FYS4150 Project 4, deadline November 19.



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Abstract

We have made a c++ program to calculate some thermodynamic variables for a square lattice using the Ising model. The variables have been used for plots and further calculations in MATLAB. The outputs have shown to be sensitive to temperature, while the initial configuration of the lattice does not matter much. The energy, magnetization, specific heat and susceptibility stay close to the same for each spin for increasing lattice sizes. The critical temperature, showing a phase shift, is found at the same temperature for our lattice sizes, and is found to be at T=2.3.

Introduction

In this project we will study phase transitions in magnetic systems. We will use the widely popular square-lattice Ising model in two dimensions, where particles interact with their nearest neighbours to get a magnetic spin of either +1 or -1. This model will be used to find a critical temperature showing a phase shift where the system goes from having magnetic moment to zero magnetic moment. We will start off by looking at the simple case of a 2x2 lattice and use this as benchmark calculations for the rest of the project. We will then move on to larger lattices, and eventually compare these results to a more realistic model where the lattice is infinite in size. In this project the emphasis will be on understanding and usage of the Metropolis algorithm, and have a look at its efficiency and precision. The codes and files used in this project can be found at https://github.com/RobinKifle/Assignment4 and https://github.com/sanderwl/Assigment-4. The last one includes the parallelization and the matlab plots.

Method

Starting of, we need to find analytical expressions for the partition function Z and the corresponding energy values E, the mean magnetization |M|, the specific heat C_v and the susceptibility χ as functions of the temperature T using periodic boundary conditions.

The partition function is given by:

$$Z = \sum_{i=1}^{M} e^{-\beta E_i}$$

Where M is the total number of configurations, β is the inverse temperature given by $\beta = \frac{1}{kT}$ with k as the Boltzmann constant. E_i is the energy for different spin settings given by:

$$E = J \sum_{\langle kl \rangle}^{N} s_k s_l$$

where J is a constant which expresses the strength of the interaction between spins. N is the total number of configurations. $\langle kl \rangle$ expresses that we only look at the nearest neighbours in the lattice. s_k and s_l are the spins of two neighbouring objects.

For now we work with a 2x2 lattice and the Ising model where the spins can only be in two states, giving a total number of $2^4 = 16$ configurations.

Table 1: Energy with periodic boundary conditions and magnetization for different spin-configurations

Spin configuration	Energy	Magnetization
$\uparrow\uparrow$	-2J	2
$\uparrow\downarrow$	2J	0
$\downarrow \uparrow$	2J	0
$\downarrow \downarrow$	-2J	-2

From the table above one can observe that the energy is only non-zero on lattices where all objects have the same spin (-8J) and when the diagonals have opposite spin (8J). The magnetization of a 2x2 lattice is only zero when half of the objects have the same spin. We can now calculate the mean energy, the mean absolute magnetization, specific heat and susceptibility with these expressions:

Partition function:

$$Z = \sum_{i=1}^{16} e^{-\beta E_i}$$

$$Z = 2e^{8\beta J} + 2e^{-8\beta J} + 12$$

$$Z = 4\cosh(8\beta J) + 12$$

Mean energy:

$$\langle E \rangle = \frac{1}{Z} \sum_{i=1}^{16} e^{-\beta E_i} E_i$$

$$\sum_{i=1}^{16} e^{-\beta E_i} E_i = -16J(e^{8\beta J} - e^{-8\beta J}) = -32J \sinh(8\beta J)$$

$$\langle E \rangle = -\frac{8J}{\cosh(8\beta J) + 3} \sinh(8\beta J)$$

Mean absolute magnetization:

$$<|M|> = \frac{1}{Z} \sum_{i=1}^{16} e^{-\beta E_i} |M_i|$$

$$\sum_{i=1}^{16} e^{-\beta E_i} |M_i| = 8e^{8\beta J} + 16$$

$$<|M|> = \frac{2e^{8\beta J} + 4}{\cosh(8\beta J) + 3}$$

Specific heat:

$$C_v = \frac{\beta}{T} (\langle E^2 \rangle - \langle E \rangle^2)$$

$$\langle E \rangle^2 = \frac{64J^2}{(\cos h(8\beta J) + 3)^2} \sinh^2(8\beta J)$$

$$\langle E^2 \rangle = \frac{1}{Z} \sum_{i=1}^{16} e^{-\beta E_i} E_i^2 = \frac{64J^2}{\cosh(8\beta J) + 3} \cosh(8\beta J)$$

$$\to C_v = \frac{64\beta J^2}{T(\cosh(8\beta J) + 3)^2} (3\cosh(8\beta J) + 1)$$

Susceptibility:

$$\begin{split} \chi &= \beta \big(< |M|^2 > - < |M| >^2 \big) \\ &< |M|^2 > = \frac{1}{Z} \sum_{i=1}^{16} e^{-\beta E_i} M_i^2 = \frac{8 e^{8\beta J} + 8}{\cosh(8x) + 3} \\ &\rightarrow \chi = \beta \big(\frac{8 e^{8\beta J} + 8}{\cosh(8\beta J) + 3} - \frac{(2 e^{8\beta J} + 4)^2}{(\cosh(8\beta J) + 3)^2} \big) \end{split}$$

Notice that the terms in C_v and χ , $\langle E^2 \rangle - \langle E \rangle^2$ and $\langle |M|^2 \rangle - \langle |M| \rangle^2$, respectively, are the variance of the energy and variance of the magnetization.

As mentioned, we will use the Metropolis algorithm. This algorithm can be described in ten steps¹

- 1. Establish an initial matrix with size $L \times L$ and compute the energy.
- 2. Position yourself at a random point in the lattice and flip one spin.
- 3. Compute the energy for this new state.

¹Hjorth-Jensen., 2015

- 4. Compute $\triangle E$
- 5. If the energy is lowered, accept the new configuration and jump to step 9
- 6. If the energy is increased, compare $w = e^{-\beta \Delta E}$ with a random number.
- 7. If the random number is bigger than w, reject the new configuration and jump back to step 2.
- 8. If the random number is smaller or equal to w, accept the new configuration.
- 9. Update expectation values
- 10. Repeat $L \times L$ times to let every lattice point get a chance to get picked.

From our simulations for a LxL lattice with L=20, L=40, L=60, L=80, L=100, we will find analytical values for the critical temperature $T_c(L)$ on different plots. We can then use these values to find a value for the critical temperature in the thermodynamic limit L $\rightarrow \infty$ using the expression:

$$T_c(L) - T_c(L = \infty) = aL^{-1/\nu}$$

Where a is a constant and we use $\nu = 1$. We set $L^{-1} = X$ and solve the equation $T_c(L) = aX + T_c(L = \infty)$ using Matlabs' least squares solver.

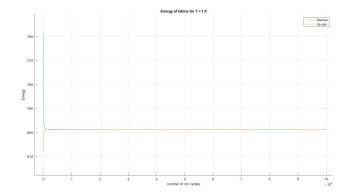
To keep the elapsed time endurable, we also parallelized the code using MPI. This allowed us to perform 8 simultaneous Monte Carlo cycles, resulting in an almost 8x faster code.

Results

Table 2: Analytical and numerical solutions for a random initial matrix

	Analytical	Numerical	MC cycles
< E >	-7.983928	-7.98472	10.000
< M >	3.994643	3.99494	10.000
C_v	0.128329	0.122007	100.000
χ	0.016043	0.015054	100.000

Starting with a 2x2 matrix with a random configuration, after only 10.000 Monte Carlo cycles the numerical solutions for the mean energy and the mean magnetization are close to the values found analytical. They are only off by a value of 10^{-3} . To get the same magnitude of agreement for the specific heat and susceptibility, 100.000 Monte Carlo cycles was needed. For fewer than 10.000 Monte Carlo cycles the values are irregular.



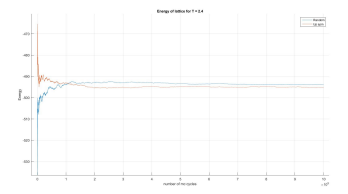
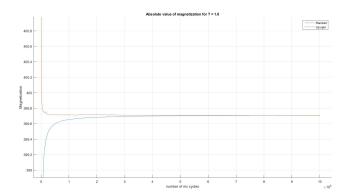


Figure 1: Energy vs. Monte Carlo cycles for random- and up-spin oriented matrix with T=1.0 and T=2.4, respectively

From figure 1 and we notice that after some amount of Monte Carlo cycles the random- and up-spin initial matrices converges to the same energy equilibriums. For T=1 the equilibrium is reached after about $2*10^4$ Monte Carlo cycles with a value of -800, or -2 per spin. The up-spin initial matrix reaches equilibrium more promptly than the random initial matrix as it starts from the steady state. For T=2.4 the energy oscillates more, and the equilibrium is reached after about 10^5 cycles with a value of -495, or -1.2375 per spin. This oscillation is expected as there is more energy in the system. At T=2.4 the up-spin initial matrix does not start from a steady state and needs more cycles before reaching the equilibrium compared to the system with T=1.



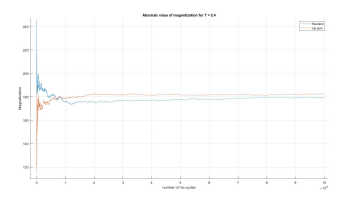
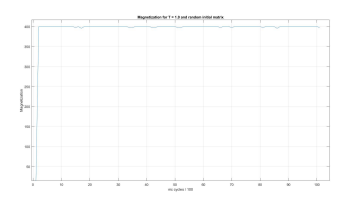


Figure 2: Absolute magnetization vs. Monte Carlo cycles for random- and up-spin oriented matrix with T=1.0 and T=2.4, respectively



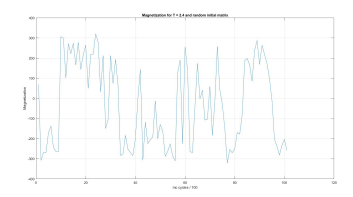
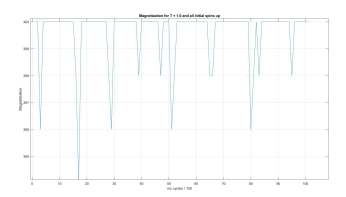


Figure 3: Mean magnetization vs. Monte Carlo cycles for random oriented matrix with T=1.0 and T=2.4, respectively



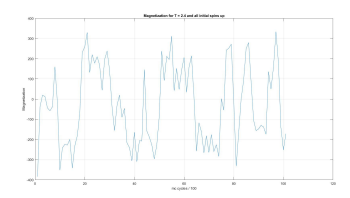


Figure 4: Mean magnetization vs. Monte Carlo cycles for up-spin oriented matrix with T=1.0 and T=2.4, respectively

In figure 2 it is noticeable that the absolute magnetization converges rapidly after the same amount of Monte Carlo cycles as the energy previously mentioned. The absolute magnetization behaves differently than the energy from figure 1, indicating that there are multiple lattice configurations giving the same energy minimum. In figure 3 and figure 4 we notice that the magnetization oscillates between positive and negative values. Comparing this with figure 1 and 2, it is noticeable that the magnetization keeps

oscillating between positive and negative values after the energy and absolute magnetization has reached equilibrium.

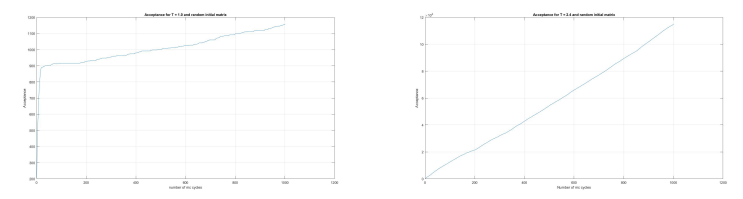


Figure 5: Acceptance vs. Monte Carlo cycles for random oriented matrix with T=1.0 and T=2.4, respectively

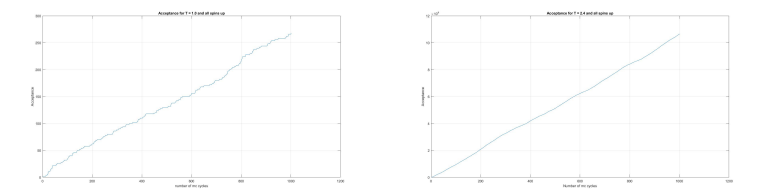


Figure 6: Acceptance vs. Monte Carlo cycles for up-spin oriented matrix with T=1.0 and T=2.4, respectively

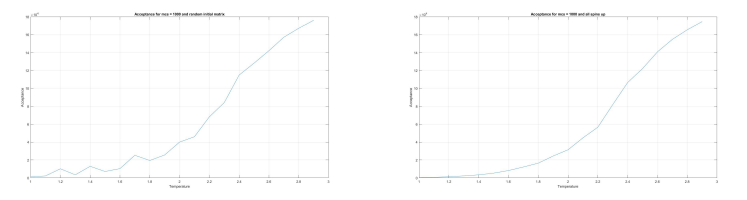


Figure 7: Acceptance vs. temperature for random- and up-spin oriented matrix, respectively

For a random initial matrix and T=1, seen in figure 5, the system start off by accepting a lot of flips before behaving linearly as the initial up-spin matrix with T=1, shown in figure 6. This is because

the random intial matrix does not start at the equilibrium. For T=2.4 there are a lot more accepted configurations. None of the matrices start at equilibrium and seem to behave the same.

In figure 7 we can clearly see an exponential growth in number of accepted configurations for increasing temperature. It seem like the number of accepted configurations might converge at the end of the graph, but we would need to run for higher temperatures to be sure about this. The up-spin initial matrix is smoother than the random initial matrix, which is expected.

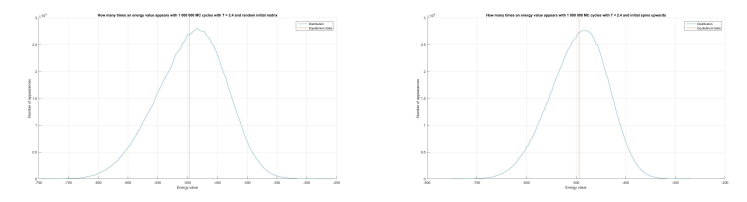


Figure 8: Number of appearances per energy value for random- and up-spin oriented matrix with T=2.4

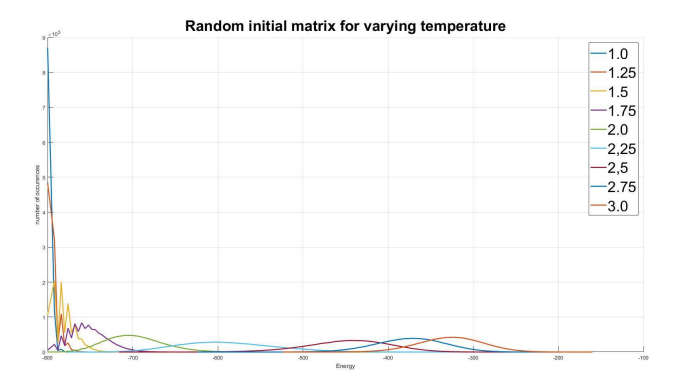


Figure 9: Probability distribution for varying temperature over 10⁶ Monte Carlo cycles

Table 3: Variance in energy for different temperatures over 10⁶ Monte Carlo cycles

Temperature	Variance
1.00	9.09764
1.25	51.7348
1.50	177.7472
1.75	478.188
2.00	1152.904
2.25	3158.504
2.50	2473.664
2.75	1700.996
3.00	1446.38

The probability distribution in figure 8 show a negative skewed distribution, with the mean value laying one the left side of the mode. From figure 9 we can clearly see that there are a higher probability for higher energies for increasing temperature. It is also noticeable that the energy is also oscillating more for higher temperatures. For T=1 there are almost only the equilibrium state. From 3 we notice that the variance peaks between T=2.5 and T=2.75. This will be discussed later.

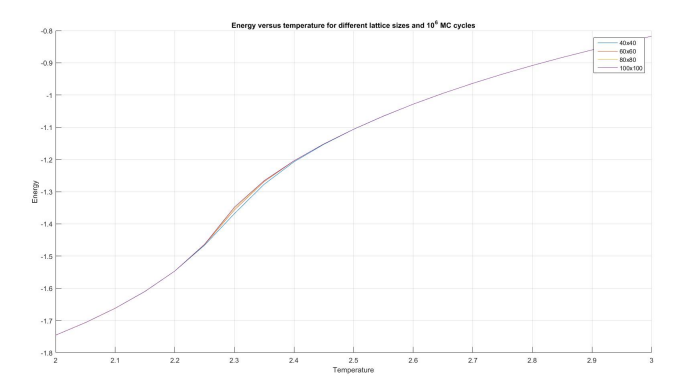


Figure 10: Mean energy vs. temperature for 10⁶ Monte Carlo cycles normalized(single lattice objects)

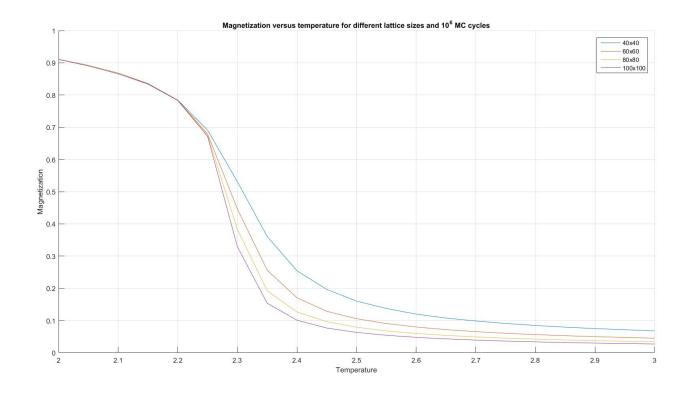


Figure 11: Mean magnetization vs. temperature for 10^6 Monte Carlo cycles normalized(single lattice objects)

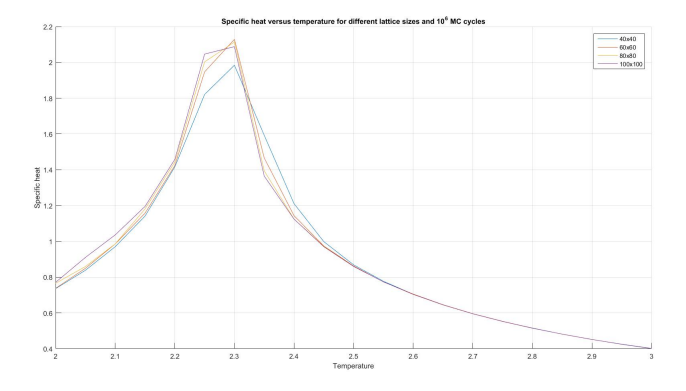


Figure 12: Specific heat vs. temperature for 10^6 Monte Carlo cycles normalized(single lattice objects)

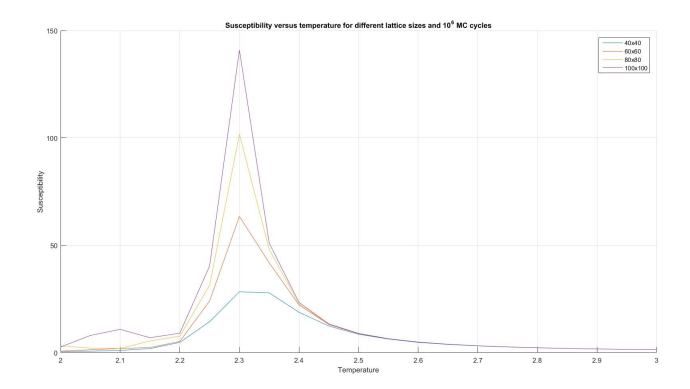


Figure 13: Susceptibility vs. temperature for 10⁶ Monte Carlo cycles normalized(single lattice objects)

Table 4: Run time for different lattice sizes with $10^6~\mathrm{MC}$ cycles

Lattice size	Time in seconds
60x60	3996.57
80x80	7081.15
100 x 100	18630.9

From figure 12 and 13 we notice peaks indicating the critical temperature at T=2.3 for all matrices. Knowing this, we notice a change in curvature in figure 10 and 11 at this same temperature. The energy goes to higher values for higher temperatures while the magnitization seem to go from having a magnetic moment to zero magnetic moment. The change in magnetization seem to become steeper for larger lattices and the susceptibility peaks higher for larger lattices.

Discussion

For a 2x2 lattice the analytical and numerical solutions correlate well after few Monte Carlo cycles.

For T=2.4 the energy oscillate more than for T=1. This is expected as energy is more active at higher temperatures.

Looking at the different magnetization plots it is noticeable that the magnetization is limited to some set values. The reasoning for this is the same as for the 2x2 lattice having few possible configurations. In the same way as for the 2x2 lattice, a larger lattice will only have a set number of values it can jump between when doing leaps between positive and negative magnetization values. The reason for the mean magnetization to keep oscillating between positive and negative values, in figure 3 and 4 for T=2.4, after the energy has reached equilibrium, seen in figure 2, is because there is a relation between the energy and the absolute magnetization, hence the mean magnetization can take these leaps between positive and negative values. This does not happen for T=1 because the temperature in the system is not high enough for the magnetization to make these leaps. This can be explained mathematically from the Boltzmann distribution:

$$w_i = e^{-E_i/T}$$

 w_i is the probability, E_i is the energy and T is the temperature. From this we can se that the probability increase with the increase of temperature for low energy values. Hence the magnetization and energy oscillates more for higher temperatures. This also explains the increase in accepted configurations from T=1 to T=2.4 shown in figure 5 and 6, which is emphasised in figure 7. A random initial matrix will behave different at the start before reaching the same state as any other initial matrix due to it stabilizing at the same values.

Looking at the probability distribution in figure 8, we notice that the probability is negative skewed, which means that the probability for lower energy values are larger than for higher energy values, which is expected.

In figure 10 and 11 we notice that the concavity changes at about T=2.3, correlating with peaks in specific heat figure 12) and susceptibility (figure 13) at T=2.3. This fits well with the fact that magnetic susceptibility is a parameter that tell us how much a extensive parameter changes when an intensive parameter increases. Hence the magnetic susceptibility peaks where the concavity of magnetization changes. We have the same correlation between specific heat and energy. We notice in figure 11 a steeper transition for larger lattices, which is also noticeable in figure 13. Following this logic we would expect an infinite slope when $L \to \infty$.

We can also see from table 3 that the variance increases until around the critical temperature before decreasing. This corresponds with the remarks of figure 10, 11, 12 and 13. The exact result for the critical temperature ² is $kT_c/J = \frac{2}{ln(1+\sqrt{2})} \approx 2.269$ with $\nu = 1$. This relates decent with our results.

Errors

Our estimates for the critical temperature is the same value for all matrix sizes. This is disappointing as this of course will give us the same value for the critical temperature in the thermodynamic limit $L \to \infty$. We used a step in temperature of 0.05. If we used a smaller stepsize we would have gotten results which would give a better estimate of the exact critical temperature. We can from table 4 see that the runtime for the lattices is quite long, even when being run on 8 processors by parallelization. Larger lattices could also give us better results, but the runtime would be awful.

Conclusion

The squared lattice Ising model has shown to be respectable for this project. Our results for the thermodynamic variables, energy, magnetization, specific heat, susceptibility and also the critical temperature fits decent with the exact values. Parallelization was found useful as it decreased the computation time.

²Lars Onsager, https://journals.aps.org/pr/abstract/10.1103/PhysRev.65.117

References

Hjort-Jensen, M., 2015. Computational physics, accessible at course github repository. 551 pages

Lars Onsager, 1944. Crystal Statistics. I. A Two-Dimensional Model with an Order-Disorder Transition