

993SM - Laboratory of Computational Physics lecture 8 - part 1 April 29, 2020

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The Ising model in the canonical ensemble

- Introduction to the Ising model
- The Ising model in the canonical ensemble: application of Metropolis Monte Carlo Method
 - Implementation in a code

M. Peressi - UniTS - Laurea Magistrale in Physics Laboratory of Computational Physics - Unit VIII

stochastic simulations of an interacting spin ensemble

We need a model for:

- energy

evolution
 (to build the trajectory / the Markov chain)

stochastic simulations of an interacting spin ensemble

We need a model for:

- energy => Ising model

evolution
 (to build the trajectory / the Markov chain)

Ising model

the simplest model of interacting spin on a lattice

$$\mathcal{H}^{spin} = -J \sum_{i,j=1}^{N} s_i s_j$$

$$s_i = \pm 1$$
 $\downarrow \qquad \uparrow \qquad \uparrow$
 $\downarrow \qquad \downarrow \qquad \uparrow$
 $\downarrow \qquad \downarrow \qquad \uparrow$
 $\downarrow \qquad \downarrow \qquad \uparrow$

But where does it come from?

Consider two fermions, $\vec{s_1}$, $\vec{s_2}$ (e.g. electrons in He) described by a hamiltonian not depending explicitly on spin:

$$\mathcal{H} = h_1 + h_2 + V_{12}$$

=> the eigenstates can be in a factorized form (symm/antisymm under particle exchange due to the Pauli exclusion principle):

$$\Psi_{+,-}(1,2) = \Psi_{+,-}^{orb}(1,2) \chi(1,2)$$

In the Hilbert spin subspace we can choose a coupled representation, referring to the spin sum S, with eigenstates labeled by $|S,S_z>$:

$$\Psi_{+,-}(1,2) = \Psi_{+,-}^{orb}(1,2) \; \chi_{S,S_z}(1,2) = \Psi_{+,-}^{orb}(1,2) \, | \, S,S_z > 0$$

Eigenstates

$$\Psi_{+,-}^{orb}(1,2)|S,S_z> = (\phi(1)\psi(2) + (-)^S\phi(2)\psi(1))/\sqrt{2} |S,S_z>$$

where $|S, S_z\rangle = \chi_{S,S_z}(\sigma_1, \sigma_2)$ have the explicit form:

$$\chi_{0,0} = \frac{1}{\sqrt{2}} \left[v_+(\sigma_1) v_-(\sigma_2) - v_-(\sigma_1) v_+(\sigma_2) \right] \quad \text{Spin singlet: antisymmetric}$$

Energy

$$<\Psi_{+,-}|\mathcal{H}|\Psi_{+,-}> = <\Psi_{+,-}^{orb}|\mathcal{H}|\Psi_{+,-}^{orb}> = <\phi|h|\phi> + <\psi|h|\psi> + J_{12}+(-1)^S K_{12}$$

= $E_0 + J_{12} + (-1)^S K_{12}$

with the Hartree and the exchange terms:

$$J_{12} = \langle \phi(1)\psi(2) | V_{12} | \phi(1)\psi(2) \rangle$$

$$K_{12} = \langle \phi(1)\psi(2) | V_{12} | \phi(2)\psi(1) \rangle$$

The Pauli principle => the energy is affected by spin even if ${\cal H}$ does not depend explicitly on it:

$$< S = 0 | \mathcal{H} | S = 0 >= E_s$$
 singlet $< S = 1 | \mathcal{H} | S = 1 >= E_t$ triplet

Hence:

$$< S = 0 |\mathcal{H}|S = 0 > - < S = 1 |\mathcal{H}|S = 1 > = E_s - E_t$$

8

$$=2K_{12}$$

Energy

The relative order of E_s , E_t depends on V_{12} (K_{12})

In He, with V_{12} repulsive Coulomb interaction,

$$E_s > E_t \quad (K > 0)$$

(a part from the ground state, when only singlet is allowed)

But present discussion is independent on the sign of $E_s - E_t$

Idea: write a model hamiltonian explicitly dependent on spin (and only dependent on spin) which gives the same difference between the energy eigenvalues of its eigenstates as the original one.

Consider the operator:

$$\sum_{12} = \vec{s_1} \cdot \vec{s_2} = \frac{1}{2}S^2 - \frac{3}{4}$$

which is diagonal on the coupled basis, with eigenvalues:

$$\langle S = 0 | \Sigma_{12} | S = 0 \rangle = -\frac{3}{4}, \quad \langle S = 1 | \Sigma_{12} | S = 1 \rangle = \frac{1}{4}$$

Consider then:

$$\mathcal{H}^{spin} = -(E_s - E_t)\Sigma_{12}$$

We have:

$$< S = 0 | \mathcal{H}^{spin} | S = 0 > - < S = 1 | \mathcal{H}^{spin} | S = 1 > = E_s - E_t$$

Heisenberg hamiltonian

$$\mathcal{H}^{spin} = -(E_s - E_t)\Sigma_{12}$$

is therefore OK! Defining: $J \equiv E_s - E_t$, we have (going back to the individual basis representation):

$$\mathcal{H}^{spin} = -J\vec{s_1} \cdot \vec{s_2}$$

Heisenberg hamiltonian

Extension to the case of several spins:

$$\mathcal{H}^{spin} = -\sum_{\substack{i,j=1\i
eq j}}^{N} J_{ij} \ \vec{s_i} \cdot \vec{s_j}$$

(see also: Ashcroft & Mermin, chap. 32, "Spin hamiltonian and Heisenberg model")

Ising model

Consider only the possibility : $s_i = \pm 1$ and nearest neighbor interaction only, with the same interaction constant J

$$\mathcal{H}^{spin} = -J \sum_{i,j=1}^{N} s_i s_j$$

J>0 (Es > Et)
$$\uparrow \uparrow$$
 spins favored => ferromagnetic case

J<0 (Es < Et) $\uparrow \downarrow$ spins favored => antiferromagnetic case

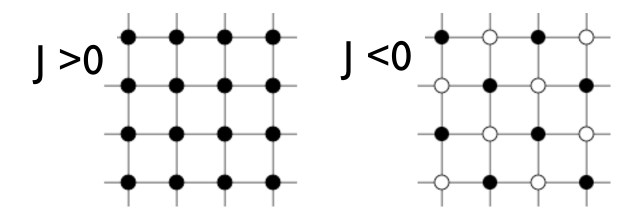
 $\downarrow \downarrow \downarrow$
 $\downarrow \downarrow$

(choosing the kind of interaction, we specify the energy model)

Ising model on a lattice

lattice containing N sites

lattice site i has associated with it a number s_i , where $s_i = +1$ for an "up" (\uparrow) spin and $s_i = -1$ for a "down" (\downarrow) spin. A particular configuration or microstate of the lattice is specified by the set of variables $\{s_1, s_2, \ldots s_N\}$ for all lattice sites.



Lowest energy state of the 2D Ising model on a square lattice with ferromagnetic (J>0) and antiferromagnetic (J<0) interactions. Solid and open circles correspond to +1 and -1 spins, respectively.

Ising model: interesting quantities

Ising model: energy

lattice containing N sites No external magnetic field:

$$E = -J \sum_{i,j=\text{nn}(i)}^{N} s_i s_j$$

(nn=nearest neighbor)

Energy in presence of an external magnetic field:

$$E = -J \sum_{i,j=\text{nn}(i)}^{N} s_i s_j - H \sum_{i=1}^{N} s_i,$$

or, better, define an average energy per spin: E/N

Ising model: magnetization

$$M = \sum_{i=1}^{N} s_i$$

"Order parameter": total magnetization, or -better-average magnetization per spin:

$$m = \frac{M}{N} = \frac{1}{N} \sum_{i=1}^{N} s_i$$

$$-1 < m < +1$$

Ising model: configurations and energy

2ⁿ different configurations for *n* spins. (microstates)

e.g. $2^4 = 16$ spin configurations for $2x^2$ lattice

For J > 0 the state of lowest energy is when all the spins are aligned. The state has macroscopic magnetization (**ferromagnetic**).

The ground state energy per spin

(ferromagnetic case, thermodynamic limit (N large), no ext. field) is:

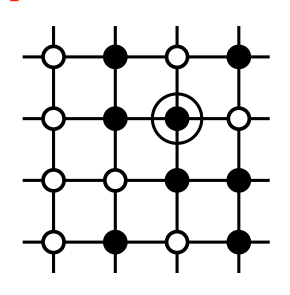
$$E_0/N = -2J$$

Ising model: dynamics?

Beside an energy model, we must define a dynamics in order to simulate the evolution of the system (i.e. to generate the trajectory in the phase space, to generate the configurations of the Markov chain)

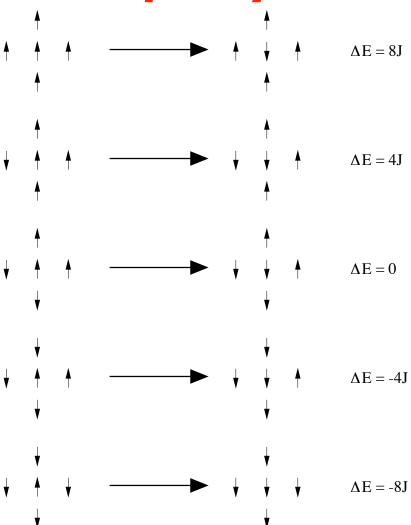
Ising model: spin flip dynamics

Consider nn interactions, choose a random spin and flip it: it's a new configuration (a microstate)



Apply **Metropolis Monte Carlo** method for evolution in the **canonical ensemble** (fix T). Evolution is driven by the **energy change** between the old and the new configuration, ΔE . Remark: Is it sufficient to calculate only ΔE , not E at each new configuration!

Ising model: spin flip dynamics



The five possible transitions of the Ising model on the square lattice with spin flip

Ising model: boundary conditions

Of course we cannot simulate an infinite system (the thermodynamic limit).

We have two choices for the simulation cell:

- free (open) boundary conditions
- periodic boundary conditions (PBC)

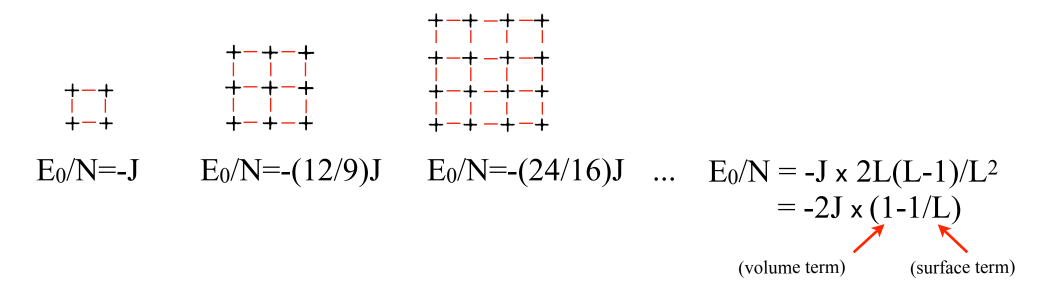
Ising model: free boundary conditions

in a N=LxL spin lattice there are 2L(L-1) nn interactions; for the ferromagnetic g.s. configuration, for instance, the energy is:

Let's count the interactions...

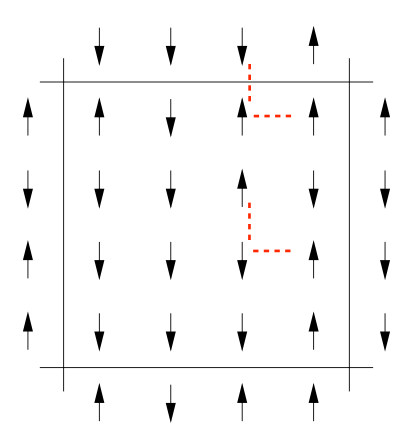
Ising model: free boundary conditions

in a N=LxL spin lattice there are 2L(L-1) nn interactions; for the ferromagnetic g.s. configuration, for instance, the energy is:



Energy per spin in the ground state converges to the value $E_0/N = -2J$ in the thermodynamic limit (with deviations $\sim 1/L$)

Ising model: PBC



The energy is a 2N-term sum: each spin interacts with its NN within the simulation cell or with the NN images

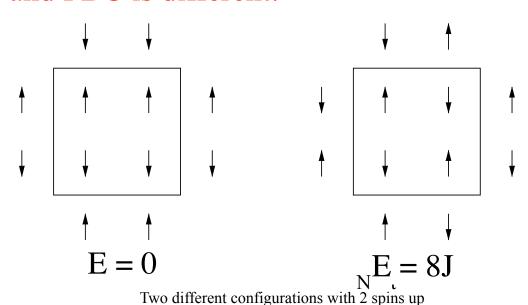
One of the 2^N possible configurations of a system of N=16 Ising spins on a square lattice. with periodic boundary conditions.

Ising model: PBC

We have always:

$$2^4 = 16$$
 spin configurations for $2x^2$ lattice_N

but the energy for each configuration in case of free boundary conditions and PBC is different:



# of spins UP	Degeneracy	Energy	Magnetisation
4	1	-8J	4
3	4	0	2
2	4	0	0
2	2	8J	0
1	4	0	-2
0	1	-8J	-4

Energy and magnetization of 16 configurations of the 2×2 Ising model with PBC

Energy per spin in the ground state is always equal to the value $E_0/N = -2J$ in the thermodynamic limit

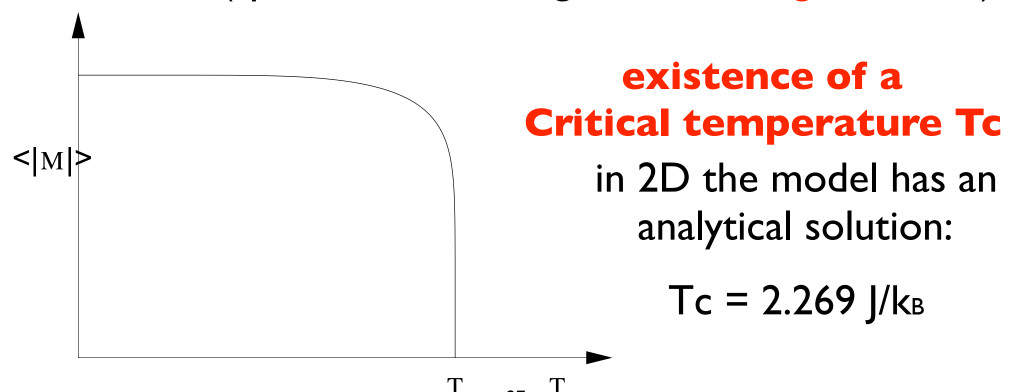
Ising model: phase transition

Low T: spin configuration minimizes energy

(if J>0: spins tend to align => high (absolute) magnetization)

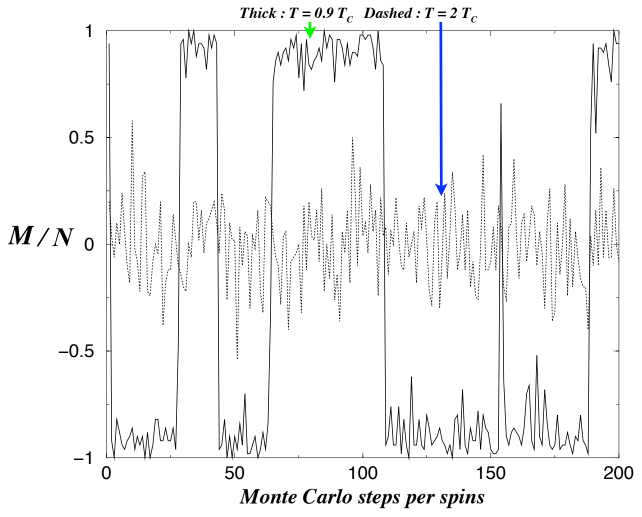
High T: spin configuration maximizes entropy

(=disorder) (spins tend to disalign => low magnetization)

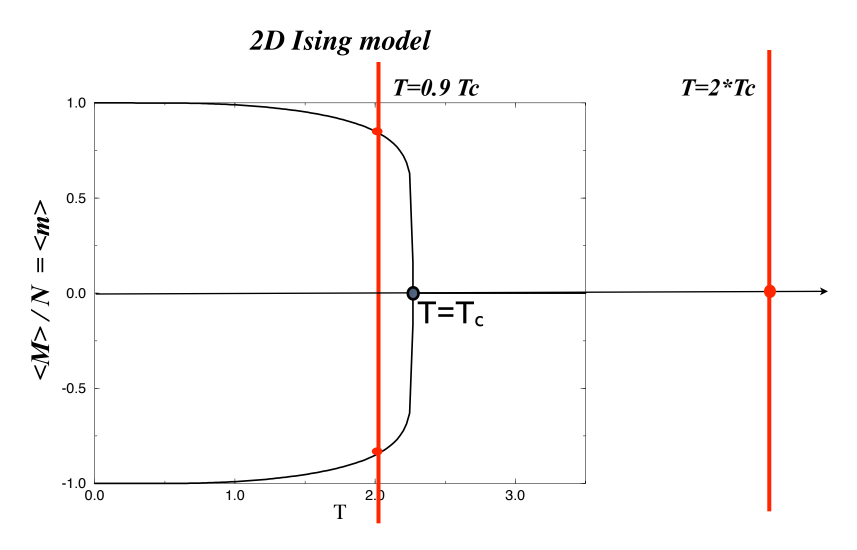


Apply **Metropolis Monte Carlo** method for evolution in the **canonical ensemble** (fix T):

Average magnetization per spin as a function of time for a lattice 10×10 and two different T

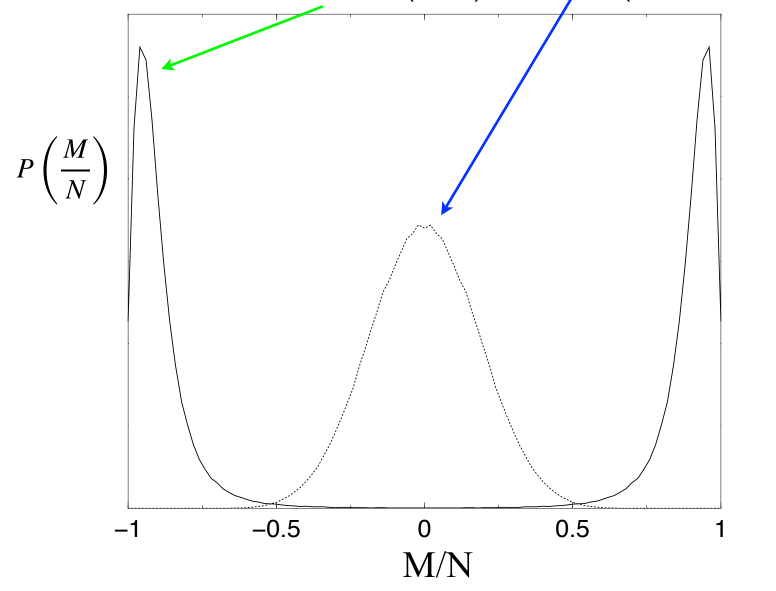


Fluctuations! do, as usual, temporal averages: < M > /N, < E > /N



Magnetisation as a function of the temperature for the 2D Ising model. (data averaged on time; red lines indicate the T values in the previous plot)

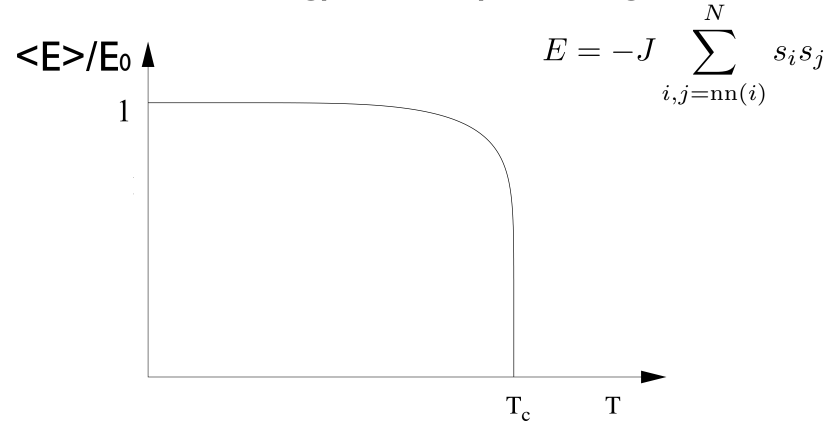
Magnetization distribution for $T < T_c$ (solid) and $T > T_c$ (dashed curve)



(data collected during time evolution, at equilibrium)

Ising model: phase transition

T_c also for energy, not only for magnetization:



and also the energy fluctuates during time evolution...

Ising model: fluctuations

Fluctuations are intrinsic to the system evolution and are important!

Linear response functions are related to equilibrium fluctuations:

(already proved):
$$C=\frac{\partial\langle E\rangle}{\partial T}$$
 , $C=\frac{1}{kT^2}\left(\langle E^2\rangle-\langle E\rangle^2\right)$

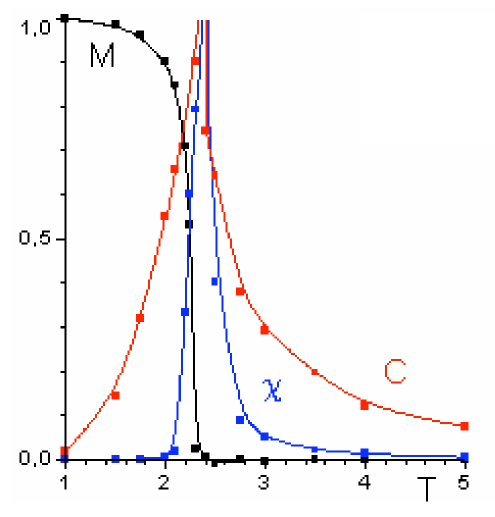
but also:
$$\chi = \lim_{H \to 0} \frac{\partial \langle M \rangle}{\partial H}$$
 , $\chi = \frac{1}{kT} (\langle M^2 \rangle - \langle M \rangle^2)$

where $\langle M \rangle$ and $\langle M^2 \rangle$ are evaluated in zero magnetic fields.

Ising model:

fluctuations and phase transition

Rapid change in <E> and <M> => singularities in C and χ



(Large fluctuations near the phase transition: Second Order phase transition)

specific heat:

$$C = \frac{\partial \langle E \rangle}{\partial T}$$

magnetic susceptibility:

$$\chi = \lim_{H \to 0} \frac{\partial \langle M \rangle}{\partial H}$$



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Implementing the Ising model in the code

Implementing the Ising model

on a 2D square lattice in the canonical ensemble zero-field, nearest neighbor interactions only

$$\mathcal{H}^{spin} = -J \sum_{i,j=1}^{N} s_i s_j \qquad s_i = \pm 1$$

Input parameters are:

- L (linear lattice dimension, which gives the number of spins: N=L*L)
- nmcs (number of total MC steps per spin)
- nequil (number of equilibration MC steps per spin)
- T (temperature of the thermal bath).

Implementing the Ising model

```
program ising
    metropolis algorithm for the ising model on a square lattice
  use common
  integer :: imcs,ispin,jspin
  real (kind = double), dimension(5) :: cum
 call initial(nequil,cum)
  ! equilibrate system
  do imcs = 1,nequil
     call metropolis()
  end do
     accumulate data while updating spins
  do imcs = 1,nmcs
     call metropolis()
     call data(cum)
  end do
  call output(cum)
end program ising
```

Ising model on a lattice

L: linear lattice dimension N = LxL: number of spins a configuration (a microstate) is the whole sequence of spins, i.e. the LxL array spin(x,y) module common integer, public, dimension(:,:), allocatable :: spin subroutine initial(nequil,cum) allocate(spin(L,L)) spin(x,y) = 1 se $s_i = \pm 1$ else spin(x,y) = -1

Ising model: magnetization

Total magnetization, or define an average magnetization per spin:

```
M = \sum s_i
compute initial magnetization
M = 0.0_{double}
                                m = \frac{M}{N} = \frac{1}{N} \sum_{i=1}^{N} s_i
do y = 1,L
   do x = 1,L
       M = M + spin(x,y)
    end do
                                             -1 < m < +1
end do
```

(Instead of the loop over x,y, write: M=sum(spin))

Ising model: energy

```
E = -J \quad \sum_{i=1}^{N} \quad s_i s_j \quad \uparrow \quad \downarrow \quad \uparrow
                                    i,j = \operatorname{nn}(i)
   compute initial energy
E = 0.0_double
do y = 1,L
   do x = 1,L
    sums = spin(x,up) + spin(right,y)
! calculate the initial energy summing all over pairs
! (for a given spin, consider only the up NN and the right NN
! - NOT the down and the left NN - : each interaction is counted once
           E = E - spin(x,y)*sums
        end do
    end do
```

Ising model: energy with PBC

do y = 1,L

```
periodic boundary conditions
       if (y == L) then
         up = 1
      else
         up = y + 1
      end if
      do x = 1,L
          if (x == L) then
             right = 1
         else
             right = x + 1
          end if
          sums = spin(x,up) + spin(right,y)
! calculate the initial energy summing all over pairs
 (gor a given spin, consider only the up NN and the right NN
! - NOT the down and the left NN - : each interaction is counted once
         E = E - spin(x,y)*sums
      end do
   end do
```

Ising model: spin flip dynamics

Choose a random spin and flip it: it's a new configuration (a microstate)

```
do ispin = 1,N
! random x and y coordinates for trial spin call random_number(rnd)
x = \text{int}(L*\text{rnd}) + 1 \qquad \qquad 1 \leq x \leq L
\text{call random_number(rnd)}
y = \text{int}(L*\text{rnd}) + 1 \qquad \qquad 1 \leq y \leq L
.....
```

Flip is: spin(x,y) = -spin(x,y)

but do it later, only if you decide to accept the flip (according to Metropolis)

Ising model: energy variations per spin flip

Evolution is driven by the **energy change** between the old and the new configuration (Metropolis MC)

```
dE = DeltaE(x,y) ← energy variation for spin(x,y) flip
call random_number(rnd)
if (rnd <= w(dE)) then ← w(dE) is e<sup>-ΔE/k<sub>B</sub>T</sup>
    spin(x,y) = -spin(x,y)
    accept = accept + 1
    ....

function DeltaE(x,y) result (DeltaE_result)
DeltaE_result = 2*spin(x,y)*(left + right + up + down)
```

Energy variations per spin flip with PBC

```
function DeltaE(x,y) result (DeltaE_result)
     periodic boundary conditions
if (x == 1) then
    left = spin(L,y)
   right = spin(2,y)
else if (x == L) then
    left = spin(L-1,y)
   right = spin(1,y)
 else
    left = spin(x-1,y)
    right = spin(x+1,y)
 end if
 if (y == 1) then
   up = spin(x,2)
    down = spin(x,L)
 else if (y == L) then
    up = spin(x,1)
    down = spin(x, L-1)
 else
   up = spin(x,y+1)
    down = spin(x,y-1)
 end if
 DeltaE_result = 2*spin(x,y)*(left + right + up + down)
```

Ising model: storage of Boltzmann's coeff.

```
! Choosing the interaction parameter J=1,
! possible energy variations per spin flip are -8,-4,0,+4,+8:
   do dE=-8,8,4
        w(dE) = exp(-dE/T)
   end do
   accept = 0
   do i = 1,5
        cum(i) = 0.0_double
   end do
```

Convenient to store the Boltzmann's coefficient for these discrete values of energy variations

The five possible transitions of the Ising model on the square lattice with spin flip

Ising model: updating energy and magnetization

```
subroutine metropolis()
     one Monte Carlo step per spin
  do ispin = 1,N
                                              DO NOT CALCULATE
                                             EVERYTHING FROM THE
     dE = DeltaE(x,y)
                                                    SCRATCH!!
     call random_number(rnd)
     if (rnd <= w(dE)) then
        spin(x,y) = -spin(x,y)
        accept = accept + 1
        M = M + 2*spin(x,y) ! factor 2 is to account for the variation:

E = E + dE ! (-(-)+(+))
     end if
                                 \Delta E is already a variation
  end do
end subroutine metropolis
```

Spin flip dynamics: how to choose spin to flip?

Random ...

```
do ispin = 1,N
! random x and y coordinates for trial spin call random_number(rnd)
x = \text{int}(L*\text{rnd}) + 1 \qquad \qquad 1 \leq x \leq L
\text{call random_number(rnd)}
y = \text{int}(L*\text{rnd}) + 1 \qquad \qquad 1 \leq y \leq L
.....
```

or ordered (sequential) ...

do
$$x = 1,L$$

do $y = 1,L$

. . .

$$spin(x,y) = -spin(x,y)$$

Spin flip dynamics: how to choose spin to flip?

- ORDERED: in some cases, it could go more slowly towards equilibrium (see later: correlation time), but it depends...
- NO appreciable differences in the statistics at equilibrium

Measuring physical quantities: how to accumulate data?

```
subroutine data(cum)
! accumulate data after every Monte Carlo step per spin
real (kind = double), dimension(5), intent (inout) :: cum
cum(1) = cum(1) + E
cum(2) = cum(2) + E*E
cum(3) = cum(3) + M
cum(4) = cum(4) + M*M
cum(5) = cum(5) + abs(M)
end subroutine data
```

After one MC step per spin for all spins:

Alternatively, do it after each MC step per individual spin...

Measuring physical quantities: how to accumulate data? Further remarks...

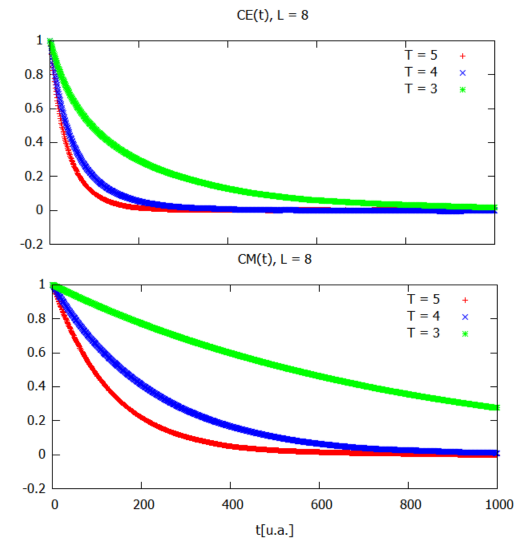
- Use statistically INDEPENDENT configurations
- Calculate therefore the CORRELATION TIME by considering the autocorrelation functions:

$$C_M(t) = \langle M(t)M(0) \rangle - \langle M \rangle^2, \quad C_E(t) = \langle E(t)E(0) \rangle - \langle E \rangle^2$$

$$(C_M(0) \propto \chi, \quad C_E(0) \propto C_V)$$

 $C_M(t) \to 0$ and $C_E(t) \to 0$ exponentially for $t \to \infty$ with a certain decay time τ : consider intervals longer than τ for statistical averages

Autocorrelation functions



(NOTE: "critical slowling down" for $T \to T_C$)

=> configurations change very slowly, and it is difficult to sample enough configurations

Measuring physical quantities: how to accumulate data? Further remarks...

- \bullet see also CORRELATION LENGTH between magnetic domains, $\zeta(T)$
- close to T_c, also the correlation length increases (spin alignments are more correlated), up to divergence

Measuring physical quantities: which errors?

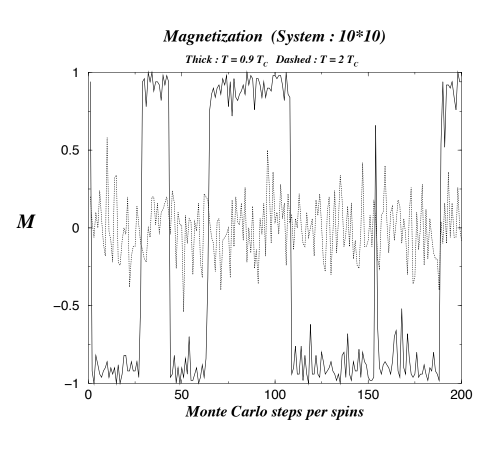
- necessary to give the ERROR ESTIMATE corresponding to the measured physical quantity !!!
- (see Tab. I of D.P. Landau, PRB 13, 2997 (1976), "Finite size behavior of the Ising square lattice")
- do also BLOCKING (called "coarse grained technique" in that paper)

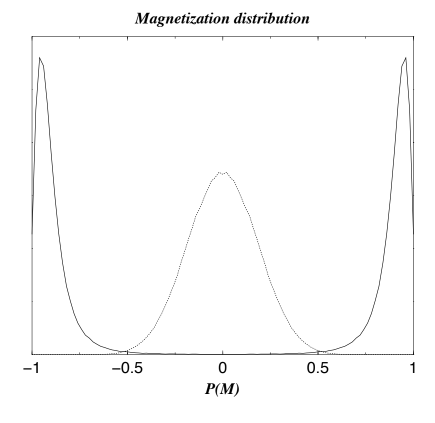
How to do efficiently simulations as a function of T?

- Sometimes EQUILIBRATION time is long...
- IDEA: for T' close to T, choose as starting point the equilibrated output of T

(see previous example, reported also on next slide)

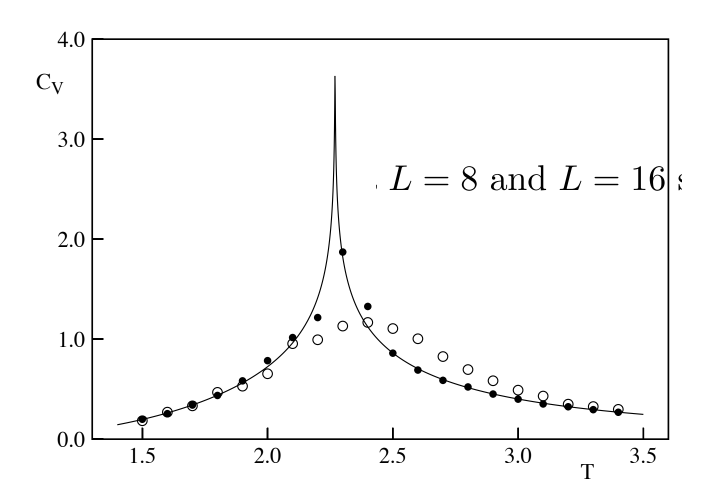
How to do efficiently simulations as a function of T?





Ising model: size problems

We cannot simulate an INFINITE system!



The temperature dependence of the specific heat C (per spin) of the Ising model

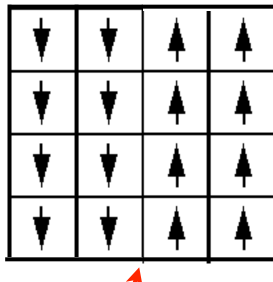
Ising model: size problems

INTERFACE EFFECTS:

example of energy for HALF UP/HALF DOWN configurations:

$$L=2$$
 $E=0$

$$L=4$$
 $E=-1$



"interface"

···· for an infinite system: E=-2 We have a ("interface") term proportional to I/L

Ising model: alternative dynamics

- in the SPIN FLIP dynamics the order parameter is not conserved (M changes during evolution)
- alternative: NN spin exchange (Kawasaki dyn.)
 (exchange two NN spins picked at random;
 M is conserved; this is equivalent to LATTICE
 GAS MODELS with fixed number of particles)

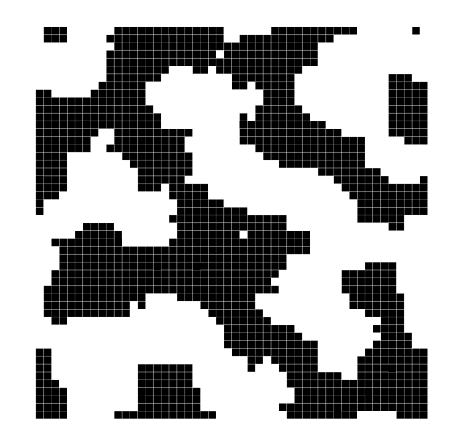
Ising model: Kawasaki dynamics

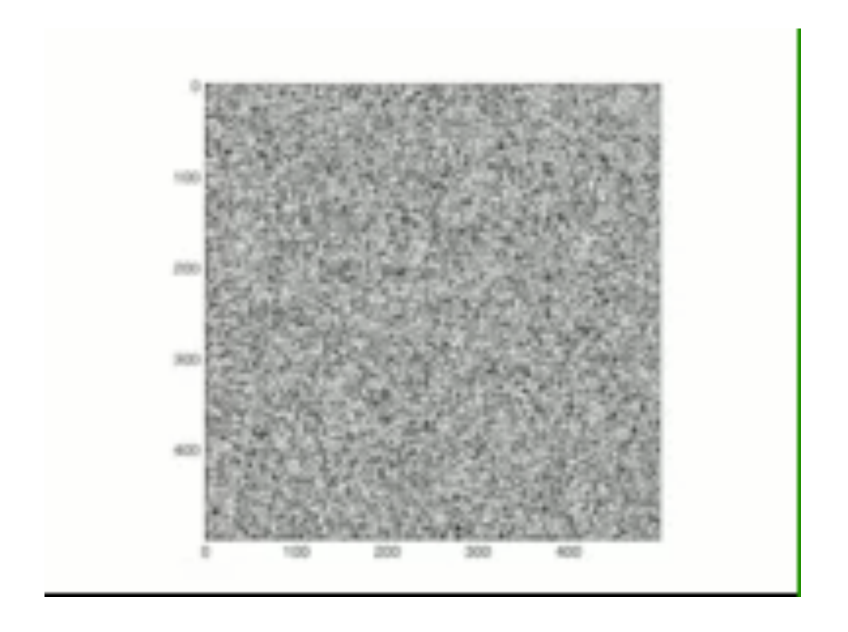
Fixed magnetization : change of thermodynamical ensemble

No modification of the equilibrium properties

except phase separation





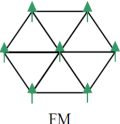


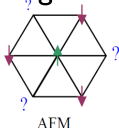
T=10, starting from random configuration

Ising model: other generalizations

- SPINS: XY, Heisenberg, Potts...
- LATTICES: Square, Triangle, Cubic, Honeycomb, Kagome....
- INTERACTIONS: Magn. Field, Antiferrom., Next Nearest Neighbor (NNN)....

Different behaviour according to the geometry and the kind of interactions. Example: frustration in the triangular antiferromagnetic Ising model:





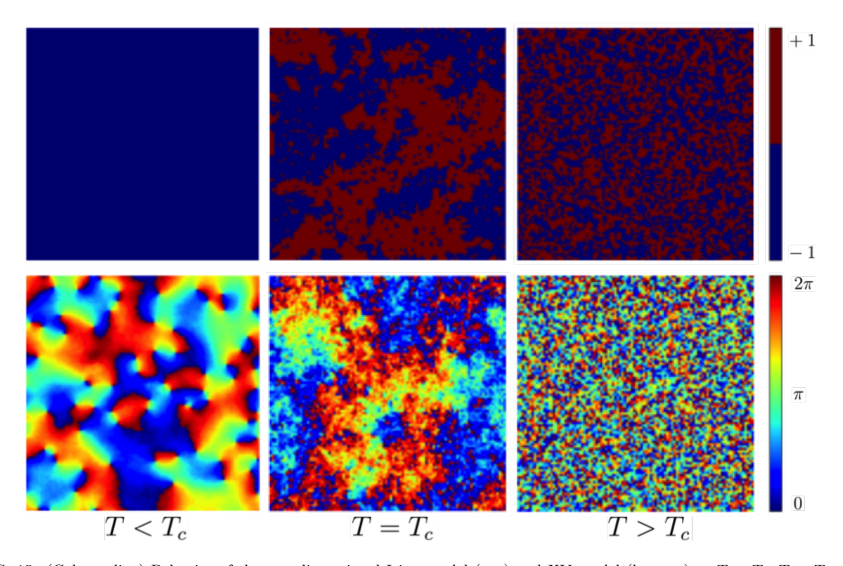


FIG. 15. (Color online) Behavior of the two dimensional Ising model (top) and XY model (bottom) at $T < T_c$, $T = T_c$ and $T > T_c$. For the Ising model, an \uparrow spin ($s_i = 1$) is represented by red and a \downarrow spin ($s_i = -1$) is represented by blue. For the XY model, because every spin is determined by θ , so $\mathbf{s}_i = (\cos \theta, \sin \theta)$, an angle of 2π is represented by red and an angle of 0 by blue. In the low-temperature phase, the Ising model exhibits spontaneous magnetization while in the XY model, vortex buddies appear (characterized by points where a continuum from blue to red, or viceversa, circle the point. It is worth to notice that these points are present by pairs with opposite circulation).

Program:

```
$\forall \text{sing} \\ \left[ \do: \percessi/comp-phys/VIII-ising/\ \text{sing}/\ \te
```

ising.f90

Exercise

(a) Choose L=30, T=2, and initially spin=±1 randomly. Calculating and plotting the instantaneous values of the energy E/N and the magnetization M/N per spin (averaged over the lattice) as a function of Metropolis-MC steps, how much time (i.e. how many nequil MC steps) is it necessary to equilibrate the system?

Hint:

- Since initially spin=±1 randomly, E/N and M/N initially will be far from the expected equilibrium average value.

First, set nequil=0 and plot instantaneous values of E/N and M/N Estimate nequil from that plot!!! Visualization is important!!!!

nequil of course depends on T and on the initial situation

Then, set *nequil* not zero and calculate the time average < E > /N and < M > /N; increasing the total *nmcs*, the two quantities should converge...

Exercise

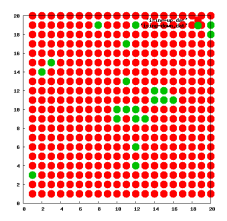
(a) Choose L=30,T=2, and initially spin=±1 randomly....

Plot a snapshot of the spin pattern: does the system appear ordered or disordered?

it should appear ordered...

p 'ising-up.dat' ps 3 pt 7, 'ising-down.dat' ps 3 pt 7

Plotting "ising-up.dat" and "ising-down.dat" which contain the coordinates of spin up and down respectively, one should get something like that:



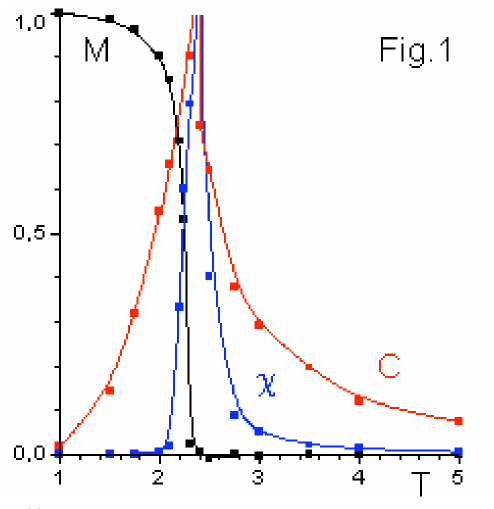
Calculate also c and χ .

Exercise

(a) Choose L=30,T=2, and initially spin=±1 randomly....

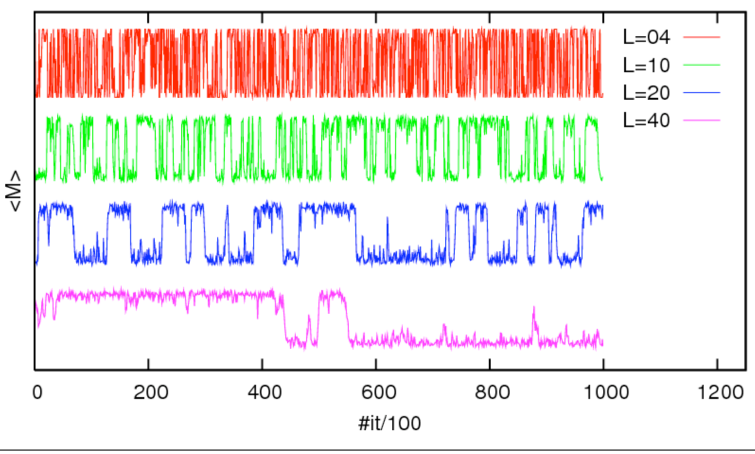
Calculate also c and χ .

(b) Choose T=I and repeat (a)...



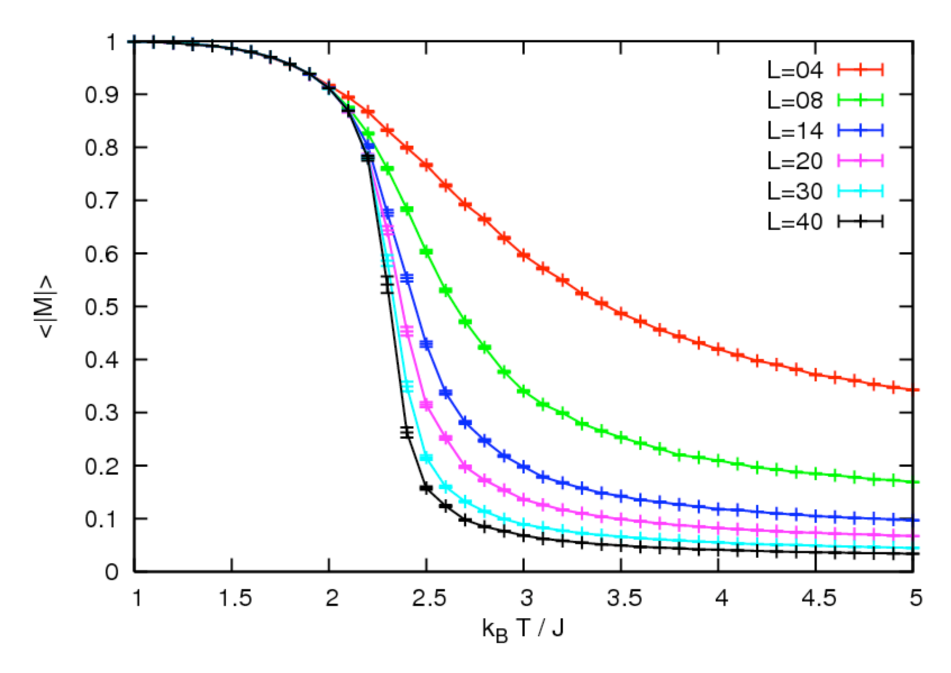
Raw data: traces, covariance and autocorrelation time

Trace: magnetization for $T=2.27J/k_B\approx T_C$ (10^5 sweeps)

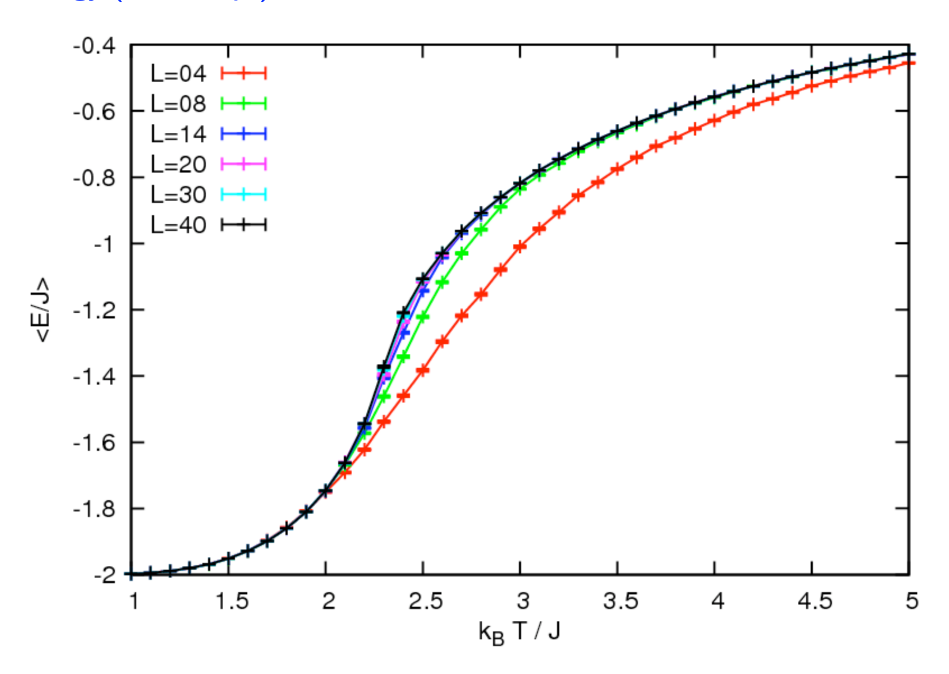


17

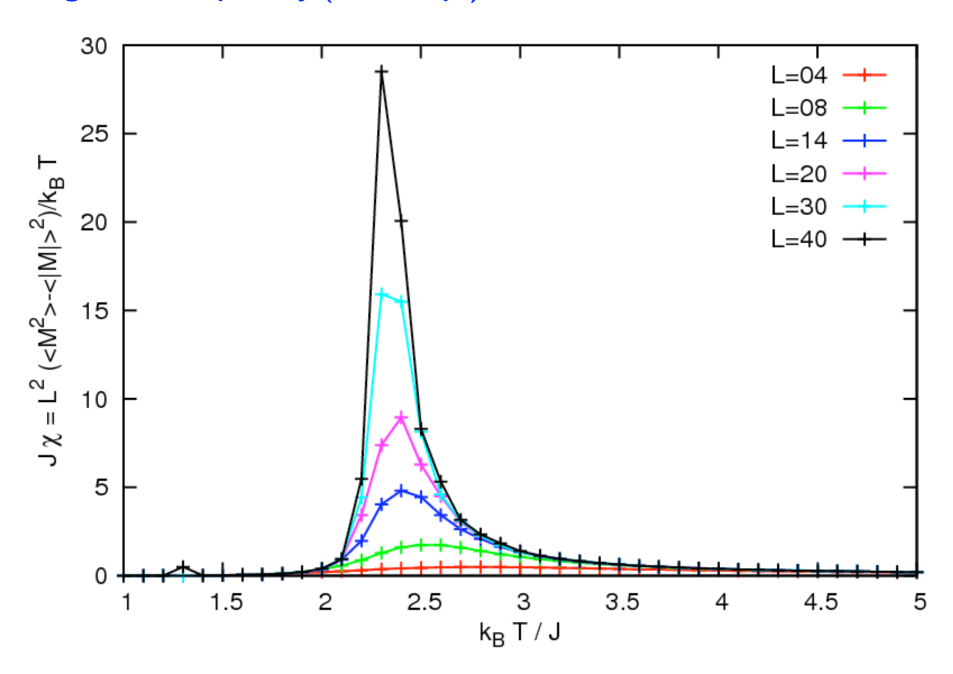
Magnetization (10^5 sweeps)



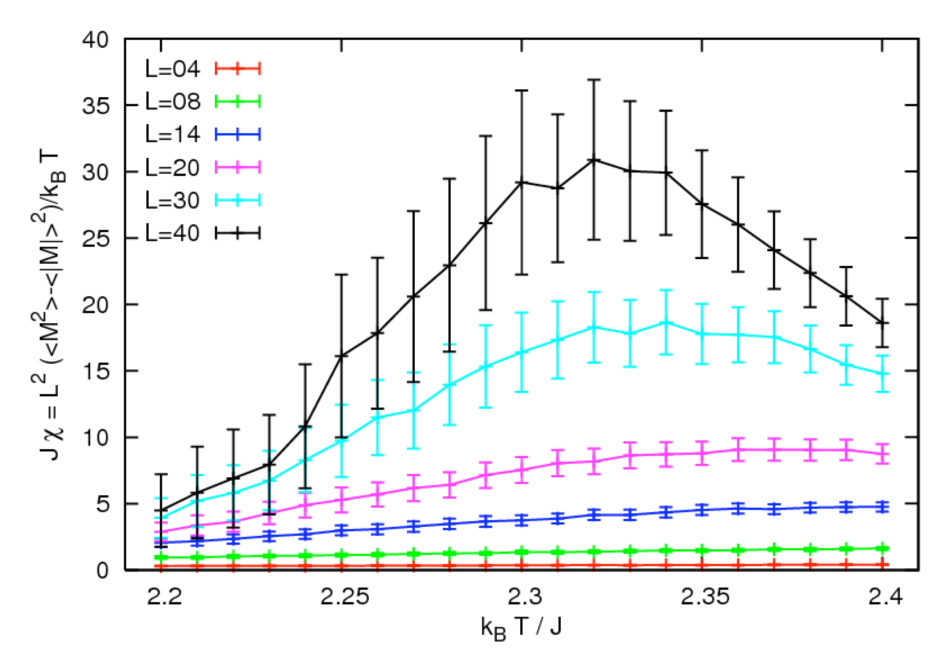
Energy (10^5 sweeps)



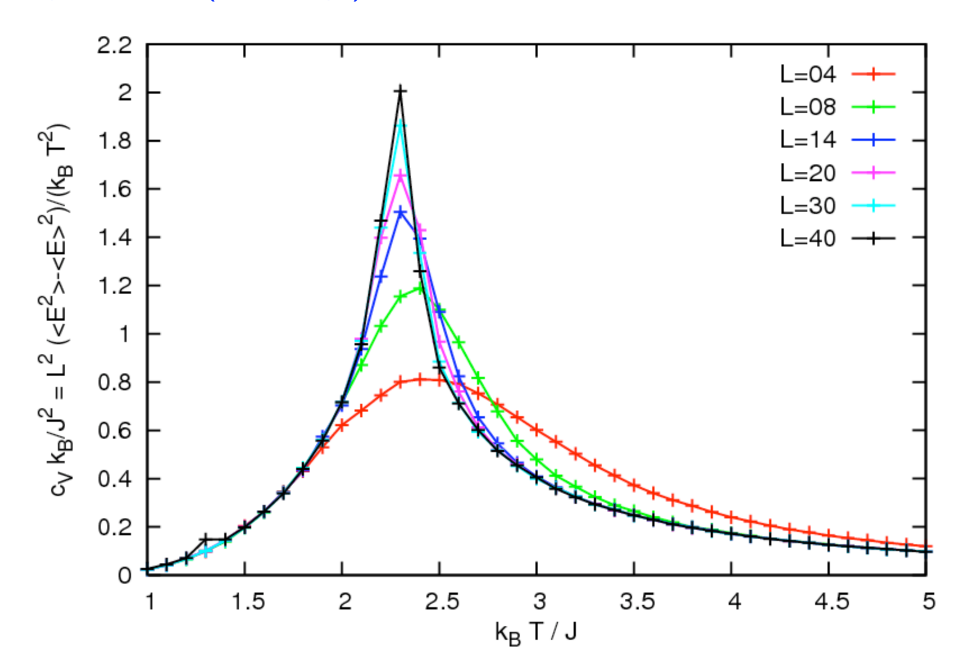
Magnetic susceptibility (10^5 sweeps)



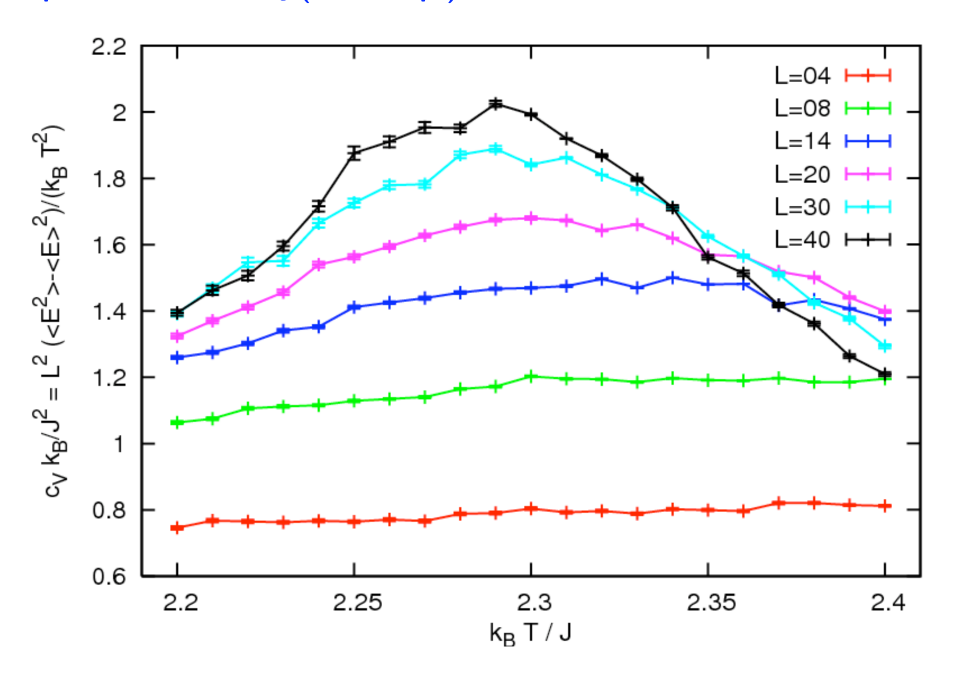
Magnetic susceptibility near T_c (10⁶ sweeps)



Specific heat (10^5 sweeps)



Specific heat near T_c (10^6 sweeps)



A few references:

Ising E 1925 Zeitschrift fur Physik 31 253–258
Bethe H 1931 Zeitschrift fur Physik 71 205
Heisenberg W 1928 Zeitschrift fur Physik 49 205

Near T_c , we can characterize the behavior of many physical quantities by power law behavior just as we characterized the percolation threshold (see Table 13.1). For example, we can write m near T_c as

$$m(T) \sim (T_c - T)^{\beta},\tag{17.22}$$

where β is a critical exponent (not to be confused with the inverse temperature). Various thermodynamic derivatives such as the susceptibility and heat capacity diverge at T_c . We write

$$\chi \sim |T - T_c|^{-\gamma} \tag{17.23}$$

and

$$C \sim |T - T_c|^{-\alpha}.\tag{17.24}$$

We have assumed that χ and C are characterized by the same critical exponents above and below T_c .

Another measure of the magnetic fluctuations is the linear dimension $\xi(T)$ of a typical magnetic domain. We expect the *correlation length* $\xi(T)$ to be the order of a lattice spacing for $T \gg T_c$. Because the alignment of the spins becomes more correlated as T approaches T_c from above, $\xi(T)$ increases as T approaches T_c . We can characterize the divergent behavior of $\xi(T)$ near T_c by the critical exponent ν :

$$\xi(T) \sim |T - T_c|^{-\nu}.$$
 (17.25)

From: Gould-Tobochnich

More precisely, the magnetisation follow a power law close to the transition only approaching from smaller T:

$$M = 0 T \ge T_c$$

$$M \sim |1 - T/T_c|^{\beta} T < T_c$$

If we use the Reduced temperature : $\Delta T = (T - T_c)/T_c$

$$C \sim |\Delta T|^{-\alpha}$$
 $M \sim |\Delta T|^{\beta} \text{ for } \Delta T < 0$
 $\chi \sim |\Delta T|^{-\gamma}$
 $\xi \sim |\Delta T|^{-\nu}$

Universalit

0.00 0.02 0.04 0.06 0.08 0.10 0.06 0.08

the critical exponents are not independent from each other, because of the following scaling laws (\bullet) :

$$\gamma = \nu(2 - \eta),$$
 $2 = \alpha + 2\beta + \gamma,$
 $\nu d = 2 - \alpha,$
 $\gamma = \beta(\delta - 1),$

so it is only necessary to know two of them to determine the others.

For the 2D Ising model:
$$egin{array}{c|c} \alpha & 0 & 0 \\ \beta & 0.125 \\ \hline \gamma & 1.750 \\ \hline \nu & 1 \\ \hline \end{array}$$

• Kerson Huang, Introduction to Statistical Physics (CRC Press)

If the heat capacity goes like: $C(T) \propto |T - T_c|^{-\alpha}$

we could plot $\frac{1}{C(T)}$ as a function of T and make a linear fit:

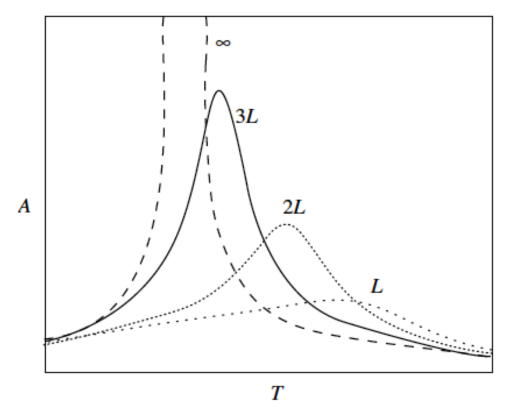
$$\frac{1}{C(T)} = a + bT$$

and similarly for the other quantities.

BUT: Because we can simulate only finite lattices, a direct fit of the measured quantities does not yield good estimates for the corresponding exponents α , ν , β , and γ => we have to take into account the finite size of the system

=> finite size scaling

The shift in the peak position of C and χ with respect to the critical temperature corresponding to the thermodynamic limit is described by: $T_c(L) - T_c(\infty) \propto L^{-\lambda}$



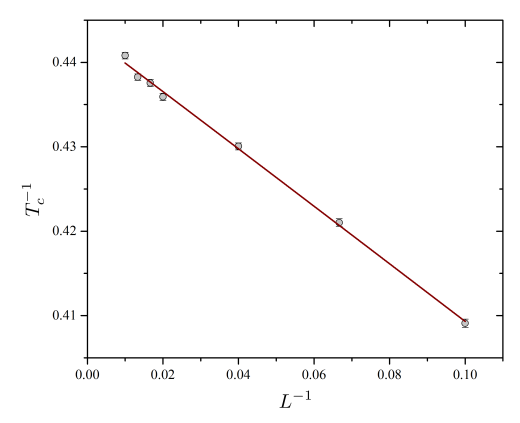


FIG. 1. Typical behaviour of a physical quantity A vs temperature close to the critical point for various system sizes. Figure taken from Thijssen⁸.

[Thijssen, Computational Physics (Cambridge University Press)]

FIG. 9. (Color online) Inverse of the critical temperature T_c^{-1} vs inverse of lattice size L^{-1} for the 2D Ising model. The curve was fitted with a power law $T_c^{-1} = T_{c_{\infty}}^{-1} - bL^{-1/\nu}$, and the critical exponent ν was determined.

• From: E. Ibarra-Garcia-Padilla et al., European Journal of Physics 37(6):065103 DOI: 10.1088/0143-0807/37/6/065103

(This is referred to percolation)

Because we can simulate only finite lattices, it is difficult to obtain estimates for the critical exponents α , β , and γ by using the definitions (17.22)–(17.24) directly. We learned in Section 13.4, we can do a *finite size scaling analysis* to extrapolate finite L results to $L \to \infty$. For example, from Fig. 17.2 we see that the temperature at which C exhibits a maximum becomes better defined for larger lattices. This behavior provides a simple definition of the transition temperature $T_c(L)$ for a finite system. According to finite size scaling theory, $T_c(L)$ scales as

$$T_c(L) - T_c(L = \infty) \sim aL^{-1/\nu},$$
 (17.27)

where a is a constant and ν is defined in (17.25). The finite size of the lattice is important when the correlation length

$$\xi(T) \sim L \sim |T - T_c|^{-\nu}.$$
 (17.28)

As in Section 13.4, we can set $T = T_c$ and consider the L-dependence of M, C, and χ :

$$m(T) \sim (T_c - T)^{\beta} \to L^{-\beta/\nu}$$
 (17.29)

$$C(T) \sim |T - T_c|^{-\alpha} \to L^{\alpha/\nu} \tag{17.30}$$

$$\chi(T) \sim |T - T_c|^{-\gamma} \to L^{\gamma/\nu}. \tag{17.31}$$

From: Gould-Tobochnich

Scaled magnetization vs unscaled T (10^5 sweeps)

