Segmenting Large Electron Microscopic Image Volumes

An Introduction to NBCR image analysis and segmentation tools



National Biomedical Computation Resource Summer Training Program @ UC San Diego

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Hands-On Session 3

Running CHM Train via chmutil on Rocce Cluster

Goals

By the end of this session, you will be able to:

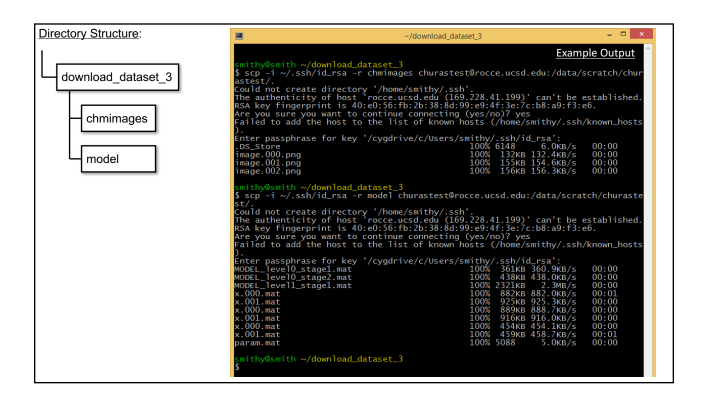
- 1. Create and run CHM Job
- 2. Download data from Rocce

1. Upload data to Rocce

- A. Download the <code>download_dataset_3.zip</code> file from https://github.com/CRBS/nbcrtrainingvm/blob/master/download_dataset_3.zip and unzip and in a terminal enter it as before.
- B. The directory will contain two items:
 - i. A directory, chmimages, which contains EM data that we will be attempting to segment with CHM
 - ii. A directory, *model*, which contains a CHM trained model generated by following steps in previous hands on session 2.
- C. Upload the *chmimages* and *model* directory to Rocce using the commands below:

```
scp - i \sim /.ssh/id\_rsa - r \ chmimages < USER > @rocce.ucsd.edu:/data/scratch/< USER > /.
```

scp -i ~/.ssh/id rsa -r model<USER>@rocce.ucsd.edu:/data/scratch/<USER>/.



2. Create CHM job

We will be running the chmutil command *createchmjob.py* to create <u>CHM</u> job on the Rocce cluster. Here is a description of the arguments used in the command invocation below

--chmbin Path to <u>CHM Singularity</u> binary (use path as shown below)

--cluster Tells tool what cluster is being used. (currently Rocce & Comet are supported)

--disablechmhisteq Disables histogram equalization of images by CHM

 Connect to Rocce, change to /data/scratch/<USER> directory, and run createchmjob.py with arguments as shown below

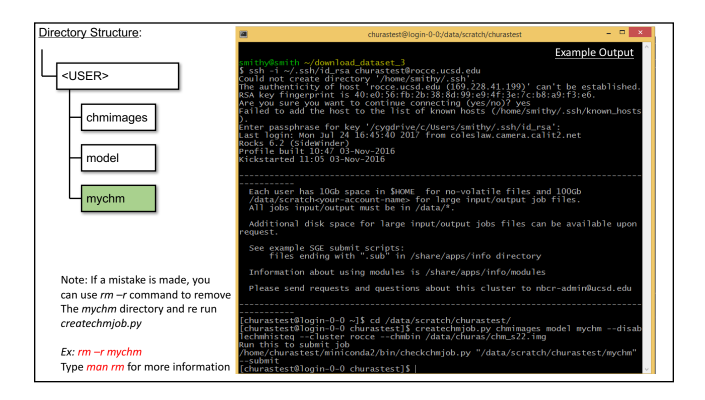
ssh –i ~/.ssh/id_rsa <USER>@rocce.ucsd.edu cd /data/scratch/<USER>

createchmjob.py chmimages model mychm --disablechmhisteq --cluster rocce --chmbin /data/churas/chm s22.img

The above command will create a new directory named mychm.

If your train job from previous the hands on ran successfully then you can use it instead by replacing the directory path *model* with *mytrain/model*. The command will also output a command to run

For more information on how to use *createchmjob.py* just run the command with no arguments or with –*h* flag (Example: *createchmjob.py* –*help*) https://github.com/CRBS/chmutil/wiki/createchmjob.py



3a. Processing loop

createchmjob.py will output a command to run. This starts the processing loop which comprises of alternating invocations of checkchmjob.py & gsub, for job submission, and gstat to see if jobs completed.

For more information visit: https://github.com/CRBS/chmutil/wiki/createchmjob.py

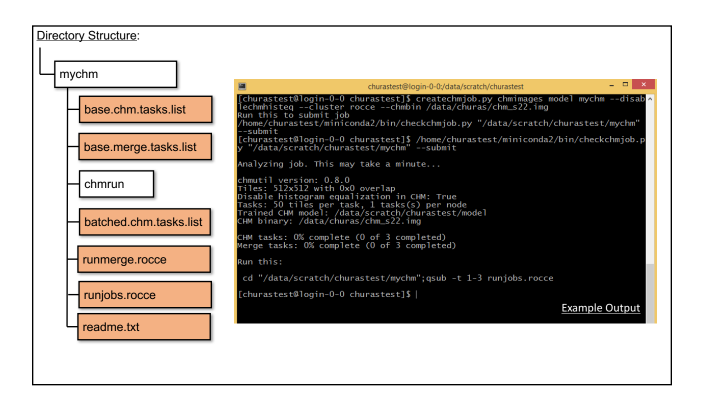
A. Lets start the processing loop by running the command output by *createchmjob.py* from the previous step which will look like the following:

/home/<USER>/miniconda2/bin/checkchmjob.py "/data/scratch/<USER>/mychm" --submit

checkchmjob.py outputs in human readable form a current completion status of the job

If –submit is passed the script will create/update batched.chm.tasks.list and batched.merge.chm.tasks.list files (described later) and output a *qsub* command to run the jobs via SGE

NOTE: The --submit flag should be omitted if any jobs are running (job is visible in *qstat*) otherwise errors will occur.



3b. Processing loop

A. When run successfully, *checkchmjob.py* – *submit* outputs a command to submit the jobs.

In the terminal copy the text line below "Run this:" and paste it in the terminal to submit the jobs.

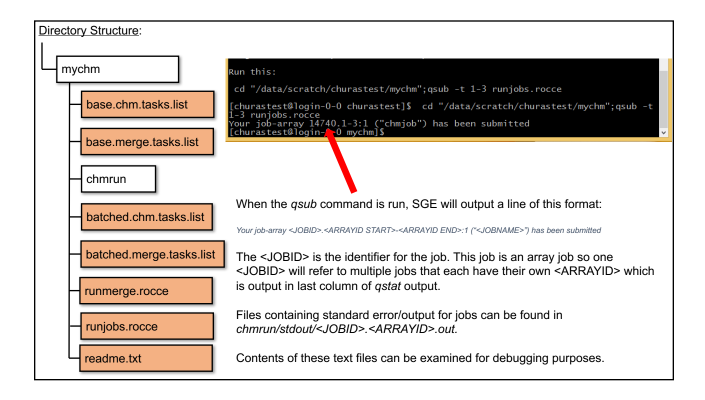
In the first invocation of *checkchmjob.py* the text to copy will look similar to the text below where we will be submitting 3 jobs using the array job feature of SGE via –*t* 1-3 option:

cd "/data/scratch/<USER>/mychm"; qsub -t 1-3 runjobs.rocce

The above command will put the terminal into mychm directory and invoke qsub to submit jobs to SGE.

The ; character lets a user put multiple commands on a single line.

Run man qsub for more information



3c. Processing loop

Completing the CHM job is now a repeat of the following operations.

1. Run *qstat* periodically and verify jobs have completed (jobs with ID should have 'c' complete state or not be listed)

astat

2. Only **AFTER** *qstat* shows jobs have finished. Run *checkchmjob.py* . *--submit* and the *qsub* command *checkchmjob.py* outputs to submit any remaining jobs. Once on a compute node CHM should only take about 3 minutes to run and the merge is just a few seconds

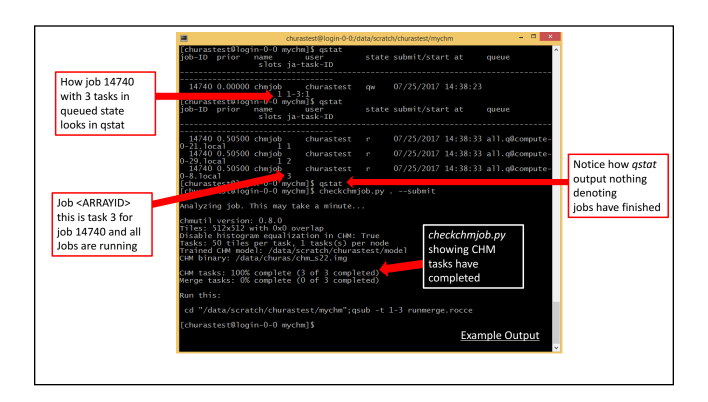
WARNING: Only add --submit to checkchmjob.py if all jobs have completed according to qstat, otherwise any running jobs will be run twice or will fail. It is safe to invoke checkchmjob.py without -submit anytime

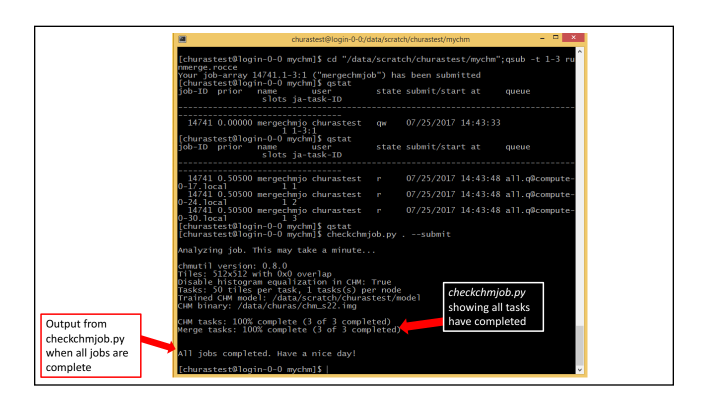
checkchmjob.py . --submit

Don't forget to run the qsub command output by checkchmjob.py

checkchmjob.py takes a directory path to job as its first argument. Since we are in mychm directory a period can be used which denotes current working directory.

For more information go to: https://github.com/CRBS/chmutil/wiki/createchmjob.py





4. Understand CHM job structure

Running createchmjob.py & checkchmjob.py --submit creates mychm directory with several files. Run Is to see a listing of files and directories.

Below is a description of the files and directories seen from the Is command

base.chm.tasks.list Main configuration file and contains CHM tasks to run

batched.chm.tasks.list Configuration file that defines how CHM tasks in base.chm.tasks.list are

batched onto individual compute nodes on the cluster

batched.merge.tasks.list Configuration file that defines how Merge tasks in base.merge.tasks.list are

batched onto individual compute nodes on the cluster

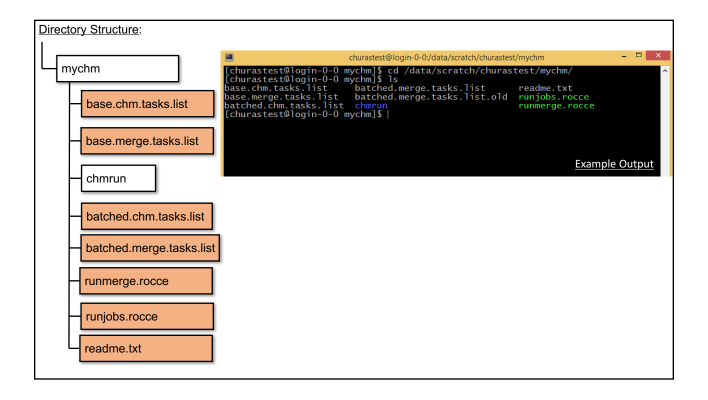
base.merge.tasks.list Configuration file for Merge phase and contains merge tasks to run

chmrun/ Directory under which all job output is written. There are several directories

within described later in this exercise

readme.txt Contains information about this job including arguments used and how to get

SGE qsub submit script to run CHM tasks on Rocce cluster runjobs.rocce runmerge.rocce SGE qsub submit script to run Merge tasks on Rocce cluster



4. Understand CHM job output structure

A. Run Is on chmrun/ to see the files and directories of CHM job output

Is chmrun/probmaps

Below is a description of the files and directories seen from the Is command

mergestdout/ Directory containing output from merge tasks. Merge tasks are directed to write to this path

via runmerge.rocce queue submit script file.

probmaps/ Directory containing final output of merged probability maps. These images are created

when the merge tasks are run.

stdout/ Directory containing output from CHM tasks. CHM tasks are directed to write to this path

via runjobs.rocce queue submit script file.

tiles/ Directory containing directories for each input image. Within each of this input image

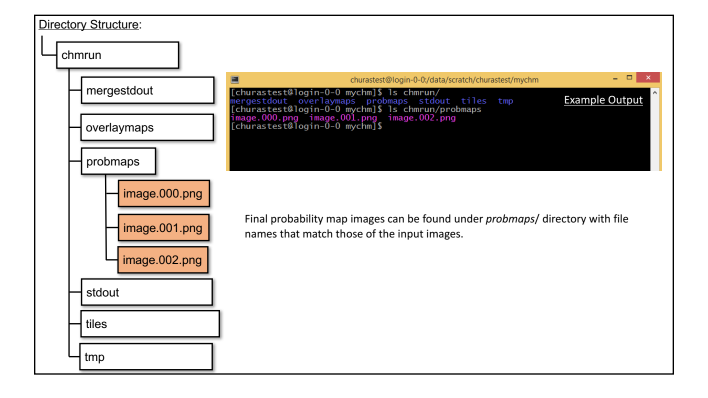
directories the CHM tasks will write partial probability maps which will be merged by merge

tasks in the SECOND phase of processing.

tmp/ Directory containing temp standard out and standard error files for chmrunner.py and

mergetilerunner.py scripts. Once the task completes this directory and files will be

removed.



9. Download data from Rocce

The merged probability map images are located in *chmrun/probmaps/<image name>* The commands below exit the terminal from Rocce cluster and use *scp* to copy back the *probmaps* directory and images within

exit
cd ~/download_dataset_3
scp -i ~/.ssh/id_rsa -r <USER>@rocce.ucsd.edu:/data/scratch/<USER>/mychm/chmrun/probmaps .

Don't forget to include the period above at the end.

For more information go to: https://github.com/CRBS/chmutil/wiki/createchmjob.py

