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# RNA Seq Walkthrough

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## 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 occurrence)
  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 **ls** 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:

```
-rwxr--r--  ? 0      0      2.8G Jan  6 14:04 1M-Ctl-
AM_R1_001.fastq.gz
-rwxr--r--  ? 0      0      3.0G Jan  6 14:04 1M-Ctl-
AM_R2_001.fastq.gz
-rwxr--r--  ? 0      0      3.0G Jan  6 14:04 1M-LPS-
AM_R1_001.fastq.gz
-rwxr--r--  ? 0      0      3.1G Jan  6 14:04 1M-LPS-
AM_R2_001.fastq.gz
-rwxr--r--  ? 0      0      3.0G Jan  6 14:04 1W-Ctl-
AM_R1_001.fastq.gz
-rwxr--r--  ? 0      0      3.3G Jan  6 14:13 1W-Ctl-
AM_R2_001.fastq.gz
-rwxr--r--  ? 0      0      3.0G Jan  6 14:13 1W-LPS-
AM_R1_001.fastq.gz
-rwxr--r--  ? 0      0      3.2G Jan  6 14:13 1W-LPS-
AM_R2_001.fastq.gz
-rwxr--r--  ? 0      0      2.1G Jan  6 14:14 2M-CSE-
AM_R1_001.fastq.gz
-rwxr--r--  ? 0      0      2.3G Jan  6 14:14 2M-CSE-
AM_R2_001.fastq.gz
```

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

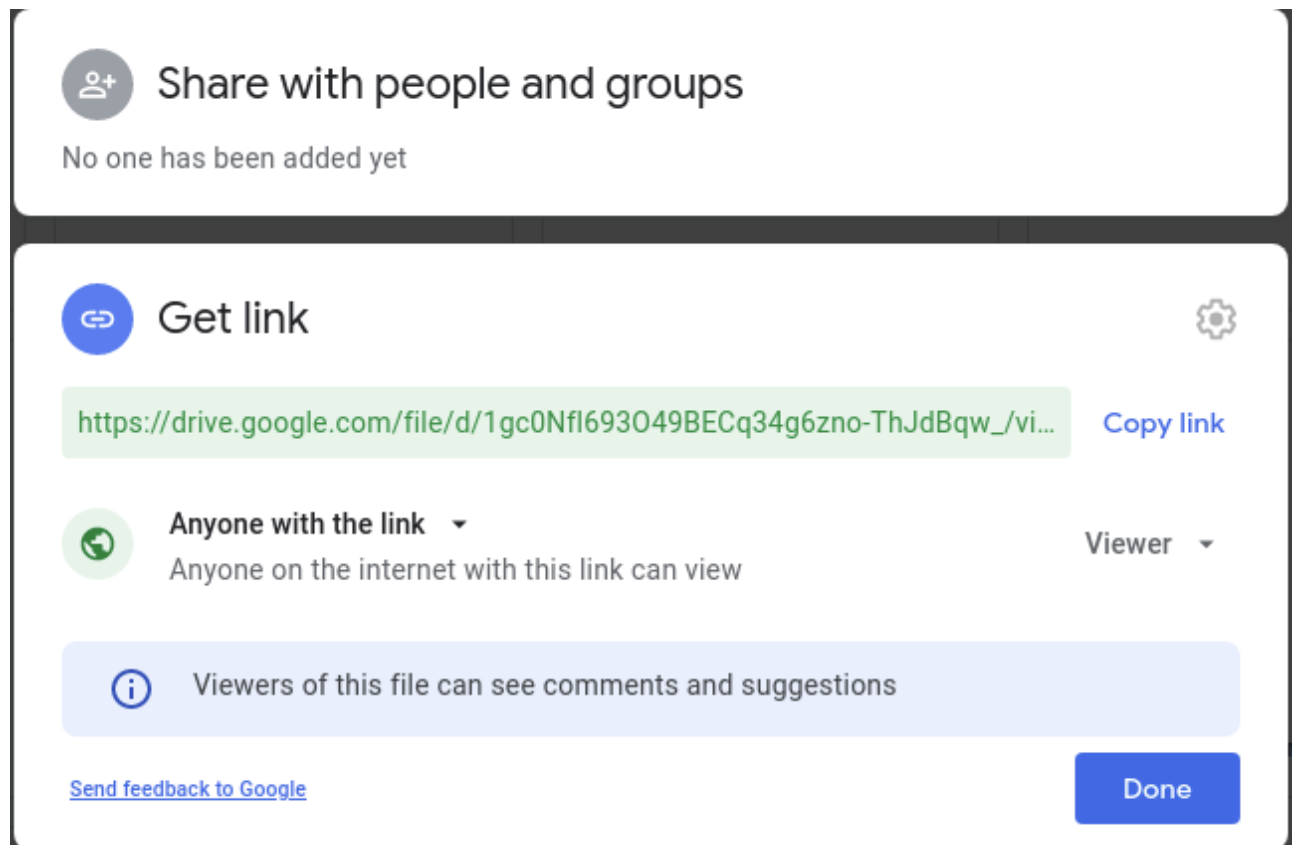
You can use this method to transfer data files from your Google Drive account to **courtyard**. It works but is somewhat clunky, and there may be a better way. For instance this method transfers one file at a time, if all relevant files could be zipped together only one command would be required.

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.

```
import gdown
url_prefix = "https://drive.google.com/uc?id="
suffix = ".fastq.gz"
filedict = {"file_1_1" : "1gc0Nfl693049BECq34g6zno-ThJdBqw_",
            "file_1_2" : "#####",
            "file_2_1" : "#####",
            "file_2_2" : "#####"}

for key, value in filedict.items():
    gdown.download(url_prefix + value, key + suffix,
quiet=False)
```

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 10000000 | gzip > reqduced_file.gz
```

This example takes the first `10000000` 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.