

Simulation of 2D 3-State Potts Model

A Metroplis Monte Carlo Approach

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Abstract

The Potts model is a generalization of the Ising model, which is a model of ferromagnetism in statistical mechanics. By using the Metropolis algorithm, a 2D 3-state Potts model is simulated under different temperatures. I studied the internal energy, magnetization, specific heat and magnetic with reference to the temperature and found that the system undergoes a phase transition at a critical temperature around 1.05.

Keywords Monte Carlo, Metropolis Algorithm, Potts Model, Phase Transition

BACKGROUND

1.a. Spin models

The Potts model is one of a group of models called spin models. In a spin model, each site on a lattice is assigned a spin variable. The spin variable can take on a discrete set of values, which are usually integers.

1.a.i. Ising model:

The Ising model is the most basic spin model. In the Ising model, the spin variable can take on only two values, +1 or -1. The Ising model is a model of ferromagnetism in statistical mechanics. The Hamiltonian of the Ising model is given by

$$H = -J \sum_{\langle i,j \rangle} S_i S_j \tag{1}$$

COURSE PROJECT

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Data Availability

Associated code repository is available on GitHub.

Honesty Statement

The author declares that the project was completed independently and without plagiarism.

where S_i is the spin variable at site i, J is the coupling constant, and the sum is over all pairs of nearest-neighbor sites.

1.a.ii. Potts model:

The Potts model is a generalization of the Ising model. In the Potts model, the spin variable can take on q different values, where q is an integer greater than or equal to 2. The Hamiltonian of the Potts model is given by

$$H = -J \sum_{\langle i,j \rangle} \delta \left(S_i, S_j \right) \tag{2}$$

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where S_i is the spin variable at site i, J is the coupling constant, and the sum is over all pairs of nearest-neighbor sites. The Kronecker delta function $\delta(S_i, S_i)$ is defined as

$$\delta(S_i, S_j) = \begin{cases} 1 \text{ if } S_i = S_j \\ 0 \text{ if } S_i \neq S_j \end{cases}$$
 (3)

If magenetic field is applied, an extra term is added to the Hamiltonian

$$H = -J \sum_{\langle i,j \rangle} \delta \left(S_i, S_j \right) - h \sum_i S_i \tag{4} \label{eq:4}$$

where h is the magnetic field strength.

1.b. Interested physical quantities in Potts model

Pratically, physicists are interested in the following physical quantities in Potts model:

1.b.i. *Internal energy:*

$$u = \frac{\langle H \rangle}{N^2} \tag{5}$$

1.b.ii. Specific heat:

$$c = \frac{k_B \beta^2 \left(\left\langle H^2 \right\rangle - \left\langle H \right\rangle^2 \right)}{N^2} \tag{6}$$

where k_B is the Boltzmann constant and $\beta=\frac{1}{k_BT}$ is the inverse temperature. 1.b.iii. *Magnetization:*

$$m = \frac{\left\langle \sum_{i} S_{i} \right\rangle}{N^{2}} \tag{7}$$

1.b.iv. Characteristic length: Define the spatial correlation function

$$C(i,j) = \left\langle S_i S_j \right\rangle - \left\langle S_i \right\rangle \left\langle S_j \right\rangle \tag{8}$$

Then we can define

$$\Gamma(k) = C(i,j)|_{|i-j|=k} \approx \frac{1}{4N^2} \sum_{i} \sum_{j \in S_i} C(i,j)$$
 (9)

where

$$S_i = \{i|i-j = \pm(k,0) \text{ or } (0,k)\} \tag{10} \label{eq:10}$$

And the characteristic length ξ is the length that $\Gamma(k)$ decays to 0. Thus the correlation length is given by

$$\Gamma(k) \propto \Gamma_0 \exp(-k/\xi), \quad k \gg 1$$
 (11)

1.c. Phase transition in Potts model

The Potts model undergoes a phase transition at a critical temperature T_* . Below T_* , the system is in a ferromagnetic phase, where the spins are aligned. Above T_* , the system is in a paramagnetic

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$$T_* = \frac{J}{k_B \ln(1 + \sqrt{q})} \tag{12}$$

where k_B is the Boltzmann constant and q is the number of spin states. In the simulation approach, the critical temperature can be determined by finding a peak in the specific heat.

2. PROBLEM SETUP

In the simulation, a 2D square lattice with periodic boundary is used as the target system. The lattice has size N=32. For simplicity, we set the coupling constant J=1 and the magnetic field strength h=0. Besides we use the reduced temperature s.t. $k_B=1$. q is set to 3, which means the spin variable can take on 3 different values, 1, 2, and 3.

The initial state of the system is randomly generated. The simulation is performed under different reduced temperatures T from 0.5 to 2.5. For each temperature, the system is evolved for 500,0000 steps. The trajectory of the system is recorded every 100 steps and thus the physical quantities are calculated with the same interval.

We selected the last half of the trajectory for analysis. The first half of the trajectory is used to ensure the system has reached equilibrium. The physical quantities are calculated by averaging over the last half of the trajectory.

3. IMPLEMENTATION OF THE SIMULATION

3.a. Setup the dependencies

In the simulation, numpy is used for the main computation. For better performance, numba is used to compile the functions just in time. multiprocessing is used to parallelize the computation, and functools.partial is used to create partial functions for the parallelization.

```
import numpy as np
from functools import partial
from numba import njit, prange, objmode, jit
import multiprocessing as mp
```

3.b. The Potts Hamiltonian

The Hamiltonian of Potts model is implemented as the following potts energy function

@njit def potts_energy(spins, J, h): # 获取形状 N = spins.shape[0]# 计算相邻自旋互作用能 E J = J * np.sum(spins[:-1, :] == spins[1:, :]) $E_J += J * np.sum(spins[:, :-1] == spins[:, 1:])$ # 计算外磁场能 $E_J += h * np.sum(spins)$ # 求和得到能量 $E = -E_J$ return E

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The implementation is straightforward. Using the slice operation, we can calculate the energy of the system by comparing the neighboring spins vertically and horizontally, and then summing up the results. The influence of the magnetic field is calculated by naively summing up all the spins. An decorator on jit is added to the function to compile it just in time. This will significantly improve the performance of the function.

3.c. The Metropolis algorithm

By splitting the Metropolis algorithm into the main loop and the operation of a single step, it is much more easier for numba to compile the code. The single step operation is implemented as the following potts mc step function

```
@niit(fastmath=True)
def potts mc step(spins, J, h, beta, q):
   # 获取形状
   N = spins.shape[0]
   # 选取随机格点
   i = np.random.randint(0, N)
   j = np.random.randint(0, N)
    # 建议新的自旋状态
    spins new = spins.copy()
    spins new[i, j] = np.random.randint(1, q + 1)
    # 直接计算能量差, 由于只改变了一个格点, 只需要计算该格点的能量差
    Eij old = 0
    Eij_new = 0
    Some code are omitted here for simplicity
   dE = Eij_new - Eij old
    # 判断是否接受
    if dE <= 0 or np.random.rand() < np.exp(-beta * dE):</pre>
       spins = spins new
    return spins
```

It is a standard implementation of the Metropolis algorithm. The only thing to note is that the energy difference dE is not calculated by comparing the energy of the new state and the old state. Instead, we only calculate the energy difference of the changed spin. The time complexity is thus reduced to O(1), which is much better than the naive implementation with time complexity $O(N^2)$. The main loop of the Metropolis algorithm is implemented naively and is not shown here. The sampled trajectories are recorded in the format of a numpy array as .npy files in the disk for further analysis.

3.d. Parallel simulation of different temperatures

The simulation of different temperatures is parallelized by using the multiprocessing module. The main idea is to create a partial function of the Metropolis algorithm with a fixed temperature. Then we can use the map function of the multiprocessing. Pool class to run the simulation of different temperatures in parallel. The implementation is shown as the following potts simulate parallel function

```
def potts_simulate_parallel(init_spin, J, h, q, k, T_series, n_steps, n_step_save=100):
   # 创建进程池
   pool = mt.Pool(8)
   # 定义部分函数
   potts mc partial = partial(
```

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```
potts mc mt,
    spins=init spin,
    J=J,
   h=h,
    q=q,
   n_steps=n_steps,
   n_step_save=n_step_save,
)
# 运行并行模拟
pool.map(potts mc partial, T series)
return 0
```

The potts mc partial function is a partial function of the potts mc mt function, which is the main loop of the Metropolis algorithm, in which parameters other than temperature are all fixed. The potts mc partial function is then mapped to the T series array, which contains the temperatures of the simulation. The potts mc partial function is run in parallel with 8 processes by using the multiprocessing. Pool class. The parallelization w.r.t. h can be implemented in a similar way.

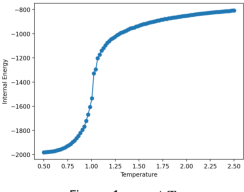
3.e. Data analysis

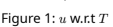
The interested physical quantities are calculated by averaging over the last half of the trajectory. The implementation is shown is the attached notebook Post-simulation.ipynb. Simply speaking, the data is loaded from the disk and then averaged over the last half of the trajectory. For internal energy, specific heat, and magnetization, it is done naively. For the characteristic length, the spatial correlation function is calculated first and then the characteristic length is calculated by fitting the correlation function after taking the logarithm. The limiting behavior around the critical temperature is also studied by fitting the data around the critical temperature using the similiar approach.

4. NUMERICAL RESULTS

4.a. Internal energy and specific heat

The internal energy and specific heat with respect to the temperature are shown in the following figure.





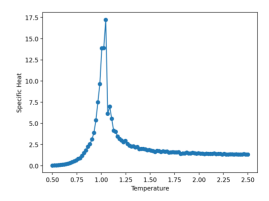


Figure 2: c w.r.t T

A peak is observed which is a obvious sign of phase transition. Before the critical temperature, the specific heat ismonotonically increasing while after the critical temperature, it is monotonically decreasing. The critical temperature is around

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$$T_* = 1.05$$
 (13)

while the theoretical value of the critical temperature is

$$T_*^t = \frac{1}{\ln(1+\sqrt{3})} = 0.995 \tag{14}$$

which is very close to the simulated value.

By using renormalisation techniques,the theory predicts that for $q \leq 4$, the ferromagnetic to paramagnetic phase transition is second order and otherwise first order. In the simulation, we found that the internal energy is a monotonically increasing function of the temperature. Besides, excluding some unnormally large values, u is almost continuous. Discontinuity is observed in the specific heat around the peak. Together with the continuity of the internal energy, we can conclude that the phase transition is **second order**, which is consistent with the theory.

4.b. Magnetization

The magnetization with respect to outer magnetic field under different temperatures is shown in the following figure

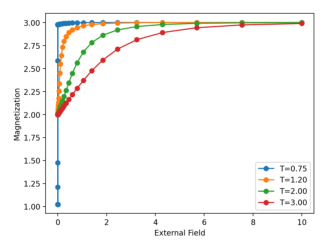


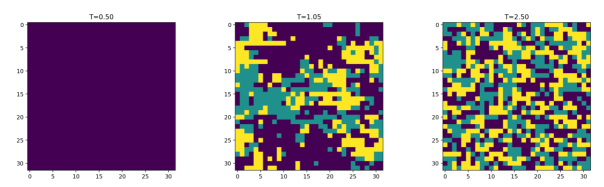
Figure 3: m w.r.t h

The dependence of the magnetization on the outer magnetic field clearly shows the magnetic property of 2D lattice before and after phase transition. When temperature is lower than T_* , the lattice is in a ferromagnetic phase, which means that once the outer magnetic field is applied, the spins will be aligned along the direction of the outer magnetic field, even the outer magnetic field is very weak. The T=0.75 line in Figure 3 shows this property. We can easily observe that the magnetization has a huge jump near h=0, and the lattice is almost fully magnetized, symbolized by the magnetization close to 3.

When temperature is higher than T_* , the lattice is in a paramagnetic phase, which means that the spins will not be aligned along the direction of the outer magnetic field unless the outer magnetic field is strong enough. The other lines in Figure 3 shows this property. In fact, we found that with the increase of temperature, the magnetization curve becomes more and more smooth, indicating the influence of the outer field becomes more obvious.

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Another thing to note is the magnetization without outer magnetic field. For ferromagnetic phase, the magnetization is close to 1, which means that the spins are almost fully aligned. On the other hand, for paramagnetic phase, the magnetization is close to 2, leading to the randomlly oriented nature of the spins. The below figure shows the states of the lattice under different temperatures.



For $T=0.5 < T_*$, the spins are almost fully aligned. For $T=1.05 \approx T_*$, the spins are mixture of oriented and random states, resulting in the jump of specific heat at the critical temperature. For $T=2.5\gg T_*$, the spins are almost fully aligned again. This is consistent with the magnetization curve and theoretical prediction.

4.c. Characteristic length

The characteristic length with respect to the temperature is shown in the following figure

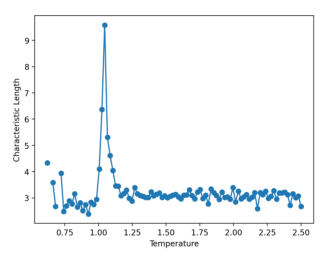


Figure 7: ξ w.r.t T

The fluctuation of the characteristic length at lower temperature is very large, indicating that the system is not well equilibrated. However, the rough trend of the characteristic length is still observable. Similar to the specific heat, the characteristic length shows a peak at the critical temperature. We found the peak and identified the corresponding temperature as the critical temperature. The critical temperature is around $T_{*}{}'=1.05$, which is the same as the critical temperature identified by the specific heat.

When temperature is very low, the characteristic length is very large, which means that the spins are highly correlated. When temperature is extrapolated to 0, the characteristic length is infinite, which means that the spins are fully aligned.

November 28, 2023 7 of 9 The correlation length at the critical temperature should be infinite theoretically. However, in the simulation, the fact is hard to observe. Since we are not able to simulate the system at the accurate critical temperature (it is even not a rational number!), the infinite correlation length is not observed.

4.d. Limiting behavior around the critical temperature

The limiting hehaviour of the specific heat and characteristic length around the critical temperature is shown in the following figure

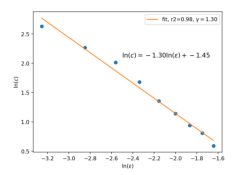


Figure 8: Limiting behaviour of c, left side

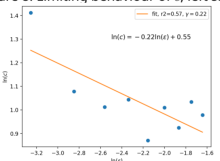


Figure 10: Limiting behaviour of ξ , left side

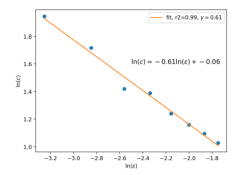


Figure 9: Limiting behaviour of c. right side

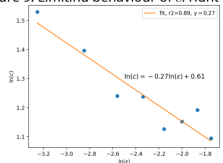


Figure 11: Limiting behaviour of ξ , right side

As a conclusion, the scaling exponents γ and δ are listed in the table below

Table 1: Scaling exponents

	Left side	Right side
γ	1.30	0.61
δ	0.22	0.27

It is necessary to emphasize the fact that the system is not well converged. Especially for the characteristic length under temperature lower than T_{\star} , the fluctuation is very large, which means that the scaling exponent is not accurate.

5. Issues and solutions

The most serious issue I encountered is the convergence of the simulation. Since the state space of the system is very large, it is very hard to ensure the system is well equilibrated. A naive solution is to increase the simulation steps. However, this will significantly increase the time cost. The interpreted nature of Python makes the situation even worse. Several tricks are used to improve the performance of the simulation and in the following section I will discuss them.

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5.a. Vectorization and correct usage of numpy

The first thing to note is that numpy is a very powerful tool for scientific computation. However, it is not a magic tool. The performance of numpy is highly dependent on the correct usage. For example, the naive implementation of the Metropolis algorithm has a time complexity of $O(N^2)$, which is very slow. By using np. roll for periodic boundary condition and np. sum for summing up the energy, the time complexity is reduced to O(1), which is much faster. Another improvement brought by usage of numpy is to bypass the global interpreter lock (GIL) and make full use of the CPU resources.

5.b. Reduction of unnecessary computation

The second thing to note is that unnecessary computation should be avoided. For example, in the Metropolis algorithm, the energy difference is calculated by comparing the energy of the new state and the old state. However, this is not necessary. Since we only change one spin, we only need to calculate the energy difference of the changed spin. This will reduce the time complexity from $O(N^2)$ to O(1). Even if the energy calculation is vectorized, it is still much slower than the current implementation.

5.c. Just-in-time compilation

The third thing to note is that numba is a very powerful tool for accelerating the computation. Although we optimize the code a lot, for Metropolis simulation tasks, the computation density is still quite low due to ubiquitous control flow, including if and for statements. Thus, the bottleneck of the simulation migrated from the computation to the control statements.

By using the @njit decorator, the whole functions will be compiled just in time. This will significantly improve the performance of the simulation. However, it is not a magic tool. Many details should be treated carefully. And the performance of numba is highly dependent on the correct usage. Many functions are not available under the most efficient nopython mode. However, by mixing the usage of nopython and object modes, the performance of the simulation can still be largely improved.

5.d. Multiprocessing and parallelization

The fourth thing to note is that multiprocessing is a very powerful tool for parallelization. In this task, we are insterested in the physical quantities w.r.t. temperature and outer magnetic field. Additionally, the MC simulation is independent for different temperatures and outer magnetic fields. Thus, we can parallelize the simulation w.r.t. temperature and outer magnetic field. By using the multiprocessing. Pool class, we can easily parallelize the simulation to take full advantage of the multi-core CPU resources. When similar strategy is used in much larger cases, it is called 'high throughput computing' and is widely used in scientific computation. Even if for each single task, the performance is not improved, sometime even worse, the total performance can be largely improved by parallelization.

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