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# Stranded Transcript Count Table Generation from Long Reads V.12

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ABSTRACT

This protocol is for comparing different samples at the transcript level, using long reads that are mapped to transcripts.

Input(s): demultiplexed and oriented fastq files (see protocol Preparing Reads for Stranded Mapping), transcript reference fasta file, annotation file

Output(s): transcript table, sorted by differential coverage, annotated with gene name / description / location

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Version created by David Eccles

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**BEFORE STARTING** 

Obtain a transcript fasta file, and an annotation file. For the mouse genome, I use the following files:

1. Transcript sequences from Ensembl; this should be the union of cDNA, CDS, and ncRNA sequences (e.g. from

mprotocols.io 10/13/2020

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#### This directory).

- 2. Annotation file obtained from <u>Ensembl BioMart</u> (Ensembl Genes -> Mouse Genes) as a compressed TSV file with the following attribute columns:
- Transcript stable ID
- Gene name
- Gene description
- Chromosome/scaffold name
- Gene start (bp)
- Gene end (bp)
- Strand

A recent version of these files can be obtained from This Zenodo Repository

## Demultiplex Reads

7

Demultiplex and orient reads as per the protocol <u>Preparing Reads for Stranded Mapping</u>. It is expected that these demultiplexed reads will be split up in the current directory, and coupled with a 'barcode\_counts.txt' file. If that's the case, the following should work:

```
for bc in $(awk '{print $2}' barcode_counts.txt);
  do ls oriented/${bc}_reads_dirAdjusted.fq.gz;
done
```

#### Example expected output:

```
oriented/BC03_reads_dirAdjusted.fastq.gz
oriented/BC04_reads_dirAdjusted.fastq.gz
oriented/BC05_reads_dirAdjusted.fastq.gz
oriented/BC06_reads_dirAdjusted.fastq.gz
oriented/BC07_reads_dirAdjusted.fastq.gz
oriented/BC08_reads_dirAdjusted.fastq.gz
```

If the 'barcode\_counts.txt' file is not present, this error will appear:

```
awk: fatal: cannot open file `barcode counts.txt' for reading (No such file or directory)
```

If one or more of the oriented read files is missing, it will look something like this:

```
oriented/BCO3_reads_dirAdjusted.fastq.gz
oriented/BCO4_reads_dirAdjusted.fastq.gz
ls: cannot access 'oriented/BCO5_reads_dirAdjusted.fastq.gz':
  No such file or directory
ls: cannot access 'oriented/BCO6_reads_dirAdjusted.fastq.gz':
  No such file or directory
oriented/BCO7_reads_dirAdjusted.fastq.gz
oriented/BCO8_reads_dirAdjusted.fastq.gz
```

### **Index Preparation**

2 Prepare a substitution matrix for barcode mapping. The default substitution matrix is swayed too much by INDELs in the barcode sequences, so here's one that I've developed using a combination of trial & error and last-train:

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```
C -18 6 -18 -12
G -7 -18 5 -18
T -18 -12 -18 6
```

cDNA.mat

[note: this is a different matrix from that used for demultiplexing and read orientation]

3 Prepare transcript index (see Guidelines for data sources). Following <u>Martin Frith's recommendation</u>, the '-uNEAR' seeding scheme is used to slightly increase sensitivity. This will generate seven additional files of the form <index name>.XXX:

lastdb -uNEAR Mus\_musculus.GRCm38.ensembl\_v98.mtr.fa Mus\_musculus.GRCm38.ensembl\_v98.mtr.fa

## Transcriptome Mapping

4 Reads are mapped to the transcriptome with LAST.

The results of that mapping can be piped through *last-split* and *last-postmask* to exclude unlikely hits, then through *'maf-convert -n tab'* to convert to a one-line-per-mapping CSV format. This CSV format is further processed to make sure that there is only one mapping per transcript-read pair.

```
mkdir -p mapped
for bc in $(awk '{print $2}' barcode_counts.txt);
  do echo "** ${bc} **";
  lastal -P 10 -p cDNA.mat Mus_musculus.GRCm38.ensembl_v98.mtr.fa <(pv
  oriented/${bc}_reads_dirAdjusted.fq.gz | zcat) | \
        last-split -n -m0.99 | last-postmask | maf-convert -n tab | \
        cut -f 2,7,10 | sort | \
        uniq | gzip > mapped/trnMapping_LAST_${bc}_vs_Mmus_transcriptome.txt.gz;
done
```

5 The result is then aggregated to sum up counts per transcript:

```
for bc in $(awk '{print $2}' barcode_counts.txt);
  do echo "** ${bc} **";
  zcat mapped/trnMapping_LAST_${bc}_vs_Mmus_transcriptome.txt.gz | \
    awk -F'\t' -v "bc=${bc}" '{print bc,$1,$3}' | sort | uniq -c | \
    gzip > mapped/trnCounts_LAST_${bc}_vs_Mmus_transcriptome.txt.gz;
done
```

Note: I've split this up into two steps (compared to previous versions of this protocol) so that an intermediate count of the total number of mapped transcripts per barcode can be done:

```
for bc in $(awk '{print $2}' barcode_counts.txt);
  do echo -n "${bc} ";
  zcat mapped/trnMapping_LAST_${bc}_vs_Mmus_transcriptome.txt.gz | \
    awk '{print $2}' | sort | uniq | wc -l;
done
```

# Annotation and Result generation

6 count\_analysis.r

Transcript counts are merged with ensembl gene annotation, then converted into wide format (one line per transcript)

 using an R script.

The transcript annotation in this case is from ensembl BioMart (see Guidelines for more details).

```
#!/usr/bin/env Rscript
library(tidyverse);
## load used barcode identifiers
bcNames <- read.table("barcode_counts.txt", stringsAsFactors=FALSE)[,2];</pre>
## load count data into "narrow" array (one line per count)
trn.counts <- tibble();</pre>
for(bc in bcNames){
   trn.counts <-
        bind rows(trn.counts,
           sprintf("mapped/trnCounts LAST %s vs Mmus transcriptome.txt.gz", bc) %>%
              read table2(col names=c("count", "barcode",
                                        "transcript", "dir")));
}
## remove revision number from transcript names (if present)
trn.counts$transcript <- sub("\\.[0-9]+$","",trn.counts$transcript);</pre>
## convert to wide format (one line per transcript)
trn.counts.wide <- spread(trn.counts, barcode, count) %>%
    mutate(dir = c("+"="fwd", "-"="rev")[dir]);
for(bd in colnames(trn.counts.wide[,-1])){
    trn.counts.wide[[bd]] <- replace na(trn.counts.wide[[bd]],0);</pre>
## load ensemble transcript metadata (including gene name)
ensembl.df <- read delim("ensembl mm10 geneFeatureLocations.txt.gz",</pre>
                          delim="\t");
colnames(ensembl.df) <-</pre>
    c("Transcript stable ID" = "transcript",
      "Gene description" = "Description",
      "Gene name" = "Gene",
      "Gene start (bp)" = "Start",
      "Gene end (bp)" = "End",
      "Strand" = "Strand",
      "Chromosome/scaffold name" = "Chr")[colnames(ensembl.df)];
ensembl.df$Description <- sub(" \\[.*$","",ensembl.df$Description);</pre>
ensembl.df$Description <- sub("^(.{50}).+$","\\1...",ensembl.df$Description);</pre>
options(scipen=15); ## don't show scientific notation for large positions
## merge ensembl metadata with transcript counts
gene.counts.wide <- inner join(ensembl.df, trn.counts.wide, by="transcript");</pre>
gene.counts.wide <- gene.counts.wide[order(-rowSums(gene.counts.wide[,-(1:8)])),];
## write result out to a file
write.csv(gene.counts.wide, file="wide_transcript_counts_LAST.csv", row.names=FALSE);
```

Downstream Workflows

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- 7 Here is a downstream workflow that carries out transcript-level differential expression analysis using <u>DESeq2</u>:
  - Creating Differential Transcript Expression Results with DESeq2

I would like to emphasise that batch effects should be considered for nanopore sequencing, given how frequently the technology changes. Make sure that at least the sequencing *library* (i.e. samples prepared in tandem on the same day from the same kit) is added into the statistical model, and try to make sure that sequencing libraries are fairly heterogeneous - replicates from a sample with skewed transcript distributions could influence the outcome of statistical tests.