## 1 Reproduce the result from paper using LAMMPS

Step 1 requires (in LJ units):

Interaction between molecules ( $\epsilon$ ) mimicked by harmonic oscillator vanished off at  $3\sigma$ . NVE with langevin. Thermostat without thinking about the on/off states and force.

Parameters: N=10, R=2, L=2, dt=0.01 in thermostat,  $r_0$ =0.5 for the bond length,  $s_s$  =2 for spring constant (bonds)  $\epsilon$ =2.5,  $\epsilon_w$ =0.3 for the LJ constant between monomers and between wall particles,  $f_{hr}$ =5 within region 1 and only acting on the first monomer,  $f_{pull}$ =1.0 is acting on the first monomer iff in region 2, 200<sup>3</sup> 3D box, ignore  $k_{on}$  = 10 and  $k_{off}$  = 1 for now.

Problems:

Lost atom (solved by discretizing the time in more detail). (Solved)

"Walls" are generated by repulsion, original method can cause diffusion. (Solved)

The positions and momentum of all atoms should be defined in data file. (Solved)

Force design: parameter  $x_{lo}$  and  $x_{hi}$  as the range of action,  $\Delta$  as scaling,  $f_{max}$  as maximum force added,  $\omega$  as frequency and t as time.  $F = \frac{tanh(\frac{x-x_{lo}}{\Delta}) - tanh(\frac{x-x_{hi}}{\Delta})}{2} f_{max}cos(\omega t)$ 

step 2: cut the LJ pairing between polymers and wall atoms at 1.12 instead of 3 to avoid overlapping around the corner; calculating the free energy change from the wall to anywhere left by  $-k_BTln(P)$  where P is subtracted from the spring we fixes by umbrella sampling.

## 2 Revise the model into 3D

(solved)

## 3 Try models in different attraction forces besides harmonic approximation

Hydrogen bonding, fene and electrostatics

- 4 Heterogeneous monomers
- 5 Design initial structures for peptides