# Critical behaviour of the surface tension in the 3D Ising model

Federico Belliardo Marco Costa

Dipartimento di Fisica Università di Pisa

August 31, 2018

### Summary

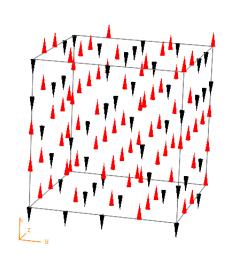
- 3D Ising models
- Definition of the surface tension
- Cluster algorithm and boundary flip
- (Notes on the implementation?)
- Estimation of the errors and autocorrelation
- Fit of the free energy
- Fit of the critical behaviour
- (Conclusion?)

### 3D Ising model

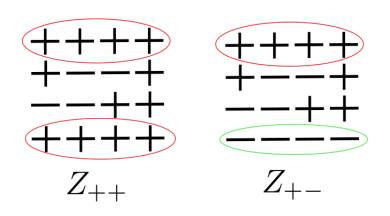
$$\mathcal{H} = -\sum_{\langle x,y \rangle} J_{\langle x,y \rangle} s_x s_y$$

 $J_{\langle x,y 
angle} = 1$  ferromagnetic

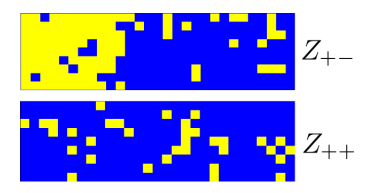
 $J_{\langle x,y 
angle} = -1$  antiferromagnetic



#### Definition of the surface tension



$$\sigma = -\lim_{L \to \infty} \frac{1}{L^2} \log \frac{Z_{+-}}{Z_{++}} \quad L \times L \times T, T = cL$$



$$\sigma = -\lim_{L \to \infty} \frac{1}{L^2} \log \frac{Z_{+-}}{Z_{++}} = \lim_{L \to \infty} \frac{1}{L^2} (F_{+-} - F_{++}) = \lim_{L \to \infty} \frac{F_s}{L^2}$$

 $\sigma = \text{interface free energy per unit area}$ 

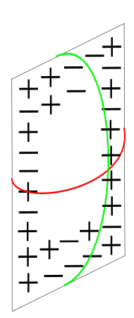
Redefinition of  $Z_{++}$  and  $Z_{+-}$ .

 $Z_{++} \rightarrow$  ferromagnetic link between top and bottom.  $Z_{+-} \rightarrow$  antiferromagnetic link between top and bottom.

Always ferromagnetic link in x and y.

Same definition for  $\sigma$ .

Periodic boundary conditions reduce the finite size effect.



### Montecarlo simulations can't measure Z!

Solution:  $J_{\langle x,y\rangle}$  between top and bottom becomes a **dinamical variable** that is summed over in Z.  $J_{\langle x,y\rangle}=1$  (periodic b.c.)  $J_{\langle x,y\rangle}=-1$  (antiperiodic b.c.) Other  $J_{\langle x,y\rangle}$  remains ferromagnetic.

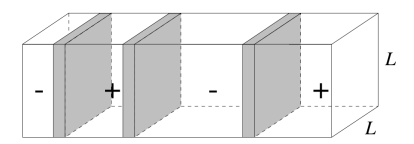
$$Z = \sum_{\{s\},J} \exp\left(\beta \sum_{\langle x,y\rangle} J_{\langle x,y\rangle} s_x s_y\right)$$

$$\frac{Z_{+-}}{Z_{++}} = \frac{\frac{Z_{+-}}{Z}}{\frac{Z_{++}}{Z}} = \frac{\langle \delta_{J=-1} \rangle}{\langle \delta_{J=+1} \rangle}$$

Ratio of measurable expectation values.

We redefine the free energy of the interface in order to improve the convergence proprieties of  $\frac{F_s}{L^2}$  to  $\sigma$  when  $L \to \infty$ . Thermodynamic limit  $\longrightarrow$  only **one** interface

For finite L multiple interface can be present. An even number for  $Z_{++}$  and odd for  $Z_{+-}$ .



 $F_s$  is the free energy of a single surface. There are  $\sim T$  different position for the interface.

$$Z_1 = T \exp(-F_s) = \exp(-F_s + \ln T)$$

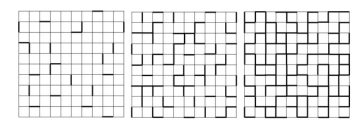
$$\frac{Z_{+-}}{Z_{++}} = \frac{Z_1 + \frac{Z_1^3}{3!} + \frac{Z_1^3}{5!} + \dots}{1 + \frac{Z_1^2}{2!} + \frac{Z_1^4}{4!} + \dots} = \tanh\left(\exp\left(-F_S + \ln T\right)\right)$$

$$F_S = \ln \left( T \right) - \ln \left( rac{1}{2} \ln \left( rac{1 + rac{Z_{+-}}{Z_{++}}}{1 - rac{Z_{+-}}{Z_{--}}} 
ight) 
ight) \quad \sigma = \lim_{L o \infty} rac{F_s}{L^2}$$

### Cluster algorithm and boundary flip

Cluster algorithms allow for simultaneous updates of large parts of the lattice. Thus reducing the autocorrelation time and the critical slowling down. Swendsen and Wang (1987).

Introduce link variables  $\sigma_{\langle x,y\rangle}=\{0,1\}$  on the lattice:



$$Z = \sum_{\{s=\pm 1\}} \exp\left(eta \sum_{\langle x,y \rangle} s_x s_y
ight) = \sum_{\{s=\pm 1\}} \prod_{\langle x,y \rangle} e^{eta s_x s_y} =$$

 $=e^{-dV\beta}$   $\left(1+\delta_{s_x,s_y}\left(e^{2\beta}-1\right)\right)=$ 

 $\{s=\pm 1\} \langle x,y \rangle$ 

 $\{s\}\ \langle x,y\rangle\ \{\sigma_{\langle x,y\rangle}=0,1\}$ 

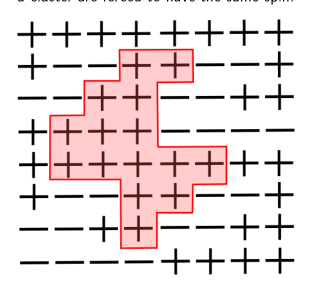
$$=e^{-dVeta}\sum\prod\sum_{\left[\left(1-\sigma_{\left\langle \mathsf{x},\mathsf{y}
ight
angle}
ight)+\sigma_{\left\langle \mathsf{x},\mathsf{y}
ight
angle}\delta_{\mathsf{s}_{\mathsf{x}},\mathsf{s}_{\mathsf{y}}}\left(e^{2eta}-1
ight)
ight]}$$

For a fixed spin configuration  $\{s\}$  the links are indipendent.

$$p_1=p\left(\sigma_{\langle x,y
angle}=0
ight)=\min\left[1,e^{-2eta s_x s_y}
ight]=\exp\left(-2eta \delta_{s_x,s_y}
ight)$$
  $p_0=p\left(\sigma_{\langle x,y
angle}=1
ight)=1-p\left(\sigma_{\langle x,y
angle}=1
ight)$  (if  $s_x s_y=1$  the weights in  $Z$  are normalized to  $e^{2eta}$ )

For fixed  $\sigma_{\langle x,y\rangle}$  only configurations of spins that satisfy the constraint  $s_x=s_y$  where  $\sigma_{\langle x,y\rangle}=1$  have a non zero probability. All configurations that satisfy the constraint have the same weight.

Definition: a **cluster** is a set of spin in the lattice path connected by links with  $\sigma_{\langle x,y\rangle}=1$ . In the ensamble all sites of a cluster are forced to have the same spin.



### Swendsen and Wang algorithm:

- Generate a link configuration  $\sigma_{\langle x,y\rangle}$  based on the current spin configuration by using probabilities  $p_0$  and  $p_1$ .
- For each cluster chosse a spin  $(s=\pm 1)$  with probability  $\frac{1}{2}$ .
- The newly generted spin configuration is the new element of the Markov chain.

The SW is **ergodic**. We know prove that it satisfy the **detailed balance**.

In the cluster algorithm we update both the spin  $\{s\}$  and the links  $\{\sigma_{x,y}\}$ . The ensamble contains both spins and link configurations:  $\{s,\sigma\}$ .

$$\frac{P\left(\left\{s_{0},\sigma_{0}\right\} \rightarrow \left\{s_{1},\sigma\right\}\right)}{P\left(\left\{s_{1},\sigma_{0}\right\} \rightarrow \left\{s_{0},\sigma\right\}\right)} = \frac{P\left(\left\{s_{1}\right\} | \left\{\sigma\right\}\right) P\left(\left\{\sigma\right\} | \left\{s_{0}\right\}\right)}{P\left(\left\{s_{0}\right\} | \left\{\sigma\right\}\right) P\left(\left\{\sigma\right\} | \left\{s_{1}\right\}\right)}$$

(this is not the detailer balance in the ensamble of  $\{s,\sigma\}$  as the lonk configurations don't get exchnged!)

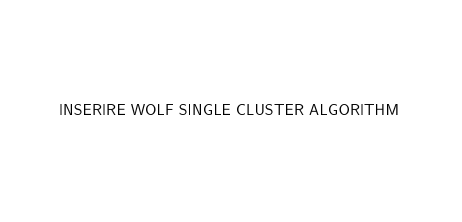
 $P\left(\{s\}|\{\sigma\}\right) = \frac{1}{2^{\#cluster}}$  if  $\{s\}$  is compatible with  $\{\sigma\}$ . Null otherwise.

Thus 
$$P(\{s_1\}|\{\sigma\}) = P(\{s_2\}|\{\sigma\}).$$

$$P(\lbrace \sigma \rbrace | \lbrace s_1 \rbrace) = \exp \left( -2\beta \sum_{\langle x,y \rangle} \delta_{s_x,s_y} \right)$$

FINIRE – In effetti questa probabilità non mi convince - rivedere.

Dire che in realtà noi usiamo il sigle cluster algorithm con un procedimento ricorsivo (implementato con uno stack) che controlla una ed una sola volta ogni link per vedere se attivarlo o meno. Costruendo così un singolo cluster che andiamo a flippare.



### Boundary flip algorithm

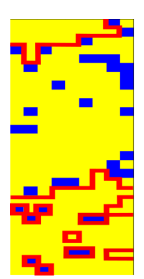
Now also the coupling between the top and the bottom is a dynamical variable to simulate:  $\{s,\sigma,J=\pm 1\}$ . At fixed  $\{s,J_{\langle x,y\rangle}\}$  the  $\sigma_{\langle x,y\rangle}$  probabilities are:

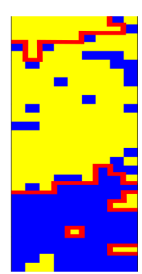
$$egin{aligned} p_1 &= p\left(\sigma_{\langle x,y
angle} = 0
ight) = \min\left[1,e^{-2J_{\langle x,y
angle}eta s_x s_y}
ight] \ p_0 &= p\left(\sigma_{\langle x,y
angle} = 1
ight) = 1 - p\left(\sigma_{\langle x,y
angle} = 1
ight) \end{aligned}$$

A link  $\sigma_{\langle x,y \rangle} = 1$  demands  $J_{\langle x,y \rangle} s_x s_y = 1$ . Thus we can flip J,  $s_x$  and all the spins connected to  $s_x$  via some link in the bulk obtaing a configuration compatible with  $\{\sigma_{\langle x,y \rangle}\}$ .

Boundary condition update. Section of a  $7 \times 7 \times 21$  lattice at temperature

 $\beta = 0.250$ 



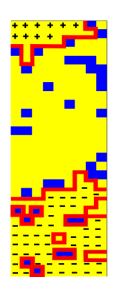


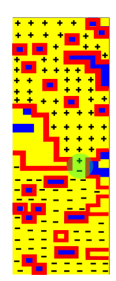
## The boundary flip generates an interface between phases!

We must check this flip of spins and J can be done without violating the constraints imposed by the bulk links  $(\sigma_{\langle x,y\rangle}=1)$  implies  $s_x=s_y$ .

Check all the clusters that contain sites of the lower surface and flip the boundary condition only if it can be done consistently on all the lattice.

Introduce an extra variable  $c_x=\pm 1$  that gets propagated in the costruction of the cluster but changes sign when crossing the boundary.





This update is ergodic. We now prove the validity of the detailed balance.

FINIRE!!!
We alternate a step of the Wolff algorithm with the boundary flip.

### Notes on the implementation

C++ for the Montecarlo and Jackknife algorithms, Python for data analysis, fits and plots.

The hot function of the simulation generates a cluster starting in a given position and exploring the neighbouring links. If a link is chosen to be  $\sigma_{\langle x,y\rangle}=1$  then the adjacent site is included in the cluster and the procedure is repeated. The extra variable  $c_x$  is also propagated.

```
stack<site> stack
stack.push(seed)
while(!stack.empty())
  site current = stack top()
  value = cluster[current]
  if(cluster[current] is incostintent) flag = 1;
  else if(cluster[current] == 0)
    cluster[current] = cluster[old];
    for (d = 0; d < 3; d++)
       for(a = -1; a < 2; a = a + 2)
       next = current + a
       Check if we are on the boundary
       if(cluster[next] == 0 \text{ and } random < p)
stack.push(next)
return flag
```

- The algorithm doesn't actually performs the spin and boundary flip it just counts for how many configuration it is possible.
- The simulation where executed with  $N=10^5, 10^6$  steps of the Markov chain.
- The first 5% of the Markov chain is ignored to avoid non termalized configurations.
- The Jackknife is executed for blocks with size starting from 10000 to 10900 with step 30.

# Estimation of the errors and autocorrelation

## Fit of the free energy - theory

## Fit of the free energy - results

### Fit of the critical behaviour

## (Conclusions?)