

Umbrella sampling / wham protocol

1. Run `ir-md-umbrella-setup-slurm.sh` and then `ir-md-umbrella-md-slurm.sh`.
2. Gather the output `*dihedral*.xvg` files in `$protein-$site-$spoint-$length-$proc/raw`.
3. Install the Grossfield Lab wham software (remember to set energy units to kJ/mol for gromacs outputs!!!): <http://membrane.urmc.rochester.edu/content/wham>, and read the documentation.
4. Run the following script (with modifications) to produce wham-2d inputs:

```
springx=0.0091385 #spring constant for CA-CB-SD-CE in units of kJ/mol/
degree^2 (while dihrestraint.itp has it in kJ/mol/rad^2!)
springy=0.0091385 #spring constant for CA-CB-SD-CE in units of kJ/mol/
degree^2 (while dihrestraint.itp has it in kJ/mol/rad^2!)
nheadings=16      #number of headings in *dihedralTime.xvg files
proc=md           #no need to change
for protein in holo.cam; do
for site in m72c; do
for spoint in n.1; do
for length in 250ps-umbrella; do
    cd $protein-$site-$spoint-$length-$proc
    mkdir processed
for center1 in $(seq 0 30 331); do
for center2 in $(seq 0 30 331); do
    awk -v nheadings=$nheadings 'NR>nheadings{print}' raw/$protein-$site-
$spoint-$length-$center1-$center2-$proc-dihedralTime1.xvg > processed/
$protein-$site-$spoint-$length-$center1-$center2-$proc-dihedralTime1-
no_heading.xvg
    awk -v nheadings=$nheadings 'NR>nheadings{print}' raw/$protein-$site-
$spoint-$length-$center1-$center2-$proc-dihedralTime2.xvg > processed/
$protein-$site-$spoint-$length-$center1-$center2-$proc-dihedralTime2-
no_heading.xvg
    awk 'FNR==NR{a[$1,1]=$1; a[$1,2]=$2;next}{print a[$1,1], a[$1,2], $2}'
processed/$protein-$site-$spoint-$length-$center1-$center2-$proc-
dihedralTime1-no_heading.xvg processed/$protein-$site-$spoint-$length-
$center1-$center2-$proc-dihedralTime2-no_heading.xvg > processed/$protein-
$site-$spoint-$length-$center1-$center2-$proc-dihedralTime-combined.xvg
    rm processed/$protein-$site-$spoint-$length-$center1-$center2-$proc-
```

```

dihedralTime1-no_heading.svg
    rm processed/$protein-$site-$spoint-$length-$center1-$center2-$proc-
dihedralTime2-no_heading.svg
    (echo processed/$protein-$site-$spoint-$length-$center1-$center2-$proc-
dihedralTime-combined.svg $center1 $center2 $springx $springy) >> holo.cam-
m72c-n.1-500ps-wham.inp
done
done
    cd ..
done
done
done
done

```

5. Run wham-2d:

```

wham-2d Px -180 180 36 Py -180 180 36 1e-12 300 0 holo.cam-m72c-n.
1-250ps-wham.inp holo.cam-m72c-n.1-250ps-wham.dat 1
Refer to manual for what each option means and whether you need to change
them: http://membrane.urmc.rochester.edu/sites/default/files/wham/doc.html

```

6. Analyze the output .dat file using 1-md/scripts/wham/wham-analysis.nb, and calculate double integrals under each dihedral combination. Consider sampling dihedral combinations with > 2% occurrence probability.