Apr 16, 2019

In devel.

Preparing Reads for Stranded Mapping

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dx.doi.org/10.17504/protocols.io.z4uf8ww



ABSTRACT

This protocol is for preparing long reads for stranded mapping, as an intermediate step for additional protocols:

- Aligning strand-oriented sequences to a transcriptome for transcript / gene counting
- Aligning strand-oriented sequences to a genome for confirmatory QC

Input(s): demultiplexed fastq files (see protocol <u>Demultiplexing Nanopore reads with LAST</u>), adapter file (containing strand-sensitive adapter sequences)

Output(s): oriented read files, as gzipped fastq files

Barcode Demultiplexing

1 Demultiplex reads as per protocol <u>Demultiplexing Nanopore reads with LAST</u>.

If this has been done, then the following command should produce output without errors:

```
for bc in $(awk '{print $2}' barcode_counts.txt); do ls reads_${bc}.fastq.gz; done
```

Example output:

```
reads_BC03.fastq.gz
reads_BC04.fastq.gz
reads_BC05.fastq.gz
reads_BC06.fastq.gz
reads_BC07.fastq.gz
reads_BC08.fastq.gz
```

If the <code>barcode_counts.txt</code> file is missing, the output will look like this:

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

If one or more of the barcode-demultiplexed files are missing, the output will look something like this:

```
reads_BC03.fastq.gz
reads_BC04.fastq.gz
reads_BC05.fastq.gz
ls: cannot access 'reads_BC06.fastq.gz': No such file or directory
ls: cannot access 'reads_BC07.fastq.gz': No such file or directory
reads_BC08.fastq.gz
```

Adapter Mapping

Prepare a FASTA file containing adapter sequences (see attached FASTA file).

__adapter_seqs.fa

3 Prepare the LAST index for the adapter file. This will generate seven additional files of the form <index name>.XXX:

```
lastdb adapter_seqs.fa adapter_seqs.fa
```

Orienting Reads

4 Map the reads to the adapter sequences. In this case it's important that the direction of mapping is also recorded, so the *cut* command selects three fields (query name [7], target name [2], mapping direction [10]):

```
for bc in $(awk '{print $2}' barcode_counts.txt);
  do echo "** ${bc} **";
  lastal -Q 1 -P10 adapter_seqs.fa <(pv reads_${bc}.fastq.gz) | \
    maf-convert -n tab | cut -f 2,7,10 | uniq | \
    gzip > adapter_assignments_${bc}.txt.gz
done
```

- 5 Reads are filtered into two groups (and one group-by-omission) based on the mapped direction of the strand-switch primer, then reverse-complemented (if necessary) to match the orientation of the original RNA strand. I use my fastx-rec.pl and fastx-rec.pl scripts for this.
 - fastx-fetch.pl
 - fastx-rc.pl

```
mkdir -p oriented
for bc in $(awk '{print $2}' barcode_counts.txt);
  do echo "** ${bc} **";
  fastx-fetch.pl -i <(zgrep 'SSP' adapter_assignments_${bc}.txt.gz | awk '{if($3 == "+"){print $2}}') <(pv reads_${bc}.fastq.gz) | \
     gzip > oriented/${bc}_reads_fwd.fastq.gz
  fastx-fetch.pl -i <(zgrep 'SSP' adapter_assignments_${bc}.txt.gz | awk '{if($3 == "-"){print $2}}') <(pv reads_${bc}.fastq.gz) | \
     fastx-rc.pl | gzip > oriented/${bc}_reads_rev.fastq.gz
done
```

6 Forward and reverse-oriented sequences are combined together to form a single group of RNA-oriented reads.

```
for bc in $(awk '{print $2}' barcode_counts.txt);
  do echo "** ${bc} **";
  pv oriented/${bc}_reads_fwd.fastq.gz oriented/${bc}_reads_rev.fastq.gz | \
    zcat | gzip > oriented/${bc}_reads_dirAdjusted.fastq.gz
done
```

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