PHYS 516: Methods of Computational Physics ASSIGNMENT 3- MC Simulation of the Ising Model

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1 Theoretical Foundation of Metropolis Algorithm

Consider a set of N states, $\Gamma_1, \Gamma_2, ..., \Gamma_N$ and let the probability to find the system in the m-th state, Γ_m , be ρ_m . Here, we prove that the probability distribution is a fixed point of the metropolis transition matrix defined below, i.e., $\Pi \rho = \rho$.

(Metropolis transition matrix)
$$\pi_{mn} = \begin{cases} \alpha_{mn} & \rho_m \ge \rho_n m \ne n \\ \frac{\rho_m}{rho_n} \alpha_{mn} & \rho_m \le \rho_n m \ne n \\ 1 - \sum_{m' \ne n} \pi_{m'n} \end{cases}$$

Here, π_{mn} are elements of the matrix Π , ρ_m are the elements of vector ρ , and α_{mn} are elements of a symmetric attempt matrix, i.e., $\alpha_{mn} = \alpha_{nm}$.

This can be proved by enforcing the Detailed Balance condition. Consider a pair of states m and n. Assuming $\rho_m < \rho_n$,

$$\pi_{nm}\rho_m = \alpha_{nm}\rho_m = \alpha_{mn}\rho_m \tag{1}$$

where the second equality holds because of the symmetric attempt. For the same case,

$$\pi_{mn}\rho_n = \frac{\rho_m}{\rho_n}\alpha_{mn}\rho_n = \alpha_{mn}\rho_m \tag{2}$$

Using equations 1 and 2,

$$\pi_{nm}\rho_m = \pi_{mn}\rho_n \tag{3}$$

The left-hand side is a flux of probability that the current state is n and that the next state is m, and the right-hand side is a flux from m to n.

In order to show that above detailed balance condition is sufficient for the unit-eigenvalue relation, sum the both side over n.

$$\sum_{n=1}^{N} \pi_{mn} \rho_n = \sum_{n=1}^{N} \pi_{nm} \rho_m \tag{4}$$

$$\implies \sum_{n=1}^{N} \pi_{mn} \rho_n = \rho_m \tag{5}$$

$$\therefore \Pi \rho = \rho \tag{6}$$

2 2D Ising Model

Ising model is used for modelling ferromagnetic materials. This model represents a lattice consisting of atoms which have quantum mechanical spin, which can be ± 1 . When these individual magnetic fields are aligned in the same direction, it gives rise to macroscopic magnetic field. This strong alignment arises from exchange interactions between electrons.

2.1 Computer Simulation

A program was written in C to simulate the 2D Ising model.

```
1 /* Monte Carlo Simulation of 2D Ising Model */
2 #include < stdio . h>
3 #include < stdlib.h>
4 | #include < time.h>
5 #include <math.h>
7 // Define global variables
8 #define L 20 //lattice size
9 int s[L][L]; //Spins s[i]/j=+-1
10 double exp dV[2][5];
11 double JdivT; //J/kBT
12 double HdivT; // H/kBT
13
14 void table_set() { //function to set up the table
15
    int sDash, sneighbor, k, l;
16
    for (k=0;k<2;k++)
       sDash = 2*k-1;
17
18
       for (1=0;1<5;1++)
19
         sneighbor = 2*l-4;
20
         \exp dV[k][1] = \exp(2*sDash*(JdivT*sneighbor + HdivT)); // check
            formula
21
22
23
  }
24
25 int main() {
26
    double runM;
27
    double sumM = 0.0, sumM2 = 0.0;
    double exp val, avgM, sigM;
28
29
    int snew, sneighbor, Sta step;
30
    int i, j, step, k, l, im, ip, jm, jp;
31
    FILE *f = fopen("Magnetization data.txt", "w");
32
33
     printf("Input J/kBT\n");
    scanf("%le",&JdivT);
34
35
36
    HdivT = 0.0;
37
    Sta step = 2000000;
```

```
38
39
     table set(); // Set up the look-up table for the exponent
        calculation
40
41
     for(i=0;i<L;i++) { //Cold start start with all spins up
        configuration
       for (j=0; j< L; j++) {
42
         s[i][j] = 1;
43
44
45
    runM=1.0*L*L;
46
47
     for(step=0; step<Sta step; step++) {</pre>
48
49
       i=rand()%L;
                      j=rand()%L;
50
       snew=-s[i][j];
51
52
       // Figure out which element of the table is to be looked up
53
       im = (i + L - 1) \% L;
54
       ip = (i + 1) \% L;
       jm = (j + L - 1) \% L;
55
       jp = (j + 1) \% L;
56
57
       k = (snew+1)/2;
58
       sneighbor = s[im][j] + s[ip][j] + s[i][jm] + s[i][jp];
59
       l = (sneighbor + 4)/2;
60
61
       //Change in Pot Energy wth flip
62
       \exp \text{ val} = \exp \text{ dV}[k][1];
63
       // Accept or reject flip conditionally
       if (\exp \text{ val} > 1.0) {
64
65
         s[i][j] = snew;
         runM += 2*snew; //update value of magnetization
66
67
       else if (rand () / (double) RAND MAX < exp val) {
68
69
         s[i][j] = snew;
70
         runM += 2*snew; //update value of magnetization
71
72
      sumM += runM;
73
      sumM2 += runM*runM;
74
       fprintf(f, "\%f \ n", runM);
75
     fclose(f);
76
    avgM = sumM/Sta step; //Mean Magnetization
77
    sigM = sqrt (sumM2/Sta step-avgM*avgM); //Standard deviation
78
79
     printf("Mean Magnetization %le, Standard Deviation %le \n",
        fabs(avgM), sigM);
80 }
```

2.2 Results

The plots below show the Mean magnetization and its standard deviation as a function of the exchange coupling, and a histogram showing the number of MC samples for every value of Magnetization. The matlab script used to generate these plots is shown below.

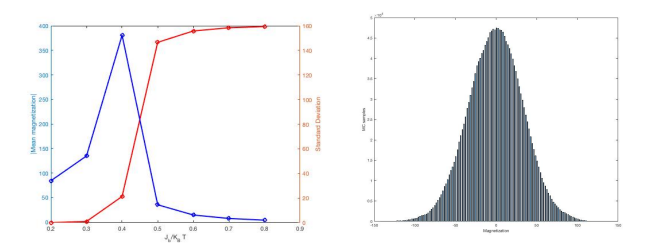


Figure 1: Plot of Magnetization and S.D vs J/K_BT , and Histogram for $J/K_BT = 0.2$

```
close all; clear all;
2
3 %% Scatter Plots
4| JDivT = [0.2:0.1:0.8];
5 \mid M = [-9.427350e-01, 1.361261e+00, 5.233627e+01, ...]
      3.662830e+02,3.894057e+02,3.959868e+02,3.984030e+02];
  yyaxis left;
  plot(JDivT, M, 'r-o', 'LineWidth', 2);
9 xlabel('J_b/K_B T'); ylabel('|Mean magnetization|');
11 sd = [3.403888e+01, 5.418066e+01, 1.523472e+02, ...
      1.449735e+01,6.180052e+00,3.360630e+00,1.993231e+00;
12
13 yyaxis right;
14 plot(JDivT, sd, 'b-o', 'LineWidth', 2);
15 xlabel('J_b/K_B T'); ylabel('Standard Deviation');
16
17 %% Histogram
18 figure()
19 A = importdata('Magnetization_data.txt', '\n', 0);
20 histogram(A);
21 ylabel('MC samples'); xlabel('Magnetization');
22 xlim([-150 150]);
```

Ising MagnetizationPlots.m