# 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 2**

Running CHM Train via chmutil on Rocce Cluster

## Goals

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

- 1. Install chmutil on Rocce Cluster
- 2. Upload training data to Rocce Cluster
- 3. Autocontrast training images
- 4. Create CHM Train job
- 5. Submit CHM Train job
- 6. Check on CHM Train job status

## 1. Install chmutil on Rocce Cluster

A. Connect to Rocce Cluster via *ssh* by opening a terminal and invoking this command:

ssh –i ~/.ssh/id\_rsa <USER>@rocce.ucsd.edu

#### 1. Install chmutil on Rocce Cluster

These instructions are taken from <a href="https://github.com/CRBS/chmutil/wiki/chmutil-on-rocce-cluster">https://github.com/CRBS/chmutil/wiki/chmutil-on-rocce-cluster</a>

A. Run these two commands to download and install Miniconda

```
qrsh
wget https://repo.continuum.io/miniconda/Miniconda2-latest-Linux-x86_64.sh
bash Miniconda2-latest-Linux-x86_64.sh –b
```

B. Run these three commands to update startup scripts with the path to Miniconda

```
echo 'export PATH="$HOME/miniconda2/bin:$HOME/.local/bin:$PATH"' >> \sim/.bash_profile echo 'export LC_ALL=C' >> \sim/.bash_profile source \sim/.bash_profile
```

C.

Run these two commands to install chmutil and verify installation by calling *createchmjob.py* 

pip install chmutil createchmjob.py –version exit

Note: The above commands only have to be run once

## 2. Upload training data to Rocce

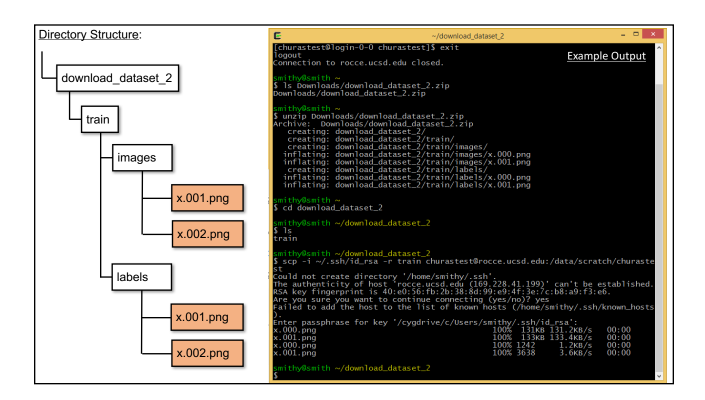
A. If still connected, disconnect from Rocce by typing exit

exit

- B. **Either** download the *download\_dataset\_2.zip* file from the BBDTC portal and unzip and enter it as before **OR** if you want, enter *download\_dataset\_1* directory where training data created from previous exercise.
- C. The directory will contain one item (more if you entered download dataset 1 folder):
  - A directory, train, which contains two directories images and labels that contain training data created in previous exercise.
- D. The training data is already in correct format, so we just need to upload it to Rocce:

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

This will recursively upload the *train* directory to /data/scratch/<USER> directory on Rocce. Note: Be sure to replace <USER> above with your Rocce username.



## 3. Autocontrast training images

When displaying images IMOD auto contrasts the data, to Train CHM it helps to apply a similar auto contrast on the images. The commands below do just that with the chmutil command *createchmimage.py* <a href="https://github.com/CRBS/chmutil/wiki/createchmimage.py">https://github.com/CRBS/chmutil/wiki/createchmimage.py</a>

A. Connect to Rocce Cluster via ssh, start an interactive job, & change directory to /data/scratch/<USER>/train/images Be sure to replace <USER> with you Rocce username.

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

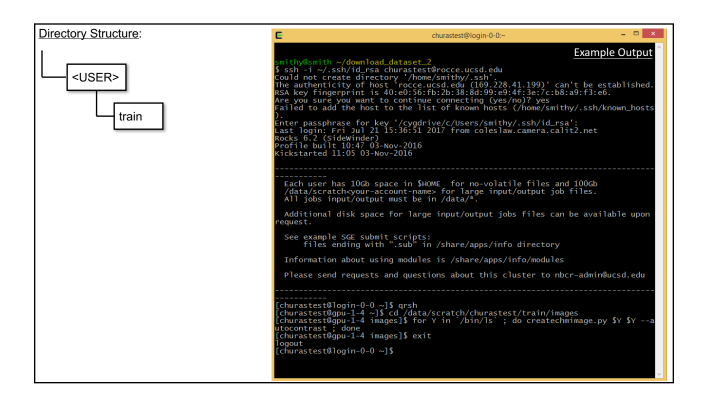
B. Use a bash for loop & chmutil command createchmimage.py to auto contrast images

for Y in `/bin/ls`; do createchmimage.py \$Y \$Y -autocontrast; done

Note: The `character is a back tick (it is usually above the tab key on keyboard)

C. Exit from interactive job

exit



#### 4. Create CHM Train job

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

--chmbin Path to CHM Singularity binary (use path as shown below)

--cluster Tells tool what cluster to generate job for. (currently Rocce & Comet are supported)

--stage CHM parameter, sets number of stages (default 2)
--level CHM parameter, sets number of levels (default 4)

--maxmem Lets caller define amount of ram in gigabytes needed by job (default 90)

A. Change to /data/scratch/<USER> and run createchmtrainjob.py with arguments as shown below

cd /data/scratch/<USER>

createchmtrainjob.py train/images train/labels mytrain –chmbin /data/churas/chm\_s22.img –cluster rocce – stage 2 –level 1 –maxmem 12

The above command will create a new directory named mytrain.

For more information on how to use *createchmtrainjob.py* just run the command with no arguments or with —h flag (Example: *createchmtrainjob.py*—help)

https://github.com/CRBS/chmutil/wiki/createchmtrainjob.py



## 5. Submit CHM Train job

Running *createchmtrainjob.py* from previous step creates a *mytrain* folder with a couple files and directories described here:

stdout/ Directory that will contain standard error and output file for training job. The file will be

named <JOBID>.out

tmp/ Directory that will temporarily hold output from CHM Train. As the job finishes this directory

will be renamed to model/

readme.txt Text file with information about job (parameters passed to createchmtrainjob.py) as well as

information on how to check on job status and get more help

runtrain.rocce Generated script to run CHM Train job on Rocce cluster. This file is passed to SGE qsub

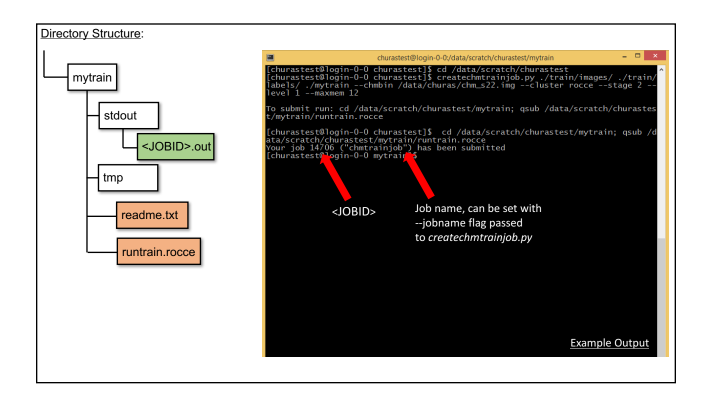
command

A. When run successfully createchmtrainjob.py outputs a command to run to actually submit the job.

In the terminal copy the text to the right of "To submit run:" and paste it in the terminal to submit the job.

The text to copy will look similar to the text below:

cd /data/scratch/<USER>/mytrain; qsub /data/scratch/<USER>/runtrain.rocce



## 6. Check on CHM Train job status

When you run the *qsub* command as done in previous step, SGE will output a line of this format:

Your job <JOBID> ("<JOBNAME>") has been submitted

The <JOBID> is the identifier for the job and is used as the file name for the file containing standard error/output of job which is stored in *stdout/<JOBID>.out*. Contents of this text file can be examined for debugging purposes.

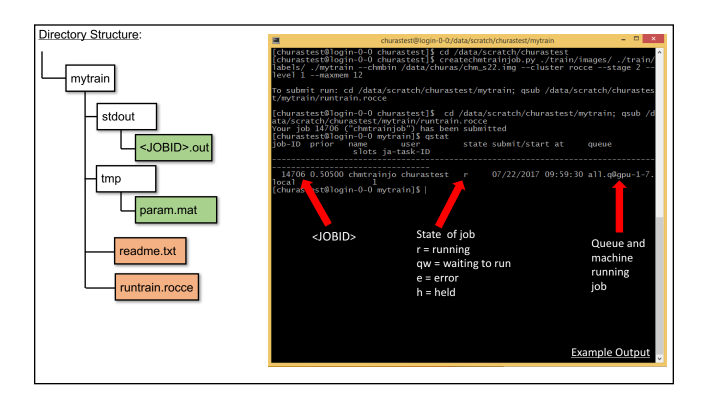
A. To check on job status run *qstat* command that is a SGE command that outputs all jobs your account is running

#### qstat

qstat will output jobs, one per row with current status.

If the job has finished or failed it will **NOT** be displayed in output.

Run man qstat for more information



## 6. Check on CHM Train job status

When *qstat* no longer displays the job, it is best to check the *stdout<JOBID>.out* file to verify correct operation.

This can be done with tail command which lets one view text files.

Is stdout tail stdout/<JOBID>.out

A successful job will have the following on the last line:

chm s22.img exited with code: 0

CHM Train writes output to tmp directory and when job completes the script renames the directory to *model*. It is a good idea to check for valid output in this *model* directory using the *Is* command

#### Is model

A successful run will have these files plus three output level.. directories:

 $param.mat,\ MODEL\_level0\_stage1.mat,\ MODEL\_level0\_stage2.mat,\ MODEL\_level1\_stage1.mat$  For more information go to:  $\frac{https://github.com/CRBS/chmutil/wiki/createchmtrainjob.py}{https://github.com/CRBS/chmutil/wiki/createchmtrainjob.py}$ 

