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Statistical geometry of packing defects of lattice chain polymer from enumeration and sequential Monte Carlo method

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Voids exist in proteins as packing defects and are often associated with protein functions. We study the statistical geometry of voids in two-dimensional lattice chain polymers. We define voids as topological features and develop a simple algorithm for their detection. For short chains, void geometry is examined by enumerating all conformations. For long chains, the space of void geometry is explored using sequential Monte Carlo importance sampling and resampling techniques. We characterize the relationship of geometric properties of voids with chain length, including probability of void formation, expected number of voids, void size, and wall size of voids. We formalize the concept of packing density for lattice polymers, and further study the relationship between packing density and compactness, two parameters frequently used to describe protein packing. We find that both fully extended and maximally compact polymers have the highest packing density, but polymers with intermediate compactness have low packing density. To study the conformational effects of void formation, we characterize the conformational reduction factor of void formation and found that there are strong end-effect. Voids are more likely to form at the chain end. The critical exponent of end-effect is twice as large as that of self-contacting loop formation when existence of voids is not required. We also briefly discuss the sequential Monte Carlo sampling and resampling techniques used in this study. © 2002 American Institute of Physics.

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I. INTRODUCTION

Soluble proteins are well-packed, and their packing densities may be as high as that of crystalline solids. $^{1-3}$ Yet there are numerous packing defects or voids in protein structures, whose size distributions are broad. The volume (v) and area (a) of protein does not scale as $v \approx a^{3/2}$, which would be expected for models of tight packing. Rather, v and a scale linearly with each other. In addition, the scaling of protein volume and cluster-radius is characteristic of random sphere packing. Such scaling behavior indicates that the interior of proteins is more like Swiss cheese with many holes than tightly packed jigsaw puzzles.

What effects do voids have? Proteins are often very tolerant to mutations, 3,6-8 which may suggest potentially stabilizing roles of voids in proteins. Voids in proteins are also often associated with protein function. The binding sites of proteins for substrate catalysis and ligand interactions are frequently prominent voids and pockets on protein structures. However, the energetic and kinetic effects of maintaining specific voids in proteins are not well understood, and the shape space of voids of folded and unfolded proteins are largely unknown.

In this paper, we examine the details of the statistical

nature of voids in simple lattice polymers. Lattice models have been widely used for studying protein folding, where the conformational space of simplified polymers can be examined in detail. Despite its simplistic nature, lattice model has provided important insights about proteins, including collapse and folding transitions, Influence of packing on secondary structure formation, Influence of packing on secondary structure formation, and designability of lattice structures. However, one drawback is that the lattice model is not well-suited for studying void-related structural features, such as protein functional sites, since it is not easy to model the geometry of voids.

In this article, we first define voids as topological defects and describe a simple algorithm for void detection in the two-dimensional lattice. We then enumerate exhaustively the conformations for all n-polymers up to n = 25, and analyze the relationship of probability of void formation, expected packing density, and compactness, as well as expected wall interval of void with chain length. To study statistical geometry of long chain polymers, we describe a Monte Carlo sampling strategy under the framework of Sequential Importance Sampling, and introduce the general technique of resampling. The results of simulation of long chain polymers up to N = 200 for several geometric parameters are then presented. We further explore the conformation reduction factor R of void formation, and describe the significant end-effect of

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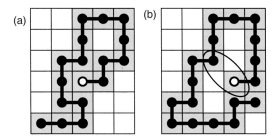


FIG. 1. Voids of polymers in a square lattice. Unfilled circle represents the first monomer. (a) A void of size 1 is formed in this 17-mer. (b) The two monomers encircled shares a vertex but not an edge of a square and are not in topological contact. The unfilled space contained within the polymer is regarded as one connected void of size 4.

void formation, as well as the scaling law of *R* and wall interval of voids. In the final section, we summarize our results and discuss effective sampling strategy for studying the conformational space of voids.

II. LATTICE MODEL AND VOIDS

Lattice polymers are self-avoiding walks (SAWs), which can be obtained from a chain-growth model. $^{26-28}$ Specifically, an n-polymer P on a two-dimensional square lattice \mathbb{Z}^2 is formed by monomers n_i , $i \in \{1,...,N\}$. The location x_i of a monomer n_i is defined by its coordinates $x_i = (a_i, b_i)$, where a_i and b_i are integers. The monomers are connected as a chain, and the distance between bonded monomers x_i and x_{i+1} is 1. The chain is self-avoiding: $x_i \neq x_j$ for all $i \neq j$. We consider the beginning and the end of a polymer to be distinct. Only conformations that are not related by translation, rotation, and reflection are considered to be distinct. This is achieved by following the rule that a chain is always grown from the origin, the first step is always to the right, and the

chain always goes up at the first time it deviates from the x-axis. For a chain polymer, two nonbonded monomers n_i and n_j are in *topological contact* if they intersect at an edge that they share. If two monomers share a vertex of a square but not an edge, these two monomers are defined as not in contact.

When the number of monomer n is 8 or more, a polymer may contain one or more void [Fig. 1(a)]. We define voids as topological features of the polymer. The complement space $\mathbb{Z}^2 - P$ that is not occupied by the polymer P can be partitioned into disjoint components,

$$\mathbb{Z}^2 - P = V_0 \dot{\cup} V_1 \cdots \dot{\cup} V_k$$
.

Here V_0 is the unique component of the complement space that extends to infinity. We call this the *outside*. The rest of the components that are disjoint or disconnected to each other are *voids* of the polymer. Because nonbonded monomers intersecting at a vertex are defined as not in contact, they do not break up the complement space. As an example, the unfilled space contained within the polymer in Fig. 1(b) is regarded as one connected void of size 4 rather than two disjoint voids of size 2. This choice is arbitrary, but is consistent with the definition of contact. A simple algorithm for void detection can be found in the Appendix.

III. VOID DISTRIBUTION BY EXACT ENUMERATION

A. Probability of forming voids and expected number of voids

The number of conformations $\omega(n)$ for an *n*-polymer up to n=25 obtained by exhaustive enumeration is shown in Table I. The numbers of conformations for polymers up to

TABLE I. Number of	conformations of	f an <i>n</i> -polymer	with different numb	er of voids	on a square lattice.

n	$\omega(n)$	$\omega_0(n)$	$\omega_1(n)$	$\omega_2(n)$	$\omega_3(n)$	$\omega_4(n)$
3	2	2	0	0	0	0
4	5	5	0	0	0	0
5	13	13	0	0	0	0
6	36	36	0	0	0	0
7	98	98	0	0	0	0
8	272	270	2	0	0	0
9	740	734	6	0	0	0
10	2034	1993	41	0	0	0
11	5513	5393	120	0	0	0
12	15037	14508	529	0	0	0
13	40617	39078	1536	3	0	0
14	110188	104566	5602	20	0	0
15	296806	280599	16088	119	0	0
16	802075	748335	53149	591	0	0
17	2155667	2002262	151052	2353	0	0
18	5808335	5327888	470386	10051	10	0
19	15582342	14222389	1325590	34287	76	0
20	41889578	37784447	3973361	131298	472	0
21	112212146	100673771	11119456	416239	2680	0
22	301100754	267136710	32479871	1471874	12293	6
23	805570061	710673806	90361878	4479355	54998	24
24	2158326727	1883960171	259195774	14946910	223458	414
25	5768299665	5005591512	717505892	44337381	862748	2132

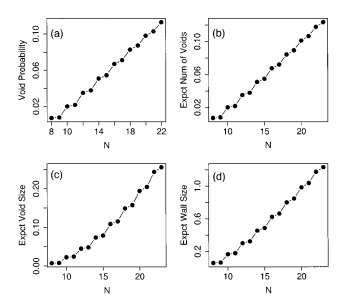


FIG. 2. Geometric properties of chain polymers by exhaustive enumeration. (a) The probability of void formation, (b) the expected number of voids contained in a polymer, (c) the expected void size, and (d) the expected wall size of voids. All these parameters increase with chain length.

n=15 are in exact agreement with those reported in Chan and Dill.¹² Table I also lists the number of conformations $\omega_k(n)$ containing k=1,2,3, or 4 voids.

The probability for a polymer to form one or more voids π_n is calculated as

$$\pi_v = \frac{\sum_{i=1}^k \omega_k(n)}{\omega(n)}.$$

The expected number of voids \bar{n}_v for a polymer is

$$\bar{n}_v = \frac{\sum_{i=1}^k \omega_k(n) \cdot k}{\omega(n)}.$$

As the chain length grows, it is clear that both π_v and \bar{n}_v increases [Figs. 2(a) and 2(b)].

B. Void size

The total size v of voids in a polymer is the sum of the sizes of all voids, namely, the total number of all unoccupied squares that are fully contained within the polymer. Let $\omega_v(n)$ be the number of conformations of n-polymer with total void size v. The expected total void size v for the v-polymer is

$$\overline{v} = \frac{\sum_{v} \omega_{v}(n) \cdot v}{\omega(n)}.$$

Figure 2(c) shows that the expected void size \bar{v} increases with chain length n.

C. Wall size of void

For a void V of size v, what is the required minimum length l(v) for a polymer that can form such a void? Equivalently, what is the size of the wall of the polymer containing void V? Here we first restrict our discussion to voids formed only by strongly connected unoccupied sites, namely, any

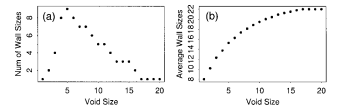


FIG. 3. Voids of fixed size in 22-mers can have different shapes and thus sometimes different wall sizes. (a) The distribution of the number of observed different wall sizes for a void depends on the size of the void. Voids of size 5 have the maximum number of different wall sizes. (b) The expected wall size for voids of different size.

neighboring two sites of a void must be sharing at least one edge of the squares. We exclude voids containing weakly connected sites, where two neighboring sites are connected by only one shared vertex [Fig. 1(b)]. For v = 1, 2, and 3, it is easy from the geometry of the voids to see that l(v) = 8, 10, and 12, respectively. However, in general l(v) also depends on the shape of the void. A void of size 4 can have five different shapes. If the void is of the shape of a 2×2 square, l(4) = 12. For the other four shapes, l(4) = 14.

For any strongly connected void, we find that the following general recurrence relationship for l(v) holds:

$$l(v) = l(v-1) + \begin{cases} 2, & \text{if } \Delta \partial V = 3 \\ 0, & \text{if } \Delta \partial V = 2 \\ -1, & \text{if } \Delta \partial V = 1 \end{cases}$$

where ∂V represents the boundary edges of void V, and $\Delta \partial V$ represents the net gain in the number of boundary edges introduced by the newly added unoccupied site. Although the number and explicit shapes of strongly connected voids of size up to 5 can be found in Ref. 29, there is no general analytical formula known for the number of shapes of a void of size v. This is related to the problem of determining the number of polyominos or animals (as in percolation theory) of a given size.

When weakly connected voids are also considered, there are more possible wall sizes for the void. For 22-mer, the number of different wall sizes observed for a void, strongly or weakly connected, at various size are shown in Fig. 3(a). Voids of size 5 have the largest diversity in wall size. This is of course due to the fixed chain length. A short chain such as the 22-mer has only a small number of ways for forming large voids. Figure 3(b) shows the average wall size for various void sizes in the 22-mer. The expected or average wall size $\overline{w}(n)$ for a void in an n-polymer can be calculated as

$$\overline{w}(n) = \sum_{v} w \cdot \frac{\omega_{v,w}(n)}{\omega_{v}(n)},$$

where v is the void size, w is the wall size of the void, $\omega_{v,w}(n)$ is the number of n-polymers containing a void of size v with wall size w, and $\omega_v(n)$ is the total number of n-polymers with a void of size v. Figure 2(d) shows that $\overline{w}(n)$ increases with chain length. Wall size and void size are analogous to the area and volume of voids in three-dimensional space.

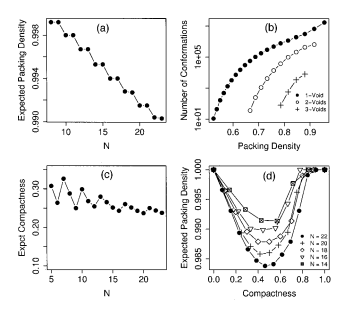


FIG. 4. Packing density and compactness are two useful parameters describing packing of chain polymers. (a) The expected packing density decreases with chain length. (b) For the 22-mer, the majority of the conformations with 1-void have a high packing density, namely, the size of the void is small. Fewer conformations are found with large voids. The same pattern is observed for conformations with 2 and 3 voids. (c) The expected compactness fluctuates but in general decreases with chain length. (d) The relationship of average packing density p and average compactness ρ for the chain polymer of length N=14-22.

D. Packing density

An important parameter that describes how effectively atoms fill space is the packing density p. In proteins, it is defined by Richards and co-workers as the amount of the space that is occupied within the van der Waals envelope of the molecule, divided by the total volume of space that contains the molecule.^{3,30} It has been widely used by protein chemists as a parameter for characterizing protein folding.³ Following this original definition, the packing density p for the lattice polymer is

$$p = n/(n+v)$$
,

when a n-polymer has a total void size of v.

The expected packing density $\bar{p}(n)$ for an *n*-polymer can be calculated as

$$\bar{p}(n) = \sum_{p} p \cdot \frac{\omega_{p}(n)}{\omega(n)},$$

where $\omega(n)$ is the number of all conformations of n-mer, $\omega_p(n)$ is the number of n-mers with packing density of p. The scaling of $\bar{p}(n)$ with the chain length n decreases roughly linearly between n=7 and n=22 [Figure 4(a)]. Because it takes at least two additional monomers to increase the size of a void by one, $\bar{p}(n)$ decreases only when n is an odd number for short chains.

Although voids are packing defects, most conformations with voids have high packing density, namely, the total size of the voids is small. Among all conformations of the 22-mer containing one void, the number of conformations increases monotonically with packing density. The lowest packing density 0.52 has only 11 conformations, whereas the highest

packing density 0.92 has the largest number (6756751) of conformations [Fig. 4(b)]. A similar relationship is found among conformations with 2 and 3 voids [Fig. 4(b)].

E. Compactness

Another important parameter that measures the packing of the lattice polymer is the number of nonbonded contacts t. It is related to the *compactness* parameter ρ , defined by Chan and Dill¹² as $\rho = t/t_{\text{max}}$, where t_{max} is the maximum number of nonbonded contacts possible for an n-polymer. Compactness ρ has been studied extensively in seminal works by Chan and Dill. ^{12,23,31} Although p is sometimes correlated with the compactness ρ , these two parameters are distinct. The relationship between compactness and expected packing density for chain polymer of length 14–22 is shown in Fig. 4. For all chain lengths, both maximally compact polymer $(\rho = 1)$ and extended polymer $(\rho = 0)$ have maximal packing density (p=1), but polymers with low packing densities have intermediate compactness on average. Polymers with ρ between 0.4 and 0.6 have lowest packing density and therefore tend to have larger void size. The explanation is simple. An extended lattice chain polymer has no voids, it therefore achieves maximal packing density of p = 1, but its compactness ρ is 0. A maximally compact polymer with $\rho = 1$ also contains no voids, its p is 1. On the other hand, nonmaximally compact polymers can have a range of packing densities.

IV. OBTAINING VOID STATISTICS FOR LONG CHAIN POLYMERS VIA IMPORTANCE SAMPLING

Sequential Importance Sampling: Geometrically complex and interesting features emerge only in polymers of sufficient length, which are not accessible for analysis by exhaustive enumeration, due to the fact that the number of possible SAWs increases exponentially with the chain length. Monte Carlo methods are often used to generate samples from all possible conformations and obtain estimates of feature statistics using those samples. However, when chain length becomes large, the direct generation of the SAWs using the rejection method (i.e., generate random walks on the lattice and only accept those that are self-avoiding) from the uniform distribution of all possible SAWs becomes difficult. The success rate s_N of generating SAWs decreases exponentially, $s_N \approx Z_N/(4 \times 3^{N-1})$. For N = 48, s_N is only 0.79%.³² To overcome this attrition problem, a widely used approach is the Rosenbluth Monte Carlo method of biased sampling.²⁶ The task is to grow one more monomer for a t-polymer chain that has been successfully grown from 1 monomer after t -1 successive steps without self-crossing, until t=n, the targeted chain length. In this method, the placement of the (t+1)th monomer is determined by the current conformation of the polymer. If there are n_t unoccupied neighbors for the tth monomer, we then randomly (with equal probability) set the (t+1)th monomer to any one of the n_t sites. However, the resulting sample is biased toward more compact conformations and does not follow the uniform distribution. Hence each sample is assigned a "weight" to adjust for this bias. Any statistic can then be obtained from weighted average of the samples. In the case of the Rosenbluth chain growth method, the weight is computed recursively as $w_t = n_t w_{t-1}$.

Liu and Chen³³ provided a general framework of Sequential Monte Carlo (SMC) methods which extend the Rosenbluth method to more general setting. Sophisticated but more flexible and effective algorithms can be developed under this framework. In the context of growing polymer, SMC can be formulated as follows. Let $(x_1,...,x_t)$ be the position of the t monomers in a chain of length t. Let $\pi_1(x_1), \pi_2(x_1,x_2),...,\pi_t(x_1,...,x_t)$ be a sequence of *target* distributions, with $\pi(x_1,...,x_n) = \pi_n(x_1,...,x_n)$ being the final objective distribution from which we wish to draw inference from. Let $g_{t+1}(x_{t+1}|x_1,...,x_t)$ be a sequence of *trial distributions* which dictates the growing of the polymer. Then we have:

Procedure SMC (n)

Draw $x_1^{(j)}$, j=1,...,m from $g_1(x_1)$ Set the incremental weight $w_1^{(j)}=\pi_1(x_1^{(j)})/g_1(x_1^{(j)})$ for t=1 to n-1for j=1 to m// Sampling for the (t+1)th monomer for the jth sample Draw position $x_{t+1}^{(j)}$ from $g_{t+1}(x_{t+1}|x_1^{(j)}\cdots x_t^{(j)})$

// Compute the incremental weight.

$$u_{t+1}^{(j)} \leftarrow \frac{\pi_{t+1}(x_1^{(j)} \cdots x_{t+1}^{(j)})}{\pi_t(x_1^{(j)} \cdots x_t^{(j)}) \cdot g_{t+1}(x_{t+1}^{(j)} | x_1^{(j)} \cdots x_t^{(j)})}$$

$$w_{t+1}^{(j)} \leftarrow u_{t+1}^{(j)} \cdot w_t^{(j)}$$

endfor Resampling

endfor

At the end, the configurations of successfully generated polymers $\{(x_1^{(j)},...,x_n^{(j)})\}_{j=1}^m$ and their associated weights $\{w_n^{(j)}\}_{j=1}^m$ can be used to estimate any properties of the polymers, such as expected void size, compactness, and packing density. That is, the objective inference $\mu_h = E_{\pi}[h(x_1,...,x_n)]$ is estimated with

$$\hat{\mu}_h = \frac{\sum_{j=1}^m h(x_1^{(j)}, \cdots, x_n^{(j)}) \cdot w_n^{(j)}}{\sum_{j=1}^m w_n^{(j)}},\tag{1}$$

for any integrable function h of interests.

The critical choices that affect the effectiveness of the SMC method are (1) the approximating target distribution $\pi_t(x_1 \cdots x_t)$, (2) the sampling distribution $g_{t+1}(x_{t+1}|x_1 \cdots x_t)$, and (3) the resampling scheme. In this study, we are interested in sampling from the uniform distribution $\pi_n(x_1 \cdots x_n)$ of all geometrically feasible conformations of length n, which we call the final objective distribution. It can also be chosen to be the Boltzmann distribution when energy function such as the HP model is introduced.

The Rosenbluth method²⁶ is a special case of SMC. Its target distributions $\pi_t(x_1 \cdots x_t)$ is the uniform distribution of all SAWs of length t. Its sampling distribution

 $g_{t+1}(x_{t+1}|x_1\cdots x_t)$ is the uniform distribution among all $n_1(x_1,\dots,x_t)$ unoccupied neighboring sites of the last monomer x_t , and the weight function is

$$w(x_1,...,x_t,x_{t+1}) = w(x_1,...,x_t)n_1(x_1,...,x_t).$$

When there is no unoccupied neighboring sites $(n_1(x_1,...,x_t)=0)$, there is no place to place the (t+1)th monomer. In this case, the chain runs into a dead end and we declare the conformation *dead*, with weight assigned to be 0. In the case of Rosenbluth method, no resampling is used.

Similarly, the k-step look ahead algorithm $^{3\overline{2},36}$ chooses $\pi_{t+1}(x_1,...,x_{t+1})$ being the marginal distribution of $\pi_{t+k}^*(x_1,...,x_{t+k})$, the uniform distribution of all SAWs of length t+k. Hence π_{t+1} is closer to the final objective distribution—the uniform distribution of all SAWs of length n. Specifically,

$$\pi_{t+1}(x_1, ..., x_{t+1})$$

$$= \sum_{x_{t+2}, ..., x_{t+k}} \pi_{t+k}^*(x_1, ..., x_{t+1}, x_{t+2}, ..., x_{t+k})$$

$$\propto n_k(x_1, ..., x_{t+1}),$$

where $n_k(x_1,...,x_{t+1})$ is the total number of SAWs of length t+k "grown" from $(x_1,...,x_{t+1})$ [i.e., with the first (t+1) positions at $(x_1,...,x_{t+1})$.] In the k-step look-ahead algorithm, the sampling distribution is

$$g_{t+1}(x_{t+1} = x | x_1,...,x_t) = \frac{n_k(x_1,...,x_t,x)}{n_{k+1}(x_1,...,x_t)}$$

It chooses the next position according to what will happen k steps later. Namely, the probability of placing the t+1th monomer at x is determined by the ratio of the total number of SAWs of length t+k grown from $(x_1,...,x_t,x)$ and the total number of SAWs of the same length t+k grown from one step earlier $(x_1,...,x_t)$. The corresponding weight function is

$$w(x_1,...,x_t,x_{t+1}) = \frac{n_k(x_1,...,x_{t+1})}{n_k(x_1,...,x_t) \cdot \frac{n_k(x_1,...,x_{t+1})}{n_{k+1}(x_1,...,x_t)}}$$
$$= \frac{n_{k+1}(x_1,...,x_t)}{n_k(x_1,...,x_t)}.$$

Although it has higher computational cost, it usually produces better inference on the final objective distribution, with less "dead" conformations. The standard Rosenbluth algorithm is a 1-step look ahead algorithm.

To compare geometric properties estimated from sequential Monte Carlo method and those obtained by exhaust enumeration, we examine the expected number of voids and expected void size for polymer of chain length 14–22. Figure 5 shows that sequential Monte Carlo can provide very accurate estimation of these geometric properties of voids. Here 2-step look ahead is used, with Monte Carlo sample size of 100 000 and no resampling is applied.

The resampling step is one of the key ingredients of the SMC method. ^{33,37} There are many cases where resampling is beneficial. First, note that it is unavoidable to have some

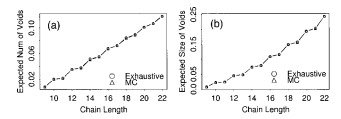


FIG. 5. Geometric properties obtained by enumeration and by Monte Carlo sampling are identical for polymers of chain length 9–22. (a) The expected number of voids, and (b) the expected size of voids. Two-step look-ahead sequential Monte Carlo sampling is used, and the sample size is 100 000.

dead conformations during the growth. These chains need to be replaced to maintain sufficient Monte Carlo sample size. Second, the weight of some chains may become so relatively small that their contribution in the weighted average [Eq. (1)] is negligible. When the variance of the weights is large, the *effective Monte Carlo sample size* becomes small. 33,37,38 Third, for a specific function h, its value may become too small (even zero) for some sampled conformations. In all these cases, efficiency can be gained by replacing those conformations with "better" ones. This procedure is called "resampling." There are many different ways to do resampling. One approach is *rejection control*, 39 which regenerates the replacement conformations from scratch. An easier approach is to duplicate the existing and *good* conformations 33 . Specifically,

Procedure RESAMPLING

Draw *jth sample from original samples $\{(x_1^{(j)},...,x_t^{(j))}\}_{j=1}^m$ with probabilities $\propto \{\alpha^{(j)}\}_{j=1}^m$ //Each sample in the newly formed sample is assigned a new weight. //*j-th chain in new sample is a copy of k-th chain in original sample.

 $w^{(*j)} \leftarrow w^{(k)}/\alpha^{(k)}$

endfor

In the resampling step, the m new samples $\{(x_1^{(*j)},...,x_i^{(*j)})_{j=1}^m$ can be obtained either by residual sampling or by simple random sampling. In residual sampling, we first obtain the normalized probability $\widetilde{\alpha}^{(j)} = \alpha^{(j)}/\Sigma \alpha^{(j)}$. Then $[m\widetilde{\alpha}^{(j)}]$ copies of the jth sample are made deterministically for j=1,...,m. For the remaining $m-\Sigma[m\widetilde{\alpha}^{(j)}]$ samples to be made, we randomly sample from the original set with probability proportional to $m\widetilde{\alpha}^{(j)} - [m\widetilde{\alpha}^{(j)}]$.

The choice of resampling probability proportional to $\alpha^{(j)}$ is problem specific. For general function h, such as the end-to-end extension $\|x_n-x_1\|$, it is common to use $\alpha^{(j)}=w_t^{(j)}$. In this case, all the samples in the new set have equal weight. When the function is irregular, a carefully chosen set of $\alpha^{(j)}$ will increase the efficiency significantly.

The method of pruning and enriching of Grassberger⁴⁰ is a special case of the residual sampling, with $\alpha^{(j)} = 0$ for the k chains with zero weight (dead conformations), $\alpha^{(j)} = 2$ for the top k chains with largest weights, and $\alpha^{(j)} = 1$ for the rest of the chains. Residual sampling on this set of α is completely deterministic. The resulting sample consists of two copies of the top k conformations (each of them having half of their original weight) and one copy of the middle n-2k chains with their original weight. The k dead conformations are removed.

In our study of the relationship between compactness and packing density, we use a more flexible resampling method. Our focus is on the packing density among all conformations within certain range of compactness. In this case, our object target distribution is the uniform distribution among all possible SAW's with compactness measure falling within a certain interval, i.e., a truncated distribution. Although compactness changes slowly as the chain grows, to grow into a long chain it is possible that the compactness of a chain evolve and cover a wide range during growth. Hence we choose the uniform distribution of all possible SAWs of length t as our target distribution at t, and only select those with the desired compactness at the end for our estimation of the packing density. In order to have larger number of usable samples (i.e., to achieve better acceptance rate) at the end, we encourage growth of chains with desirable compactness through resampling. Specifically,

Procedure RESAMPLING (m,d,c_t)

// m: Monte Carlo sample size, d: steps of looking-back. // c_t : targeting compactness.

 $k\leftarrow$ number of dead conformations.

Divide m-k samples randomly into k groups.

for group i=1 to k

Find conformations not picked in previous d steps. //Pick the best conformation P_j , for example P_i —polymer with $\min |c-c_t|$

Replace one of k dead conformations with P_j Assign both copies of P_j half its original weight. endfor

Here d is used to maintain higher diversity for resampled conformations.

Most polymers sampled by sequential Monte Carlo with two step look-ahead but no resampling are well-extended with few voids, as shown in Figs. 6(a) and 6(b). In Fig. 6(b), the majority of the conformations have a higher packing density. They have small compactness (<0.5) and large packing density. There are not enough compact conformations. As a result, a small number of samples are accepted at the end whose compactness falls within the desired interval of higher than 0.5. By using the resampling step described above, we were able to generate more samples near the desired compactness value of 0.6 [Fig. 6(c)]. Figure 6(c) is a pure histogram of compactness in the observed samples, without regarding the weight of the samples. Here resampling is applied at each sequential Monte Carlo growth step. Figure 6(d) shows that the resampling technique is also very effective in shifting the samples to small packing density values,

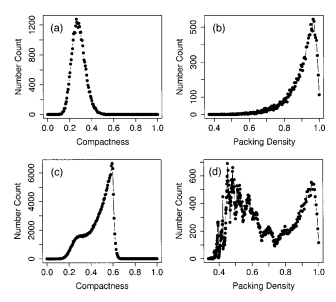


FIG. 6. The distribution of configurations of polymers obtained by the sequential Monte Carlo method can be adjusted by resampling. (a) Histogram of conformations at different compactnesses generated without resampling. The compactness of the majority of the conformations is less than 0.5. (b) Histogram of conformations at different packing density generated without resampling. The number of conformations with packing density below 0.8 is small. (c) After applying the resampling technique favoring compactness of 0.6, the majority of the conformations have compactness between 0.5 and 0.6. (d) Resampling can also be applied to generate conformations with low packing densities with voids. Sample size of 100 000 is used in all calculations

hence improve the inferences. Here resampling favoring lowest packing density is applied every 5 growth steps.

V. VOID DISTRIBUTION OF LONG CHAINS

We apply the techniques of sequential Monte Carlo with resampling to study the statistical geometry of voids in long chain polymers. Each Monte Carlo simulation starts with a sample size of 200 000, and we take the averaged values of 20 simulations. Resampling is carried out every 5 steps in the process of the chain growth. Figure 7(a) shows that the probability of void formation increases with the chain length. At chain length 105-110, about half of the conformations contain voids. At chain length 200, the standard deviation (8.5 $\times 10^{-3}$) is maximum. The expected number of voids [Fig. 7(b)] increases linearly with chain length. Similar linear scaling behavior is also observed in proteins.4 The expected wall size of void and void size also increase with chain length [Figs. 7(c) and Fig. 7(d)]. The expected packing density is found to decrease with chain length, which is consistent with the scaling relationship of void size and chain length shown in Fig. 7(d). The compactness ρ of chain polymer has been the subject of several studies. ^{12,41} The asymptotic value of ρ we found is 0.18, slightly different from that reported in Ref. 41 (ρ = 0.16), and is within the range of 0.16–0.24 reported in Ref. 12. Different resampling strategies are applied where dead conformations are removed and other conformations with the targeted property is duplicated. Resampling favors conformations with small radius-of-gyration in Figs. 7(a)-7(e), and conformations with large weight in Fig. 7(f).

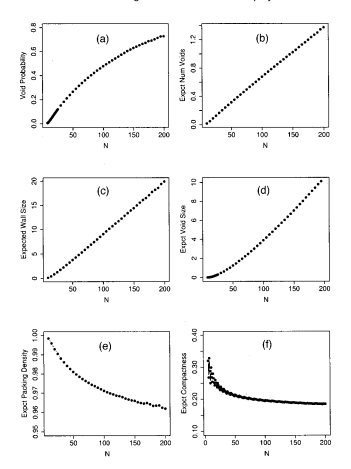


FIG. 7. Geometric properties of lattice polymers of different lengths, estimated by the sequential Monte Carlo method with the 2-step look-ahead and resampling technique. (a) The probability of void formation increases with chain length. Standard deviation ($\le 8.5 \times 10^{-3}$) increases slowly with the length. The expected number of voids (standard deviations $\le 1.6 \times 10^{-2}$) (b) and wall size (standard deviations ≤ 0.25) (c) are linearly correlated with chain length. (d) The expected void size increases with chain length (standard deviations $\le 8.3 \times 10^{-3}$). (e) The expected packing density decreases with chain length (standard deviations $\le 7.5 \times 10^{-4}$). (f) The expected compactness decreases with chain length and reaches an asymptotic value of $\rho = 0.18$ (standard deviations $\le 5.7 \times 10^{-4}$).

To explore the relationship of packing density p and compactness ρ , we use sequential Monte Carlo with 2-step look-ahead to sample 200 000 conformations, each with an appropriate weight assigned. This is repeated 20 times, and the weighted average values of the packing density at various compactness for chains with 60-100 monomers are plotted (Fig. 8). The compactness value corresponding to the minimum packing density seems to have shifted from 0.462 for the 22-mer by enumeration to above 0.5 for the 100-mer as obtained by sampling. However, the overall pattern of p and ρ found by Monte Carlo is very similar to the pattern found by enumeration for polymers with $N \le 22$ [Fig. 4(d)]. Data shown in Figs. 8 and 4(d) are not redundant. Rather, they complement each other and together provide a full picture of the relationship of ρ and p for chain length between N= 14-22 and N = 30-100. The lowest packing density p_{\min} of these lattice polymers occur at compactness $\rho = 0.525$. However, an accurate estimation of the value of asymptotic p_{\min} as $N \rightarrow \infty$ requires simulation of longer chains.

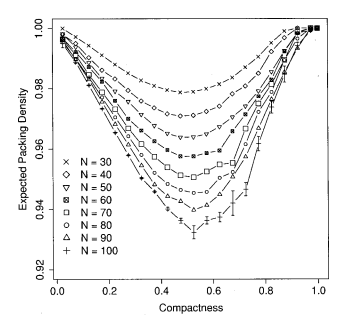


FIG. 8. The relationship of expected packing density and compactness for the long chain polymer. These data are estimated by the sequential Monte Carlo method using the 2-step look-ahead and a sample size of $20 \times 200\,000$ with resampling. Resampling is designed to favor compactness at specified values. The expected packing density calculated by averaging from the 20 runs has the largest standard deviations for the 100-mer, which are shown in the figure.

The accuracy of geometric properties of long chain polymers estimated by Monte Carlo can be assessed by the standard deviation obtained from multiple Monte Carlo runs.

VI. END EFFECTS OF VOID FORMATION

What is the effect of void formation on the size of conformational space? We consider the conformational reduction factor of voids. Following Refs. 12, 23, 31, we define the conformational reduction factor due to the constraint of a void as

$$R(n;i,j) = \frac{\omega(n;i,j)}{\omega(n)},$$

where $\omega(n;i,j)$ is the number of conformations that contains a void beginning at monomer (i) and ending at monomer (j), and $\omega(n)$ is the total number of conformations of n-polymers. R(n;i,j) reflects the restriction of conformational space due to the formation of a void with wall interval of k=|i-j|. Figure 9(a) shows a 24-mer with one void that starts at i=4 and k=19. Unlike self-contacts or self-loops, which was subject of detailed studies by Chan and Dill, 12,23,31 all conformations analyzed here must contain a void. The polymer shown in Fig. 9(b) with a large loop has no void, and such polymers do not contribute to the numerator of R.

Figure 10(a) shows the reduction factor R calculated by enumeration for voids at different starting positions with wall intervals k=7, 9, and 11. There are clearly strong endeffects: The reduction factor of voids of the same wall interval depends on where the void is located. R decreases rapidly as the void moves from the end of chain towards the middle. Void formation is much more preferred at the end of chain.

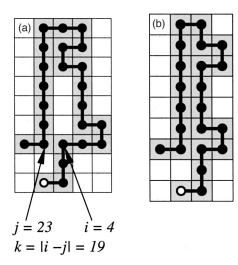


FIG. 9. The starting position of a void and its wall interval. (a) This 24-mer has a void that starts at i=4 and ends at j=23. Its wall size is k=19. (b) This polymer has a contact-loop but contains no void.

Similar end effects of void formation are also observed for the 50-mer sampled by sequential Monte Carlo [Fig. 10(b)]. Because the exact total number of conformations of the 50-mer $\omega(50)$ is unknown, we plot the value of $R \times$ constant, where constant is common to all data points at different starting positions and wall intervals. Our interest is how R changes its relative values.

The end-effect of voids has the same origin as the end-effect of self-contact, which has been extensively studied by Chan and Dill. 12,23,31 Because of the effect of excluded volume, sterically it is less hindering to form a void at the end of a polymer. When a void is formed, the conformational

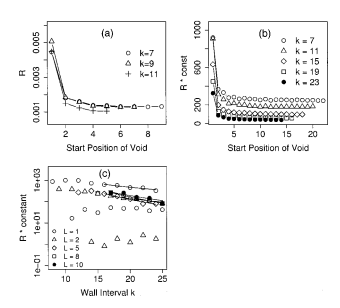


FIG. 10. The end-effect of void formation on conformational reduction. (a) Conformational reduction factor R when voids are formed in a 22-chain as examined by enumeration. R depends on the starting position and the wall interval of void. (b) Conformational reduction factor R up to a normalizing constant when voids are formed in a 50-chain as sampled by sequential Monte Carlo (standard deviations \leq 6.2). (c) Scaling of conformational reduction factor R and the wall interval k at different initiation position L for 50-chain (standard deviations \leq 6.4).

space of the k+1 monomers between monomer i and j, as well as the two tails become restricted. When the void is formed at the chain end, only one tail is subject to conformational restriction.

Void formation is different from self-contact. When monomer i and j form self-contact, it may involve the formation of a void, but it is also possible that there will be no unfilled space between i and j. When a void is formed beginning at monomer i and ending at monomer j, some monomers between i and j will have unsatisfied contact interactions. Self-contacting loops have been analyzed in previous theoretical studies^{42,43} which have been confirmed by exhaustive enumeration.^{23,31} An important study of the role of loop formation in understanding the disulfide bond and protein folding can be found in Ref. 44. Compared to nonbonded self-contact, the effect of conformational reduction is more pronounced for void formation. For the twodimensional lattice, the ratios between reduction factors of self-contact at chain end and mid-chain of a sufficiently long polymer are 1.3, 1.4, 1.5, and 1.6 for k=3, 5, 7, and 9, respectively,²³ whereas the ratios for voids at the chain end and the symmetric midpoint of the N=22 polymer are 3.4, 4.0, and 4.4. for k = 7, 9, and 11.

We now consider the power-law dependence of R(N;i,j) on the wall interval k=|i-j|. In the study by Chan and Dill, 31 the scaling exponent ν of the reduction factor R and loop length k=|i-j| for $R(N;i,j)\approx k^{-\nu}$ is found to be dependent both on k and the location of the cycle in the chain. The values of ν for self-contact range from 1.6 when k=N to 2.4 when the loop is in the middle of a long chain with two long tails. Because void formation involves at least 8 monomers, its scaling behavior is less amenable to exhaust enumeration, and application of Monte Carlo sampling is essential. Based on estimations from Monte Carlo simulation of void formation in the 50-mer, the value of ν depends on the void initiation position l_0 from the end of the polymer chain. ν ranges from 1.4 ± 0.2 for $l_0=1$ to 3.0 ± 0.2 for $l_0 = 8$ [Fig. 10(c)]. Our results show that the scaling exponent of R with k = |i - j| for void formation is similar to that of the self-contacting loop. This scaling exponent also depends on the location of the void. The exponent ν is estimated from the nonlinear least square regression fit of the data using the Gauss-Newton algorithm as implemented in the GNU package R. A cautionary note is that the estimation of standard error of ν is accurate asymptotically only for large samples, 45 and therefore maybe overly optimistic in our case, where the number of data points is very small. An accurate estimation of confidence interval of ν for small sample nonlinear regression is beyond the scope of this work.

VII. CONCLUSION

In this work, we have studied the statistical geometry of voids as topological features in two-dimensional lattice chain polymers. We define voids as unfilled space fully contained within the polymer, and have developed a simple algorithm for its detection. We have explored the relationship of various statistical geometric properties with the chain length of the polymer, including the probability of void formation π_v ,

the expected number of voids \bar{n}_v , the expected void size \bar{v} , the expected wall size of voids \bar{w} , packing density p, and the expected compactness ρ . Our results show that for chains of >105-110 monomers, at least half of the conformations contain a void. At about 150 monomers, there will be at least one void expected in a polymer. The expected wall size scale linearly with the chain length, and about 10% of the monomers participate in the formation of voids. We formalize the concept of the packing density for lattice polymers. We found that both the packing density and compactness decrease with chain length. The asymptotic value of compactness ρ is estimated to be 0.18.

We have also characterized the relationship of packing density and compactness, which are two parameters that have been used frequently for studying protein packing. Our results indicate that packing density reaches minimum values between compactness 0.4-0.6. The effects of voids are studied by analyzing the conformational reduction factor R of void formation. We found that there is a significant endeffect for void formation; the ratio of R at chain end and at midchain may be twice as large as that of the R factor for contact loops, where the formation of voids is not required.

In this study, we have applied sequential Monte Carlo sampling and resampling (SMC) techniques to study the statistical geometry of voids. SMC is essential for exploring the geometry of long chain polymers. The origin of SMC can be traced back to the work of Rosenbluth and Rosenbluth,²⁶ where the idea of placing the current monomer with probability dictated by the outcome of future steps was first formulated. Grassberger was the first to apply the technique of resampling by weight to grow the chain polymer.³³ Independently, Liu and Chen provided the general framework of Sequential Monte Carlo, which unifies for the first time the techniques of delayed sampling (look-ahead of future steps) and resampling by arbitrary statistical property. Although SMC has had many applications in science and engineering, 46 we report in this paper the first application of SMC for sampling a variety of rare events in growing polymers. Specifically, we make concrete novel applications of SMC in sampling rare conformations with the prescribed value of radius of gyration and compactness. Because the general framework of the SMC has not been described before for growing chain polymer, we also provide in this publication details of the validation of the method.

Sequential Monte Carlo allows the generation of an increased number of conformations with a variety of interesting characteristics. For example, we can replace dead conformations with existing conformations of highest weight, or conformations with highest compactness, or with the smallest radius of gyration. Figure 6 provided examples where the distribution of conformations of polymers obtained by SMC can be adjusted by resampling. In Fig. 11, we further elaborate on the general flexibility of SMC for sampling that targets on a variety of statistical properties, as well as the important fact that SMC maintains properly weighted samples, which is essential for any statistical inference.

Figure 11(a) shows the histograms of conformations of 100-mer at different packing density generated without resampling. Figure 11(c) shows the histograms of conformations

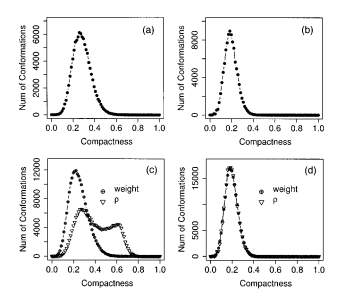


FIG. 11. Histograms of conformations of 100-mers generated by sequential Monte Carlo with and without resampling at different compactness. (a) Histogram of conformations at different compactness generated without resampling. (b) Weighted histogram of conformations generated without resampling, which is proportional to the distribution of all geometrically feasible 100-mers. (c) Histograms of conformations at different compactness when resampling is applied. (d) The weighted histograms of conformations under different resampling are in excellent agreement with each other and with that when no resampling is applied.

when resampling by weight and resampling by compactness ρ are used. To resample by weight, dead conformations are replaced with conformations of the highest weight. To resample by compactness, dead conformations are replaced with conformations of the lowest compactness. Note that the total number of surviving conformations that reach a chain length of 100 is much higher than without resampling. Resampling by compactness generates many more conformations with higher compactness. In Figs. 11(c) and 11(d), resampling is applied to every step of the chain growth process. Unlike Fig. 6(d), where resampling favors the compactness value of $\rho = 0.6$, resampling in Fig. 11(c) favors samples with the highest value of compactness ρ . Other resampling schemes are possible, e.g., resampling by radius-of-gyration, by packing density. During resampling, the number k of dead conformations at each step of growth is identified and these are replaced with conformations of interest from k randomly divided groups. These conformations must have not been resampled in the previous four steps of the growth process to maintain sample diversity. Both histograms where resampling is used deviate from that of Fig. 11(a). Resampling by weight shifts the peak of the conformations to below 0.2, and resampling by compactness turns the histogram into bimodal. The latter produces a lot more conformations with compactness $\rho > 0.4$.

SMC sampling and resampling use biased samples since conformations are generated with a probability different from that of the target distribution. The bias is dictated by the different method of resampling and different choices of the number of steps of look-ahead in sequential Monte Carlo. An essential component of a successful biased Monte Carlo sampling is the appropriate weight assignment to each

sample conformation. This is necessary because we need to estimate the expected values of the parameter such as packing density and void size under the target distribution of all geometrically feasible conformations. In Fig. 11(a), where each of the 100 000 starting conformations is generated by a two-step look-ahead without resampling, not every conformation is generated with the same probability and therefore is assigned a different weight accordingly. Figure 11(b) shows the weight-adjusted histogram, which is indicative of the probability density function at different compactness for the population of all geometrically feasible 100-mers. Figure 11(d) shows that when weights are incorporated and the area of the histogram normalized to the final number of surviving conformations, the weighted distributions of conformations using different resampling techniques have excellent agreement with the weighted distribution when no resampling is used [Fig. 11(b)]. All weighted histograms are normalized so the total area equals to the total number of surviving conformations reaching 100-mer. This example shows that by incorporating weights, the target distributions can be faithfully recovered even when the sampling is very biased.

Although sequential Monte Carlo sampling is very effective, the estimation of parameters associated with rare events remain difficult. In Fig. 10 where the conformational reduction factor R is plotted at various void initiation positions and wall interval lengths, voids starting at position 1 but with odd wall intervals ($k \in \{11,13,\ldots,25\}$) are much rarer, and it is unlikely that sequential Monte Carlo sampling with limited sample size can provide a large enough effective sample size for the accurate estimation of scaling parameters ν , where $R(N;i,j) \approx k^{-\nu}$.

In this study, we are interested in the statistics of void geometry, and our target distribution is the uniform distribution of all conformations of length n. With the introduction of an appropriate potential function and alphabet of monomers such as the HP model, 11,34,35 we can study the thermodynamics, kinetics, and sequence degeneracy of chain polymers when voids are formed in polymers. In these cases, our target distributions will be chain polymers under the Boltzmann distribution derived from the corresponding potential functions.

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APPENDIX: VOIDS DETECTION IN TWO-DIMENSIONAL LATTICE POLYMER

To detect voids in a polymer, we use a simple search method. For an $l \times l$ lattice, we start from the lower-left corner. Once we found an unoccupied site u, we use the breadth-first-search (BFS) method to identify all other unoccupied sites that are connected to site u. These sites are grouped together and marked as "visited." Collectively they represent one void in the lattice. We continue this process until all unoccupied sites are marked as visited:

```
Algorithm VOIDDETECTION (lattice, l) v=0 // Number of voids for i=1 to l for j=1 to l if site (i,j) is unoccupied and not visited v\leftarrow v+1 Mark (i,j) as visited. BREADTHFIRSTSEARCH(lattice, (i,j)) Update the size of void (i,j) endif endfor
```

Details of BFS can be found in algorithm textbooks such as Ref. 47.

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