# Geometric Properties of Ising model on an Interacting Self-Avoiding Walks

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This is an abstract and that's really abstract.

### I. INTRODUCTION

The model of self-avoiding walks is one of extensively studied examples of linear polymers. Moreover, it is the simplest model to study critical behaviour while other models polymer chains have different states in the thermal equilibrium in various solvent conditions. By adding interaction between nearest neighbors of monomers of the walk, we are allowed to study phase transition fixed between solvent conditions, so the given polymer in the thermal equilibrium become extended in good-solvent conditions and collapsed in poor-solvent one. This tricritical nature was described in Ref. [1].

The impact of close-range interaction was studied precisely in models of magnetic polymers, where interaction between monomers became more complex after each monomer carried a spin and strength of nearest neighbors coupling became variable. This is so called Ising Model on a SAW conformation. In Ref. [2], the model was complicated by adding external magnetic field and all conclusions about magnetic properties were by comparing with mean-field model. However, there are some geometric properties which impact of magnetic properties is not clear while studying of require more brute methods.

In previous studies [3], it was established that Ising model on the self-avoiding walk conformations (SAWs) has a continious type of phase transition. In this work, we continue study geometric properties of this model and compare them with "papent" models and its modifications, such as Ising model on the rectangular lattice[4] and two-dimensional interacting self-avoiding walks exactly in their respective critical regions. We suggest that models with similar geometric properties will also have same magnetic propertries, what we suggest to observe in comparing values of Binder cumulants in the  $\theta$ -transition of models with the equal values of asphericities.

# II. MODELS AND METHODS

In the paper we consider several models: the first one is Ising model on interacting self-avoiding walk from the [3], on three different lattices: 2D-square lattice, 3D-square lattice and 2D-triangle lattice. The main difference between square and triangle lattice in defining two additional diagonal monomers on lattice as nearest too. Considering the case of lack of outer magnetic field in this work, the Hamiltonian of the model of fixed conformation u with length N and strength of nearest-neighbors

interaction J reads:

$$H_{u,N,\{\sigma\}} = -\sum_{\langle i,j\rangle} J\sigma_i\sigma_j, \quad i,j \in u, \ |u| = N$$
 (1)

The summation runs through spins involved in conformation and only with the nearest neighbors.

The second model considered in this paper is the Ising model on the rectangular lattice from the [4]. Simulated lattices has  $L \times rL$  spins and the Hamiltonian is calculated through interaction between all spins and their nearest neighbors respectively:

$$H_{L,r,\{\sigma\}} = -\sum_{\langle i,j\rangle} J\sigma_i \sigma_j \tag{2}$$

Here the *i*-th spin of the lattice has a pair of coordinates from  $[1..L] \times [1..rL]$ . For comparing magnetic properties of models with similar geometric ones we also define shape factors, such as gyration tensor of system with N points  $w_i$  [5]:

$$Q_{N,\alpha\beta} = \frac{1}{N} \sum_{i=1}^{N} (w_{i,\alpha} - w_{c,\alpha})(w_{i,\beta} - w_{c,\beta})$$
 (3)

where N is length of the system (number of monomers in conformations of Ising-ISAW models or number of spins in the lattice in rectangular Ising), and  $w_{i,\alpha}, w_{i,\beta}$  are the coordinates of i-th point of conformation.  $w_{c,\alpha}, w_{c,\beta}$  are coordinates of the center of system (so,  $Q_{N,xx}$  and  $Q_{N,yy}$  can be defined as mean squares of coordinates of the points of the model in the cartesian coordinate system with the center in the center of model). Eugen values q1, q2 of given tensor can be interpreted as  $Q_{N,xx}$  and  $Q_{N,yy}$  in the coordinate system of eugen vectors, or more important - as square of semi-axes of ellipse of inertia of given system. The proportion of them for systems with length N will be[5]:

$$r = \sqrt{\frac{\langle q_1 \rangle_N}{\langle q_2 \rangle_N}} \tag{4}$$

Eugen values q1, q2 are also used in enumerating another important shape factor - mean asphericity[5]:

$$\mathcal{A} = \left\langle \frac{(q_1 - q_2)^2}{(q_1 + q_2)^2} \right\rangle_N \tag{5}$$

The compared magnetic property of our models is the fourth order cumulant of the magnetization of the Binder cumulant, defined as [4]:

$$U_4 = 1 - \frac{\langle m^4 \rangle}{3\langle m^2 \rangle^2} \tag{6}$$

Where  $\langle m^4 \rangle$  and  $\langle m^2 \rangle$  are mean fourth and second order of mean magnetization per spin respectively.

We also need to define mean proportion of monomers with fixed number i of nearest neighbors  $\langle n_i \rangle$ , which is counted directly for every monomer in every simulated conformation of walk.

We are interested in comparing models in their respective critical regions. For each structure, critical temperatures of Ising models are known as [3, 4, 6]:

Structure	lattice	$T_c$
ISAW conformation	Square	1.198(9)[3]
ISAW conformation	Cubic	$1.90 \pm 0.02[6]$
Regular lattice	Rectangular	$2/\ln{(1+\sqrt{2})[7]}$

TABLE I. Known values of critical temperature of different modifications of Ising-ISAW model and normal Ising on the rectangular lattice

lattice	$T_c$
Square	1.4986(12) [8]
Cubic	1.318(29)[8]

TABLE II. Known values of critical temperature of different modifications of ISAW model

## III. RESULTS

### A. Mean Asphericity and Critical Cumulant

We attempted to learn how magnetic properties of Ising-like models depend on their geometrical ones and to define their comparability in critical region, where observable values of models don't depend on the length of conformation N. The idea is to compare critical cumulants  $U_4$  (6) of both models of Ising having equal asphericities. Both models are considered to have open boundary conditions (OBC). As we know, in the Ising model on rectangular lattice shape factors like aspect ratio r are the parameters, not observable values. Therefore, we can find (2) value of the aspect ratio of lattice for any asphericity  $\mathcal{A}$  (5) (see fig. 2). For rectangular lattice, the leading correction term to the asphericity behave like  $A^*(r) - A(r,L) \propto 1/L^2$ . Moreover, we know that value of Binder cumulant in Rectangular Ising[4] in critical region depends on aspect ratio r.

Using Monte-Carlo simulations with steps of Wolff algorithm [9], we enumerated Binder cumulant  $U_4$  of

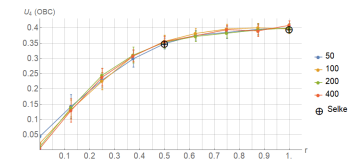


FIG. 1. Critical cumulant  $U_4(6)$  of Ising model on a rectangular lattice with open boundary conditions as function of aspect ratio r with side length L=50 (blue), 100 (yellow), 200 (green) and 400 (red). Black markers define values from [4]

Rectangular Ising model with aspect ratio r and open boundary conditions in  $\theta$ -point (see table I). Our results matched with known values from Ref. [4].

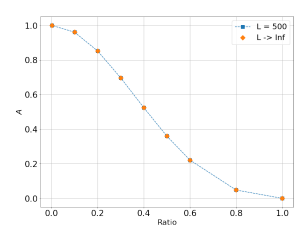


FIG. 2. Asphericity as function of aspect ratio r of the rectangular lattice with side length =500 and approximate values for rectangular lattice with infinitely long side

We enumerated values of asphericity (5) of Ising-ISAW model on 2D-square lattice in its critical region I for lengths N=1000-4900. (See figure 3) For simulations of this model we used method described in [3]. Vertical lines on figures define borders (according to statistical errors of known values from [6], our value from [3] is comparable with it) of critical regions of Ising-ISAW (red lines) and ISAW models (black line). Horizontal line define value of critical asphericity of ISAW model, which is known from Ref. [5].

We took mean values of asphericity of Ising-ISAW model in the borders of critical region and in the point of the best crossing of plots where we observe phase transition according to our numerical results. All these points are marked as black in zoomed figures 4. Our following

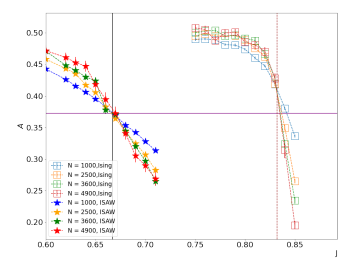


FIG. 3. Asphericity of Ising-ISAW (empty squares) and ISAW-only models (stars) as function of J=1/T, varying lengths of conformations N=1000 (blue), 2500 (yellow), 3600 (green) and 4900 (red)

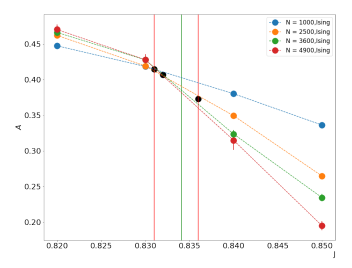


FIG. 4. Asphericity of Ising-ISAW model as function of zoomed in the critical region (red vertical lines, according to Ref. [6] and green vertical line, according to Ref. [3]), varying lengths of conformations N=1000 (blue), 2500 (yellow), 3600 (green) and 4900 (red)

steps was to pick up values of aspect ratio, so the Rectangular Ising had the same asphericity and to enumerate critical cumulant of the model with the same shape factors. As it seen from fig. 2, it is enough to use lattice with length N=500 for picking up the aspect ratio. For simulations we used cluster update based on Wolff algorithm [9] on a rectangular lattice with the same length.

As a result, comparison with critical cumulant of Ising-ISAW model, which was enumerated in [3] ( $U_4 = 0.308(8)$ ) showed significant mismatch of values, which means that we had not took into account some other

Ising-ISAW				
J	$\mathcal{A}$	r	$U_4$ Rectangular	
0.831	0.415	0.465	$0.340 \pm 0.006$	
0.832	0.4072	0.47	$0.343 \pm 0.006$	
0.836	0.373	$0.490 \pm 0.002$	$0.348 \pm 0.006$	

TABLE III. Values of critical cumulant for Ising model on rectangular lattice with mean asphericity related to Ising-ISAW model in its critical region

geometrical properties - for example, which will be considered in the next part - proportions of monomers with different quantities of nearest neighbors. It is obvious that in Ising model on the rectangular lattice most of monomers located inside the lattice and have 4 nearest neighbors, while monomers spread around the perimeter of the lattice have at least 2 (corners) and 3 nearest neighbors. Proportions in Ising-ISAW conformations are completely different.

#### B. Bulk

In this section we studied proportions of monomers with fixed numbers of nearest neighbors for Ising-ISAW and ISAW models on 3D-square and 2D-triangular lattices. Monomers of all four modified models can have from 2 to 6 close-range energy connections and some types of monomers, according to number of connections they have, can be interpreted similarly: for example, it is obvious that parts of conformations with monomers with only two nearest neighbors represent 1D-conformations or chains whatever lattice was used in observed model. And the opposite - regions where monomers have maximum number of close-range connections represent densely packed areas deep inside the Talking about cubic lattice, other types of monomers can be also interpreted: monomers with only three neighbors define "corners" on conformation, while only monomers on a cube edge (or it also can be on a isolated plane from other part of globule) can have four neighbors. Presence on five neighbors belongs to the monomers on a surface of conformation. Unfortunately, we cannot make similar interpretations for conformations on a triangular lattices.

# 1. Ising model on a SAW-conformation

Using method of MC simulations from Ref. [3], we enumerated proportions of monomer with 2-6 nearest neighbors in Ising-ISAW model on a three-dimensional square (or cubic) lattice.

As it is seen from left part of Figure 5, increasing the strength of nearest-neighbors interaction J leads to conformation becoming more dense as proportions of monomers with higher numbers of close-range connections significantly increases after  $\theta$ -transition located between blue vertical lines (see Table I).

We also repeated MC simulations for two-dimensional triangular lattice. Unlike the 3D-square lattice, the 5-th and 6-th possible neighbors located on the same plane as the first four, so conformations are expected to be more dense on this lattice.

Critical region of the model on a triangular lattice was not enumerated yet. But as it was suggested, density of conformations becomes higher as nearest-neighbors interaction J strengthen. Moreover, proportion of monomers with six neighbors on a triangular lattice is far higher than on a cubic one. It is also significant that "triangular" conformations with no inner interaction have almost twice shorter one-dimensional chains than conformations on a cubic lattice in the same conditions.

#### 2. ISAW model

To understand how the presence of magnetic properties affects on density of models, we also performed Monte-Carlo simulations on a parental model of self-avoiding walks on the same lattices. Results was similar to Ising-ISAW model, as suggested from "parent" model. Figure 5 shows that geometric aspects of phase transitions in ISAW model manifect earlier than in Ising-ISAW ones, despite the fact that the ISAW's  $\theta$ -point (black vertical line) is significantly above the Ising-ISAW one (blue vertical dashed lines) (see Table II).

### IV. DISCUSSION

#### V. ACKNOWLEDGMENTS

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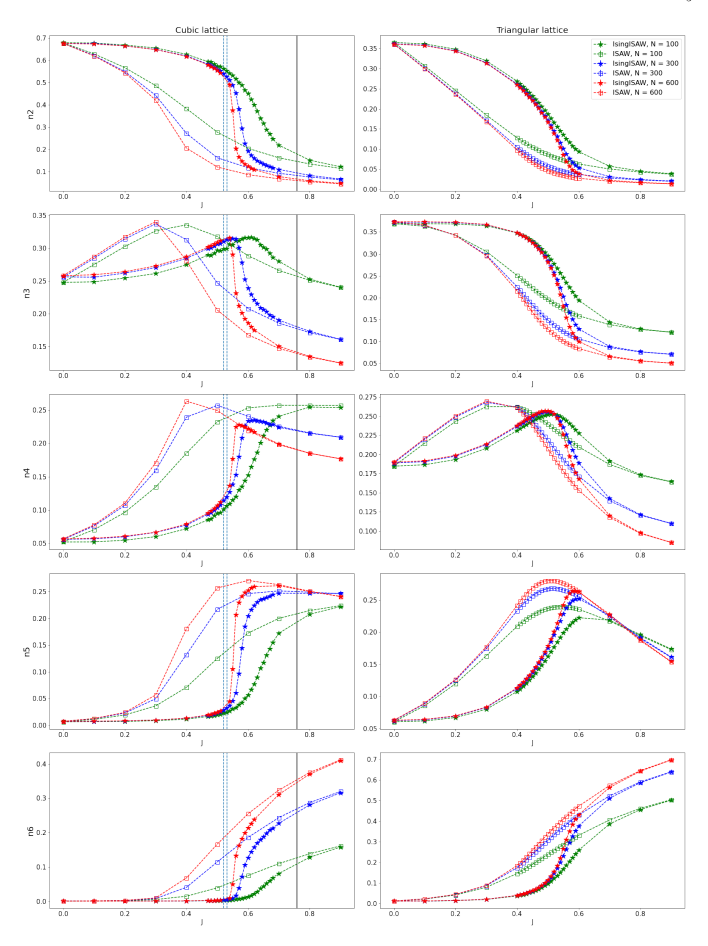


FIG. 5. Fractions of monomers of Ising-ISAW model (stars) and ISAW model (open squares) on a cubic lattice (left column) and 2D-triangle lattice (right column) with 2-6 nearest neighbors as function of J with length of conformations N=100 (green), 300 (blue) and 600 (red)