# Simulation of Liquid Argon

# Vo Chau Duc Phuong

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#### 1 Introduction

Using the FORTRAN, I simulate the liquid Argon in the Lennard-Jones potential between the particles. The input value into the code will be these parameters and variables:

```
integer :: n = 864, bc = 1, tr = 0 !bc: periodic boundary, tr: poor-mans algorithm real, allocatable :: dens(:,:,:) !crucial parameters real, parameter :: T = 94.4, kb= 1.380649*10.**(-16.) real, parameter :: m = 39.95*1.67*10.**(-24), sigma = 3.4*10**(-8.) real, parameter :: ep = 120.*kb, rc = 2.25, L = 10.229, halfL = 10.229/2. !time parameters integer :: nt, ntrelax real, parameter :: tmin = 0.0, tmax = 100.0, trelax = 10.0 !in ps real :: dt = 0.01 !ps
```

The first line is specify the number of particles, use the parameter and poormans thermal-stat or not. (in this case, no thermal-stats).

Second line is the define of the density matrix, contains the position and velocity of all particles.

Crucial parameter will be in the unit of cgs. The time parameters as shown.

```
subroutine Initial() implicit none integer :: i, j, k, p real :: spacing, v_stddev, rand1, rand2, Tp, E allocate(dens(n,3,2)) ntrelax = int(trelax/dt); nt = int((tmax - tmin)/dt) dt = dt*1.0e-12*(ep/m)**0.5*1./sigma
```

```
! Initialize densitions randomly within the box
spacing = L / (int(n**(1.0/3.0)) + 1)! Adjust spacing to avoid boundary
p = 0
do i = 0, int(n**(1.0/3.0))
do j = 0, int(n**(1.0/3.0))
do k = 0, int(n**(1.0/3.0))
p = p + 1
if (p > n) exit
dens(p,1,1) = i * spacing
dens(p,2,1) = j * spacing
\mathrm{dens}(p,3,1) = k * \mathrm{spacing}
end do
if (p > n) exit
end do
if (p > n) exit
end do
v_stddev = sqrt(kb * T / m)
do i = 1, n
do j = 1, 3
call random_number(rand1); call random_number(rand2)
dens(i,j,2) = v_stddev * sqrt(-2.0 * log(rand1)) &
* \cos(2.0 * 3.14159265359 * rand2)
end do
end do
do i = 1, 3
dens(:, i, 2) = dens(:, i, 2) - sum(dens(:, i, 2))/n
Tp = sum(dens(:,:,2)**2.) * m / (3.0 * n * kb)
dens(:,:,2) = dens(:,:,2) * sqrt(T / Tp)
dens(:,:,2) = dens(:,:,2) / (ep/m)**0.5! Convert to dimensionless units
end subroutine
```

First, I generate the position of the particles in a lattice to avoid the very strong at the near interaction of Leornard-Johnes potential

## 2 Method

## 3 Results and Discussion