

Canadian Bioinformatics Workshops

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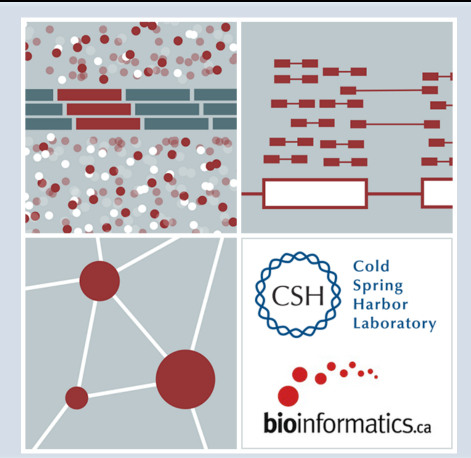
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English French

RNA-Seq Module 3 Expression and Differential Expression (lecture)

Malachi Griffith, Obi Griffith, Fouad Yousif High-Throughput Biology: From Sequence to Networks March 20-26, 2017



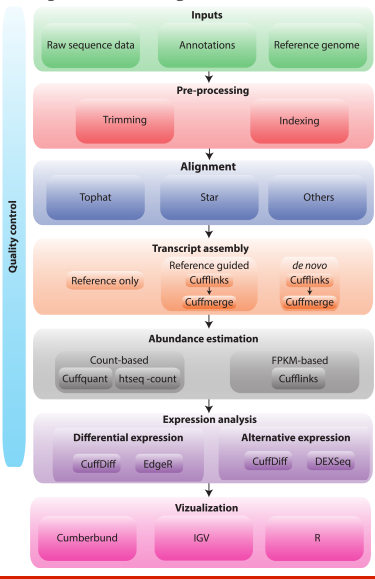




Learning Objectives of Tutorial

- Generate gene/transcript expression estimates with StringTie
- Perform differential expression analysis with Ballgown
- Summarize and visualize results
 - Ballgown
 - Old school R methods

RNA-seq Analysis Flow Chart



4-i. Generate expression estimates

- The alignment SAM/BAM files generated in the previous step will now be used by StringTie to calculate expression estimates
 - For all transcripts on the target chromosome
- For this step options '-G' and '-e' are used
 - '-e' forces StringTie to calculate expression values for known transcripts
 - To discover novel transcripts with StringTie you should:
 - Not use the '-e' o '-G' option. De novo transcript assembly and estimation will be performed. (we will try this in Module 4) OR ...
 - Use the '-G' option only. Known transcripts will be used as a 'guide', but novel transcripts will also be predicted.
- This step will generate one isoform and one gene expression file for each library
 - Expression values are reported as 'FPKM', or 'Fragments Per Kilobase of exon per million fragments Mapped'
 - Where each 'fragment' corresponds to a read-pair mapped to the genome

4-i. Generate expression estimates (Optional Alternatives)

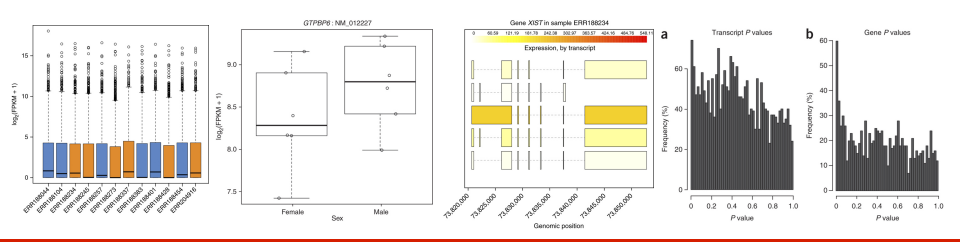
- Another alternative we will explore is a count-based method
 - We will use a program called htseq-count
 - Requires name-sorted SAM file
 - We will count at the gene level (transcript-level is also possible)
- In the end we will have two expression estimates for each sample
 - HISAT2/StringTie
 - HISAT2/Htseq-count

4-ii. Perform differential expression analysis

- In this step we will use Ballgown to:
 - Combine expression estimates from our 6 libraries into more convenient files
 - Combine expression estimates across replicates
 - Compare UHR vs. HBR and identify significantly differentially expressed genes and isoforms (transcripts)
- Note that these commands can get quite complicated when you have replicates
 - The positioning of spaces and commas, and grouping of libraries matters!
- Comparisons
 - Compare UHR vs. HBR using all replicates, for known (reference only mode) transcripts

4-iii. Summarize and visualize results

- In this step we will run the R package Ballgown to visualize our expression and differential expression results.
 - See online tutorial for details
 - https://github.com/alyssafrazee/ballgown
 - http://bioconductor.org/packages/release/bioc/html/ ballgown.html

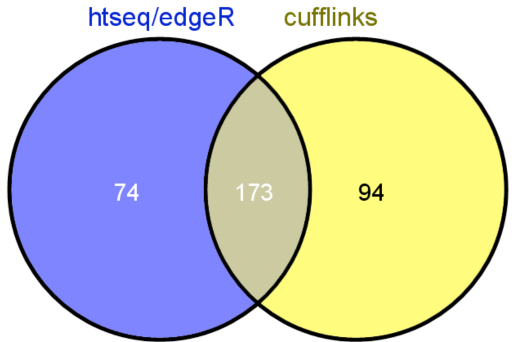


Summarize and visualize results (optional)

- In this step we will use R to summarize and visualize the results of the previous steps
- Explanation of the R commands is provided in the online wiki
- Examples of the tasks performed:
- Examine the expression estimates
 - How reproducible are the technical replicates?
 - How well do the different library construction methods correlate?
 - Visualize the differences between/among replicates, library prep methods and tumor versus normal
- Examine the differential expression estimates
 - Visualize the expression estimates and highlight those genes that appear to be differentially expressed according to Ballgown
 - Generate a list of the top differentially expressed genes

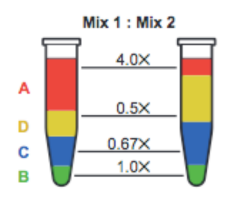
Perform differential expression analysis with edgeR using htseq output (optional)

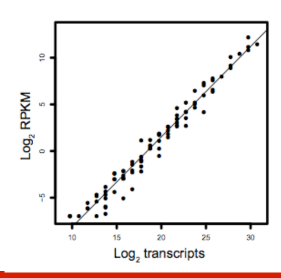
- Make use of raw counts generated by htseq-count
- Load into R and process with edgeR package
- Compare significantly differentially expressed genes from two methods

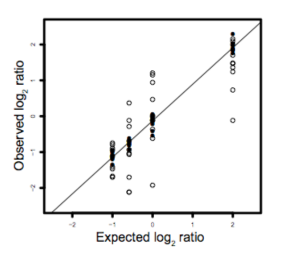


Analysis of ERCC spike-in expression and differential expression (optional)

- https://tools.lifetechnologies.com/content/sfs/manuals/ cms 086340.pdf
- Lower Limit of Detection
- Dynamic Range (dose response)
- Fold-change response (DE)







We are on a Coffee Break & Networking Session