Pores in a two-dimensional network of DNA strands - computer simulations

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January 11, 2022

Abstract

Formation of random network of DNA strands is simulated on a two-dimensional triangular lattice. We investigate the size distribution of pores in the network. The results are interpreted within theory of percolation on Bethe lattice.

PACS numbers: 05.10.Ln, 87.14.Gg, 87.15.Aa
Keywords: DNA junctions; DNA networks computer simulations

1 Introduction.

Besides their extremely important biological function, molecules of DNA are useful in many areas, as biotechnology, nanotechnology and electronics. Possibility of design of molecules of exactly known composition and length opened an area for using DNA for construction of different structures. Appropriate choice of DNA sequences allows to obtain not only linear chains but also branched DNA molecules. These synthetical branches are analogs of branched DNA molecules (Holliday junctions) discovered in living cells. Addition of sticky ends to such junctions enables a construction of complex molecular systems. Holliday junctions occurring naturally are present between two symmetrical segments of DNA strands, and because of that, the branch point is free to migrate. Such a behaviour is not advantageous when DNA strands are used for a construction of networks. Elimination of symmetry in synthetically made DNA molecules fixes the branch points location. Synthetical junctions give an opportunity for building complex structures from branches prepared specially for given purposes; this enables to construct junctions with different numbers of arms. Addition of sticky ends to three- or four-armed branches allows to construct two-dimensional lattices, and using six-armed branches – for a construction of 3D lattices [1, 2]. Different kinds of branched molecules can be connected with linear strands or to each other. Appropriate choice of DNA and other allows for a construction of lattices of specific parameters.

One of the most important problems occurring during the network formation is that the angles between the arms of branched junctions are variable. This problem can be solved by using branches with four different sticky ends. Besides a stabilization of the edges, this solution eliminates improper connections between arms. Second important problem with DNA network formation is that the polymer strands are flexible. They can cyclize on itself what prevents further
growth of the system \cite{2}. To construct periodic lattices, such as e.g. crystals, which can be used in nanoelectronics, polymers must be rigid. Construction of crystals is possible since DNA double-crossover motifs were developed \cite{2}. As the DNA double helix width is about 2 nm and one helical repeat length is about 3.4 nm, lattices made from DNA strands can be useful in the emerging nanotechnology. It is possible to use DNA to form crystals with edge length of more than 10 $\mu$m and a uniform thickness of about 1-2 nm \cite{4}. One of main advantages of DNA structures is a possibility of their modifications by binding different ions and groups of ions or fluorescent labels \cite{3,2}. This is useful in electronic devices where nanometer-scale precision is extremely important.

Here we are interested in a design of random networks of DNA molecules. This aim is motivated as follows: the above mentioned limitations and conditions of the production of DNA lattices and crystals can be found to be too severe and in consequence too costful for some applications, where periodic structures are not necessary. In this case, properties of networks of DNA can be relevant for these potential applications of various two-dimensional structures, e.g. of polymers and liquid crystals \cite{6,7}. On the other hand, process of formation of a random network is close to some model processes of current interest for theories of disordered matter \cite{8}. Then, interpretation of our numerical results allows to comment to what extent assumptions of these theories are minimal. As it is discussed below, our results suggest that some conclusions of the percolation theory in uncorrelated Bethe lattice can apply to correlated structures.

We are going to investigate the size distribution of empty holes (pores) in a random network. This aim should allow to evaluate the network in a role of a kind of molecular sieve \cite{9}. Whilst disorder in such a sieve is obviously a disadvantage when compared to periodic structures \cite{2}, it seems to us worthwhile to evaluate consequences of this disorder.

In subsequent section the model is described and some details are given on its numerical realization. In Section III we show the results on the size distribution of the pores, obtained for a triangular two-dimensional lattice. Short discussion closes the text.

2 Model

In our model we initially consider two kinds of molecules: linear DNA strands and DNA-junctions, which connect ends of the strands. The latter can be of three or four arms; they are called Y- and X-junctions. We assume that the length of the linear molecules is much larger, than the size of the junctions. Also, we assume that the number of junctions is so much larger than the number of strands, that \textit{i}) practically any collision of two ends of the strands leads to their fixed connection, \textit{ii}) the junctions do not influence the motion of the strands in other way, than as in \textit{i}). Summarizing, our assumptions on the DNA-junctions (enough small and enough numerous) allow to neglect them during the simulation.

To speed up numerical calculations, the algorithm is based on a periodic lattice. It is straightforward to apply the square lattice in the case of X-junctions, and the triangular lattice in the case of Y-junctions. The length of the linear DNA strand is chosen to be an integer multiple of the size of the lattice cell.
An additional assumption is that, no more than arbitrary chosen number \( M \) of different DNA strands are placed at the same lattice node, at the same time. This limits the density of the system from above, to reproduce the steric effect. We have only three model parameters: the length \( L \) of the strands, their number \( N \), and the maximal number \( M \) of strands at one lattice cell. A strand cannot rotate, but only moves along its axis. The direction of the latter is random, with equal probabilities \( 1/2 \) at each time step. Initial set of the strand positions is selected to be entirely random (positions and orientations), with only the above mentioned limitation of the density. We choose periodic boundary conditions, to limit an influence of the borders. Subsequent positions are controlled by the assumption that in the one cell can not be more molecules that arbitrary chosen number \( M \). Possible orientations of the strands are horizontal or vertical in the case of square lattice. In the case of triangular lattice, three orientations of the strands are equally possible, along all three sides of the triangle. If the number of ends of the strands at a given cell of the lattice is equal to or more than two (but not more than \( M \), which is forbidden by the code), these strands become connected to a DNA-junction and to each other and they cannot move anymore. If at any time step two different strands are placed at the same cells only one of them, chosen randomly, can be connected to the DNA-junction. In one simulation step all molecules which are not immobile can move with equal probability to the right or left if they are oriented horizontally, and up or down if they are vertical in the case of square lattice. In the case of triangular lattice the strands can move in two possible directions, in accordance with their orientation. At each time step each strand can move by the length equal to the size of the one lattice cell. A movement is accepted if, as its consequence, the number of strands at all lattice cells is not more then \( M \).

All results are obtained as average value from ten independent simulations, and the lattice size was 512\( \times \)512. Each simulation was made for 100 time steps. Duration of the simulation was chosen so as to a number of free strands was in range of few percent (2-3). In all simulations we studied distribution of the pores, i.e. the lattice nodes which are not occupied by the DNA strands as a function of the model parameters. The size distribution of the pores was investigated using the Hoshen-Kopelman algorithm [10].

### 3 Results

The results presented in Figs. 1-4 are obtained for the case of triangular lattice with \( M = 3 \). The number \( N \) of molecules introduced to the system varies from \( 10^4 \) (Fig. 1) to \( 4 \times 10^4 \) (Fig. 4). The plot shown in Fig. 1 shows that there are large pores in the system, but their number is exactly one for each size \( s \); in other words, the system is diluted. For larger \( N \) we see that the pore size distribution is close to a power law, i.e. \( N_g(s) \propto s^{-\tau} \), with \( \tau \approx 1.6 \) for \( N = 2 \times 10^4 \) and \( \tau \approx 2.0 \) for \( N = 3 \times 10^4 \). This is seen in Figs. 2 and 3. A further increase of the number \( N \) of molecules leads to a cutoff of the scale-free character of the plot (Fig. 4). The size of maximal pores is then reduced from 100 or more to about 30.

Similar results are obtained for other strand lengths \( L \). As a rule, for low values of \( N \) the shape of the function \( N_g(s) \) is close to the one in Fig. 1. For larger \( N \) we see a power-law function, as in Figs. 2 and 3. When \( N \) increases
further, large pores vanishes. This common behaviour is summarized in Fig. 5, where a kind of phase diagram is presented. Assuming that the scale-free power-law function is proper at only one value of $L_c(N)$, where the concentration of the strands is at the percolation threshold, we marked approximate position of this threshold with a schematic solid line. This plot is qualitative: the only conclusion from our data is that $L_c$ decreases with $N$.

4 Discussion

As it was noted in the Introduction, our results appear to be close to what could be expected for an uncorrelated percolation in Bethe lattice [8]. For $M = 3$, the percolation threshold is $p_c = 1/(M - 1) = 0.5$. By symmetry, we can interchange occupied cells and empty cells. Then, the pore distribution for $p > p_c$ is equivalent to the cluster size distribution below the percolation threshold, i.e. for $p < p_c$, where the giant component is absent. At the percolation threshold, we expect that the average number of strands at the network nodes is $p_cM = 1.5$. Actually, we obtain the average number of strands equal to 1.39, 1.48, 1.52 and 1.53 for $L = 7$ and $N = 1, 2, 3$ and $4 \times 10^4$, respectively. This means that, keeping the analogy with the uncorrelated percolation in the Bethe lattice, we are close to the percolation threshold. Theoretical value of the exponent $\tau$ in the two-dimensional Bethe lattice is 2.5 [8]. Taking into account our numerical errors, this is not in contradiction with our evaluations. Moreover, the cluster size distribution above the percolation threshold in the Bethe lattice theory fulfils the equation $N_g(s, L)/N_g(s, L_c) \propto \exp(-cs)$. This formula is checked in Fig. 6 with data from Figs. 3 and 4. Again, the obtained data are not in contradiction with theory.

However, our network is neither uncorrelated, nor the Bethe lattice. Cor-

Figure 1: Pore size distribution $N_g(s)$ for $L = 7$, $N = 10^4$, $M = 3$. 

![Figure 1: Pore size distribution $N_g(s)$ for $L = 7$, $N = 10^4$, $M = 3$.](image)
Figure 2: Pore size distribution $N_g(s)$ for $L = 7$, $N = 2 \times 10^4$, $M = 3$.

Figure 3: Pore size distribution $N_g(s)$ for $L = 7$, $N = 3 \times 10^4$, $M = 3$. 
Figure 4: Pore size distribution $N_g(s)$ for $L = 7$, $N = 4 \times 10^4$, $M = 3$.

Figure 5: Phase diagram (L,N) for $M = 3$. Different kinds of plots of $N_g(s)$ are marked with different signs. In the case of small $N$, a part of plot for large $s$ is a horizontal line at $N_g = 1$, as in Fig.1; this kind of plot is marked by ($\times$). The scale-free power functions, as in Figs. 2 and 3; this kind of plot is marked by ($\bullet$). Finally, for the largest $N$, in many cases we observe a curve with a cutoff, as in Fig.4; this kind of plot is marked by ($\triangle$). A hypothetical percolation threshold is marked by a solid line.
relations appear because the strands are allowed to move until finally attached to some other strands. From this point of view, the system is analogous to the diffusion-limited aggregation (DLA), where the density of the growing cluster decreases in time. The difference is that in DLA, the cluster grows from some nucleus, which is absent in our simulation. On the other hand, loops are not prevented here, unlike in the Bethe lattice. The similarity of our results to the latter model allows to suspect that in random structure near the percolation threshold the contribution to the cluster size distribution from accidental loops remains small.

The same calculations are performed also for the two-dimensional square lattice. Here the power-law like character of the pore size distribution is less convincing. It seems likely that this additional difficulty is due to the fact that the triangular lattice is closer to the off-lattice model than the square lattice. In general, the lattice introduces numerical artifacts, which can be expected to disturb the system for small values of $L$. In the limit of large $L$, it is only the anisotropy of the lattice which is different from the ideal off-lattice limit. This numerically induced anisotropy allows for three orientations for the triangular lattice and two orientations for the square lattice. In an off-lattice calculation, the number of orientations would be infinite.
References

[1] Seeman N.C., TIBTECH 17 (1999) 437
[2] Seeman N.C. et al., Nanotechnology 9 (1998) 257
[3] Ito Y., Fukusaki E., J.Mol.Catal.B:Enzymatic 28 (2004) 155
[4] Niemeyer Ch.M., Current Opinion in Chemical Biology 4 (2000) 609
[5] Mao Ch. et al., Nature 397 (1999) 144
[6] Clarke S. M. et al., Macromol.Chem.Phys. 198 (1997) 3485
[7] Sheming Lu et al., Cryst. Res. Technol. 39 (2004) 89
[8] Stauffer D., Aharony A., Introduction to Percolation Theory, 2nd edition, Taylor and Francis, Routledge 2003
[9] Polarz S., Smarsly B., J.Nanoscience and Nanotechnology 2 (2002) 581
[10] Hoshen J., Kopelman R., Phys. Rev. B 14 (1976) 3428