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# MG\_HW2: Downloading SRA data using the SRA toolkit Version

## **James Thornton**

#### **Abstract**

The Sequence Read Archive (SRA) is a database for biological sequence data and is maintained by the National Center for Biotechnology Information (NCBI). Sequence files can be obtained by using the SRA Toolkit. This protocol provides the steps necessary to use the SRA toolkit to get sequence data in fastg format.

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## **Guidelines**

**SRA Toolkit Documentation** 

## **Before start**

Login to the UA hpc. This protocol will begin in your home directory.

## **Protocol**

## Step 1.

Make sure you have /rsgrps/bh\_class/bin in your path:

If you don't, you will get an error message like this:

2016-09-12T17:15:17 prefetch.2.4.4 int: path not found while resolving tree - cannot get cache location for SRR1647046

cmd COMMAND

\$ 00

\$ nano .bashrc

export PATH=/rsgrps/bh class/bin:\$PATH

\$ source .bashrc

export PATH=/rsgrps/bh\_class/bin:\$PATH is copied into .bashrc. Then save and quit nano to source it.

## NOTES

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This step allows you to execute the executable files found in /rsgrps/bh\_class/bin. Executable files appear green on the HPC.

## Step 2.

Utilize the "prefetch" command from the SRA toolkit to get your SRA file.

#### cmd COMMAND

\$ prefetch SRR1647145

NOTE: make sure to use your SRR number

#### NOTES

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Your SRR numbers are found in the <u>google drive sheet</u> shared with the class under column 'M'. There should be a total of 8 files you need to download. See next step on how to download multiple files at once.

#### **ANNOTATIONS**

#### Amy Hudson 19 Sep 2016

I'm getting a warning after my prefetch SRR... statement:

warn: lock exists while copying file - Lock file /home/u2/amyrhudson/ncbi/public/sra/SRR1647109.sra.lock exists: download canceled

any ideas?

## Step 3.

You can pass 'prefetch' multiple arguments to download all data files at once:

#### cmd COMMAND

\$ prefetch SRR1647238 SRR1647240 SRR1647144 SRR1647260 SRR1647239 SRR1647236 SRR1647237 NOTE: make sure you use your SRR numbers .

## **P** NOTES

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Rather than copying and pasting each file name, you can use Unix to help you!

# make a file with the list of SRR files copied from the excel spread sheet

% nano list

# use the translate command to convert new lines to spaces. Note the space in the second set of quotes.

```
% tr '\n' ' ' < list
```

# then copy the line with the file names separated by space into the prefetch command, as below.

## Step 4.

The .sra files will be stored in /ncbi/public/sra

Move into that directory, then make sure all 8 files are present:

```
cmd COMMAND
$ cd ~/ncbi/public/sra
$ ls

\times EXPECTED RESULTS
```

SRR390728.sra SRR1647238.sra SRR1647240.sra SRR1647144.sra SRR1647260.sra SRR1647239.sra SRR1647236.sra SRR1647237.sra

## Step 5.

Convert the .sra file into fastq format using the fastq-dump command from the SRA toolkit. All files can be converted in one command by passing fastq-dump all files with the .sra extension.

```
cmd COMMAND
```

- \$ fastq-dump \*.sra
- \*.sra defines all files with a .sra extension NOTE: make sure you are in ~/ncbi/public/sra when you execute this command.

#### **EXPECTED RESULTS**

Read 2533849 spots for SRR1647144.sra Written 2533849 spots for SRR1647144.sra Read 3649566 spots for SRR1647145.sra Written 3649566 spots for SRR1647145.sra Read 3051288 spots for SRR1647236.sra Written 3051288 spots for SRR1647236.sra
Read 1856522 spots for SRR1647237.sra
Written 1856522 spots for SRR1647237.sra
Read 492203 spots for SRR1647238.sra
Written 492203 spots for SRR1647238.sra
Read 1191553 spots for SRR1647239.sra
Written 1191553 spots for SRR1647239.sra
Written 1191553 spots for SRR1647240.sra
Written 1527542 spots for SRR1647240.sra
Written 1527542 spots for SRR1647240.sra
Read 39872 spots for SRR1647260.sra
Written 39872 spots for SRR1647260.sra
Read 14342395 spots total
Written 14342395 spots total

#### Step 6.

Now check to see you have 8 .fastq files, 1 for each .sra file. Make a /rsgrps/bh\_class/<user>/fastq directory. Where you will replace <user> with your github id. Then move all of the all fastq files there for later use.

```
cmd COMMAND
ls
mkdir -p /rsgrps/bh_class/bhurwitz/fastq
mv *fastq !$
cd !$
ls
```

use mkdir -p to create all directories listed. In this case, we are creating bhurwitz (my user id) and the fastq directories. Note that you should use your github id here, so we can track your user id easily, and so it is consistent with your homework. Note that I am using !\$ to use the argument from the last command line.

```
EXPECTED RESULTS
```

SRR390728.fastq SRR1647238.fastq SRR1647240.fastq SRR1647144.fastq SRR1647260.fastq SRR1647239.fastq SRR1647236.fastq SRR1647237.fastq

#### Step 7.

Do a read count using seqmagick.

```
cmd COMMAND
```

\$ seqmagick info ./\*.fastq --input-format fastq > readcounts.txt
seqmagick info will generate sequence statistics for all fastq files found in the current directory
and redirect the output into a file.

#### **EXPECTED RESULTS**

```
name alignment min_len max_len avg_len num_seqs ./SRR1647144.fastq FALSE 1 300 247.30 2533849 ./SRR1647145.fastq FALSE 4 300 257.64 3649566 ./SRR1647236.fastq FALSE 1 302 254.69 3051288 ./SRR1647237.fastq FALSE 2 302 273.31 1856522
```

./SRR1647238.fastq FALSE 2 302 258.27 492203 ./SRR1647239.fastq FALSE 2 300 255.07 1191553 ./SRR1647240.fastq FALSE 4 302 270.61 1527542 ./SRR1647260.fastq FALSE 8 302 176.50 39872

#### Step 8.

Input the seqmagick 'num\_seqs' results for each sample into a summary table entitled 'Table 1'. The format should be:

## Table 1:

sample name	num_seqs
SRR1647144	2533849
SRR1647145	3649566
SRR1647236	3051288
SRR1647237	1856522
SRR1647238	492203
SRR1647239	11191553
SRR1647240	1527542
SRR1647260	39872

## Step 9.

Add metadata for each sample to your summary table. This information can be found in columns K (Occlusion\_s), Q (Age\_s), R (Sex\_s), S (Site\_Categories), T (Site\_Symbol\_s), V (TimePoint\_s), and X (Visit\_Date\_s) from the google doc. The resulting table should look like this:

#### Table 1:

sample name	num_seqs	occlusion_s	age_s	ssex_s site_categories	site_symbol_s	stimepoint_s	visit_date_s
SRR1647144	2533849	Intermittently_Occluded	24	Female Rarely_Intermittently_Moist	Ac	2	8/19/13
SRR1647145	3649566	Occluded	24	Female Moist	Ax	2	8/19/13
SRR1647236	3051288	Exposed	24	Female Sebaceous	Fh	2	8/19/13
SRR1647237	1856522	Exposed	24	Female Rarely_Intermittently_Moist	Pa	2	8/19/13
SRR1647238	492203	Occluded	24	Female Sebaceous	Ra	2	8/19/13
SRR1647239	11191553	Exposed	24	Female Sebaceous	Sc	2	8/19/13
SRR1647240	1527542	Occluded	24	Female Moist	Tw	2	8/19/13

SRR1647260 39872 Occluded 24 Female Moist Um 2 8/19/13

## NOTES

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Metadata is any information that describes the context of the sample. For example, age, sex, location, timepoint etc. is metadata.

# Step 10.

Add "Table 1" to your google doc under the tables section.