

# LAMMPS gcmc with umbrella

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I test liquid-gas coexistenc in LAMMPS with my custom `gcc/umbrella` fix. I take a small LJ system in a volume  $V = 8^3\sigma^3$  and test it at the  $\beta\mu$  given by MC

I place a quadratic umbrella with weight

$$w(n) = \exp \left[ -\frac{k(n - n_0)^2}{2T} \right]$$

where  $n$  is the number of particles. It is centered at  $n_0 = 150$  with  $k = 0.0005$ .

I run `lammmps` in a hybrid GCMC coupled to a constant temperature thermostat (which should correctly take into account the change of particles).

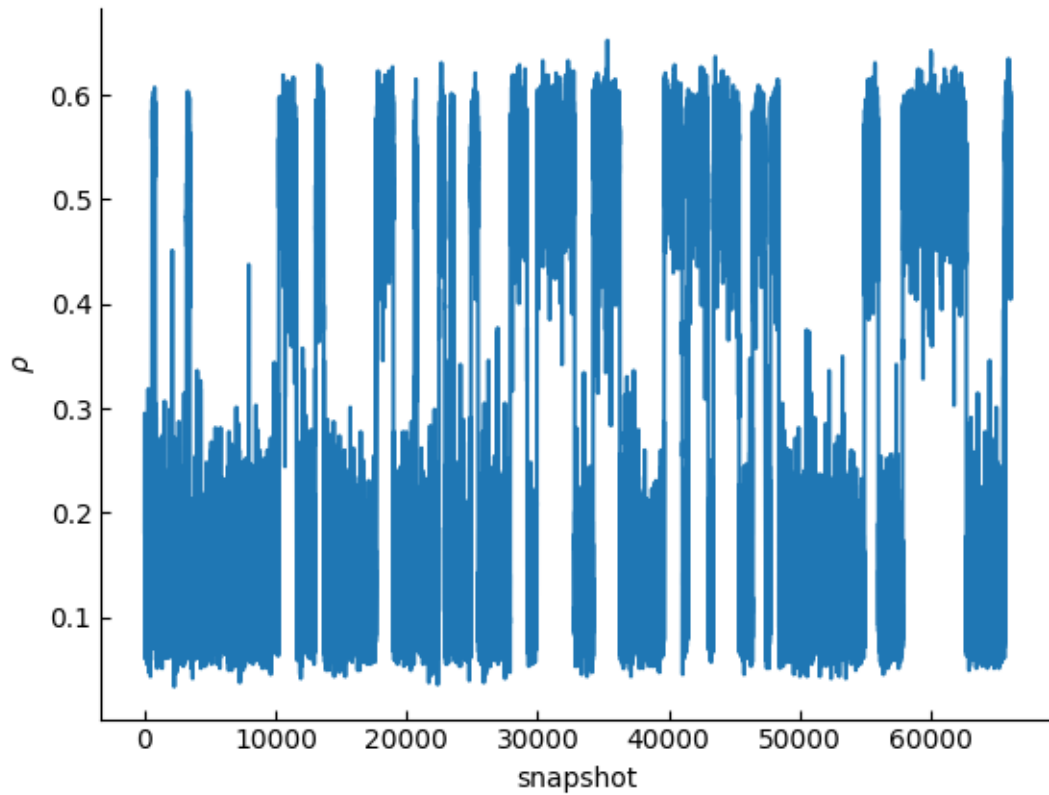
First I load the outputs for teh density.

```
import numpy as np
import matplotlib.pyplot as plt
```

```
data = np.loadtxt("n-betamu-3.092-T1.txt")
```

```
plt.plot(data[:,1])
plt.xlabel("snapshot")
plt.ylabel(r"$\rho$")
```

```
Text(0, 0.5, '$\rho$')
```



The simulation correctly jumps from one phase to the other.

I can check the PDFs and apply an unbiasing.

```
# consider the pdf and the umbrella
Vol = L**3
rhos = np.linspace(0,data[:,0].max()+1)/Vol
# print(rhos, rhos.max())
h,edges = np.histogram(data[:,1],bins=rhos, density=True)
rho = edges[:-1]+0.5*(edges[1]-edges[0])
F = np.log(h)

plt.plot(rho,F,label='raw')

# umbrella
k, n0 = 0.0005,150
L = 8.0

T = 1.0
```

```
umbrella = - 0.5/T*k*(rho*Vol-n0)**2
```

```
plt.plot(rho,F-umbrella,label='corrected for the umbrella')
```

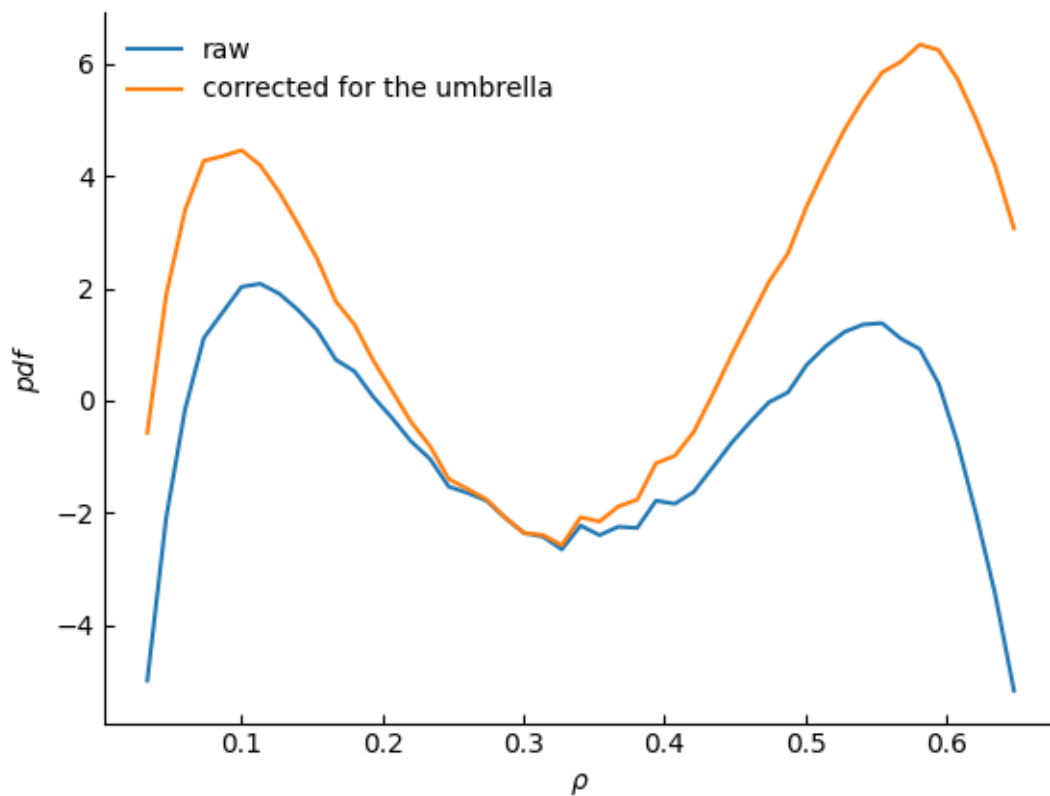
```
plt.xlabel(r"$\rho$")
```

```
plt.ylabel(r"$pdf$")
```

```
plt.legend()
```

/var/folders/lt/p1n6bb0j4\_173x9548b473jh0000gn/T/ipykernel\_71458/1092576372.py:7: RuntimeWarning

```
F = np.log(h)
```



The peaks are unequal. Maybe I can estimate a correction to the chemical potential?

```
def objective(dmu, original_F=None, rho=None, mid=0.3, Volume=1.):
```

```
    N= rho*Volume
```

```
    F = original_F+dmu*N
```

```
    val = (F[rho<mid].max()-F[rho>mid].max())**2
```

```
    return val
```

```
# objective(0,F-V, rho=rho,V=V)
from scipy.optimize import minimize_scalar

res = minimize_scalar(objective,args=(F-umbrella,rho,0.3,Vol))
res.x
betamu = -3.092
print(f"The correction dmu is {res.x} which is {res.x/betamu} of the original value")
```

The correction dm<sub>u</sub> is -0.0076565803405382795 which is 0.0024762549613642558 of the original value

```
plt.plot(rho,F-umbrella+res.x*rho*Vol,label='corrected chemical potential')
plt.xlabel(r"$\rho$")
plt.ylabel(r"$pdf$")
plt.legend()
```

