## Part 5

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#### 0.0.1 Submitted by: Anushka and Yashasvee

### 0.0.2 Answer 5: Autocorrelation time for the absolute magnetization

This is only for part 5. Sorry for the confusion.

```
[55]: # Necessary modules
      %matplotlib inline
      import matplotlib.pyplot as plt
      import numpy as np
      import random as r
      from random import choice
      import sympy
      # We now work to generalize the Ising model to 2-Dimensions and or that the
       \hookrightarrow lattice size becomes (N_x) \times (N_y) where we
      # assume periodic boundary conditions in both, x and y, directions. It is given
       ⇔that we use the same coupling constant in
      # both directions.
      # Number of sweeps for thermalization for every value of J,h
      n_{therm} = 20000
      # Number of measurements for each J,h
      n_meas = 100
      # External magnetic field
      h = 0
```

```
skm = np.fft.ifft2(s, norm = 'forward')
          return np.fft.fft2(skm*skp, norm = 'backward')/s.size**2
[57]: J_c = 0.44068 \# Critical J
[58]: \# To calculate change in energy after one spin flips at position (x,y)
      # This is the general formula and we have already defined h=0 above
      def energy_flip(s,J,h,x,y,n):
            Here, the s is the spin, x and y are th positions of the 2 D lattice,
       \hookrightarrow J, n, and h are the same variables as defined
            before.
            The function calculates the energy after the spin at one site is flipped.
          return 2 * s[x][y]*(J*(s[((x+1)%n)][y] + s[((x-1)%n)][y] + s[x][((y+1)%n)]_{U}
       \hookrightarrow+ s[x][((y-1)\%n)]) + h)
      def energy(s,J,h,x,y,n):
          return -J*((s[((x+1)\%n)][y] + s[((x-1)\%n)][y] + s[x][((y+1)\%n)] +_{\sqcup}
       \Rightarrows[x][((y-1)\%n)])*s[x][y])-h*s[x][y]
[59]: # fft of sx and sy
      def variables(n,J,h,obs):
      # Defining local arrays
          m = np.array([]) # Magnetization
          E = np.array([]) # Energy
          m absolute = np.array([]) # Absolute value of magnetization
          prob = np.array([]) # Probability information
      # Now we need to assign spins to the site and because this is a 2D lattice, ___
       we have to keep in mind the dimensions
         x and y and do that. This has been done as follows:
      # The variable (s) is the spin here and n is some random integer
          s = [[choice((+1,-1)) \text{ for } x \text{ in } range(n)] \text{ for } y \text{ in } range(n)]
          for j in range(n_therm):
                 Assigning a random integer value to x and y
               x = np.random.randint(n)
              y = np.random.randint(n)
      # We need to define the change of energy after flipping the lattice site_
       \rightarrow (x,y) picked randomly. We call the defined
```

# print(skp)

```
function energy flip to calculate that.
              delta_energy = energy_flip(s,J,h,x,y,n)
              if delta_energy < 0:</pre>
                  s[x][y] *= -1
                                       # Condition to accept the spin flip
              else:
                  if np.random.uniform(0,1) <= np.exp(-delta_energy):</pre>
                      s[x][y] *= -1
          for i in range(n_meas):
              for j in range(n**2): #sweeping the lattice
                  x = j \% n
                  y = j // n
                  delta_energy = energy_flip(s,J,h,x,y,n)
                  if delta_energy < 0:</pre>
                      s[x][y] *= -1
                                            # Condition to accept the spin flip
                  else:
                      if np.random.uniform(0,1) <= np.exp(-delta_energy):</pre>
                           s[x][y] *= -1
                          prob = np.append(prob,1.) # Accept.
                      else:
                           prob = np.append(prob,0.) # We reject the other values.
                        Measurements
              if obs == "m":
                  m=np.append(m,np.mean(s))
                  obs=np.mean(m)
              if obs == "E":
                  temp = np.array([])
                  for x in range(n):
                      for y in range(n):
                           temp = np.append(temp,energy(s,J,h,x,y,n))
                  E=np.append(E,np.mean(temp))
                  obs=np.mean(E)
              if obs == "m_absolute":
                  m_absolute=np.append(m_absolute,np.absolute(np.mean(s)))
                  obs=np.mean(m_absolute)
          return obs
[60]: # plt.figure(figsize=(12,10))
      # plt.title("C(s)", fontsize=12)
      # plt.grid()
```

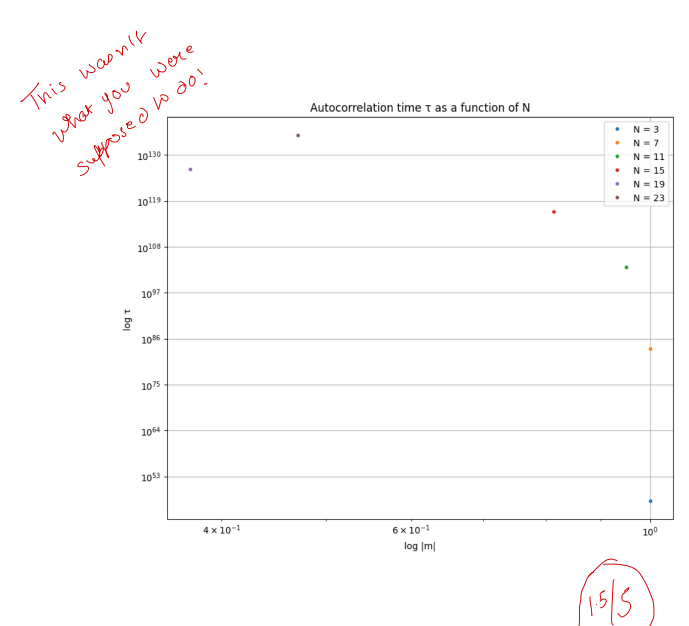
# plt.plot(C,r)
# plt.legend()

```
[61]: def int_autocorrelation(N):
    for z in range(1,100):
        tau_int = N**z
    return tau_int

tau_int3 = int_autocorrelation(3)
tau_int7 = int_autocorrelation(7)
tau_int11 = int_autocorrelation(11)
tau_int15 = int_autocorrelation(15)
tau_int19 = int_autocorrelation(19)
tau_int23 = int_autocorrelation(23)
```

Autocorrelation time as a function of N

```
[63]: # Now plotting the required variables
      J = 0.5
                  # Keeping J fixed at a certain value > J_c
      h = 0
      # We are supposed to take different values of N (N_x = N_y), so we define as
      ⇔follows:
      mag_3 = [variables(3,J,h,"m_absolute")]
      mag_7 = [variables(7,J,h,"m_absolute")]
      mag_11 = [variables(11,J,h,"m_absolute")]
      mag 15 = [variables(15,J,h,"m absolute")]
      mag_19 = [variables(19,J,h,"m_absolute")]
      mag_23 = [variables(23,J,h,"m_absolute")]
      # For different values of N:
      plt.figure(figsize=(10,8))
      plt.title("Autocorrelation time as a function of N", fontsize=12)
      plt.grid()
      plt.ylabel("log
      plt.xlabel("log |m|")
      plt.loglog(mag_3, tau_int3,'.', mag_7, tau_int7,'.', mag_11,tau_int11,'.',u
       →mag_15, tau_int15,'.', mag_19, tau_int19,'.', mag_23, tau_int23,'.')
      plt.legend(("N = 3", "N = 7", "N = 11", "N = 15", "N = 19", "N = 23"))
      plt.show()
```



# 0.0.3 Answer 5 : Dynamical variable

Port 6 not othern bred! []: