

# CRSP: Comparative RNA-seq Pipeline

Python Requirement: Python version  $\geq 3.0$

You will need to have the following programs installed and in your PATH

- cd-hit-est (<https://github.com/weizhongli/cdhit>)
- NCBI-BLAST+ (<ftp://ftp.ncbi.nlm.nih.gov/blast/executables/blast+/LATEST/>)
- RSEM (<http://deweylab.github.io/RSEM/>)
- Bowtie (<http://bowtie-bio.sourceforge.net/>)

## Workflow (Examples)

1. Merge multiple transcriptome assemblies, prepending a label to contigs from each assembly to avoid name collisions

```
./src/crsp_merge_assemblies.py assembly_1 ./User_Assembly_Files/Trinity_assembly_1.fasta \  
assembly_2 ./User_Assembly_Files/Trinity_assembly_2.fasta \  
> merged_assembly.fasta
```

2. Create a non-redundant transcriptome assembly using cd-hit-est

```
cd-hit-est -M 200000 -T 0 -i merged_assembly.fasta -o non_redundant_assembly.fasta -c 1 \  
> reduce_assembly.log
```

3. Create a BLAST database for a comparative reference protein set [please see the folder: Blastdb\_Protein]

CRSP (default) has build the mouse protein blastdb (the users can build their own blastdb)

```
makeblastdb -in ./Blastdb_Protein/Screened_Mouse_Protein_with_UniqueID.fa \  
-dbtype prot \  
-out ./Blastdb_Protein/Mouse_protein_blast_index \  
-title mouse_protein \  
-logfile mouse_protein.makeblastdb.log
```

4. Run BLAST+ locally on the non-redundant assembly and the comparative protein set

```
blastx \  
-num_threads 96 \  
-db ./Blastdb_Protein/Mouse_protein_blast_index \  
-outfmt 6 \  
< non_redundant_assembly.fasta \  
> mouse_protein.blast \  
2> mouse_protein.blast.log
```

5. Extract the best BLAST hit for each contig

```
./src/crsp_blast_tophit.py < mouse_protein.blast > mouse_protein.tophits
```

6. Create a mapping from contigs to comparative reference proteins with the given e-value threshold

```
./src/crsp_tophits_to_map.py -e 0.00001 < mouse_protein.tophits > contig_to_mouse_protein.map
```

## 7. Prepare an RSEM reference using the non-redundant assembly

```
rsem-prepare-reference \  
  --bowtie \  
  non_redundant_assembly.fasta \  
  rsem_reference \  
  &> rsem_prepare_reference.log
```

## 8. Compute contig expression levels using RSEM | The example is for paired end reads. For single end reads please see RSEM website: <http://deweylab.github.io/RSEM/>

```
rsem-calculate-expression \  
  --bowtie-n 2 \  
  --no-bam-output \  
  --paired-end \  
  ./User_RNASeq_Files/Sample_1.R1.fastq \  
  ./User_RNASeq_Files/Sample_1.R2.fastq \  
  rsem_reference \  
  Sample_1 \  
  &> rsem_calculate_expression.log
```

## 9. Map contig expression levels to comparative reference protein expression levels

```
./src/crsp_map_abundance_estimates.py \  
  contig_to_mouse_protein.map \  
  < Sample_1.genes.results \  
  > Sample_1.proteins.results
```

## 10. Map protein expression levels to gene symbol expression levels

```
./src/crsp_map_abundance_estimates.py \  
  ./Blastdb_Protein/Mouse_Protein_UniqueID_to_Symbol.map \  
  < Sample_1.proteins.results \  
  > Sample_1.gene_symbols.results
```

## Output File:

Sample\_1.gene\_symbols.results

## Citation

A Nile Grass Rat Transcriptomic Landscape Across 22 Organs By Ultra-deep Sequencing and Comparative RNA-seq pipeline (CRSP)(Submitted)

## Contact

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