## **Omics Big Data**

Spring 2018

## Project 1, week 6

Which genome annotation set is the best for the rice reference genome?

The goal of this project is to evaluate the quality of the three genome annotation datasets for the rice genome.

There are three commonly used genome annotations for the reference rice genome. You can access them from

http://rice.plantbiology.msu.edu/ (known as MSU rice genome annotation, Release 7),

http://rapdb.dna.affrc.go.jp (known as RAP DP) and

ftp://ftp.genomics.org.cn/pub/ricedb/rice\_update\_data/bgf\_genes/irgsp/\_ (known\_as\_BGI\_ricedatabase).

You can also find these annotation files from the server /share/home/ccwei/courses/2018/omics/proj1/gff.

We have collected <u>226 RNA-seq datasets for rice</u>. You can access some of them at /share/home/ccwei/courses/2018/omics/proj1/rice\_rna. The rice reference genome is also given to you at /share/home/ccwei/courses/2018/omics/proj1/IRGSP-1.0 genome.fasta.

You are expected to evaluate two rice genome annotation data by these RNA-seq data. Please pick at least 5 RNA-seq data, and align these RNA-seq data to the rice genome and check how many of the genes/transcripts can be validated by at least 2 RNA-seq datasets. You can use any sequencing alignment (or mapping) tools to align these RNA-seq data back to the rice genome. You are expected to report the description of steps in your analysis pipeline, the reasons why you choose these tools for each step, together with the following results.

- 1. The total number of RNA-seq reads for your analysis;
- 2. The number of reads that can be aligned to the rice genome (with the criteria you use) for each RNA-seq dataset;
- 3. The number of reads that can't be aligned to the rice genome;
- 4. For all genome annotation sets,
  - a) How many of the aligned reads locate inside an exon, i.e., not include exon junctions;
  - b) How many of the aligned <u>reads include exon junction</u>;
  - c) How many genes are validated with at least two exons with their junctions supported with at least k reads, where k=1, 5, 10 and 50;
  - d) the percentages of exons in the rice genome that can be covered by at least k reads, where k=1, 5, 10, and 50;
- 5. (bonus) You are assigned with a rice chromosome. In order to get a full credit of this project, you need to have results on at least this chromosome. You will get some bonus if you finish your analysis for more than one chromosome.

(input the assignment of chromosomes to students here )

Chr11	Chr5
Chr3	Chr8
Chr10	Chr9
Chr5	Chr2
Chr6	Chr9
Chr7	Chr6
Chr1	Chr2
Chr8	Chr12
Chr4	Chr7
Chr1	

时间

年 月 日

This is a real research project. The lecturer does not know the result either. Therefore, you need to give detailed information about each step in your analysis so that your result can be reproduced by others if it is needed.

Students are encouraged to form a team of two or three (at most three), and submit a report for the whole team instead of a report for each individual. However, you have to describe the contribution of each individual in the report if you choose to work as a team instead.

Note: You are expected to use the task management system OpenLava to submit all your computation jobs. Please don't run your job in the management node. A brief introduction for OpenLava system is in our server (not the course webpage) under the directory /share/home/ccwei/OpenLava. Please download it to your own computer and try to use the task management system for all the project.

## Turning in your project work Submit your results through the course website at http://cgm.sjtu.edu.cn/index/pub/courses/2018/omics/courseG/login.html before 10AM, May 9, 2018. The result includes your project report and your source codes for the project. -----cut-----here-----独立作业承诺: (请选择一个, 并签名) 1. 本人, \_\_\_\_\_, 保证本次作业由自己独立完成。 签名 时间 年 月 日 或者 2. 本人,\_\_\_\_\_,保证本次作为和\_\_\_\_\_\_同学讨论后,由自己独立完成。 讨论内容包括 签名