MapQ Tutorial

Last Updated: Aug. 21, 2019

Greg Pintilie gregp@slac.stanford.edu

Overview

- MapQ is a UCSF Chimera plugin that allows calculation and visualization of Q-scores, which measure resolvability of atomic features in cryoEM maps using a fitted model.
- This tutorial covers the following topics:
 - 1. Get the software (UCSF Chimera and the MapQ plugin)
 - 2. Getting an example map and model
 - Feel free to use your own map and model here
 - 3. Use the MapQ interface to calculate and visualize Q-scores for amino acid residues and solvent atoms.
- The plugin has been tested on MacOS (10.13.16), Linux (Ubuntu 14.04), and Windows 10 with Chimera 1.12, 1.13 and 1.14.

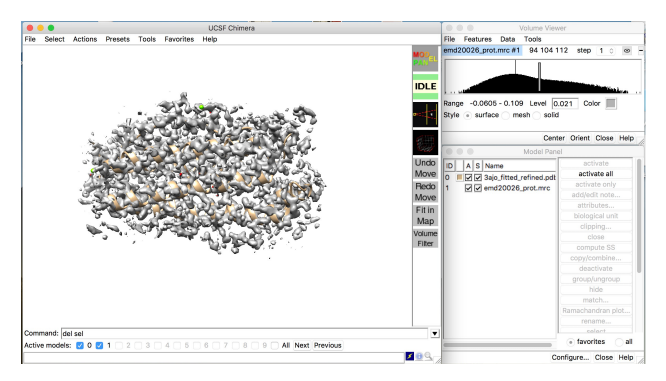
1. Getting the Software

- Download Chimera for your system from
 - o https://www.cgl.ucsf.edu/chimera/download.html
 - o Either production or daily release should work.
- Download and install the MapQ plugin
 - o https://github.com/gregdp/mapq
 - o Follow the Install instructions at the above link.

2. Loading the map and model

- Download the map and model from the following link:
 - o https://github.com/gregdp/mapq/blob/master/data/QScore_Apoferritin_Tutorial.zip
 - Once downloaded, unzip the file. It should contain two files, a map and a model:
 - Emd20026_prot.mrc this is a portion of the entire Apoferritin map representing a single protein and solvent atoms around it.
 - 3ajo_fitted_refined.pdb this is the X-ray model PDB:3ajo which was first rigidly fitted to the cryoEM map, and then refined using Phenix.real space refine
- Open Chimera
- Load the map:

- o In Chimera, from File menu, select Open, navigate and select the map (emd20026 prot.mrc), click Open
- Load the model:
 - In Chimera, from File menu, select Open, navigate and select the model
 (3ajo fitted refined.pdb), click Open
- You should see something like this in the Chimera window, and two dialogs should pop up, 'Model Panel, and 'Volume Viewer'
 - o If these dialogs go away, they can be opened again from the top menu:
 - Tools -> General Controls -> Model Panel
 - Tools -> Volume Data -> Volume Viewer



Volume Viewer

- o This shows the Cryo-EM maps that are open (emd_20026_prot.map).
- o For efficiency, the 'step' may be "2" initially, however this shows less detail. Select 1 to see the full detail.
- The map is shown in the main window (titled UCSF Chimera) as an 'Iso-surface', i.e. a surface that has the same map value at all the points on it. The range and histogram of the values in the map are shown in the Volume Viewer dialog.
- O Drag the bar on the histogram from left to right (with the mouse left button), and see what happens to the surface shown in the main window.
- You can click the 'eye' icon to show/hide the map in the main window (useful when dealing with multiple maps).

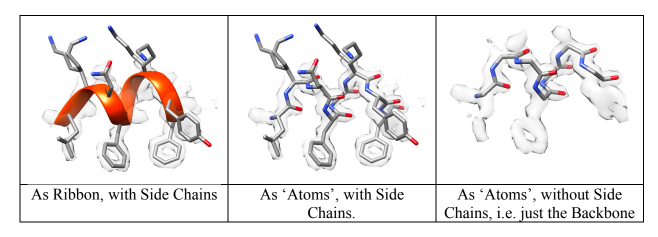
3. Using MapQ to calculate and visualize Q-scores

• For this part, we will use the 'MapQ' tools/plugin

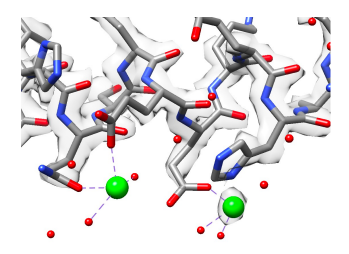
- Open ModelZ dialog:
 - o Tools menu -> Volume Data -> MapQ
- The ModelZ Dialog looks like this:



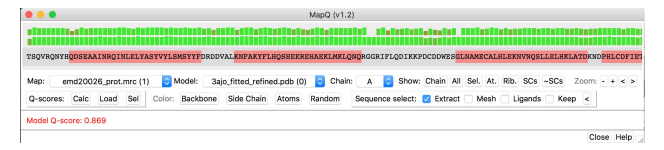
- A map, model, and chain are already selected.
 - o If not, or to select a new map/model/chain, click on the respective fields just under the sequence.
- The sequence for the selected chain is shown
 - o The sequence is made up of amino acid residues (e.g. Tyrosine, T, Serine, S, Glutamine, Q, etc.)
- Residues that are in 'loops' are shown with gray background.
- Residues that are in 'helices' are shown with reddish background.
- Select an entire loop/helix (Ctrl+Click+Drag on the sequence).
 - o This selects just that part of the protein, and,
 - o The map around the selection will be extracted and shown.
- With each part selected, click 'At.', 'Rib.', 'SCs', '~SCs' to show/switch between seeing Atoms, Ribbon, Side Chains, No-Chains. This will show the protein in different ways, as illustrated below.



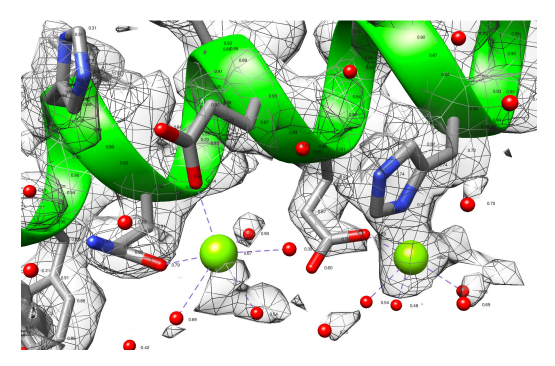
• Make sure 'Ligands' is checked in the MapQ dialog (to the right of 'Sequence Select:'). Then, select the entire second helix. The selection and extracted map will now include nearby 'ligand' atoms, which in this case are water and Mg atoms as below:



- Note how at lower thresholds you can see contours around some of the Mg and water (O) atoms. These atoms were placed in the X-ray model, and are also resolved in the cryoEM map.
- To calculate Q-scores, you can press either:
 - o "Calc" button
 - This will calculate Q-scores for all atoms in the selected model and chain. Select 'All' in the Chain field to calculate for all atoms in the model.
 - For this protein, it will take about 4 min to calculate the Q-scores for each atom. You can see the progress and ETA at the bottom of the main Chimera window. To cancel, you can press the X button.
 - o "Load" button
 - Once the Q-scores are calculated once, a new file will appear in the folder of the map, which is the same as the original model, but will have Q-scores stored in the B-factor column.
 - Pressing this button will look for this file and load it instead of calculating Q-scores again.
 - o "Sel" button
 - This will calculate Q-scores only for atoms that are selected (with Ctrl+Left Click – press the up arrow to expand selection to residue, secondary structure, or entire chain).
 - It typically takes just a few seconds to calculate the Q-score for each atom, so this is faster than calculating Q-scores for an entire model.
- After calculating Q-scores using the **Calc** button (or loading them with the **Load** button), you will see the area above the sequence show a bar graph of Q-scores for each residue.
 - On the bottom are backbone Q-scores, on the top are side chain Q-scores:



- Because this map is very good, you will see most residues in this case have good Q-scores the bars for these are green.
 - A couple have lower (red). These tend to be residues which are more dynamic (e.g. Asp, Glu), or portions of loops.
- To visualize Q-scores on the model, you can press a button to the right of the Color: label:
 - Backbone colors the model ribbon by backbone score (green is high, red is lower Q-score).
 - Side Chain colors ribbon by side chain score.
 - o Atoms shows a label with the Q-score next to each atom, as below
 - o Random colors each chain a random color



- If you select the '**Keep**' checkbox, when selecting a new part of the sequence, the previous selection is also included in what is shown.
 - o So for example, you can extract/show two parts of the model at the same time.
 - Press the '<' button to undo the last selection (helpful if the new selection was not what you were looking for).

 Using this mode, we can show, for example, the GLU 27 residue which is close to the Mg on the right, while keeping the previous selection:

