Y [4036497, 4100320] + | ENSG00000229308
 ENSG00000229308
                      Y [25378300, 25394719] - | ENSG00000228786
 FNSG00000228786
                    Y [25482908, 25486705] + | ENSG00000240450
 ENSG00000240450
 FNSG00000231141
                      Y [25728490, 25733388] + | ENSG00000231141
                  entrezid gene_biotype seq_coord_system
     gene_name
   <character> <character> <character> <character> 
 ENSG00000278847 RP11-414C23 1
                                              lincRNA
                                                          chromosome
                                             lincRNA
 ENSG00000231535 LINC00278 100873962
                                                          chromosome
 FNSG00000229308 AC010084 1
                                              lincRNA
                                                          chromosome
 ENSG00000228786 LTNC00266-4P
                                              lincRNA
                                                          chromosome
 FNSG00000240450
                   CSPG4P1Y
                                   114758
                                             lincRNA
                                                          chromosome
 FNSG00000231141
                       TTTY3 474148:114760
                                            lincRNA
                                                          chromosome
symbol
   <character>
 ENSG00000278847 RP11-414C23.1
 ENSG00000231535 LINC00278
 ENSG00000229308 AC010084 1
```

Available filters

- For genes: GeneidFilter, GenenameFilter, EntrezidFilter, GenebiotypeFilter, (SymbolFilter).
- For transcripts: TxidFilter, TxbiotypeFilter.
- For exons: ExonidFilter, ExonrankFilter.
- NEW: for proteins: ProteinidFilter, UniprotidFilter, UniprotdbFilter, UniprotmappingtypeFilter, ProtdomidFilter.
- Based on chromosomal coordinates: SeqnameFilter,
 SeqstrandFilter, SeqstartFilter, SeqendFilter,
 GRangesFilter: condition can be within or overlapping.
- Multiple filters are combined with a logical AND.
- Each filter supports 1:n values, =, != and also a *like* condition.

Building annotation databases

- Option A): from GTF/GFF files or AnnotationHub.
- Example: create an EnsDb using AnnotationHub.

- Option B) (preferred): using the Ensembl Perl API:
 - fetchTablesFromEnsembl and makeEnsemblSQLiteFromTables.
 - Fetches also protein annotations.
 - Requirements: Perl, Bioperl, Ensembl Perl API.

MySQL backend

 <u>Example</u>: listEnsDb list all available databases, useMySQL to insert an EnsDb to a MySQL server.

```
library(RMySQL)
dbc <- dbConnect(MySQL(), host = "localhost", user = "anonuser", pass = "")</pre>
## list all available EnsDb databases.
listEnsDbs(dbc)
Loading required package: DBI
                   dbname
                               organism ensembl_version
1 ensdb acarolinensis v83 acarolinensis
     ensdb byulgaris v86
                              bvulgaris
                                                      86
3 ensdb_dmelanogaster_v86 dmelanogaster
                                                      86
       ensdb hsapiens v75
                               hsapiens
                                                      75
## Connect to a database
dbc <- dbConnect(MySOL(), host = "localhost", user = "anonuser", pass = "",</pre>
                  dbname = "ensdb_dmelanogaster_v86")
edb 2 <- EnsDb(dbc)
## To insert an EnsDb to a MySQL: useMySQL
db_my <- useMySQL(edb, host = "localhost", user = "anonuser", pass = "")</pre>
```

Enables a central, MySQL-based annotation server.

Protein data: fetch protein data from an EnsDb

Example: add protein columns to the columns parameter.

```
## Get all genes with a C2H2 Zinc finger domain and
## return all of their Uniprot IDs
pfam <- ProtdomidFilter("PF13912")</pre>
genes(edb, filter = pfam, return.type = "DataFrame",
    columns = c("gene_name", "uniprot_id"))
DataFrame with 583 rows and 4 columns
gene name uniprot id
                            gene id protein domain id
     <character> <character>
                                <character>
                                                  <character>
     AC002310.11
                      B7Z5R0 ENSG00000261459
                                                      PF13912
      AC092835 2 A0A087WIV0 FNSG00000233757
                                                      PF13912
   CTD-2006C1.13 F5H0A9 ENSG00000267179
                                                      PF13912
581
          7SCAN9 015535 FNSG00000137185
                                                      PF13912
582
          ZSCAN9 A0A024RCK9 ENSG00000137185
                                                     PF13912
583
          7SCAN9
                      E9POL7 ENSG00000137185
                                                      PF13912
```

Protein data: fetch protein data from an EnsDb

Example: use proteins method to specifically fetch protein data.

```
## Return the protein annotation as a AAStringSet:
prts <- proteins(edb, filter = GenenameFilter("ZBTB16"),</pre>
                 columns = c("tx_id", "tx_biotype"),
                 return.type = "AAStringSet")
prts
 A AAStringSet instance of length 5
   width sea
                                                           names
     673 MDLTKMGMIQLONPSHPTGLLCK...GHKPEEIPPDWRIEKTYLYLCYV ENSP00000338157
[1]
Γ27
     115 MDLTKMGMIOLONPSHPTGLLCK...QAKAEDLDDLLYAAEILEIEYLE ENSP00000437716
Г37
     148 MDLTKMGMIOLONPSHPTGLLCK...OASDDNDTEATMADGGAEEEDR ENSP00000443013
[4]
     673 MDLTKMGMIOLONPSHPTGLLCK...GHKPEEIPPDWRIEKTYLYLCYV ENSP00000376721
Γ51
      55 XGGLLPQGFIQRELFSKLGELAV...GEQCSVCGVELPDNEAVEQHRVF ENSP00000445047
## Additional columns are available as mools:
mcols(prts)
DataFrame with 5 rows and 4 columns
                              tx_biotype protein_id gene_name
           tx id
     <character>
                             <character>
                                             <character> <character>
1 ENST00000335953
                          protein_coding ENSP00000338157
                                                              ZBTB16
2 FNST00000544220
                          protein coding ENSP00000437716
                                                              7BTB16
                          protein_coding ENSP00000443013
3 ENST00000535700
                                                              ZBTB16
4 FNST00000392996
                          protein coding ENSP00000376721
                                                              ZBTB16
5 ENST00000539918 nonsense mediated decay ENSP00000445047
                                                              ZBTB16
```

- Pbase: (Laurent Gatto and Sebastian Gibb): provides classes and functions for the analysis of protein sequence data in proteomics experiments.
- The Proteins object: container for proteins and peptide ranges within the AA sequences.
- <u>Example</u>: fetch a Proteins object for all ZBTB16 proteins including their protein domains.

```
## load Pbase - we need the "ensembldb" branch.
library(Pbase)

## Fetch proteins including protein domains for ZBTB16
prts <- Proteins(edb, filter = GenenameFilter("ZBTB16"))

## Amino acid sequence:
aa(prts)</pre>
```

Loading required package: Rcpp Loading required package: Gviz Loading required package: grid

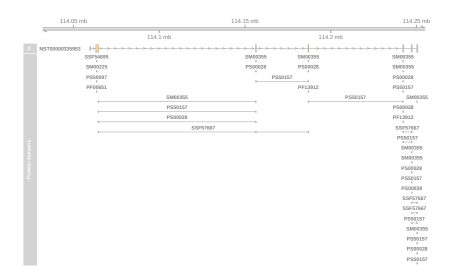
This data.table install has not detected OpenMP support. It will work but slower in single th

Example: fetch a Proteins object for all ZBTB16 proteins including their protein domains (continued).

```
## Peptide features:
pranges(prts)
IRangesList of length 5
$FNSP00000338157
IRanges object with 36 ranges and 3 metadata columns:
                          width I
                                        protein id protein domain source
      start
                  end
  <integer> <integer> <integer> |
                                     <character>
                                                              <character>
  PS50157
                602
                          629
                                      28 | ENSP00000338157
                                                                            pfscan
  PS50157
                490
                          517
                                      28 | ENSP00000338157
                                                                            pfscan
                          657
  PS50157
                630
                                      28 | ENSP00000338157
                                                                            pfscan
                                                                               . . .
  SM00355
                432
                          454
                                      23 | ENSP00000338157
                                                                             smart
  SM00355
                574
                          596
                                      23 | ENSP00000338157
                                                                             smart
  SM00225
                 34
                           126
                                      93 | ENSP00000338157
                                                                             smart
  interpro accession
 <character>
  PS50157
                   IPR007087
  PS50157
                   TPR007087
  PS50157
                   TPR007087
  SM00355
                   TPR015880
  SM00355
                   TPR015880
  SM00225
                   TPR000210
```

Example: use ensembldb to map peptide features to the genome.

```
## Map all protein domains to the genome
gen_map <- mapToGenome(prts, edb)</pre>
## Plot the results for the first protein (transcript)
txid <- gen_map[[1]]$tx_id
## Get the gene region track for the first transcript
tx <- getGeneRegionTrackForGviz(edb, filter = TxidFilter(txid))</pre>
## Add a protein ID column
map_1 <- gen_map[[1]]</pre>
map 1$id <- names(map 1)</pre>
## Plot using Gviz
library(Gviz)
plotTracks(list(GenomeAxisTrack(),
                 GeneRegionTrack(tx, name = "tx"),
                 AnnotationTrack(map_1, groupAnnotation = "id",
                                 just.group = "above",
                                 name = "Protein domains")),
           transcriptAnnotation = "transcript")
```



Things not covered

ensembldb provides full AnnotationDbi support.

Res <- select(edb, keys="SKA2", keytype="GENENAME")

17

17

Get all data for the gene SKA2

chromosome 83257441

chromosome 83257441

• Example: use AnnotationDbi's select method to fetch annotations: keys can be a character list of IDs or a list of filter objects.

```
head(Res. n=3)
 ENTREZID
                  EXONID EXONIDX EXONSEQEND EXONSEQSTART
                                                        GENEBIOTYPE
   348235 FNSF00001324111
                                 59155269
                                             59155131 protein_coding
2 348235 ENSE00003636954 2 59131367 59131281 protein_coding
3 348235 ENSE00003478713 3 59119495
                                             59119319 protein_coding
  GENEID GENENAME GENESEOEND GENESEOSTART INTERPROACCESSION ISCIRCULAR
1 ENSG00000182628
                   SKA2 59155269
                                      59109951
                                                     TPR026762
2 ENSG00000182628 SKA2 59155269 59109951
                                                    IPR026762
3 ENSG00000182628
                   SKA2 59155269 59109951
                                                     IPR026762
 PROTDOMEND PROTDOMSTART PROTEINDOMAINID PROTEINDOMAINSOURCE
                                                              PROTEINID
        115
                              PF16740
                                                   pfam ENSP00000333433
        115
                              PF16740
                                                 pfam ENSP00000333433
                              PF16740
        115
                                                   pfam ENSP00000333433
   PROTEINSEOUENCE
1 MEAEVDKLELMFQKAESDLDYIQYRLEYEIKTNHPDSASEKNPVTLLKELSVIKSRYQTLYARFKPVAVEQKESKSRICATVKKTMNM
2 MEAEVDKLELMFOKAESDLDYIOYRLEYEIKTNHPDSASEKNPVTLLKELSVIKSRYOTLYARFKPVAVEOKESKSRICATVKKTMNM.
3 MEAEVDKLELMFOKAESDLDYIOYRLEYEIKTNHPDSASEKNPVTLLKELSVIKSRYOTLYARFKPVAVEOKESKSRICATVKKTMNM
 SEOCOORDSYSTEM SEOLENGTH SEONAME SEOSTRAND SYMBOL TXBIOTYPE TXCDSSEOEND
                            17 -1 SKA2 protein_coding 59155163
     chromosome 83257441
```

-1 SKA2 protein_coding 59155163 -1 SKA2 protein_coding 59155163

Things not covered

- Easy integration of UCSC and Ensembl annotations: use seqlevelsStyle to change chromosome naming scheme in EnsDb.
- Example: How to integrate Ensembl based annotation with UCSC data?

```
## Get chromosome names, they are "Ensembl-formatted"
head(seglevels(edb))
Γ17 "1" "10" "11" "12" "13" "14"
## Get genes on chromosome Y, UCSC style.
genes(edb, filter=SegnameFilter("chrY"))
GRanges object with 0 ranges and 6 metadata columns:
  seqnames ranges strand | gene_id gene_name entrezid gene_biotype
     <Rle> <IRanges> <Rle> | <character> <character> <character> <character>
  seq_coord_system
                        symbol
       <character> <character>
 seginfo: no seguences
## Solution: change the chromosome naming style:
sealevelsStvle(edb) <- "UCSC"
## Get chromosome names
head(seqlevels(edb))
```

4□ > 4同 > 4 = > 4 = > ■ 900

[1] "chr1" "chr10" "chr11" "chr12" "chr13" "chr14"

Warning massaga.

Finally

Thank you for your attention!

https://github.com/jotsetung/EuroBioC2016-ensembldb.git