## Overview

Advances in technology have made large numbers of assembled genomes available. Submitted assemblies selected for NCBI Reference Sequence project (RefSeq) are annotated by NCBI's genome annotation pipeline and made available through the NCBI FTP site (<a href="ftp://ftp.ncbi.nlm.nih.gov/genomes/refseq/">ftp://ftp.ncbi.nlm.nih.gov/genomes/refseq/</a>). The hierarchical organization of the FTP directories, by taxonomic groups and then by species and assemblies, facilitates finding and downloading data for individual genome assemblies, but is not well suited for downloading data for all assemblies from a broad taxonomic group. In this document, we will describe a Linux shell command-based workflow that allows us to take advantage of the assembly\_summary.txt file, found in the genomes FTP site for representative taxonomic groups, to extract URLs for files or directory of interest, and use these URLs and wget to download selected sequences or all data files for these sets of genome assemblies.

## **Genomes FTP Directory Structure**

The /genomes/refseq and /genomes/genbank directories organize available data by large taxonomic groups, i.e., archaea, bacteria, fungi, invertebrate, plant, protozoa, vertebrate\_mammalian, vertebrate\_other, and viral (last is RefSeq only, details at: <a href="http://www.ncbi.nlm.nih.gov/genome/doc/ftpfaq/">http://www.ncbi.nlm.nih.gov/genome/doc/ftpfaq/</a>). Each group level directory contains an assembly\_summary.txt file with details on the latest versions of assemblies available for that group. This file also contains many fields of metadata, useful in identifying genome assemblies of interest, as well as the URLs for the subdirectories from which the data files can be downloaded. For a detailed description of the file structure, see ftp://ftp.ncbi.nlm.nih.gov/genomes/README\_assembly\_summary.txt.

NCBI organizes the genomes data files with a consistent directory hierarchy and naming convention: ACCN\_NAME/ACCN\_NAME\_X.Y.gz, where ACCN is the assembly's accession.version, NAME is the assembly name, and X.Y are sequence type and file type. Taking advantage of this consistency and information from assembly\_summary.txt for each taxonomic group, workflows below use common Linux shell utilities to process the information into direct FTP URLs for genomic sequences or full subdirectory content download.

### **Use Cases**

### Case 1: Get all the genomic sequence files for the fungal RefSeq assemblies

Under the /genomes/refseq directory of the NCBI FTP site, available data are grouped by large taxonomic groups, i.e., archaea, bacteria, fungi, invertebrate, plant, protozoa, vertebrate\_mammalian, vertebrate\_other, and viral, each with its own assembly\_summary.txt file that provides detailed information of available assemblies along with the URLs for those subdirectories in the 20th column. The workflow consists of two steps, collecting and modifying the FTP URLs for the desired file format (genomic FASTA sequences), and downloading the relevant files using the collected URLs as input.

# Step 1. Collect and modify the FTP URLs to point to the genomic.fna.gz files The command line is a pipe symbol– linked set:

The "\" is a Linux shell command to indicate that the command line continues in the next line. We use it to break the linked commands into distinctive steps so we can clearly see and discuss each sub-step:

- The first **curl** command simply gets the specified assembly\_summary.txt file and passes its content to the next step with a pipe ("|", instead of displaying it in console).
- The second command uses the **awk** utility to separate each line's content by tab (FS="\t"), skip header line (!/^#/) and print out the value of the 20th column (print \$20), and passes the output to the next step with pipe ("|").

**NOTE**: if you need the assembly submitted to GenBank, you will need to change the curl command's "refseq" to "genbank" and change sed command's "GCF" to "GCA" since their accession initials are different.

## **Use Cases (cont.)**

• The third command uses **sed** to modify the extracted URL string that points to an assembly directory to point to the "\_genomic.fna.gz" file instead. Specifically, with the pipe ("|") as delimiter, it first matches the URL into substrings using regular expression matching and captures them using parentheses (**s|(ftp://ftp.ncbi.nlm.nih.gov/genomes/all/)(GCF\_.+)|**), then reconstructs the string (\1\2) for the path and add another directory level (/) and a specific file name (\2\_genomic.fna.gz). This modifies the existing URL to point to the \_genomic.fna.gz file for that assembly. The last part (>genomic\_file) redirects the output into a file named genomic file (partially shown below).

ftp://ftp.ncbi.nlm.nih.gov/genomes/all/GCF\_000002945.1\_ASM294v2/GCF\_000002945.1\_ASM294v2\_genomic.fna.gz ftp://ftp.ncbi.nlm.nih.gov/genomes/all/GCF\_000149845.2\_SJ5/GCF\_000149845.2\_SJ5\_genomic.fna.gz

#### Step 2. Use the output from step 1 as input to wget to download

The second step is very simple. The command (below left) calls the wget utility and passes the output from step 1 as an argument to the "—input-file" switch. It will iterate through the FTP URLs and pull down those files from NCBI's FTP site to the working directory (partially listed, below right).

wget --input genomic\_file

-rw-r--r-- 1 samd sdesk 3989616 Dec 31 10:52 GCF\_000002945.1\_ASM294v2\_genomic.fna.gz -rw-r--r-- 1 samd sdesk 3492573 Dec 31 10:52 GCF\_000149845.2\_SJ5\_genomic.fna.gz

The command gunzip \*.gz will unpack them all to regenerate the FASTA files for further downstream need. For better file management, first move \*.gz files to a new directory so they are isolated from other files.

## Case 2: Get the directories and their contents for all the fungal RefSeq assemblies

If we wish to completely mirror and archive the files for this group or organisms, we can modify the above commands to download the directories along with all their contents (excluding subdirectories).

## Step 1. Collect and modify the FTP URLs to get only the directory name

curl 'ftp://ftp.ncbi.nlm.nih.gov/genomes/refseq/fungi/assembly\_summary.txt' | \ awk '{FS="\t"} !/^#/ {print \$20"/"}' > genomic\_directory ftp://ftp.ncbi.nlm.nih.gov/genomes/all/GCF\_000149845.2\_SJ5/ ftp://ftp.ncbi.nlm.nih.gov/genomes/all/GCF\_000150505.1\_SO6/

We can modify the command from Case 1 by dropping the sed command, and modify the awk slightly to get the directory URLs (left). Last two lines are example output.

## Step 2. Pass the filte to wget to pull down the directories and their contents

wget -r --no-parent --no-host-directories --cut-dirs=2 --level=1 \
--input-file=genomic directory

- --input-me-genomic\_unectory
- -r: recursively works through the directory
- --no-parent: ignores the parent directory
- --no-host-directories: saves the files without prepending the NCBI FTP URL
- --cut-dirs=2: saves the files without creating intermediate directories
- --level=1: works only at that level of directory
- --input-file=value: sets directory input to the file specified by value

We use the wget command (left) to get all the directories and their files. The command is more complex than in Case 1, so we explain the meaning of each command line arguments separately below the command. This will pull all the directories and their contents down to your Linux box, so make sure you have enough disk space to handle them.

**NOTE**: For users on PCs without Linux or cygwin access, we can do the same first steps described above in a different way. The example at the end uses inline Perl commands (below). It allows us to generate the same outputs as on Linux, and then use the same wget commands for the second steps to download. The PC port of wget is available from: <a href="https://eternallybored.org/misc/wget/">https://eternallybored.org/misc/wget/</a>

perl -e "use LWP::Simple;  $file=get(\t ftp.-/ftp.ncbi.nlm.nih.gov/genomes/refseq/fungi/assembly_summary.txt\"); while (<math>file= \t ftp.+)(GCF.+?)\$  {print \$1, \$2, \\\, \$2, \\\_genomic.fna.gz\n\\";}\" > fungi\_genomic\_files perl -e "use LWP::Simple;  $file=get(\t ftp.-/ftp.ncbi.nlm.nih.gov/genomes/refseq/fungi/assembly_summary.txt\"); while (<math>file= \t ftp.-+GCF.+?$ )\s/g){print \$1, \\\\n\\";}\" > fungi\_directory

Warning: Type all example commands instead of copying/pasting since hidden characters may break their execution.