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# RNA Seq Walkthrough

## Introduction

Purpose: provide a step-by-step, end-to-end RNA Seq analysis walkthrough.

#### **Environment:**

• All commands and scripts are made to run on the UCSC courtyard server, but should work in any bash environment.

OS: Red Hat 4.8.5-39
Python3: 3.6.8
R: 3.6.0

- If running on Windows try the Linux Subsystem for Windows or a Docker instance. Either of these can be used to create Linux environments on Windows (albeit with some drawbacks). I suspect any of the recent Ubuntu releases (16.04, 18.04, 20.04) should work.
- If using courtyard (or any remote server) it's **highly recommended** that you start commands in a screen environment. This is a very simple step and provides at least two substantial advantages:
  - 1. A disruption to your ssh connection will not result in termination of whatever you're running. (Execution can take many hours to days so this is a likely occurence)
  - 2. You can open multiple windows in the same terminal and switch between them, running different commands in each.

In the examples below I will always be working in a screen window, however, I will only demonstrate how to start one in section 0.0.

More on using screens.

The Data: I will be using paired-ended 151 bp Illumina sequence data starting in fastq.gz format.

## Step 0: Getting the Data

In cases where it seems to make sense I will include the generic command followed by the command I am using for actual data as an example. Otherwise I will just show the command I am using.

#### 0.1 From an SFTP Server

0. Connect to courtyard, start a screen and navigate to your directory

```
$ ssh <username>@courtyard.gi.ucsc.edu
$ ssh emalekos@courtyard.gi.ucsc.edu
# enter password at prompt
$ screen
```

```
$ cd <directory/to/transfer/data/to>
$ cd /public/groups/carpenterlab/people/emalekos/
```

1. Connecting to data storage on remote server

```
$ sftp <username>@<hostname>
$ sftp emalekos@sftp.genewiz.com
# enter password at prompt
```

2. We are now in an sftp environment (\$ -> sftp). Navigate to data folder, use 1s to list available directories

```
sftp> ls
30-422456969

# Now I navigate to the folder I found above
sftp> cd/30-422456969
sftp> ls
00_fastq

# Found another folder, I'll navigate into that.
# This time I find the data files.
sftp> cd 00_fastq/
sftp> ls -lh
```

If you are in the data folder you should see something like:

-rwxrr ? 0 0 2.86 Jan 6 14:04 1M-Ctl- M_R1_001.fastq.gz -rwxrr ? 0 0 3.06 Jan 6 14:04 1M-Ctl- M_R2_001.fastq.gz -rwxrr ? 0 0 3.06 Jan 6 14:04 1M-LPS- M_R1_001.fastq.gz -rwxrr ? 0 0 3.16 Jan 6 14:04 1M-LPS- M_R2_001.fastq.gz -rwxrr ? 0 0 3.06 Jan 6 14:04 1W-Ctl- M_R1_001.fastq.gz -rwxrr ? 0 0 3.36 Jan 6 14:13 1W-Ctl- M_R2_001.fastq.gz -rwxrr ? 0 0 3.06 Jan 6 14:13 1W-LPS- M_R1_001.fastq.gz -rwxrr ? 0 0 3.26 Jan 6 14:13 1W-LPS- M_R2_001.fastq.gz -rwxrr ? 0 0 2.16 Jan 6 14:14 2M-CSE- M_R1_001.fastq.gz -rwxrr ? 0 0 2.36 Jan 6 14:14 2M-CSE-				
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	\M_R1_001.fastq.g	Z		
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1-12-001114064192	\M_R2_001.fastq.g	Z		

1. To copy the files from this server to your workspace use the get command. Here I give some examples

```
# To copy a single file
sftp> get 1M-Ctl-AM_R1_001.fastq.gz

# To copy all files
sftp> get *

# To copy all files ending in 'fastq.gz'
sftp> get *fastq.gz

# To copy all files starting with 'LPS' somewhere in the middle
sftp> get *LPS*

# To copy all files starting with 'LPS' and with '_R1' somewhere
before the end
sftp> get LPS*_R1*
```

**NOTE:** '\*' performs wildcard expansion - it can fill in for any characters and is useful for pattern matching. When you use it it's good to check which patterns it's actually matching by double tapping the Tab key. This will list everything that matches. If you see the files you want press Enter to execute.

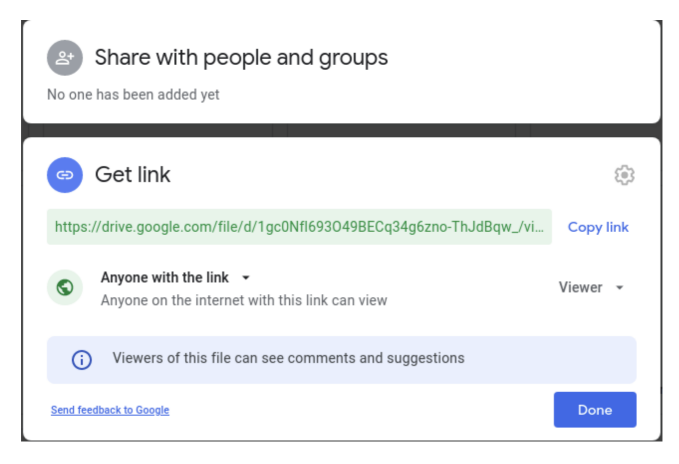
### 0.2 From Google Drive

You can use this method to transfer data files from your Google Drive account to courtyard. It works but is somewhat clunky, and there may be a better way. For instance this method transfers one file at a time, if all relevant files could be zipped together only one command would be required.

1. We will use the python package gdown. The first time you use this you will have to install it

```
pip3 install gdown --user
```

- 2. Get the share link for your Gdrive file
  - IMPORTANT make sure the the file is accessible to "Anyone with the link"
  - The relevant part of the link is the string of characters between /d/ and /view/
  - In the example below this is what we want: 1gc0Nf1693049BECq34g6zno-ThJdBqw\_



## 3. Run with python3

Now you can start a python3 session and copy the files over

```
$ python3

# Now in python shell
>>> import gdown
>>> url="https://drive.google.com/uc?
id=1gc0Nfl693049BECq34g6zno-ThJdBqw_"
>>> output="desired_filename.fastq.gz"
>>> gdown.download(url, output, quiet=False)
```

• Alternatively you can open a text editor and write a script like this one.

After changing the entries in filedict to your desired filenames and corresponding links, run with:

```
python3 file.py
```

## 0.3 Make Smaller Files for Pipeline Practice (OPTIONAL)

You may want to practice running through the pipeline with a reduced dataset. This would allow you to troubleshoot much more quickly than if you tried with all of your data at once. We can make some reduced, but still functional fastq.gz files with the following command

```
zcat <file.in> | head -n <# of lines> | gzip > <file.out>
zcat full_file.fastq.gz | head -n 100000000 | gzip > reqduced_file.gz
```

This example takes the first 100000000 lines of the input file (or the first 2500000 fastq entries). The resulting gzipped files is ~200 MB. Adjust <# of lines> as you see fit, but make it divisible by 4 to avoid cutting off fastq entries.