Process Single Cell RNA-Seq reads using scruff

Zhe Wang 2017-11-25

scruff is a toolkit for processing single cell RNA-seq fastq reads. It does demultiplexing, alignment, Unique Molecular Identifier (UMI) correction, and transcript counting in an automated fashion and outputs expression table. This vignette provides a brief introduction to the scruff package by walking through the demultiplexing, alignment, and UMI-counting of a built-in example dataset.

Quick Start

Stepwise Tutorial

Load Example Dataset

The scruff package contains a example single cell RNA-seq dataset examplefastq which is stored as a list of ShortReadQ objects. Each object has 20,000 sequenced reads. They can be accessed via variable examplefastq and exported to current working directory as fastq.gz files. scruff only accepts fastq/fastq.gz files for demultiplexing so these objects have to be stored in fastq.gz format first.

```
library(scruff)

# write fastg.gz files to working directory
for (i in seq_len(length(examplefastq))) {
    writeFastq(examplefastq[[i]], names(examplefastq)[i], mode = "w")
}
```

Demultiplex and Assign Cell Specific Reads

Now the fastq files are saved in current working directory and are ready to be demultiplexed. scruff package provides built-in predetermined sample annotating table exampleannot and barcodes example for demultiplexing the example dataset. In the example fastq files, the barcode sequence of each read starts at base 6 and ends at base 11. The UMI sequence starts at base 1 and ends at base 5. They can be set via bc.start, bc.stop, and umi.start, umi.stop parameters. By default, reads with any nucleotide in the barcode and UMI sequences with sequencing quality lower than 10 (Phred score) will be excluded. The

following command demultiplexes the example fastq reads and trims reads longer than 50 nucleotides. The command returns a summary data.table containing the sample barcodes, filenames, number of reads, sample name, file paths, etc. By default, the cell specific demultiplexed fastq.gz files are stored in ../Demultiplex folder.

```
demultiplex.res <- demultiplex(
    exampleannot,
    examplebc,
    bc.start = 6,
    bc.stop = 11,
    bc.edit = 1,
    umi.start = 1,
    umi.stop = 5,
    keep = 50,
    min.qual = 10,
    yield.reads = 1e+06,
    out.dir = "../Demultiplex",
    cores = 2,
    overwrite = TRUE,
    verbose = TRUE
)</pre>
```

Alignment

scruff has a wrapper function align.rsubread which is a wrapper function to align in Rsubread package. It aligns the reads to reference sequence index and outputs sequence alignment map files. For demonstration purpose, the built-in mitochondrial DNA sequence from GRCh38 reference assembly GRCh38_MT will be used to map the reads. First, a Rsubread index for the reference sequence needs to be generated.

```
ref <- "Homo_sapiens.GRCh38.dna.chromosome.MT.fa"</pre>
index <- "GRCh38_MT"</pre>
# write reference sequence to "./Homo_sapiens.GRCh38.dna.chromosome.MT.fa".
write.table(GRCh38_MT,
            ref,
            quote = FALSE,
            sep = "\t",
            row.names = FALSE,
            col.names=FALSE)
# NOTE: Rsubread package does not support Windows environment.
library(Rsubread)
# Create index files for GRCh38_MT. For details, please refer to Rsubread user manual.
buildindex(
  basename = index,
  reference = ref,
  indexSplit = FALSE,
  memory = 16000
```

The following command maps the fastq files to GRCh38 mitochondrial reference sequence GRCh38_MT and returns the file paths of the generated sequence alignment map files. By default, the files are stored in BAM format in ../Demultiplex folder.

```
# get the paths to demultiplexed fastq.gz files
fastq.paths <- demultiplex.res[!(is.na(cell_num)), fastq_path]

# map the reads
align.res <- align.rsubread(
   fastq.paths,
   index,
   format = "BAM",
   out.dir = "../Alignment",
   cores = 4,
   overwrite = TRUE,
   verbose = TRUE
)</pre>
```

UMI correction and Generation of Expression Table

Example GTF file GRCh38_MT_gtf will be used for feature counting. Currently, scruff applies the union counting mode of the HTSeq Python package. The following command generates the UMI corrected expression table for the example dataset.

Collect gene annotation information from Biomart

Gene annotation information is needed before visualization of the data quality. In this tutorial, Ensembl Biomart database is used for the query and collection of gene annotation information. The following codes retrieve human gene names, biotypes, and chromosome names from latest version of biomart database.

```
# get gene names, biotypes, and chromosome names from biomart
host = "www.ensembl.org"
biomart = "ENSEMBL_MART_ENSEMBL"
dataset = "hsapiens_gene_ensembl"
GRCh = NULL
# get gene IDs
```

```
# exclude ERCC spike-ins and last two rows in expression table
geneID <- expr[!(gene.id %in% c("reads_mapped_to_genome",</pre>
                                "reads mapped to genes")) &
                !grepl("ERCC", expr[,gene.id]), gene.id]
# Get feature annotation data
ensembl <- biomaRt::useEnsembl(biomart = biomart,</pre>
                                dataset = dataset,
                                host = host,
                                GRCh = GRCh)
biomart.result <- data.table::data.table(biomaRt::getBM(</pre>
  attributes = c("ensembl_gene_id", "external_gene_name",
                  "gene_biotype", "chromosome_name"),
 filters = "ensembl_gene_id",
 values = geneID,
 mart = ensembl))
# remove rows containing duplicate gene IDs
biomart.result <- biomart.result[!base::duplicated(biomart.result,</pre>
                                                     by = "ensembl_gene_id"), ]
# Reorder/insert rows so that rows of Biomart query result match
# rows of the expression matrix
biomart.result <- data.table::data.table(biomart.result[</pre>
  match(expr[!(gene.id %in% c("reads_mapped_to_genome",
                             "reads_mapped_to_genes")) &
             !grepl("ERCC", expr[, gene.id]), gene.id],
        biomart.result$ensembl_gene_id),])
```

Visualization of QC metrics

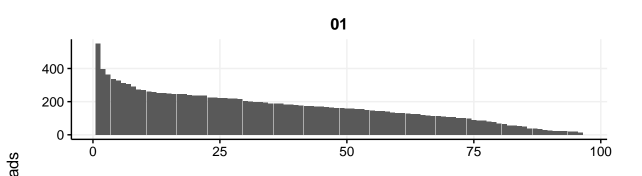
In order to plots data quality, a QC metrics table needs to be generated. collect.qc function collects and summarizes data quality information from results from previous steps and returns a QC metrics table for plotting purpose.

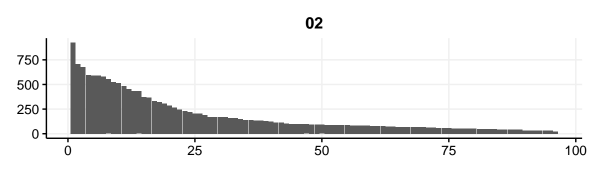
```
qc <- collect.qc(demultiplex.res, align.res, expr, biomart.result)</pre>
```

QC metrics can be visualized by running plot.qc function.

[[1]]

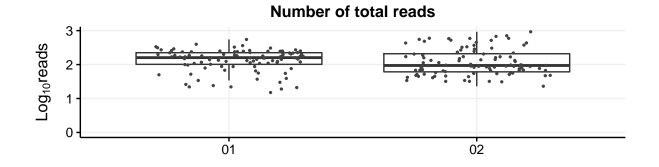
Reads per cell

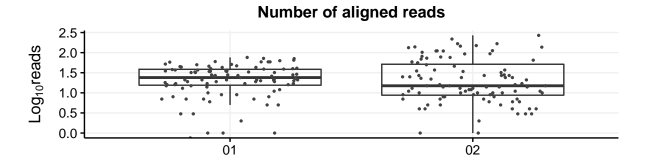


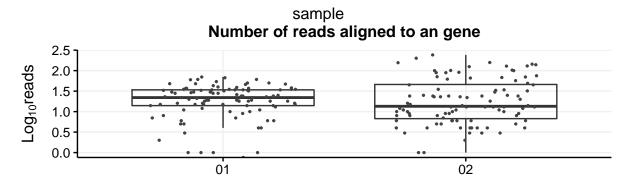


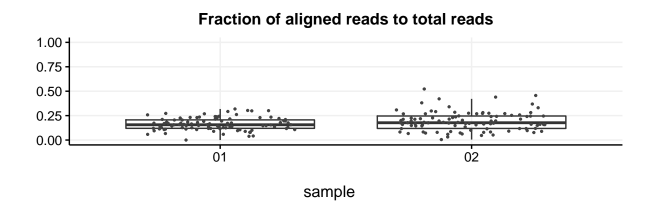
Cells in descending order

[[2]]

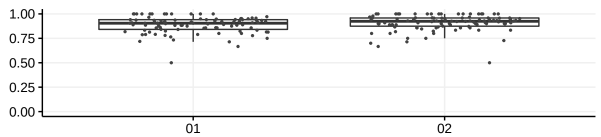










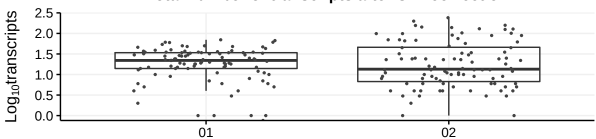


Fraction of reads aligned to an gene out of tatal number of reads

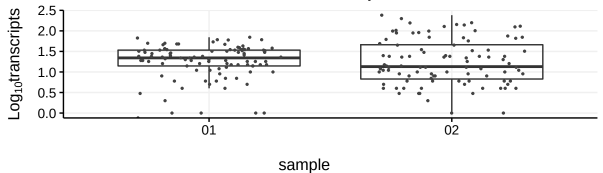


sample

Total number of transcripts after UMI correction



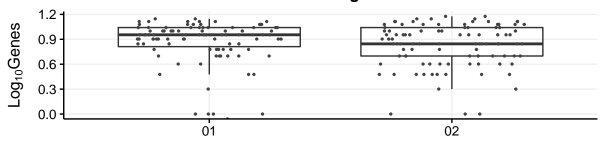
Number of mitochondrial transcripts after UMI correction







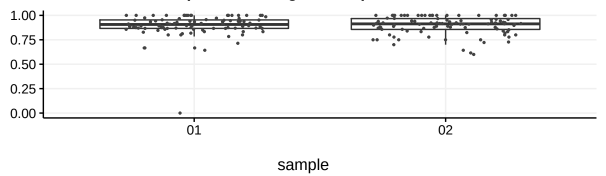
Number of genes



sample Fraction of protein coding genes



Fraction of protein coding transcripts after UMI correction





Genes detected divided by total number of reads sequenced per million



sample