

DIGITAL PHYSICS

# Quantum Monte Carlo algorithm for spins systems

Etienne CAMPHUIS  
Barthélémy MEYNARD  
PIGANEAU  
X 2016  
December 2018, 7<sup>th</sup>

# Table des matières

<b>1</b>	<b>Introduction</b>	<b>2</b>
<b>2</b>	<b>Local Algorithm</b>	<b>4</b>
2.1	Description . . . . .	4
2.2	Domain of validity . . . . .	4
2.2.1	Winding number . . . . .	4
2.2.2	Straight line only can be flipped . . . . .	5
2.3	First results . . . . .	6
2.4	Autocorrelation function . . . . .	8
2.5	Limits . . . . .	10
<b>3</b>	<b>Loop Algorithm</b>	<b>11</b>
3.1	Failure of the local algorithm . . . . .	11
3.1.1	Winding number . . . . .	11
3.1.2	Acceptance rate . . . . .	11
3.1.3	Decorrelation . . . . .	11
3.2	The loop update . . . . .	11
3.2.1	Example . . . . .	11
3.2.2	Formalism . . . . .	13
3.3	Encoding the loop algorithm . . . . .	16
3.3.1	Parameters . . . . .	16
3.3.2	Methods . . . . .	17
3.4	Results . . . . .	19
3.4.1	Case $J_x = J_z$ . . . . .	19
3.4.2	Case $J_x >  J_z $ . . . . .	22
<b>4</b>	<b>Conclusion</b>	<b>24</b>
<b>5</b>	<b>Bibliography</b>	<b>24</b>

# 1 Introduction

The aim of this project is to encode an algorithm able to compute the mean energy at fixed temperature of a spin chain modeled by the XXZ model. This model is described by the following Hamiltonian :

$$H = J_x \sum_i (S_i^x S_{i+1}^x + S_i^y S_{i+1}^y) + J_z \sum_i S_i^z S_{i+1}^z$$

FIGURE 1 – Hamiltonian of the XXZ model

The basic idea of this original World Line approach is to split the XXZ Hamiltonian into a set of independent two sites problems. The way to achieve this decoupling, is with the use of a path integral and the Trotter decomposition. First we write :

$$H = \underbrace{\sum_n H^{(2n+1)}}_{H_1} + \underbrace{\sum_n H^{(2n+2)}}_{H_2}$$

FIGURE 2 – Separation of the Hamiltonian

One may verify that  $H_1$  and  $H_2$  are sums of commuting (*i.e.* independent) two sites problems. Hence, on their own  $H_1$  and  $H_2$  are trivially solvable problems. However,  $H$  is not. To use this fact, we split the imaginary propagation  $\exp(-\beta H)$  into successive infinitesimal propagations of  $H_1$  and  $H_2$ . Here  $\beta$  corresponds to the inverse temperature, with  $\beta = m\_trotter * \Delta\tau$ . The path integral is then a  $2D$  representation, with horizontally the space, *i.e.* the spin chain, and vertically the imaginary time, *i.e.* the Trotter decomposition, with  $2m\_trotter$  steps. Actually, as both  $H_1$  and  $H_2$  acts on half of the spin pairs, the path integral can be represented by a checker board. The bold lines follow the time evolution of the up spins and empty sites, with respect to the world lines, correspond to the down spins. A full time step  $\Delta\tau$  corresponds to the propagation with  $H_1$  followed by  $H_2$ . Periodic boundary or open border conditions are chosen in the spatial direction. In the imaginary time direction, periodic boundary conditions are mandatory for the imaginary time decomposition to work. The weights for a given world line configuration is the product of the weights of tiles listed in the figure 3. Note that, although the spin-flip processes come with a minus sign, the

overall weight for the world line configuration is positive since each world line configuration contains an even number of spin flips. A further description is given in the article by Assaad and Evertz.

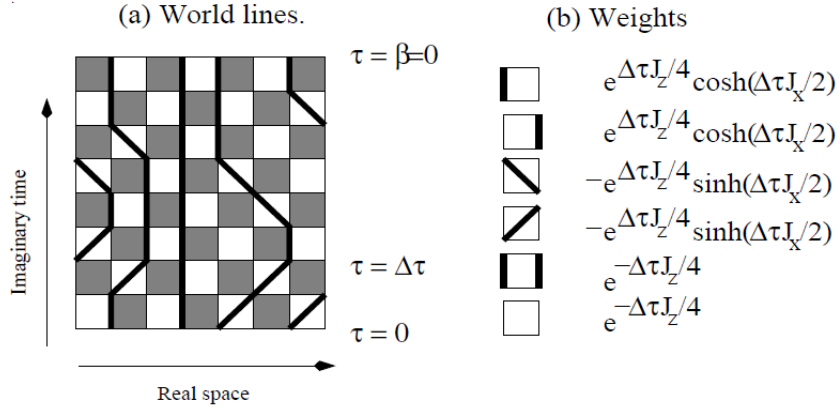


FIGURE 3 – 2D Path integral with the associated weights of the tiles ; Here  $m\_trotter = 4$  and  $L = 8$

**Computation of the weights** For a given configuration, the weight can be expressed with this formalism :

$$Z = \sum_w \Omega(w)$$

$$\Omega(w) = \langle \sigma_1 | e^{-\Delta\tau H_1} | \sigma_{2m} \rangle \cdots \langle \sigma_3 | e^{-\Delta\tau H_1} | \sigma_2 \rangle \langle \sigma_2 | e^{-\Delta\tau H_2} | \sigma_1 \rangle$$

where  $w$  defines the states  $|\sigma_1\rangle \cdots |\sigma_{2m}\rangle$ .

FIGURE 4 – Formula of the weight of a given configuration

**Computation of the energies** The following formula give the energy of a given configuration :

$$E(w) = -\frac{1}{m} \frac{\partial}{\partial \Delta\tau} \ln \Omega(w).$$

FIGURE 5 – Formula of the energies of a given configuration

## 2 Local Algorithm

### 2.1 Description

This section focuses on a way to move through the configurations. The local Algorithm need to implement two distinct moves :

- ★ The splitline move : This move is supposed to inverse a complete line. To be implemented we firstly chose a spin, then check if the all the spins upward have the same value, and in that case invert all of them.

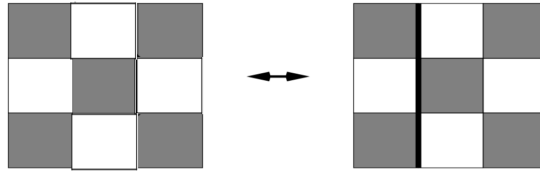


FIGURE 6 – Splitline move

- ★ The kink move : This move move represents the effect of the quantum spins. The part  $J_x$  and  $J_z$  create variation from the usual spin model. It is taken in account with this move, which flip the 4 spins around a black case, if the two left spins are the same and are the opposite of the two right spins.

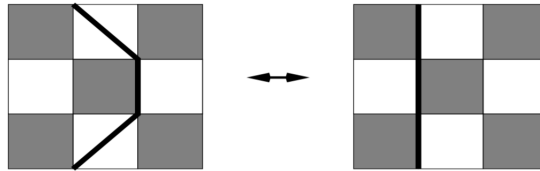


FIGURE 7 – kink move ie local update in our code

### 2.2 Domain of validity

#### 2.2.1 Winding number

It can not deal with periodic boundary conditions. Indeed, because the updates are either local or along a temporary line, the winding number is fixed. The winding number is an integer counting the number of period that the world-line takes to return to the same position.

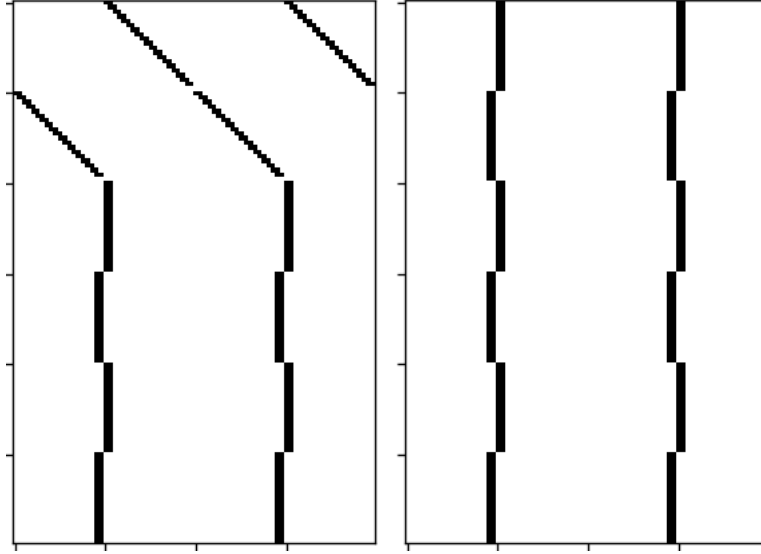


FIGURE 8 – Configurations with  $W = 2$  (left) and  $W = 1$  (right)

### 2.2.2 Straight line only can be flipped

After a few attempts to split any world-line we only had wrong results. Indeed we realized it caused an error in the detailed balance. If we start from the following situation 1 and split the world line 3 we get to the situation 2.

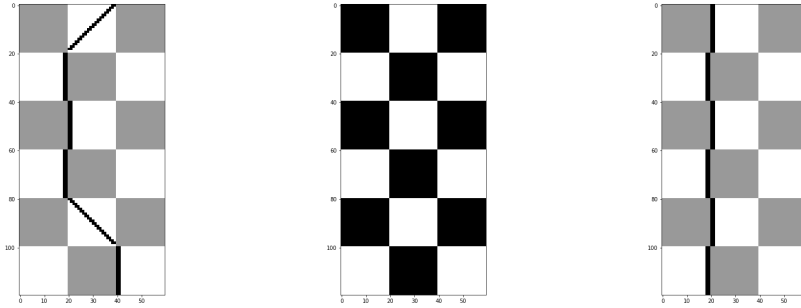


FIGURE 9 – situation 1 (left), situation 2 (middle), situation 3 (right)

From the situation 2 it is not possible to come back at the situation 1 with a simple splitline. Indeed such move only creates situation 3. To do so, a splitline and a kink move is mandatory. The detailed balance is in this case unbalanced. The solution is to accept a flip on a straight world-line only.

To code a Monte Carlo algorithm, we then had to condensed the two moves in one. We called it *stoch\_move(threshold)*, this algorithm has a probability  $P = threshold$  to make a splitline move, and  $P = 1 - threshold$  to make a kink move.

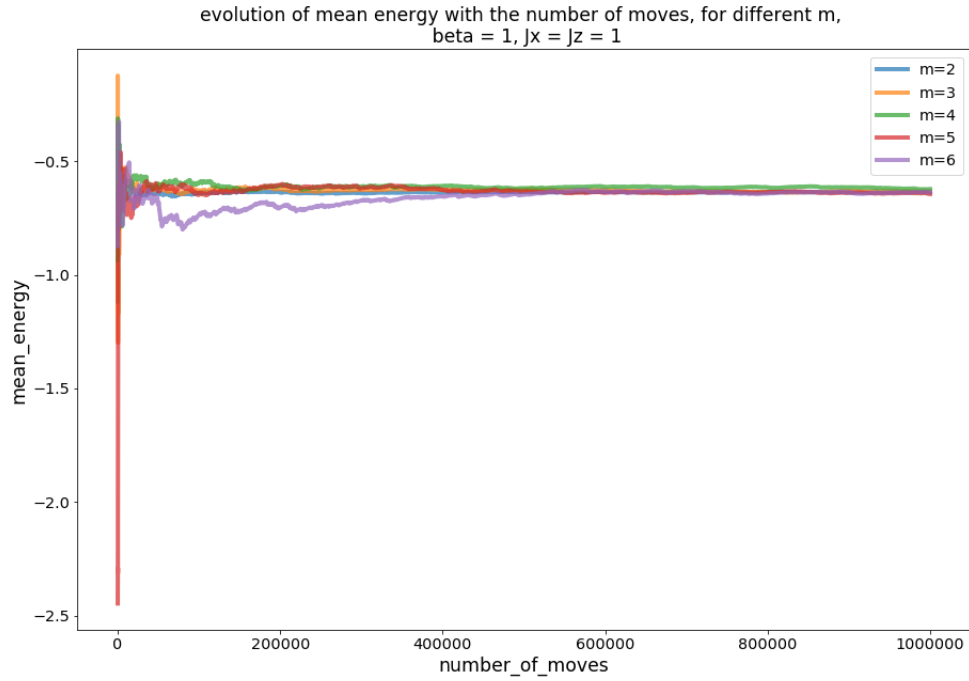
## 2.3 First results

For the proper parameter the algorithm enable to compute the mean energy. In the following we will focus on the case  $J_x = J_z = 1, n\_spins = 4, \beta = 1$ . Firstly we made a few run of the montecarlo algorithm, with a small *length\_cycle*(10) and a small number of measures(100 000), to have a first look.

The code produces the following results :

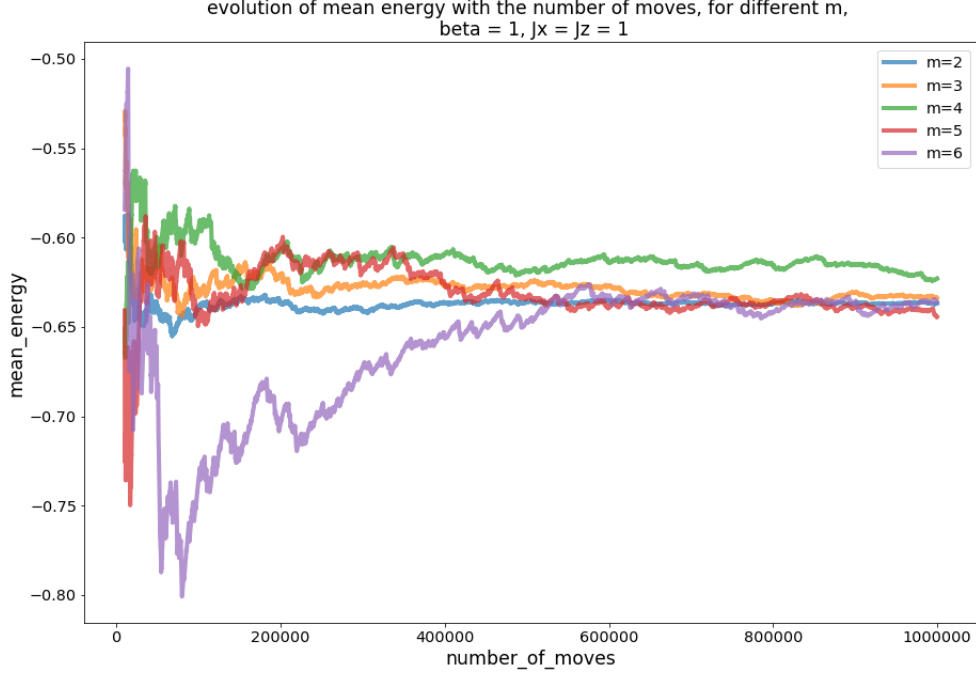
- ★ theoretical energy : -0.6330214400416617
- ★ Energy for m=2 : -0.6368988448758103 +/- 0.002502566964381992
- ★ Energy for m=3 : -0.6338973315707276 +/- 0.0027341143579975723
- ★ Energy for m=4 : -0.6229915669089899 +/- 0.002819998026568314
- ★ Energy for m=5 : -0.6445715838659634 +/- 0.0029559595617620286
- ★ Energy for m=6 : -0.6358614671234866 +/- 0.0029930079421800816

At the first glance it seems that the algorithm give a good approximation of the result. We firstly wanted to have an idea of the evolution of the mean energy with the number of measures.



We clearly feel that the 5 results converge to the same limit around  $-0.63$ . We also observe that the case  $m = 6$  takes more moves to converge. The beginning of the curves doesn't have much meaning. It has a high dependence regarding the initial configuration. We can have a better look on the end of the graph :





This graph enables us to see two things more clearly. First, the number of moves necessary for the mean to converge rises with  $m$ . Second, the choice of  $m\_trotter$  seems to involve a bias in the mean. While  $m\_trotter$  rises, the number of configurations follows and the weight changed by the *local\_update* move decreases.

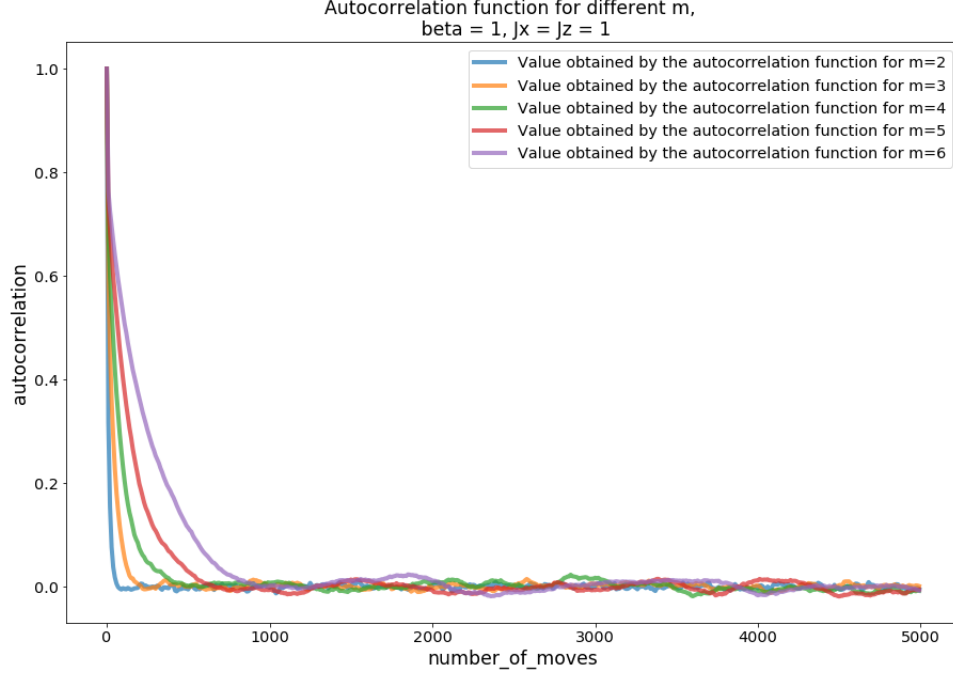
## 2.4 Autocorrelation function

To have a better idea of the length between two measures, we computed the autocorrelation function. This function is defined for an observable  $Q$  by the following formula :

$$A_Q(\tau) = \frac{\langle Q_k Q_{k+\tau} \rangle - \langle Q_k \rangle^2}{\langle Q_k^2 \rangle - \langle Q_k \rangle^2}$$

Mean being on the  $k$ .

We computed the autocorrelation function for 5000 moves and the result is clear.



We can observe that the correlation between two states requires more and more moves to disappear when  $m$  rises. This is due to two factors : firstly when  $m$  rises the system get bigger and therefore you need more accepted moves to de-correlate, secondly the weights of the kinks decrease which means that less and less moves are accepted.

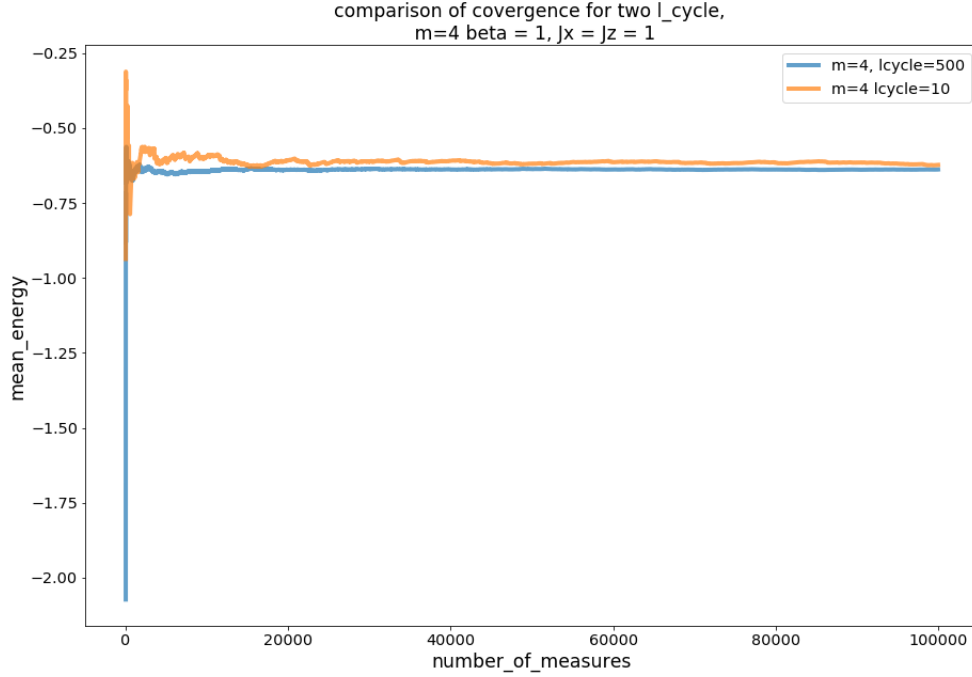
If we look at these functions more precisely, we can find the proper cycle length, which guarantees that two measures are not correlated. We find :

- ★  $m = 2$  : length of de-correlation 50
- ★  $m = 3$  : length of de-correlation 200
- ★  $m = 4$  : length of de-correlation 500
- ★  $m = 5$  : length of de-correlation 800
- ★  $m = 6$  : length of de-correlation 1000

The following results are found :

- ★ Energy for  $m=2$  :  $-0.6393578437601745 \pm 0.002507393542860798$
- ★ Energy for  $m=3$  :  $-0.6387051296013677 \pm 0.002749339051345642$
- ★ Energy for  $m=4$  :  $-0.6380826437580377 \pm 0.0028694731132348273$

We can compare the evolution of the mean energy for the two different length cycle



## 2.5 Limits

Several problems remains with this algorithm. Firstly as we have seen, it can only works on open border condition. Moreover It get very expensive to compute this Monte Carlo algorithm when we are not in the proper condition. Indeed the the weight to do a kink is proportional to  $\sinh^2(\Delta\tau \frac{Jx}{2})$ . Consequently when  $\beta$  gets smaller or  $m\_trotter$  bigger, the moves proposed are mostly rejected, and the Monte Carlo becomes highly inefficient. Unfortunately to get precise results we need to make m big and so  $\Delta\tau$  small. To conclude this idea only works for low precision estimation and high  $\beta$ .

## 3 Loop Algorithm

In this section, we will discuss the loop algorithm, introduced in the Assaad and Evertz article page 12.

### 3.1 Failure of the local algorithm

The previous algorithm is not reliable for the three following reasons.

#### 3.1.1 Winding number

It can not deal with periodic boundary conditions. Indeed, because the updates are either local or along a temporary line, the winding number is fixed. The winding number is an integer counting the number of period that the world-line takes to return to the same position.

#### 3.1.2 Acceptance rate

The rate of acceptance of the move is highly dependant on the parameters. For instance, should ...

#### 3.1.3 Decorrelation

The decorrelation between the configuration is rather slow as shown in the previous part. It can go up to several hundred of moves. This expands the time of computation necessary to reach a suitable result.

### 3.2 The loop update

#### 3.2.1 Example

The idea of Assaad and Evertz is to map the periodic boundary conditions model onto another one called "six-vertices model". Thanks to the result obtained on this model, we are now able to implement an update which allows fast de-correlation, high acceptance rate and changing winding number. Let us imagine each tile is replaced by its representation in the six-vertices model which is described in figure 10.

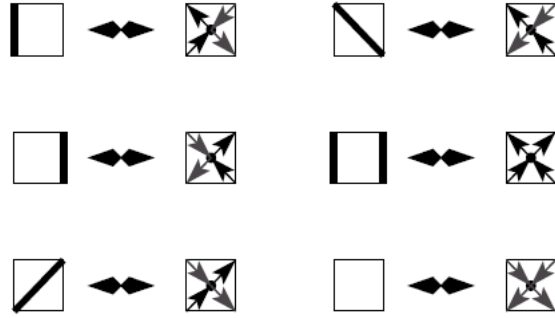


FIGURE 10 – Rule to change a tile in a vertex plaquette

The rule is the following : each spin up is transformed into an arrow going up ; each spin down is transformed into an arrow going down. Given this transformation one is now able to imagine loops on the pattern. One starts from a given spin and a given vertex plaquette, and go along a chosen arrow in the same direction. If one does not choose two times the same arrow, he will come back to the first spin and thus create a loop. This mechanism is illustrated by the figure 11. The described loop "goes through" the right side to come back to the left side thanks to the spatial periodic boundary conditions.

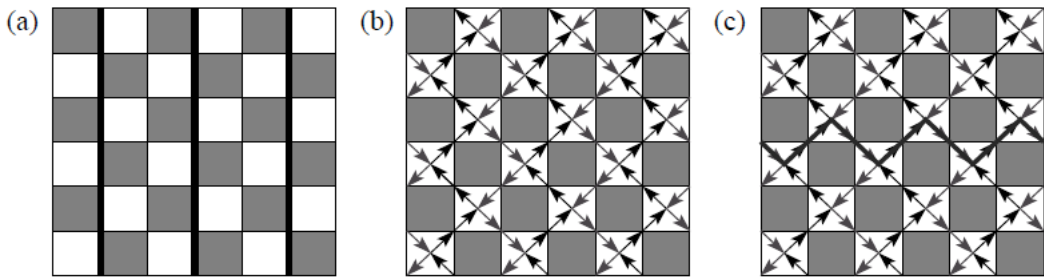


FIGURE 11 – Illustration of the creation of a loop

With the chosen loop, one can now flip the spins along the loop, which corresponds to changing the direction of all the arrows. This generates a new configuration. In the described one in figure 11 and figure 12, the winding number is changed from 1 to 3. One can observe that even this quite simple update (only one loop has been studied), the new configuration seems less dependent that any other that would have been generated thanks to the local algorithm.

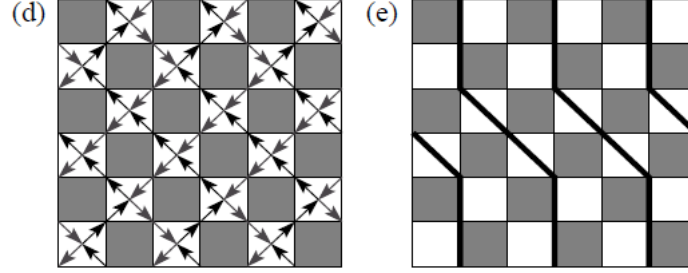


FIGURE 12 – Update by flipping the spins along the loop

### 3.2.2 Formalism

*For the quantum Monte-Carlo algorithm, the detailed balance must be fulfilled. For a given tile in the configuration  $S$ , and for any other tile  $S'$ , the detailed balance is given by equation (1)*

$$W(S)P(S \rightarrow S') = W(S')P(S' \rightarrow S) \quad (1)$$

In order to make a move on the configuration, one has to change all the tiles in vertex plaquettes, then to choose loops and to choose whether to flip the spins along it or not. The process is clearer if one creates a virtual configuration called graph. For each tile, the associated vertex plaquette will be replaced by a graph-tile representing the choice of the arrows. The choices are either the loop goes vertically, either diagonally, either horizontally along the tile. Actually, there is another graph possible which switch all the spins on the tile, but it will not be considered here. Summing all the graph will create loops on the configuration. Figure 13 is an example of total graph

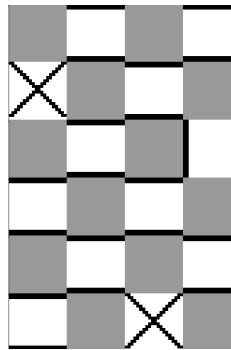


FIGURE 13 – Example of graph ; one can verify that all loops are closed

First of all, one needs to know which graphs can be reached by a given tile. Let us give an example for the tile with all the spins down, the white tile. The associated vertex plaquette is the one with all the arrow heading down. If the loop arrives from the upper left side of the tile, it can either go to the bottom left or to the bottom right. Thus, only 2 graphs are allowed : either a vertical graph or a cross graph. The illustration is given in figure 14 for all tiles, one can verify the association following the previous logic applied to the white tile.

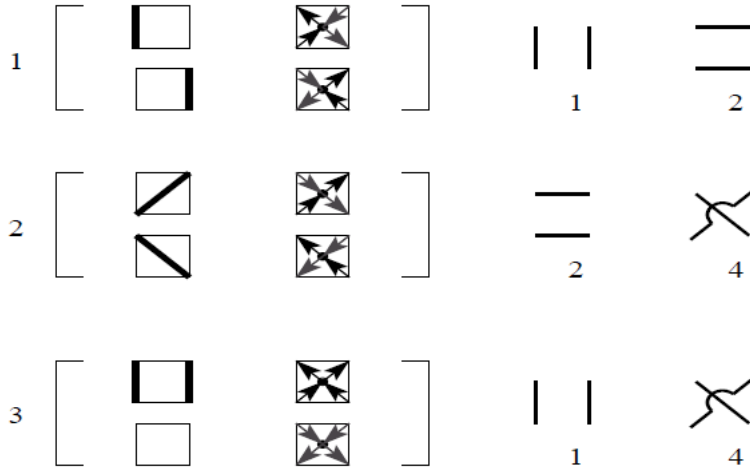


FIGURE 14 – Possible graphs for tiles

However, depending on the interactions and the temperature, the graphs will not be equally possible. One has to choose weights for each of them. Let us call  $W(S)$  the weight of the tile  $S$  and  $W(S, G)$  the weight of the graph  $G$  for a tile  $S$ . The equation (2) is required to have a total probability equal to one, and then be sure that every tile is replaced by a graph.

$$\sum_G W(S, G) = W(S) \quad (2)$$

So we choose a graph from a given plaquette with probability

$$P(S \rightarrow (S, G)) = \frac{W(S, G)}{W(S)} \quad (3)$$

The detailed balance written with the graph transition

$$\begin{aligned} W(S) \sum_G P(S \rightarrow (S, G)) P((S, G) \rightarrow (S', G')) = \\ W(S') \sum_{G'} P(S' \rightarrow (S', G')) P((S', G') \rightarrow (S, G')) \end{aligned} \quad (4)$$

By using equation (3) is can be rewritten

$$\sum_G W(S, G) P((S, G) \rightarrow (S', G)) = \sum_{G'} W(S', G') P((S', G') \rightarrow (S, G')) \quad (5)$$

Then one can choose the different parameters to make the computation easier. For instance, the weights of the graph is set constant, not dependent on the spin configuration. This is easier to encode and clearer for the detailed balance :

$$\forall G, \forall (S, S'), W(S, G) = W(S', G) = V(G) \quad (6)$$

Moreover, with the use of the heat-bath algorithm for flipping, the probability to change the spin from  $S$  to  $S'$  given a choice of graph  $G$  can be set to  $P((S, G) \rightarrow (S', G)) = \frac{W(S', G)}{W(S', G) + W(S, G)}$ . Thus the detailed balance is verified and the code is easy to encode. Indeed

$$\begin{aligned} P((S, G) \rightarrow (S', G)) &= \frac{W(S', G)}{W(S', G) + W(S, G)} \\ &= \frac{V(G)}{V(G) + V(G)} \\ &= \frac{1}{2} \end{aligned} \quad (7)$$

The equation (7) means that each loop is flipped with probability  $1/2$ .

To set the weights of the graphs  $V(G)$ , equations (2) and (6), with the use of the notations of figure 14 give

$$\begin{cases} W(S = 1) = V(G = 1) + V(G = 2) \\ W(S = 2) = V(G = 2) + V(G = 4) \\ W(S = 3) = V(G = 1) + V(G = 4) \end{cases} \quad (8)$$

This set of equation (8) has the solution :

$$\begin{cases} V(G = 1) = \frac{1}{2}[W(S = 1) + W(S = 3) - W(S = 2)] \\ V(G = 2) = \frac{1}{2}[W(S = 1) + W(S = 2) - W(S = 3)] \\ V(G = 4) = \frac{1}{2}[W(S = 2) + W(S = 3) - W(S = 1)] \end{cases} \quad (9)$$

Then, the weights  $W(S)$  are

$$\begin{cases} W(S = 1) = \exp(\Delta\tau J_z/4) \cosh(\Delta\tau J_x/2) \\ W(S = 2) = \exp(\Delta\tau J_z/4) \sinh(\Delta\tau J_x/2) \\ W(S = 3) = \exp(-\Delta\tau J_z/4) \end{cases} \quad (10)$$



The weights of each graph is now set. All the problem is characterized and the implementation is possible. However, the *Metropolis* algorithm is valid only for positive weights for  $W(S)$  and  $V(G)$ . This fixes conditions on  $J_z$  and  $J_x$ . With the condition  $V(G = 2) > 0$  :

$$\begin{aligned} \exp(\Delta\tau J_z/2) \sinh(\Delta\tau J_x/2) + \exp(\Delta\tau J_z/2) \cosh(\Delta\tau J_x/2) - 1 &> 0 \\ \exp(\Delta\tau J_z/2) \exp(\Delta\tau J_x/2) &> 1 \\ J_x &> -J_z \end{aligned} \tag{11}$$

With the condition  $V(G = 4) > 0$  :

$$\begin{aligned} \exp(\Delta\tau J_z/2) \sinh(\Delta\tau J_x/2) + 1 - \exp(\Delta\tau J_z/2) \cosh(\Delta\tau J_x/2) &> 0 \\ \exp(-\Delta\tau J_z/2) &> \exp(-\Delta\tau J_x/2) \\ -J_z &> -J_x \\ J_x &> J_z \end{aligned} \tag{12}$$

The final condition is then

$$J_x > |J_z| > 0 \tag{13}$$

If ever  $|J_z|$  is higher than  $J_x$ , the algorithm can be implemented with another type of graph, called  $G = 3$ , that propagates the loop along all the arrows, which means all the spins on the tile are flipped. This is a graph accessible by all the vertex plaquettes, which allows a new system (8) and a supplementary degree of freedom to set the weights  $V(G)$ . This option will not be considered in our algorithm.

### 3.3 Encoding the loop algorithm

The whole methods will be encoded in an algorithm

#### 3.3.1 Parameters

First, we create a class containing the necessary parameters.

- ★ The interactions  $J_x$  and  $J_z$
- ★ The number of spins  $n\_spins$
- ★ The division of the imaginary time  $m\_trotter$  and  $\Delta\tau$
- ★ The representation of the spins  $spins$
- ★ The pattern configuration  $pattern$
- ★ The graph configuration  $total\_graph$

- ★ The list of energies depending on the concerned tile *energymatrix*
- ★ The list of tile weights depending on the concerned tile *weightmatrix*
- ★ The list of arrays imaging the tiles *cases*
- ★ The list of arrays imaging the graphs *graphs*
- ★ The graph weights depending on the graphs  $w_{11}, w_{12}, w_{22}, w_{24}, w_{31}, w_{34}$

### 3.3.2 Methods

Here are the methods used in the Quantum Monte Carlo algorithm.

- ★ *total\_energy()* : Thanks to array combination and the use of masks, this methods extract the energy of each tile on the pattern then sum those energies.
- ★ *weight()* : Thanks to array combination and the use of masks, this methods extract the weight of each tile on the pattern then make the product of those weights to return the full weight of the combination.
- ★ *spins\_to\_pattern()* : With the use of array combination, this method computes the pattern from the spin configuration. At each coordinate  $(i, j)$  is a number corresponding to the tile.
- ★ *createimage()* : Returns an array imaging the total world-line pattern.
- ★ *tile\_in\_graph(pos)* : Chooses a graph for the tile in position  $(pos[0], pos[1])$  with probability  $\frac{V(G)}{W(S)}$ .
- ★ *set\_total\_graph()* : Goes over the whole parameter *total\_graph* and choose for each position a graph thanks to the previous method *tile\_in\_graph(pos)*. An example of transformation is given in the figure 15.

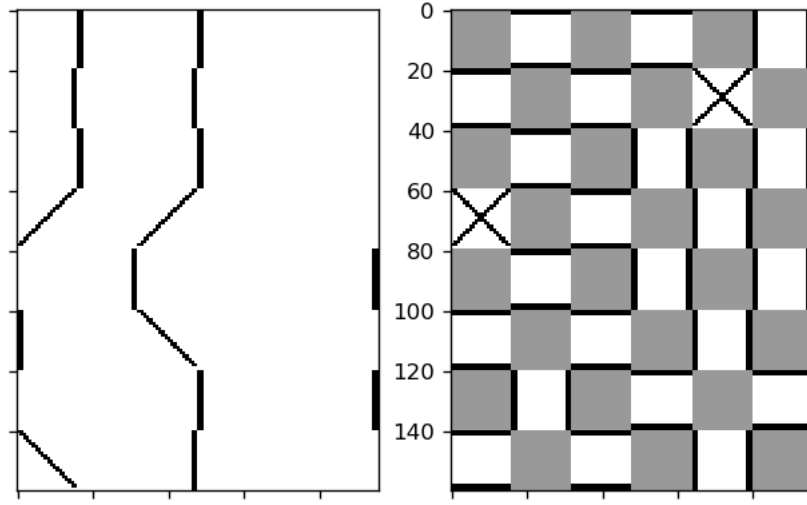


FIGURE 15 – Example of transition from a given pattern to a graph

- ★ *find\_next(pos, graph)* : This method is used to get the loops from the graph. The *pos* and *graph* parameters corresponds respectively to the position of a spin and the one of a graph. The method returns a new spin position and a new graph position after propagating the loop through the given *graph* from the *spin* position. For instance, if the graph is  $G = 1$ , *i.e.* vertical, and the spin is on the upper left side on the graph, the method will return the spin just below the first spin and the graph down left the first graph.

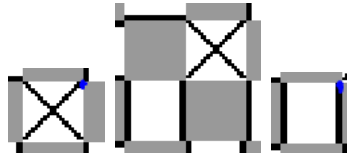


FIGURE 16 – Ensemble of two graphs ; the left graph with the blue spin position will give the right graph with the blue spin position through *find\_next()*

- ★ *find\_loops()* : This method will find all the loops and flip them with probability  $\frac{1}{2}$ . It goes over all the spin, and when any spin is not already in a loop, it uses the method *find\_next()* to know the following spin in the associated loop, thus until it goes back to the same spin. At each step, it stores the spin to know which spins have already been treated. When it finds a not treated spin, it chooses with probability  $\frac{1}{2}$  whether to flip all the spin along it or not.
- ★ *creategraph()* : Returns an array imaging the total graph configuration.

- ★ *pattern\_to\_string()* : Transforms the *spins* into a string to allow the script *check\_loop\_ergodicity.py* to check if the algorithm is ergodic, i.e. that all configurations are reachable.
- ★ *QMC\_step()* : This method applies a step on the configuration. First, it computes the pattern from the spins. Then, it uses the computed pattern to set the *total\_graph*. Then, thanks to the *find\_loop()* method it changes the spins to a new configuration.
- ★ *Quantum\_Monte\_Carlo(n\_warmup, n\_cycles, length\_cycle)* : This method uses the metropolis algorithm to compute the mean energy. First, the initial state with all spins down is "warmed up" by *n\_warmup* cycles. Then, there are *n\_cycles* cycles of *length\_cycle* steps. For each cycle, the energy is measured once. The parameter *length\_cycle* is used to get rid of the correlation between two successive configuration. This parameter can be optimize.

## 3.4 Results

### 3.4.1 Case $J_x = J_z$

**Parameters** For this case only, the graph  $G = 4$  is not considered, because it's weight is null.

**Validity of the algorithm** This case allowed us to check that our algorithm worked well with simple parameters. We used our code *ExactComputation* to know the theoretical values for the following parameters :  $J_x = J_z = -1$ ,  $m\_trotter = 10$ ,  $N\_spins = 8$ . The results for the Quantum Monte Carlo are computed thanks to the script *testloop.py*.

$\beta$	Theoretical mean energies	Monte Carlo computed mean energies
0.1	-0.146105	$-0.175369 \pm 0.045031$
0.2	-0.283938	$-0.318095 \pm 0.052067$
0.3	-0.412980	$-0.428188 \pm 0.047555$
0.4	-0.532973	$-0.550819 \pm 0.046009$
0.5	-0.643891	$-0.614586 \pm 0.039800$
0.6	-0.745901	$-0.775494 \pm 0.040258$
0.7	-0.839321	$-0.858818 \pm 0.039029$
0.8	-0.924584	$-0.912076 \pm 0.034779$
0.9	-1.002198	$-0.960219 \pm 0.035408$
1.0	-1.072715	$-1.081081 \pm 0.035624$

This computation confirms that our code runs and produces the good results for parameters with  $J_x = J_z$ . This was tested for fairly few *n\_cycles* in the Metropolis algorithm (10000), however it is sufficient.

**Ergodicity** In order to test our algorithm, we counted the number of different patterns that could be reached. If the algorithm is ergodic, it reaches all those patterns, that we are able to count. The script *check\_loop\_ergodicity.py* allowed us to verify this property.

**Autocorrelation** Let us show on figure 17 the mean energy depending on the number of moves. We use the measures produced by the script *compute\_x=z=1\_beta1.py*. Those are lists of  $1e7$  measures for  $J_x = J_z = 1$  and  $\beta = 1$ , but only the first 1000 moves are shown, it is on this time scale that the mean energy varies a lot. It can be seen that depending on the value of  $m\_trotter$ , the mean energy converges towards a different limit. This will be studied in the following paragraph.

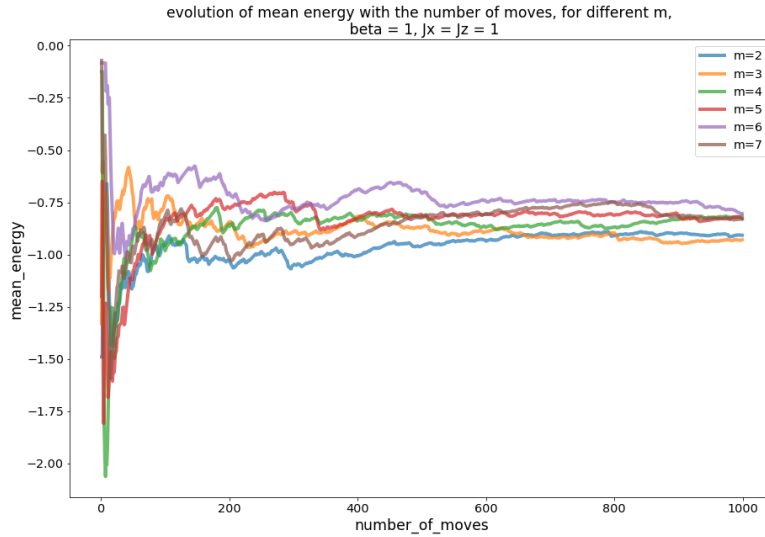


FIGURE 17

The script *loop\_result.ipynb* delivers the figure 17 and 18. The following figure directly computes the autocorrelation in the same way as in the local algorithm. One can observe that the autocorrelation is quite small, with an advised *length\_cycle* at 10.

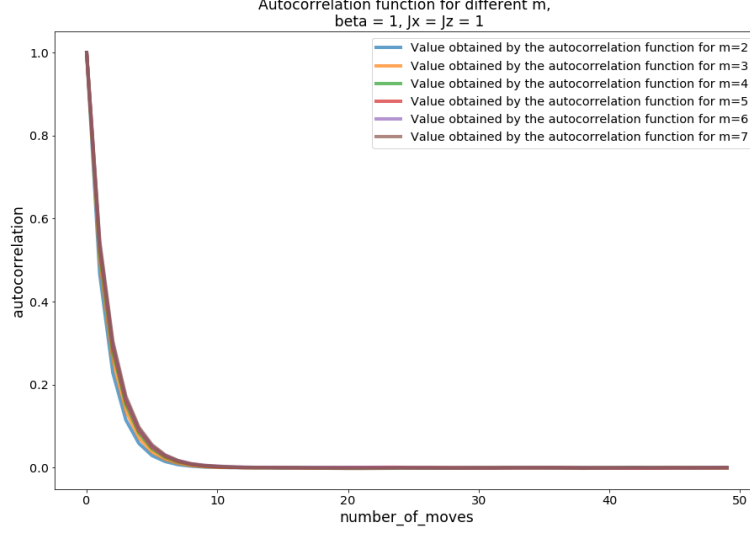


FIGURE 18

**Result dependence on  $m$**  For a given number of spins and a given division of the imaginary time, there is a given number of configuration of the spins. Each has a weight. The Quantum Monte Carlo computation of the energy returns an energy than can be predicted for low values of  $n\_spins$  and  $m\_trotter$ . Those predictable energies were given to us by M. Ferrero. The parameters are :

$$\begin{aligned}
 J_x = J_z = 1, \beta = 1, n\_spins = 4 \\
 m \in [2, 3, 4, 5, 6, 7] \\
 \Delta\tau \in [\frac{1}{2}, \frac{1}{3}, \frac{1}{4}, \frac{1}{4}, \frac{1}{5}, \frac{1}{6}, \frac{1}{7}, ]
 \end{aligned} \tag{14}$$

The following figure 19, given by the script *show\_ThvsComp.py*, shows the theoretical values for the energy on the orange dashed line. The energies are then computed by the loop algorithm with  $1e7$  measures. The error-bars are shown.

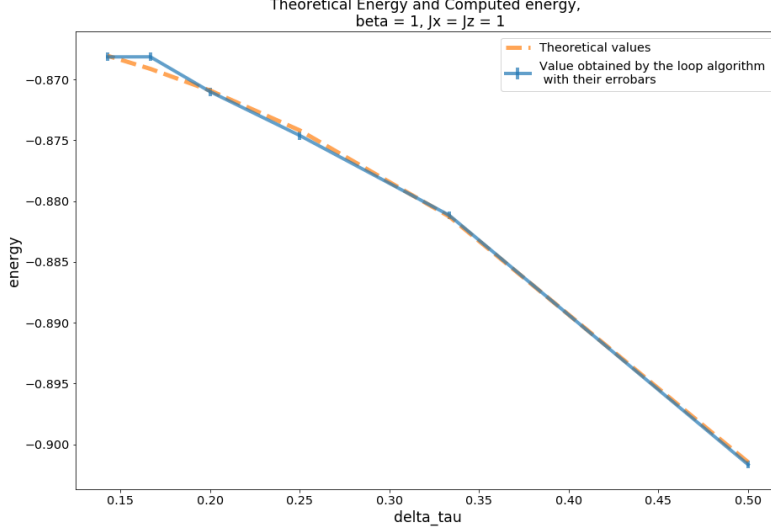


FIGURE 19

### 3.4.2 Case $J_x > |J_z|$

**Parameters** For this case, all graphs are considered. *During the encoding, because we faced difficulties, we tried to implement the code with another rule of repartition of the graphs' weights  $V(G)$  such as the graph  $G = 4$  was accessible only from the diagonal tiles  $S = 2$  (i.e.  $W(S = 3, G = 4) = 0$ ). This algorithm did not work, and the article written by Wessel that proposed such a repartition was wrong. Indeed, in this particular case, the detailed balance is not verified, as through the graph  $G = 4$ , the tiles  $S = 2$  are transformed to  $S = 3$ , and tiles  $S = 3$  can not reach tiles  $S = 2$ .*

**Validity of the algorithm** We used our code *ExactComputation* to know the theoretical values for the following parameters :  $J_x = 1.0$ ,  $J_z = 0.5$ ,  $m\_trotter = 10$ ,  $N\_spins = 8$ . The results for the Quantum Monte Carlo are computed thanks to the script *testloop.py*.

$\beta$	Theoretical mean energies	Monte Carlo computed mean energies
0.1	$-0.114286$	$-0.122148 \pm 0.014636$
0.2	$-0.231759$	$-0.255334 \pm 0.014792$
0.3	$-0.351769$	$-0.310946 \pm 0.013171$
0.4	$-0.473585$	$-0.487812 \pm 0.014690$
0.5	$-0.596416$	$-0.619067 \pm 0.014897$
0.6	$-0.719437$	$-0.747027 \pm 0.014502$
0.7	$-0.841814$	$-0.834654 \pm 0.013837$
0.8	$-0.962735$	$-0.936949 \pm 0.014312$
0.9	$-1.081434$	$-1.122287 \pm 0.014313$
1.0	$-1.197217$	$-1.149319 \pm 0.013671$

**Other results** Due to time of computation, major results will be given during the oral.



## 4 Conclusion

Thanks to the power of the loop algorithm, one can compute the mean energy with a very low *length\_cycle*. Quite rapidly, as shown in figure 17, the mean energy converges towards the limit, which is very close to the real value. This algorithm is reliable.

## 5 Bibliography

- ★ Assaad, Evertz, *World line and determinantal Quantum Monte Carlo methods for spins, phonons, and electrons*
- ★ Sandvik, *Monte Carlo simulations in classical statistical physics*