## Calculation of Q-scores from Command Line

First, download and install UCSF Chimera and the MapQ plugin: https://github.com/gregdp/mapq

Then, Q-scores can be calculated from the command line using a command which varies slightly by platform, and whether to use multiprocessing:

Using only a single processor:

### Mac Command:

[path to Chimera]/Contents/MacOS/chimera --nogui --silent --nostatus [path to map] [path to model] [path to Chimera]/Contents/Resources/share/mapq/mapq run.py

### Mac Example:

```
~/Desktop/Chimera.app/Contents/MacOS/chimera --nogui --silent --nostatus ~/_data/mmetrics/emd_20026/emd_20026.map ~/_data/mmetrics/emd_20026/3ajo.pdb ~/Desktop/Chimera.app/Contents/Resources/share/mapg/mapg run.py
```

#### Unix Command:

```
[path to Chimera]/bin/chimera --nogui --silent --nostatus [path to map] [path to model] [path to Chimera]/share/mapq/mapq_run.py
```

### Unix Example:

```
~/_mol/Chimera/bin/chimera --nogui --silent --nostatus ~/Dropbox/_data/Ferritin/k175/k175_fi.mrc ~/Dropbox/_data/Ferritin/k175/3ajo_f.pdb ~/_mol/Chimera/share/mapq/mapq_run.py
```

Using multi-processing (the number of processors/cores will be automatically detected and used – this may be set as an option in the future):

#### Mac Command:

[path to Chimera]/Contents/MacOS/chimera --nogui --silent --nostatus [path to map] [path to model] [path to Chimera]/Contents/Resources/share/mapq/mapq\_run\_MP.py [number of processors]

### Mac Example:

```
~/Desktop/Chimera.app/Contents/MacOS/chimera --nogui --silent --nostatus ~/_data/mmetrics/emd_20026/emd_20026.map ~/_data/mmetrics/emd_20026/3ajo.pdb ~/Desktop/Chimera.app/Contents/Resources/share/mapq/mapq_run_MP.py 6
```

# Unix Command:

```
[path to Chimera]/bin/chimera --nogui --silent --nostatus [path to map] [path to model] [path to Chimera]/share/mapq/mapq_run_MP.py [number of processors]
```

#### Unix Example:

```
~/_mol/Chimera/bin/chimera --nogui --silent --nostatus ~/Dropbox/_data/Ferritin/k175/k175_fi.mrc ~/Dropbox/_data/Ferritin/k175/3ajo_f.pdb ~/_mol/Chimera/share/mapq_run_MP.py 6
```

• Please note that [path to model] is required when using the multi-processing script, and should be an absolute path, even if the script is run from the directory where the model is located. This is so that per-atom Q-scores can be written to a PDB file in the same folder (see below).

The output should be something like this (will vary slightly depending on whether multi-processing was used or not):

```
Q Scores:
 - map: emd_20026.map
 - mol: 3ajo.pdb, chain: _all_
 - sigma: 0.50
 - mind: -0.061, maxd: 0.109
 - atoms to do: 1714
 - path: /Users/greg/Desktop/Chimera.app/Contents/Resources/share/mapqp.py
 - on mac
[...]
 - done, time: 70.172916
 - done, time: 1 min, 10.2 sec
Saving pdb with Q-scores: /Users/greg/_data/mmetrics/emd_20026/3ajo__Q_emd_20026.pdb
O for 413 res...
Model Q-score: 0.776
                       Est.Res.(A)
Chain Avg.Q-score
chain A (hoh) 0.506 3.45
chain A (mg) 0.510 3.42
chain A (protein)
                       0.820 1.69
      Avg.Q-score
                        Est.Res.(A)
Protein 0.820
MG 0.510 3.42
HOH 0.506 3.45
                       1.69
Avg. Q score: 0.776
```

- The score for the entire model is on the third last line (0.776).
- The other few last lines summarize the scores by atom type: protein, nucleic acids, and other, e.g. Mg, Hoh (water atoms).
- Per-atom Q-scores are stored in the pdb file [model name] Q [map name] in the B-factor column.