# RNASeq

Efforts to sequence the transcripts expressed in a cell or organism.

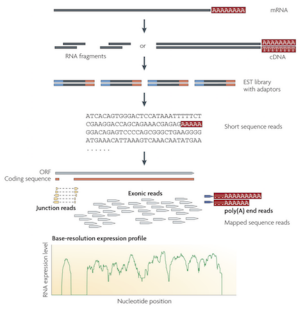
#### **Statistics**

Measuring gene expression

#### Resources

• See the StatQuest Video on RPKM/FPKM/TPM to better understand how the statistics can be used to evaluate gene expression.

Using techniques to extract the Wang et al. Nat Rev Genetics. 2009.



doi:10.1038/nrg2484

Figure 1. A typical RNA-Seq experimen

# Multiple approaches to understanding the transcriptome

- 1. Genome sequenced, align RNAseq reads to genome
- 2. de novo Assembly of mRNA into transcripts
- 3. Quantify gene expression from reads aligned to genome or transcripts

### Reads to Genome mapping

It is important to note that aligning sequences to the genome when there are introns requires dealing with introns. So splice-aware alignements are needed in some cases.

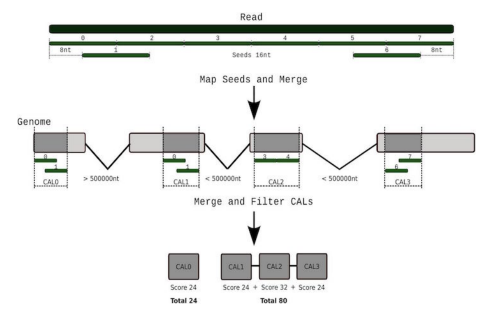


Figure 1: SpliceAlign

Tarraga et al 2017. DNA Research.10.1093/dnares/dsv039

#Reads to Genome mapping

Challenges: mRNA is spliced, genome contains introns

Splice-aware short read aligners. Speed and accuracy tradeoffs \* Tophat + Bowtie – this is old don't use \* HISAT2 \* GMAP/GSNAP \* STAR

## Need to Quantify expression

- Count reads overlapping exons
- Table of total read counts per gene
- Normalize counts for gene length and sequencing library depth
- $\bullet\,$  Gene expression then is FPKM Fragments per Kilobase per Millions of reads
- Tools: htseq-count, stringtie
- SubRead
- BEDtools
- R tools with iRanges

#### Evaluating expression differences

Statistical tools for evaluating gene expression differences

- Ballgown bioconductor package
- DESeq bioconductor package
- edgeR bioconductor package

#### Alternative approach for Quantifying

Compare reads to **Transcripts** instead of Genome \* Kalisto and Sailfish are common tools \* Bray et al 2016 "Near-optimal probabilistic RNA-seq quantification" doi:10.1038/nbt.3519 \* Patro et al 2014 "Sailfish enables alignment-free isoform quantification from RNA-seq reads using lightweight algorithms" doi:10.1038/nbt.2862

## Alignment free quantification

Usage: kallisto quant [arguments] FASTQ-files

Required arguments:

-i, --index=STRING Filename for the kallisto index to be used for

quantification

-o, --output-dir=STRING Directory to write output to

Optional arguments:

--bias Perform sequence based bias correction
-b, --bootstrap-samples=INT Number of bootstrap samples (default: 0)

--seed=INT Seed for the bootstrap sampling (default: 42)

--plaintext Output plaintext instead of HDF5
--fusion Search for fusions for Pizzly
--single Quantify single-end reads

--single-overhang Include reads where unobserved rest of fragment is

predicted to lie outside a transcript
--fr-stranded Strand specific reads, first read forward
--rf-stranded Strand specific reads, first read reverse

-1, --fragment-length=DOUBLE Estimated average fragment length

-s, --sd=DOUBLE Estimated standard deviation of fragment length

(default: -1, -s values are estimated from paired
end data, but are required when using --single)

-t, --threads=INT Number of threads to use (default: 1)

--pseudobam Save pseudoalignments to transcriptome to BAM file
--genomebam Project pseudoalignments to genome sorted BAM file

-g, --gtf GTF file for transcriptome information

(required for --genomebam)

```
-c, --chromosomes Tab separated file with chromosome names and lengths (optional for --genomebam, but recommended)
```

Note this won't quite work to copy and paste.

/master/Rscripts/kallisto profile rf stranded.R

```
#!/usr/bin/bash

module load kallisto
ln -s /bigdata/gen220/shared/data-examples/rnaseq/kallisto/S_cerevisiae_ORFs.fasta
ln -s
kallisto index -i Scer.idx S_cerevisiae_ORFs.fasta
cat samples.tsv | while read ACC COND REP
do
   OUT=output/$COND.$REP
kallisto quant -t 8 --single -l 300 -s 20 -i Scer.idx -o $OUT data/${ACC}_1.fastq.gz
done
Go see /bigdata/gen220/shared/data-examples/rnaseq/kallisto
See also https://github.com/stajichlab/C_lusitaniae_DHED1_RNAseq/blob
```

### Denovo assembly

```
Trinity Assembler for RNASeq
```

```
$ module load trinity-rnaseq
$ module switch perl/5.22.0
$ Trinity --seqType fq --left reads_1.fq --right reads_2.fq --CPU 8 --max_memory 20G
```

#### **ORF** identification

Once we have assembled the transcriptome, want to find genes in there.

#### TransDecoder

• Finds Open Reading Frames in mRNA transcripts

```
$ module load transdecoder
$ TransDecoder.LongOrfs -t target_transcripts.fasta
```

## RNAseq read mapping

```
Using HISAT2 for RNAseq read mapping * S_cerevisiae.fasta.gz * S_cerevisiae.gff3.gz
```

Download those files.

```
# start an interactive session
srun -N 1 -n 4 -p short --mem 16gb --pty bash -l
module load hisat2
# uncompress
gunzip S_cerevisiae.gff3.gz S_cerevisiae.fasta.gz
# build index
hisat2-build S_cerevisiae.fasta yeast
# run search
ln -s /bigdata/gen220/shared/data-examples/rnaseq/yeast_rnaseq/*.gz .
hisat2 -x yeast -1 SRR3396381_1.fastq.gz -2 SRR3396381_2.fastq.gz -S SRR3396381.sam -p 4
module load samtools
samtools view -Ob -o SRR3396381.bam SRR3396381.sam
samtools sort -o SRR3396381.sort.bam SRR3396381.bam
samtools index SRR3396381.sort.bam SRR3396381.bam
samtools flagstat SRR3396381.sort.bam
Get counts
Subread - http://subread.sourceforge.net/
module load subread
GENOME=S_cerevisae.fasta
GFF=S_cerevisae.gff3
OUTFILE=SRR3396381.tab
INFILE=SRR3396381.sort.bam
featureCounts -g gene id -T $CPUS -G $GENOME -s 0 -a $GFF -o $OUTFILE \
-F GTF $INFILE
Template for Projects
Here's a template for RNASeq analyses
https://github.com/biodataprog/RNASeq_template Click on 'Use this template'
- you can create your own version of this.
It will prompt you give it a name.
Go to the command line to download.
git clone yourname/YourRNASeqAnalysis.git
Edit samples.csv to describe names of some experiments SRR3396381.
Download data in input folder.
```

Download or us download script to get genome files (need to put a genome FASTA file in the folder). If want to do kallisto will need a mRNA file of transcriptome.