PH: 322 Molecular Simulation Homework 2

1. Consider the angle bending potential of the following form

$$E_{\theta} = \frac{1}{2} \sum_{angles} K_{\theta} (\theta - \theta_{o})^{2}$$

Derive the expression for force on the three atoms making the angle bending term.

Due date: 5th March

2. Consider the exponential-6 potential given by following form

$$V_{vdw} = Ae^{-\alpha r} - \left(\frac{C}{r}\right)^6$$

Plot this potential as a function of r (inter-particle separation) and discuss its behavior for very small values of r

- 3. For a Lennard-Jones system in NVE ensemble plot the RMS fluctuation in energy as a function of time step (δt) of integration. From this comment on how large time step one can choose for simulation.
- 4. Remember we discussed the use of switching functions in the context of truncating and shifting the non-bond potential. We discussed the use spline functions as well as various order polynomial switching functions. For non-bond LJ interactions please compare the energy accuracy and time efficiency of cubic spline as well as higher order (5th and 7th order) polynomial switching functions.
- 5. Show that the heat capacity is related to the fluctuation in total energy through the following relation

$$C_V = \frac{\left\langle U^2 \right\rangle - \left\langle U \right\rangle^2}{k_B T^2}$$

From your NVT MD simulation calculate the specific heat of the Lennard-Jones system at two densities (one in gas phase and another in liquid phase).

6. Using the MD code (supplied) compute the equation of state for LJ fluid at T=1.0, T=2.0 and T=0.9. Take a system of 500 LJ particles.