RNA-Seq Demo on Galaxy

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*August 12, 2015*
Our RNA-Seq Demo Data

We will be assembling the DNA Polymerase protein units from the H37Rv strain of *Mycobacterium tuberculosis*, the causative agent of TB, also known as the consumption.

The raw reads originated from the Short Read Archive on NCBI. The accession number for the set is SRX212035.

This dataset consists of paired-end, ~75bp RNA-Seq reads.

Cristobal Rojas, *La miseria* (1886)
Let's get some sequence data

Galaxy allows users to publish their data to share with each other.

Let’s start with “Shared Data” at the top. Then select Data Libraries from the menu.

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Let’s get some sequence data

Choose Workshop Data.

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Expand folder

Check both boxes

Import the Data sets to current history.

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Data set is imported – Click on Analyze Data to return.
Step 1: Assess the Quality of Inputs

We will first get an idea of the quality of our input data sets.

The FastQC tool will produce graphical output that makes it easy to gauge the characteristics of the data – quality, patterns, biases, gc content etc.

Choose the left or right reads file and run it. Compare your results with your neighbors'.

Pro tip: Use the search bar to find tools

(I added a space at the end here)
Step 1: Assess the Quality of Inputs

The input data usually declines in quality as the reads progress.

The quality score is assigned by the sequencing machine as it reads each base. It is a rough estimate of how ambiguous the signal is.

Sequence: ATGCAT
Quality Score: 39 38 23 19 3 3

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Step 2: Trim Input Sequences

We’ve determined that the input data sets need some work before they are used in downstream processes. We’ll use the FASTQ quality trimmer by sliding window to trim reads based on quality score.

Run this tool for both input data sets.

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Step 3: Rinse, Repeat

Now that the files are trimmed, we will re-assess their quality. If necessary, keep trimming away until you are satisfied with the input files.

I renamed my trimmed files to help me keep them straight.
Step 3: Rinse, Repeat

Pictured are the left and right reads after trimming is complete. These will do!
Step 4: Assembly

Next we will put the reads together to create a complete picture of the actively transcribed genes of the sample organism.

Trinity is a de novo assembler that has been optimized for use on Mason. We will use it to assemble our reads.

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It finished! We’re done, right?

An assembler solves a computer problem of putting together a puzzle from tiny pieces. The output of the assembler is a guess – but we don’t know how accurate it is. We could look at:

- Basic stats of the assembly – “Contigs”
  - Number of “Contigs” vs. Expected Number
  - N50 – a weighted average
  - Average Length
  - Max Length
- Check contigs against known genes with Blast
Step 5: Assessing Quality of Assembly

Important statistics for assembly quality:
Contig Length Distribution

Assemblies will typically produce a number of complete contigs representing whole transcripts, and a large number of partial transcripts. This biases the average contig length toward the low end. The N50 is a measure weighted by total sequence length in the assembly.

Total Sequence: 100bp

Contigs, sorted by length

Histogram of isotig lengths

N50 = 9

Average Contig Length = 4.5

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Step 5: Assessing Quality of Assembly

Getting these stats in Galaxy:

Run assemblystats to get a summary and histograms of your contig length distribution.

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Step 6: Getting more data

Right-click **HERE** and choose “Copy link location”.

The TB accession can be found **at NCBI**.
Step 7: Check Against Database

For this step, we’ll check to see how well our assembled transcripts compare to what we already know.

Use this step to give a rough annotation of genes, to make sure that your transcripts are from nuclear genes, or to gauge how complete your sequence is.
Step 7: Check Against Database

We will use Blastx to search the database for our genes.

Use blast database from history.

Make sure to choose Pairwise HTML output for readability.
**Step 7: Check Against Database**

We see the expected genes as the top hits!

We could limit the number of hits depending on output desired.

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Step 8: Differential Expression

We will use RSEM and EBSeq to calculate differential gene expression.

First we need to build a reference using RSEM prepare reference.
Step 8: Differential Expression

Next, gene counts will be produced using RSEM. Make sure to use the prepared library and the paired end reads. Do not create a BAM file.
Step 8: Differential Expression

We will treat this sample like a real set with three replicates. The notation for the condition is tricky – C1 and C2 are sample names, and it assumes the data is in tabular with one gene name column and all other columns are counts.
Step ..?

RNA-Seq is a very versatile technology. You can use the data for:

- Gene discovery based on transcripts
- Genome evidence – introns, exons, junction
- Gene expression patterns in multiple samples
- SNP calling/other variants
- Protein divergence between samples

We have gotten to the assembly step, but there is a lot to learn about the data now that it is put together. A foundation in the use of Galaxy coupled with Indiana University resources will enable you to reach these goals.
Fin

Thanks for watching!
Questions and comments:
Email help@ncgas.org