### 22.01.15

#### **Tutorial from the book**

- Rerun the simulation from the beginning, this time using different folders for each step of the simulation
- simulation started

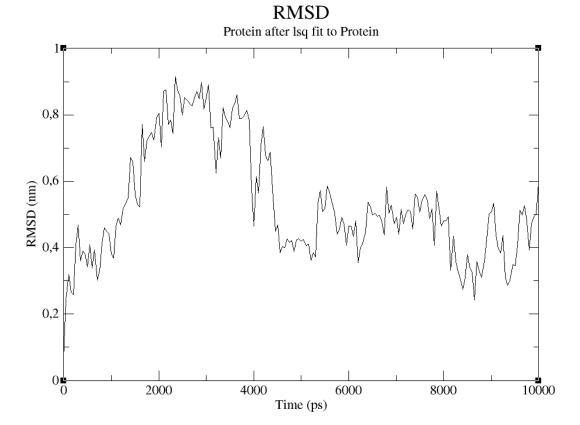
### **Trajectory Analysis of simulation from 20.01.15**

- Simulation of the 2x Heptad
- · Connect via ssh

```
ssh hvoehri@hvoehri.mpibpc.intern
```

• RMSD calculation performed as follows, selecting two times 1 for protein

```
g_rms -s ../em.tpr -f traj.part0001.xtc
xmgrace rmsd.xvg
```

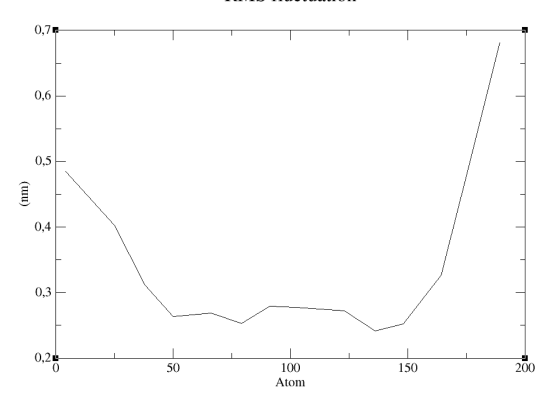


RMSD

• Comparing fluctuations of Ca-atoms with g\_rmsf tool

```
g_rmsf -s md.tpr -f traj.part0001.xtc -o rmsf.xvg -oq bfaq.pdb
xmgrace rmsf.xvg
```

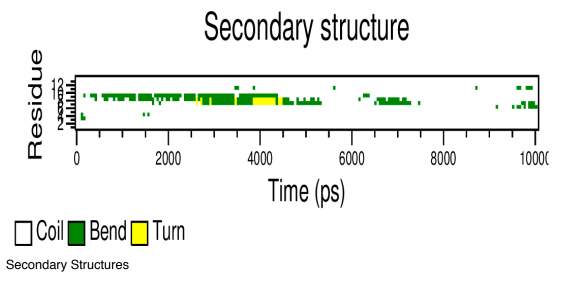
#### RMS fluctuation



RMSF of Ca atoms

• Analysis of Secondary structures

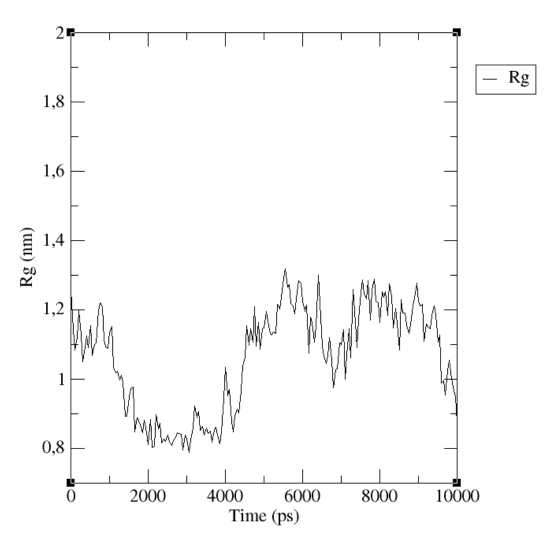
```
do_dssp -s md.tpr -f traj.part0001.xtc -ver 1
xmp2ps -f ss.xpm
gv plot.eps
```



Radius of Gyration

```
g_gyrate -s md.tpr -f traj.part0001.xtc
```

# Radius of gyration

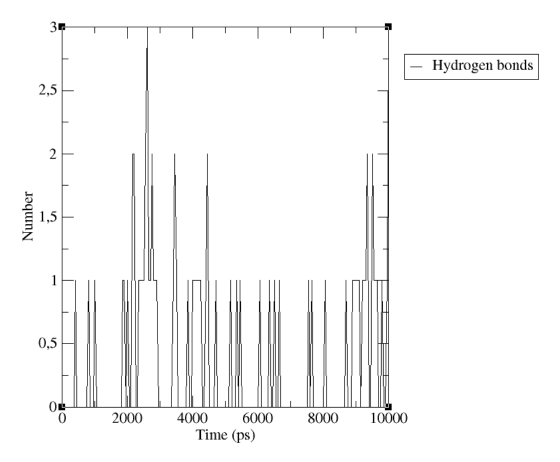


Radius of gyration

Number of H-bonds between protein and protein itself

```
g_hbond -s md.tpr -f traj.part0001.xtc -num hbnum.xvg
```

## Hydrogen Bonds



Number of H-bond formed by the protein itself