# Series 2

## Genomics and bioinformatics - Week 3 - September 27, 2012

In today's session, you will use publicly available genome sequence and annotation data for a particular species to extract some biological information about that species.

If you do not have a working copy of Python and R on your computer, please go through last week's tutorial before starting this exercise.

# 1 Getting data with UCSC

### 1.1 Visualizing genome data

- 1. Go to the UCSC Genome Browser and select the Mus musculus genome.
- 2. Visualize the most recent assembly (mm10) of mouse chromosome 18.
- 3. Scroll down to "Mapping and Sequencing Tracks" and load the GC percent track.

### 1.2 Downloading genome data

Copy the "sequence" file chr18.fa and "annotation" files chr18.gtf, chr18\_mod.txt for mouse chr18 from the USB keys provided by us.

Note: Actually, the .fa files for mouse can be downloaded from the UCSC Downloads page: http://hgdownload.cse.ucsc.edu/goldenPath/mm9/chromosomes/, and the .gtf file for the whole mouse genome can be downloaded from ENSEMBL: ftp://ftp.ensembl.org/pub/release-64/gtf/mus\_musculus/.

# 2 Manipulating data with Python

- 1. Load the .fa file for chr18 and extract the sequence.
- 2. Determine the length of the sequence (see len).
- 3. Calculate the number of As, Gs, Cs and Ts in the sequence.
- 4. Compute the GC content of the chromosome.
- 5. Plot GC content along mouse chr18 using an appropriate window (bin) sizes (use matplotlib).
- 6. Write the start and end coordinates of each bin and it's corresponding GC content to a file, as follows: binStart <tab> binEnd <tab> GC\_content

Note: if you feel confident, this is a good occasion to save time trying the Biopython library: from Bio import seqIO # then use seqIO.read() from Bio.SeqUtils import GC # then use GC() Note: list methods such as count, append, etc. may be useful.

# 3 Manipulating data with R

#### 3.1 Exons

The .gtf file is a tab-delimited file, with the following column headers: chromosome source feature start end score strand frame attributes.

- 1. Load the .gtf file for chr18 in R.
- 2. Extract the rows corresponding to exons from the feature column to a new table.
- 3. Compute exon sizes, and attach them to the table in a new column "exonSize".
- 4. Plot the exon size distribution for chr 18.

#### 3.2 Genes

The modified annotation file chr18\_attributes.txt is also a tab-delimited file, with the following column headers:

chromosome source feature start end score strand frame gene\_id transcript\_id exon\_number gene\_name gene\_biotype transcript\_name protein\_id .

- 1. Load the modified .txt annotation file for chr18 in R.
- 2. Find out the ID and name of the gene containing,
  - a) the longest exon, b) most number of exons.
- 3. List all the intron-less genes in the chromosome.

#### 3.3 GC content

- 1. Load the file (table) generated by your python script into R.
- 2. Recreate the GC content plot for chr18 in R.

### 3.4 Some useful features for this exercise

- which, length, max, hist
- Loops and conditions: for, if, in
- Conversions: as.vector, as.numeric, as.data.frame, as.factor, float, int,...