**The main pipeline for the manuscript**

**1. De novo transcriptome assembly**

**I created two different assembly; one from C3 homografts and another from C4 homografts.**

**To do so, I created two sample list file contatining the samples that were used for the assembly construction:**

**Sample list table for C3 homografts:**

| **Tissue #** | **Replication** | **Left** | **Right** |
| --- | --- | --- | --- |
| HM.Root | R.C3\_C3.R1 | 78062\_1\_paired.fastq.gz | 78062\_2\_paired.fastq.gz |
| HM.Root | R.C3\_C3.R2 | 78064\_1\_paired.fastq.gz | 78064\_2\_paired.fastq.gz |
| HM.Root | R.C3\_C3.R3 | 78066\_1\_paired.fastq.gz | 78066\_2\_paired.fastq.gz |
| HM.Shoot | S.C3\_C3.R1 | 78086\_1\_paired.fastq.gz | 78086\_2\_paired.fastq.gz |
| HM.Shoot | S.C3\_C3.R2 | 78088\_1\_paired.fastq.gz | 78088\_2\_paired.fastq.gz |
| HM.Shoot | S.C3\_C3.R3 | 78090\_1\_paired.fastq.gz | 78090\_2\_paired.fastq.gz |

**Sample list table for C4 homografts:**

| **Tissue #** | **Replication** | **Left** | **Right** |
| --- | --- | --- | --- |
| HM.Root | R.C4\_C4.R1 | 78080\_1\_paired.fastq.gz | 78080\_2\_paired.fastq.gz |
| HM.Root | R.C4\_C4.R2 | 78082\_1\_paired.fastq.gz | 78082\_2\_paired.fastq.gz |
| HM.Root | R.C4\_C4.R3 | 78084\_1\_paired.fastq.gz | 78084\_2\_paired.fastq.gz |
| HM.Shoot | S.C4\_C4.R1 | 78104\_1\_paired.fastq.gz | 78104\_2\_paired.fastq.gz |
| HM.Shoot | S.C4\_C4.R2 | 78106\_1\_paired.fastq.gz | 78106\_2\_paired.fastq.gz |
| HM.Shoot | S.C4\_C4.R3 | 78108\_1\_paired.fastq.gz | 78108\_2\_paired.fastq.gz |

**1.1. Trinity run for De novo transcriptome assembly:**

Trinity --seqType fq --samples\_file "Homo.C3.sample.txt" --max\_memory 40G --CPU 14

**Note: I did not know whether the read files were strand specific or not, so that's why I ran it first without --SS\_lib\_type, and then examine the strand specificity in the next step.**

**1.2. Examine Strand Specificity of RNA-Seq Reads**

**Note: I did not know if the reads were sequenced stranded or not.**

So, first, I ran Trinity without --SS\_lib\_type option and then I used the guidence in this link to find if they are stranded or not (<https://github.com/trinityrnaseq/trinityrnaseq/wiki/Examine-Strand-Specificity>).

**First, align your reads back against your Trinity assembly. Bowtie2 works well for this:**

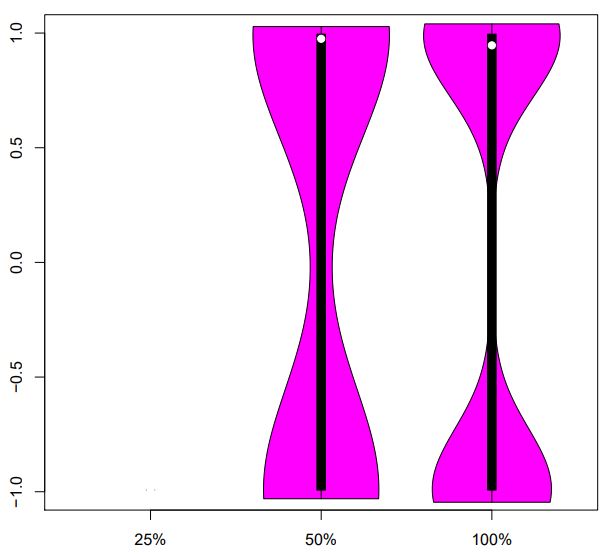
/usr/lib/trinityrnaseq/util/misc/run\_bowtie2.pl --target Trinity.fasta --left 78080\_1\_paired.fastq --right 78080\_2\_paired.fastq | samtools view -Sb - | samtools sort - -o bowtie2.coordSorted.bam mkdir Strand\_specificity\_dir mv \*.bt2 /mkdir Strand\_specificity\_dir cd /mkdir Strand\_specificity\_dir

**Then, examine the distribution of strand-specificity - looking at the distribution of orientations for the first read of paired-end fragment reads.**

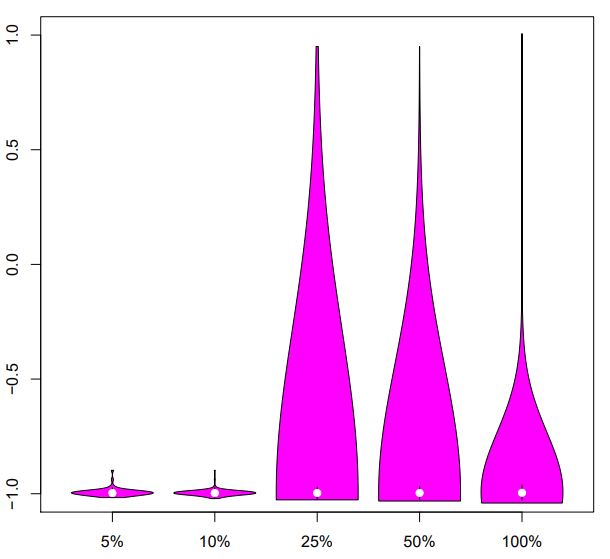
/usr/lib/trinityrnaseq/util/misc/examine\_strand\_specificity.pl bowtie2.coordSorted.bam So, when I realized that it is stranded, I renamed the trinity output dir and ran the analysis against cd ../ rm trinity\_out\_dir unstrand\_trinity\_out\_dir

**1.2.1. An unstranded example.**

This is from my C4 assembly that I made it without --SS\_lib\_type

**1.2.2.Stranded example.**

This is from C3 assembly that I made it with --SS\_lib\_type

**1.3. Run Trinity as Stranded reads:**

Trinity --seqType fq --samples\_file "Homo.C3.sample.txt" --SS\_lib\_type RF --max\_memory 40G --CPU 14

**2. Assembly quality assessment**

**2.1. Assessing the read content of the transcriptome assembly**

**2.1. RNA seq Read Representation**

In order to comprehensively capture read alignments, we run the process below. Bowtie2 is used to align the reads to the transcriptome and then we count the number of proper pairs and improper or orphan read alignments. Related link: <https://github.com/trinityrnaseq/trinityrnaseq/wiki/RNA-Seq-Read-Representation-by-Trinity-Assembly>

**First, build a bowtie2 index for the transcriptome:**

bowtie2-build Trinity.fasta Trinity.fasta

Then perform the alignment to just capture the read alignment statistics.

bowtie2 -p 10 -q --no-unal -k 20 -x Trinity.fasta -1 reads\_1.fq -2 reads\_2.fq \ 2>align\_stats.txt| samtools view -@10 -Sb -o bowtie2.bam

**The output from C3 homograft was copied here:**

28489039 (100.00%) were paired; of these: 3042513 (10.68%) aligned concordantly 0 times 10653739 (37.40%) aligned concordantly exactly 1 time

14792787 (51.92%) aligned concordantly >1 times

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3042513 pairs aligned concordantly 0 times; of these:

377855 (12.42%) aligned discordantly 1 time

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2664658 pairs aligned 0 times concordantly or discordantly; of these:

5329316 mates make up the pairs; of these:

1064874 (19.98%) aligned 0 times

1270360 (23.84%) aligned exactly 1 time

2994082 (56.18%) aligned >1 times

98.13% overall alignment rate

**Interpration:**

Bowtie2 found 3042513 read pairs that it couldn't align concordantly and looked for discordant alignments for them. It could align 377855 of them discordantly, leaving 2664658 still unmapped. It then tried to align each of the reads in those pairs separately (i.e., as singletons), which worked for about 79% of them.

**So, I could say 88% (37.40 + 51.92) mapped or 98.13% in overall that is a good percentage and can say the quality of assembly is good.**

A typical Trinity transcriptome assembly will have the vast majority of all reads mapping back to the assembly, and ~70-80% of the mapped fragments found mapped as proper pairs (yielding concordant alignments 1 or more times to the reconstructed transcriptome).

**2.2. Transcriptome Contig Nx and ExN50 stats**

Below we describe Trinity toolkit utilities for computing contig Nx statistics (eg. the contig N50 value), in addition to a modification of the Nx statistic that takes into consideration transcript expression (read support) data, which we call the ExN50 statistic.

**The 'Gene' Contig Nx Statistic**

Based on the lengths of the assembled transcriptome contigs, we can compute the conventional Nx length statistic, such that at least x% of the assembled transcript nucleotides are found in contigs that are at least of Nx length. The traditional method is computing N50, such that at least half of all assembled bases are in transcript contigs of at least the N50 length value.

**The following script in the Trinity toolkit will compute these values for you like so:**

TRINITY\_HOME/util/TrinityStats.pl Trinity.fasta

**2.2.1 Nx stat for C3:**

**#**

**Counts of transcripts, etc.**

**#**

Total trinity 'genes': 47158

Total trinity transcripts: 88253

Percent GC: 40.69

**#**

Stats based on ALL transcript contigs:

**#**

Contig N10: 3620

Contig N20: 2849

Contig N30: 2374

Contig N40: 2037

Contig N50: 1752

Median contig length: 911

Average contig: 1179.64

Total assembled bases: 104106958

**#**

**Stats based on ONLY LONGEST ISOFORM per 'GENE':**

**#**

Contig N10: 3493

Contig N20: 2715

Contig N30: 2258

Contig N40: 1929

Contig N50: 1644

Median contig length: 627

Average contig: 988.83

Total assembled bases: 46631287

**2.2.2 Nx stat for C4:**

**#**

**Counts of transcripts, etc.**

**#**

Total trinity 'genes': 46304

Total trinity transcripts: 83692

Percent GC: 40.37

**#**

Stats based on ALL transcript contigs:

**#**

Contig N10: 3792

Contig N20: 2998

Contig N30: 2515

Contig N40: 2159

Contig N50: 1864

Median contig length: 956

Average contig: 1236.91

Total assembled bases: 103519264

**#**

**Stats based on ONLY LONGEST ISOFORM per 'GENE':**

**#**

Contig N10: 3615

Contig N20: 2825

Contig N30: 2353

Contig N40: 1999

Contig N50: 1704

Median contig length: 620

Average contig: 1005.51

Total assembled bases: 46558986

**3.Trinity Transcript Quantification**

**My goal to perform this step:**

This step will give me the expression level as count, FPKM, and TPM. I will run this for both homo root and shoot against their assembly for both C4 and C3. Then I will run it for both hetero root and shoot against their assemblies for both C4 and C3. And then I can run DEG analysis for both root and shoot that how transcriptome profile was changed in root or shoot when they are homografted and heterografted.

| **Tissue#** | **Condition** | **Assembly** |
| --- | --- | --- |
| Root | C3.C3 | C3.assembly |
| Root | C3.C4 | C3.assembly |
| Root | C4.C4 | C4.assembly |
| Root | C4.C3 | C4.assembly |
| Shoot | C3.C3 | C3.assembly |
| Shoot | C4.C3 | C3. assembly |
| Shoot | C4.C4 | C4.assembly |
| Shoot | C3.C4 | C4.assembly |

**3.1. Estimating Transcript Abundance**

There are now several methods available for estimating transcript abundance in a genome-free manner, and these include alignment-based methods (aligning reads to the transcript assembly) and alignment-free methods (typically examining k-mer abundances in the reads and in the resulting assemblies).

In Trinity, we provide direct support for running the alignment-based quantification methods RSEM, as well as the ultra-fast alignment-free method kallisto and 'wicked-fast' salmon.

The Trinity software does not come pre-packaged with any of these software tools, so be sure to download and install any that you wish to use. The tools should be available via your PATH setting (so, typing 'which kallisto'on the linux command line returns the path to where the tool is installed on your system).

If you have multiple RNA-Seq data sets that you want to compare (eg. different tissues sampled from a single organism), be sure to generate a single Trinity assembly and to then run the abundance estimation separately for each of your samples.

**3.2. RSEM method**

**Just prepare the reference for alignment and abundance estimation:**

/usr/lib/trinityrnaseq/util/align\_and\_estimate\_abundance.pl --transcripts Trinity.fasta --est\_method RSEM --aln\_method bowtie --trinity\_mode --prep\_reference

**Run the alignment and abundance estimation (assumes reference has already been prepped, errors-out if prepped reference not located.)**

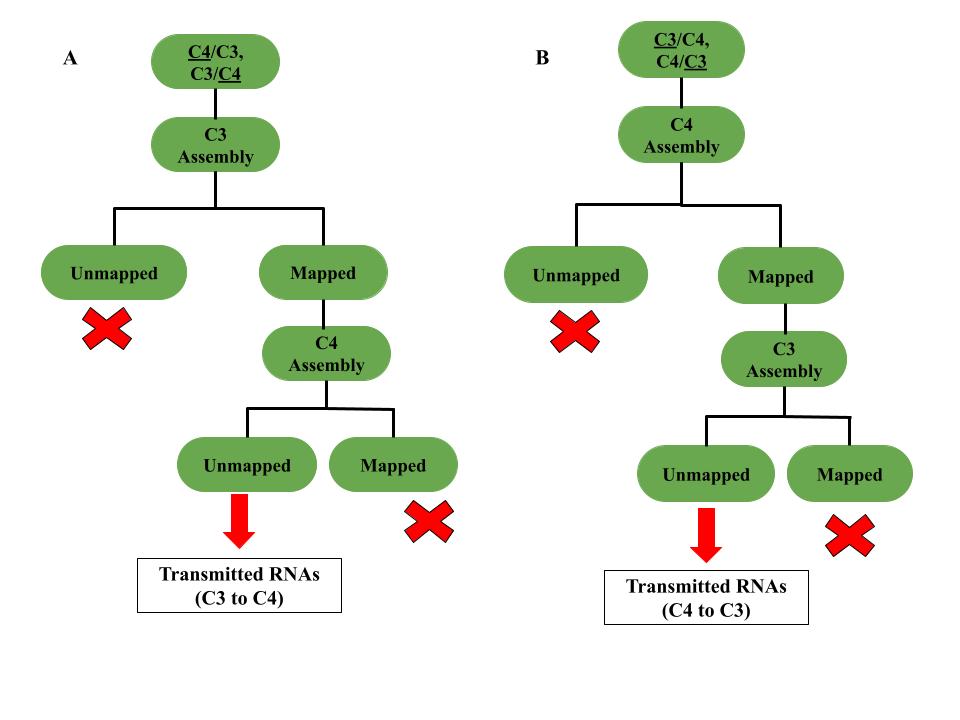
/usr/lib/trinityrnaseq/util/align\_and\_estimate\_abundance.pl --transcripts Trinity.fasta --gene\_trans\_map Trinity.fasta.gene\_trans\_map --seqType fq --samples\_file "" --SS\_lib\_type RF --est\_method RSEM --aln\_method bowtie --trinity\_mode --output\_dir rsem\_outdir

**Note: I used --samples\_file option so, do not need run the --left and --right options and run the command for each condition. It will store output in the folder with condition name.**

**If you have strand-specific data, be sure to include the '--SS\_lib\_type' parameter.**

**It is useful to first run 'align\_and\_estimate\_abundance.pl' to only prep your reference database for alignment, using '--prep\_reference', and then subsequently running it on each of your sets of reads in parallel to obtain sample-specific abundance estimates.**

**PART 2: FINDING TRANSMITTED RNAs**

In this section I will perform different mapping to find the RNAs that are transmitted in heterografts: 

**Note: Gene (contig) will be reported as transmitted RNA that it exists in all 3 replicates.**

**The commands were used for this part:**

**1. create index for the assembly**

bwa index TrinityC3.fasta

**2. Alignment to find the RNAs from C4 which mapped to C3 assembly:**

bwa mem TrinityC3.fasta /home/mahnaz/C4.C3.transcriptome/trimmed.files/Variant\_calling/transmitted\_C3\_to\_C4/root\_C4/78074\_1\_paired.fastq /home/mahnaz/C4.C3.transcriptome/trimmed.files/Variant\_calling/transmitted\_C3\_to\_C4/root\_C4/78074\_2\_paired.fastq | samtools view -o 78074.C4R1\_vs\_C3assembly.bam | samtools fastq -F4 78074.C4R1\_vs\_C3assembly.bam > matched.78074.C4R1\_vs\_C3assembly.fastq It runs an error that can't find the 78074.C4R1\_vs\_C3assembly.bam, but if it runs for a while, a truncated 78074.C4R1\_vs\_C3assembly.bam file will be created and I stopped the run and run it again, it solves the error.

**3. We need to align the mapped reads from step above to C4 assembly to be find the unmapped reads to C4, these reads can be transmitted from C3 grafts or artifacts, so, that's why I consider genes that identified by all 3 replicates. Anyway, we need an index for C4 assembly as well.**

bwa index TrinityC4.fasta

**4. And then, align the mapped reads to C4 to get the unmapped reads:**

bwa mem TrinityC4.fasta /home/mahnaz/C4.C3.transcriptome/trimmed.files/Variant\_calling/transmitted\_C3\_to\_C4/matched.78074.C4R1\_vs\_C3assembly.fastq | samtools view -o matched.78074.C4R1\_vs\_C3assembly.bam | samtools fastq -f4 matched.78074.C4R1\_vs\_C3assembly.bam > unmatched.78074.C4R1\_vs\_C4assembly.fastq

**5. Lastly, run the RSEM step for the output from step4 against C3 assembly (I put the outputs in the directory of 3.1 and run the command below). Actually, I used the index file from part 3.1:**

/usr/lib/trinityrnaseq/util/align\_and\_estimate\_abundance.pl --transcripts Trinity.fasta --gene\_trans\_map Trinity.fasta.gene\_trans\_map --seqType fq --samples\_file "" --SS\_lib\_type RF --est\_method RSEM --aln\_method bowtie --trinity\_mode --output\_dir rsem\_outdir