

Chemical Energy Wise Decomposition (CHEWD)

CHEWD plugin can be downloaded from

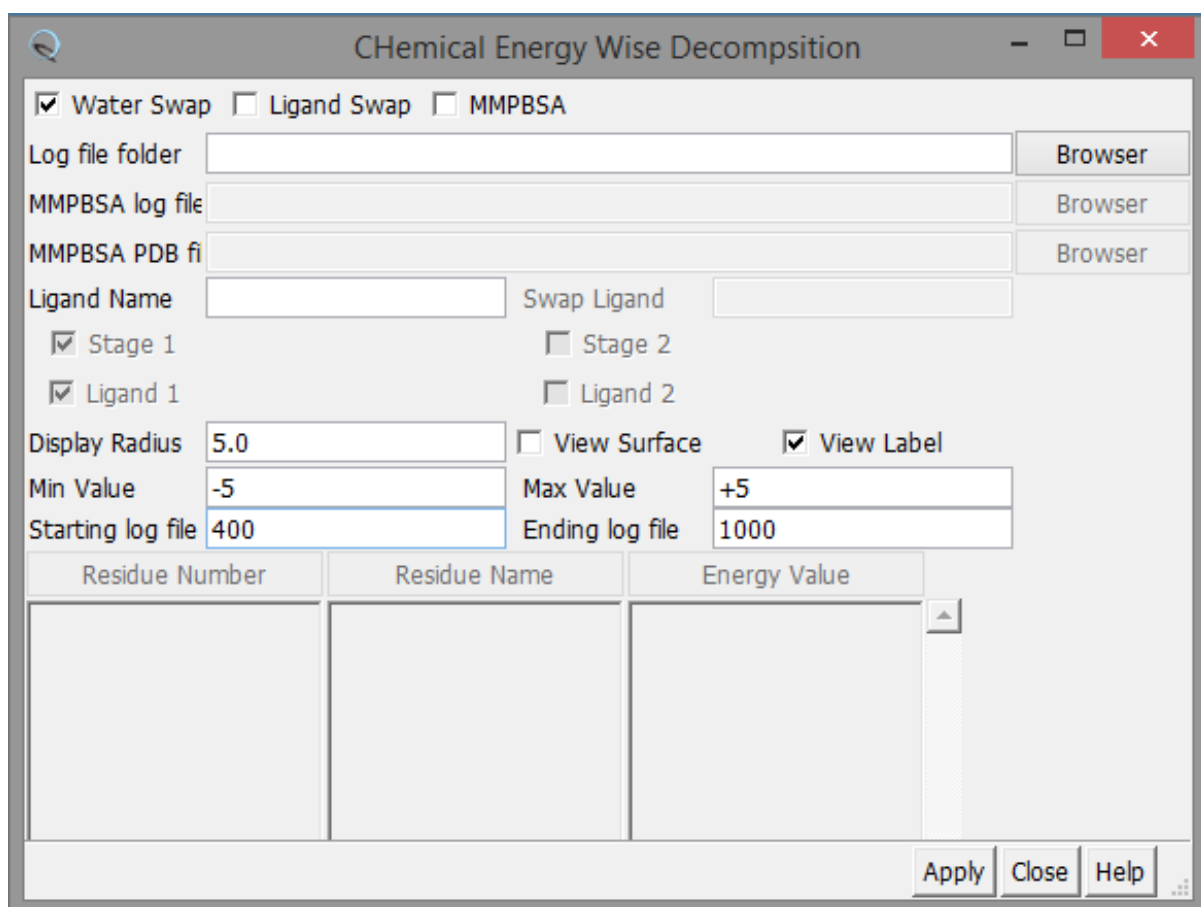
<https://github.com/saadraza128/CHEWD>

For installation:

```
cd $CHIMERA/share
```

```
tar -zxvf /path/to/EnergyVisualizer.tgz
```

The plugin is located in the menu Tools | Utilities | Sire Energy Visualizer. Clicking on this option will open the below GUI dialog:

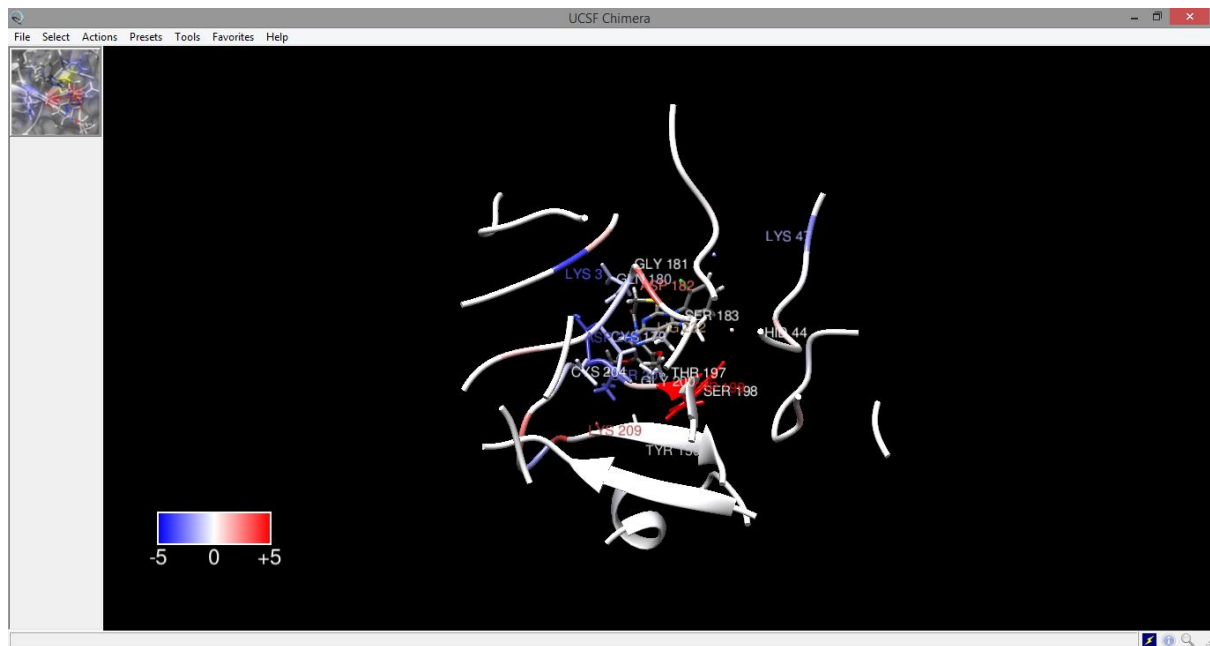


Next, make sure that the Water Swap box is checked at the top of the dialog. This puts the plugin into Waterswap mode. Next, set the Ligand Name according to your topology file (In this example we set it to LIG).

[illegible]

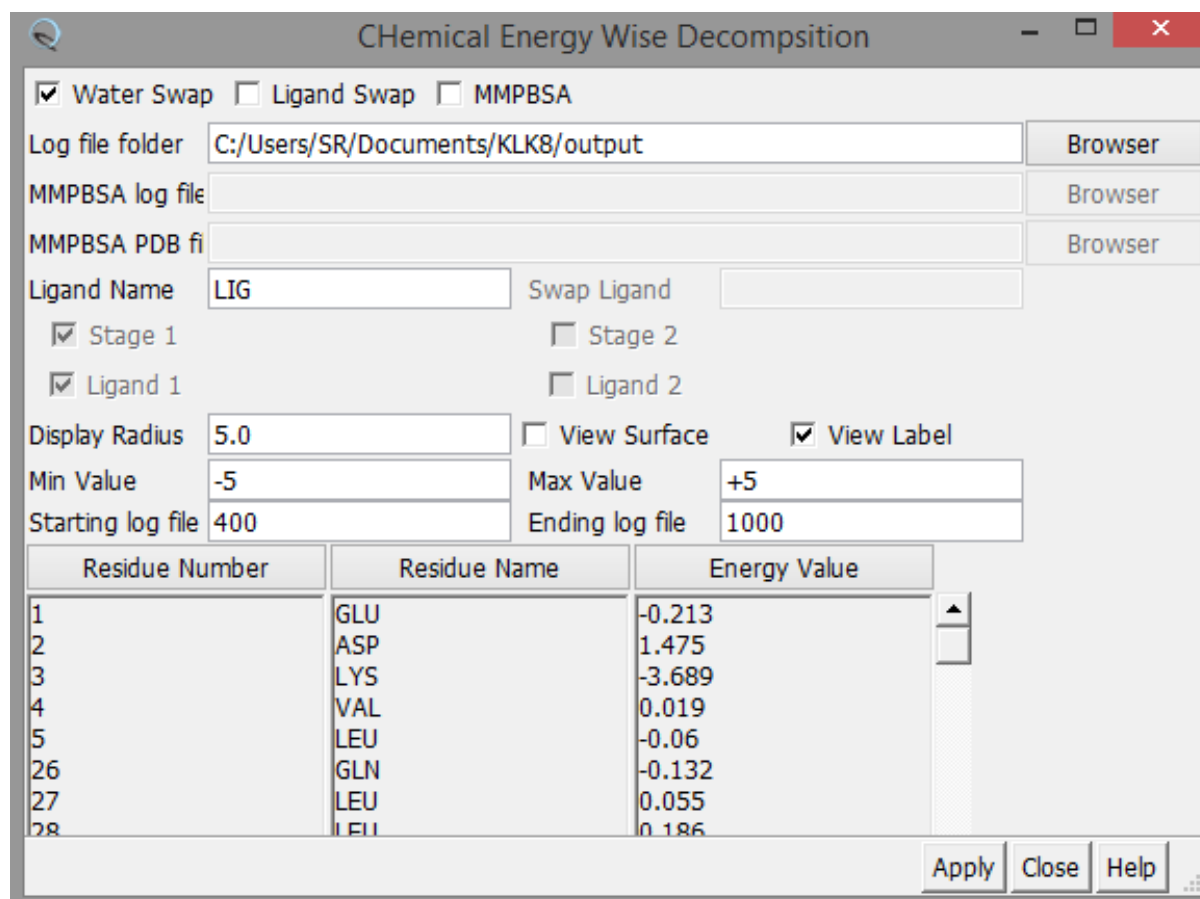
Now click Apply. This will take a while to load up all of the output.

Once loaded, you should see that the main chimera window looks something like this;



This shows a cutout of the full protein-ligand-water system, showing only the protein and ligand atoms that were mobile during the waterswap calculation (everything within 15 angstroms of the swapped ligand). The residues are colour-coded according to their preference for either ligand or water. Red shows a preference for water, with stronger colours indicating a stronger preference. Blue shows a preference for ligand, again with a stronger colour implying a stronger preference.

The exact values of these free energy component preferences are shown in the Sire Energy Visualizer dialog, which looks something like this;



Chemical Energy Wise Decomposition

☒ Water Swap ☐ Ligand Swap ☐ MMPBSA

Log file folder:

MMPBSA log file:

MMPBSA PDB file:

Ligand Name: Swap Ligand:

☒ Stage 1 ☐ Stage 2

☒ Ligand 1 ☐ Ligand 2

Display Radius: ☐ View Surface ☒ View Label

Min Value: Max Value:

Starting log file: Ending log file:

Residue Number	Residue Name	Energy Value
1	GLU	-0.213
2	ASP	1.475
3	LYS	-3.689
4	VAL	0.019
5	LEU	-0.06
26	GLN	-0.132
27	LEU	0.055
28	LEU	0.186