Ergodicity and slowing down in glass-forming systems with soft potentials: no finite-temperature singularities

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Abstract

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Ergodicity and slowing down in glass-forming systems with soft potentials: No finite-temperature singularities

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The aim of this paper is to discuss some basic notions regarding generic glass-forming systems composed of particles interacting via soft potentials. Excluding explicitly hard-core interaction, we discuss the so-called glass transition in which a supercooled amorphous state is formed, accompanied by a spectacular slowing down of relaxation to equilibrium, when the temperature is changed over a relatively small interval. Using the classical example of a 50-50 binary liquid of $N$ particles with different interaction length scales, we show the following. (i) The system remains ergodic at all temperatures. (ii) The number of topologically distinct configurations can be computed, is temperature independent, and is exponential in $N$. (iii) Any two configurations in phase space can be connected using elementary moves whose number is polynomially bounded in $N$, showing that the graph of configurations has the small world property. (iv) The entropy of the system can be estimated at any temperature (or energy), and there is no Kauzmann crisis at any positive temperature. (v) The mechanism for the super-Arrhenius temperature dependence of the relaxation time is explained, connecting it to an entropic squeeze at the glass transition. (vi) There is no Vogel-Fulcher crisis at any finite temperature $T>0$.

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I. INTRODUCTION

It is not uncommon to read in papers devoted to the glass transition a statement of the type “it is well known that glass-forming systems lose ergodicity.” It is even more common [1] to use fits of the $\alpha$ relaxation time $\tau_\alpha$ in such systems to the Tamam-Vogel-Fulcher formula

$$\tau_\alpha \propto \exp\left[\frac{AT}{T - T_{VF}}\right],$$

(1)

which indicates a belief that the relaxation time actually diverges at some temperature $T_{VF}>0$. Related to these issues is the concept of the Kauzmann temperature [2], which is a finite temperature $T_K>0$ where the extrapolated entropy appears to vanish. In this paper we argue that these related questions should be addressed with care; we wish to clarify which of these issues can actually appear in glass-forming systems, and which of them are only a consequence of inadequate simulations or interpretations, or even of a confusion of questions.

In some parts this paper is a review, or an outgrowth of our own efforts to understand basic concepts of glass formation, which became more transparent in some of our earlier work [3–5], and whose findings we combine in this paper. We offer, however, in this paper a number of new results as is explained below. Glasses and their formation have occupied researchers for many decades, with ideas being first developed on theoretical bases [6], and, in more recent years, being studied extensively with the help of computer simulations [7–10]. Such simulations vary from very realistic models to toy models with simplified dynamics, but correspondingly faster simulations. We can summarize our findings in the following logical structure.

1. One has to distinguish between large (but finite) systems as compared to infinite systems. Infinite systems pose difficult conceptual problems, because in this case, density and close packing are not tightly related (any small lowering of density will allow for arbitrarily large voids, and the jamming problems disappear). We will therefore focus our discussion only on systems with a finite, but arbitrary, number of particles. In this view an Avogadro number of particles is finite, and it is important to state this fact.

2. There is an essential difference between systems or particles with a hard core and systems of particles interacting with soft repulsive potentials. In the first case, there is obviously a density where the particles cannot move any more (or at least not all of them can be moved). This poses interesting problems of jamming, or contact geometry. These issues have been studied in depth by Stillinger and his school, with a careful analysis of different types of “movability” [11]. We point out that these problems, while very interesting from the geometrical point of view (see also the book by Conway and Sloane [12]), do not really address the questions of what a generic glass is. We will thus focus attention to systems with a finite number of particles, and with soft potentials.

3. There is another issue that deserves attention: time. Many concepts of glasses are based on the notion that, if something does not happen before a given time, then it will never happen. Such reasoning is unacceptable from a theoretical point of view, since a natural time scale does not exist. We will argue that in systems with a finite number of particles interacting via soft potentials there is no singularity in the relaxation time at any positive temperature. In particular, such systems undergo a glass “transition” in the sense that their relaxation time increases without limit as temperatures are lowered, but they remain ergodic. Observing the consequences of ergodicity may necessitate waiting for an unbounded, but still finite, time. At any temperature larger than $T=0$ all of phase space is available to the system, all the configurations are dynamically connected; the configura-
tional entropy can be computed, yielding a finite result for any given finite energy (or finite temperature). We stress and reiterate that this paper deals with systems having a finite number of particles, which are interacting via soft potentials and are being observed for an unbounded time.

One can exemplify the discussion with the help of many simple models, and for concreteness we choose the classical example of a glass-forming binary mixture of particles interacting via a soft 1/r 12 repulsion with a “diameter” ratio of 1.4. More or less related models can be found in [13–18]. This two-dimensional model had been selected for simulation speed and, more importantly, for the ease of interpretation. We refer the reader to the extensive work done on this system [3,4,7,8,19]. It shows that it is a bona fide glass-forming liquid meeting all the criteria of a glass transition. In short, the system consists of an equimolar mixture of two types of particles with diameters σ2 = 1.4 and σ1 = 1, respectively, but with the same mass m. The three pairwise additive interactions are given by the purely repulsive soft-core potentials

\[ u_{ab} = \varepsilon \left( \frac{\sigma_{ab}}{r} \right)^{12}, \quad a, b = 1, 2, \]  

where σ_{ab} = σ_a and σ_{ab} = (σ_a + σ_b)/2. The cutoff radii of the interaction are set at 4.5σ_{ab}. The units of mass, length, time, and temperature are m, σ1, τ = σ1/√m/ε, and T = ε/k_B, respectively, with k_B being Boltzmann’s constant.

It has been shown that for temperatures T > 0.5 the system behaves like a liquid. For temperatures lower than 0.5 the system begins to slow down; the correlation functions stop decaying exponentially, and can be fitted within reason to a stretched exponential form; see Fig. 1. The time constant (or so-called α relaxation time) can be fitted to the Vogel-Fulcher form (1) for temperatures not too close to T_VF. The model shows the expected behavior of the specific heat at the temperature range that is considered the glass transition (i.e., where the relaxation time increases rapidly), and the super-Arrhenius dependence of the viscosity on the temperature. In short, the system appears as a good example to consider to elucidate the fundamental issues that concern us here.

In Sec. II we explain how to discretize the configurational space using Voronoi tessellations and their dual Delaunay triangulations. We generalize the discussion of [5] to include systems in three dimensions and binary mixtures (colored graphs). In three dimensions triangulations are replaced by tetrahedrizations, and some subtleties of the relation between them and Voronoi tessellations are explained. The Delaunay graphs offer a convenient basis of the demonstration of ergodicity in two or three dimensions at any finite temperature. We also review the mathematical results concerning the size of this phase space (exponential in the number of particles N) and the fact that its graph has the small-world property in the sense that any configuration can be reached from any other using a polynomial number of steps. This review is supplemented with an extension to the case of binary mixtures which concern us here. In Sec. III we discuss the configurational entropy, and explain a new result, i.e., that there is never a Kauzmann crisis in this system. Finally in Sec. IV we identify what is a typical relaxation process in this system and present the fundamental reason for the super-Arrhenius dependence of the relaxation time on the temperature. This is due to the fast decrease in the concentration of some quasispecies, leading to an entropic squeeze. The new result is that the Vogel-Fulcher fit is untenable, since the only divergence in relaxation time can appear at T = 0. In Sec. V we summarize the paper and offer concluding remarks.
II. VORONOI CONSTRUCTION AND ERGODICITY

A. Voronoi tessellation and Delaunay triangulation

We begin by discussing the possible configurations of the system in a systematic way, using the time-honored Voronoi polygon construction [7]. It associates with every configuration of the particles a subdivision of position space into polygons, one per particle. These polygons will also be called cages. The polygon associated with any particle contains all points closer to that particle than to any other particle. The edges of such a polygon are the perpendicular bisectors of the vectors joining the centers of the particles (actually the coordinates of the point particles we consider). As had been noted in [7,8], when periodic boundary conditions are used, the average number of sides of the polygons is exactly 6. This follows because the Euler characteristic on the torus is 0: \( \chi = V - E + F = 0 \), where \( V \), \( E \), and \( F \) are, respectively, the numbers of vertices (corners), edges, and faces in the given polyhedron. This constraint is important since in fact there is no limit to the number of nearest neighbors around a given particle, even in two dimensions. We can always organize as many particles as we want in a circle around a given particle, and increase the number of neighbors by increasing the radius of the circle. In reality such wasteful arrangements are costly, and in simulations at low temperatures we find typically only pentagons, hexagons, and heptagons.

Typical examples of such Voronoi tessellations for two different temperatures are shown in Fig. 2. Since there are two types of particles (small or large, or blue and red, respectively), in order to have a mapping between the particle positions and the number of sides in the polygons of the Voronoi tessellation, we distinguish between polygons having a small and a large particle in their center. Thus, a coloring scheme of cells will take into account not only the number of sides, but also the type of the particle (big or small) in each cell. The tessellation obtained without distinguishing the type of particles will be referred to as “colorless” and so, below, will be the resulting triangulation. When distinguishing the two types of particles, the tessellation and its triangulation will be referred to as colored.

Note that in a generic configuration the Voronoi decomposition has the property that exactly three lines meet at a point. Accordingly, the dual graph of such a decomposition is a triangulation. Such triangulations are called Delaunay triangulations. They are obtained by simply connecting the centers of those particles that are separated by one edge of the polygon. In other words, the vertices of the triangles coincide with the particles at the centers of the Voronoi cells. Let us remark that this construction can also be done in three dimensions (or more), where now instead of edges of polygons we have faces of polyhedra and four faces will meet at a point. In this case, the dual graphs are now made of tetrahedra. In three dimensions we will call the graphs obtained in this way a Delaunay tetrahedrization. It is noteworthy that there is an important difference between two and three dimensions. In two dimensions (2D), the points of any triangulation can be arranged in such a way in position space that the triangulation is indeed a Delaunay triangulation, as described above, that is, the dual of a Voronoi decomposition.

However, in 3D there exist tetrahedrizations which are not duals of Voronoi tessellations, or, in other words, cannot be obtained starting with particles spread in three dimensions, mapped onto a Voronoi tessellation and then tetrahedrized ([21]).

Having outlined the description of configurations in this model (and in fact all models of this general type), we consider now the question of dynamic accessibility, viz., ergodicity. The model under consideration is Hamiltonian, but in simulations it is coupled to a heat bath of temperature \( T \). There are several methods for achieving such a coupling, leading to the study of this model in either \( N,V,T \) or \( N,p,T \) ensembles. In either ensemble the mean energy per particle is fixed, but there can be arbitrarily large fluctuations in the energy. It is precisely these fluctuations which are the source of ergodicity. The energy is partially kinetic and partially potential. By ergodicity, one means that the natural motion of
the system can reach every point in phase space. For systems having one type of (indistinguishable) particles, it is not important to look at particles exchanging positions; indeed, every point in phase space can be obtained by moving particles within their cages and translating cages as is necessary. In contrast, when there is more than one type of particle (here we have two types), ergodicity means that particles must also be able to change their relative positions in all imaginable ways. It is therefore useful in both cases to distinguish between thermal motion of individual particles rattling in their own cages, without really changing their relative positions, and large-scale movements and exchanges of particles that change the macroscopic configuration. Only the second class of movements is important in terms of the configurational entropy of the system, which arises from counting different configurations, for which the thermal motion does not matter. For example, in systems with hard-core interactions, there are various levels at which motion can be restricted, as we have mentioned before. But here we consider soft particles.

Since we are starting from a problem with particles that exist in a concrete space, we can focus on the restricted class of Delaunay graphs (which actually coincides in 2D with all triangulations, as we have explained earlier). In other words, in two (three) dimensions the natural phase space for our system is the set of all Delaunay triangulations (tetrahedrizations).

In two dimensions the elementary change in the Voronoi tessellation is obtained by a so-called $T1$ process as seen in Fig. 3. (This operation has various names in the literature: Gross-Varstedt move, Pachner move, flip [22–26].) Note that the operation of a $T1$ process in the Voronoi tessellation translates to a flip in the Delaunay triangulation: see Fig. 3.

To simplify things further, we consider instead of the triangulations of the torus the triangulations $T$ of the sphere. The simplification is that the genus is 0, and that more is known about the combinatorics of triangulations of the sphere than of the torus. Given $N$, corresponding to the number of particles in the original model, we let $T$ denote a triangulation of the sphere with $N$ nodes and we let $T_{N,0}$ denote the set of all such triangulations. By this, one means the set of all combinatorially distinct rooted simplicial three-polytopes. In particular, a triangulation should not have any "double edges." We further refine the definition, by distinguishing two types of nodes in the triangulation: We first number the nodes from 1 to $N$ and then define two types of nodes. Those with even index are the “small particles” and those with odd index the “large” ones. This means that the triangulation has the same number of large and small nodes (up to a difference of 1). We will also call the two types of nodes two colors.

We shall call the odd nodes blue and the even ones red and will refer to the triangulations as colored triangulations. Once the colors are assigned, the numbers are again forgotten. The set of all colored triangulations with $N$ nodes will be called $T_N$. This is our phase space and the dynamics is mapping points in this phase space to other points. Having understood that, we can now ask the major question of this section, the question of ergodicity: Is every point in phase space accessible through a sequence of motions that are flips? The answer is not obvious from the outset. For example in two dimensions one can consider a triangulation that includes locally a form as in Fig. 4. Obviously, a form like this is stuck, since one cannot do any flip without doubling one of the links. (One can see this easily by noticing that the configuration in Fig. 4 is a valid triangulation of the sphere, and it is the unique triangulation with four points.) The question of ergodicity is even less trivial in three dimensions, where there is a whole class of unflippable tetrahedrizations. However, they are not duals of Voronoi. But the relevant question (in any dimension) is whether the triangulations (tetrahedrizations, etc.) which are duals of Voronoi tessellations are indeed all connected in one large component. The answer is nevertheless “yes” in both two and three dimensions as we explain next.

**B. Demonstration of ergodicity**

One can attack this problem in two ways, the first being more physical and the second more mathematical. The physical argument is trivial. Suppose that the system is efficiently coupled to a heat bath, such that the energy is unbounded. Since the particles are soft, and the energy is not bounded, they can be moved around each other in any way one can imagine. We stress again that, while this needs perhaps a lot of energy, the large fluctuations of the energy will guarantee that this will eventually happen, rarely, but surely. And hence ergodicity is obvious. The only care one must observe is that moves through degenerate situations (four lines meeting at a Voronoi vertex) must be avoided. Unfortunately, this simple
argue that the question is more intriguing when formulated in terms of triangulations alone, since we have seen in Fig. 4 that a local configuration in which three links emanate from a node cannot be flipped. Could it be that the triangulation is so imbricated that in fact none of the edges can be flipped? Indeed, for four particles this is exactly what happens, but then the phase space consists of exactly one triangulation and thus there is no need to move any link. In 1936, Wagner [27] showed that a finite number of flips will transform any triangulation of the sphere into a “Christmas tree,” which is the configuration shown in Fig. 5. See also [28] for a discussion of several related issues. Since one can undo the flips, this implies immediately that any two triangulations can be connected by a sequence of flips going through the Christmas tree. There is abundant literature on this question [21,24] which also plays a certain role in the classification of threemanifolds.

C. The size of phase space

The possible states of our system of triangulations with \( N \) nodes is the set \( T_N \) of all possible colored triangulations. The set \( T_N \) has, as we will see, a number of elements which grows like \( C^N \) for some constant \( C > 1 \). It is thus a discrete space with a finite number of states. To describe the dynamics of flipping in a geometric way, one should view this set as the dynamical graph \( \mathcal{G} \), whose nodes are now the elements of the set \( T_N \) [not to be confused with the nodes (particles) of any triangulation \( T \)] and two of its nodes are linked if one can be reached from the other by a flip. (This makes an undirected graph, since one can flip back and forth.) The reader should note that there are two graphs in this discussion: Each triangulation is a graph with \( N \) nodes, and \( 3N - 6 \) links (by Euler’s theorem), while the graph \( \mathcal{G} \) has about \( C^N \) nodes, and about \( 3N - 6 \) links per node. This last statement follows because in every state of \( T_N \) one can choose which of the \( 3N - 6 \) links of the triangulation \( T \) one wants to flip. However, there will, in general, be somewhat fewer links that are candidates for flipping, because whenever there is a node of degree 3 in the triangulation \( T \) its links cannot be flipped (a tetrahedron is unflippable).

Finally, given any two elements in \( T_N \), that is, any two triangulations with \( N \) nodes, we will show below that \( O(N^3) \) flips are sufficient to walk on the graph \( \mathcal{G} \) from one to the other. Thus, the diameter of the graph \( \mathcal{G} \) is at most \( O(N^3) \) while it has \( O(1)^N \) vertices. This means that \( \mathcal{G} \) has the smallworld property [29]. It has also a small clustering coefficient, since there are very few triangles in the graph \( \mathcal{G} \) (it is difficult to get from a triangulation back to the same triangulation with three flips).

In the remainder of this section, we spell out these statements. They are well known for uncolored graphs, so the only task is to prove them for the colored graphs; this is the added result of this section. The method of proof can be taken from [5].

We first state two known results for the set \( T_{N,0} \) of uncolored triangulations.

**Lemma 1** [23,24,30]. The number of elements in \( T_{N,0} \) is asymptotically

\[
\left( \frac{256}{27} \right)^{N-3} \frac{3}{16\sqrt{6\pi N}}.
\]

The distance between any two uncolored triangulations is at most \( 6N - 30 \) flips.

For the case of colored graphs, with \( N_{\text{red}} = N_{\text{blue}} + c \) and \( c \in \{0,1\} \), that is, about equal numbers of red and blue nodes, one has the following lemma.

**Lemma 2**. The number of elements in \( T_N \) is asymptotically bounded above by

\[
2^N \left( \frac{256}{27} \right)^{N-3} \frac{3}{16\sqrt{6\pi N^3}}
\]

and below by the expression (3). The distance between any two colored triangulations in \( T_N \) is bounded by

\[
C_1 N^2 + C_2
\]

flips with some universal constants \( C_1 \) and \( C_2 \).

We note that the phase space as defined here is (obviously) independent of the temperature. We can thus conclude that the present classical model of glass formation does not suffer from any issue of loss of ergodicity. Accordingly, it should have a valid statistical mechanics at any temperature \( T > 0 \). Next we show that indeed its configurational entropy \( S \) never suffers any finite-temperature crisis, and there is no Kauzmann temperature where \( S \to 0 \).

III. CONFIGURATIONAL ENTROPY

A. Statistical mechanics

Needless to say, the configurations discussed in the previous section have different energies and therefore the probability to see any particular one can be strongly dependent on the temperature. To discuss the temperature dependence of the configurational entropy of this system, we need to review the statistical mechanics that was introduced for this system in [4]. The basis of the analysis is again the Voronoi tessellation.
of quasispecies increases to 42 (six, seven, and eight for each pentagon, hexagon, and heptagon, respectively). While double and a bit more precise, this more involved statistical mechanics does not shed more light on the issues of principle that interest us in the present paper, and therefore we do not discuss such improvements any further.

Coming back to the simpler variant, we note that in the free energy the value of \( U \) is \( \sum_{i=1}^{6} N_i e_i \). The \( pV \) term is simply \( p \sum_{i=1}^{6} N_i \Omega_i \). Lastly, we need to estimate the entropy term. In principle, this should be computed from the number of possible complete tilings of the area by \( N_i \) cells of each type with its given area and shape, subject to the Euler constraint \( \sum_{i=1}^{6} N_i = 6N \), where \( E_i \) is the number of edges of the \( i \)th polygon. This is a formidable problem. A useful estimate can be obtained by considering the area only, and filling space starting with the largest objects, then the next largest, and so on, until the smallest are fitted in. We do this by dividing the (remaining) volumes into boxes, and studying the combinatorial filling of these boxes. Denoting the possible number of boxes to fit the largest cells by \( N_1 \), then the number of boxes available for the second largest cell by \( N_2 = (V - N_1 \Omega_1) / \Omega_2 \), etc., the number of possible configurations \( W \) is

\[
W = \prod_{i=1}^{6} \frac{N_i!}{N_i^! (N_i - N_i)!}.
\]

Using the abbreviation \( x_i = N_i / N \), we compute directly \( x_i = c_i \Omega_i / \sum_{i=1}^{6} c_i \Omega_i \) where \( c_i \) is the number concentration of each defect. We can now compute \( S = k_B \ln W \) and write \( G \) together with a Lagrange multiplier for the Euler constraint,

\[
G = \sum_{i=1}^{6} N_i e_i + p \sum_{i=1}^{6} N_i \Omega_i + \lambda \sum_{i=1}^{6} N_i E_i + T \sum_{k=1}^{6} N_k \ln x_k + x_k (1 - x_k) \ln(1 - x_k)).
\]

The chemical potential \( \mu_i = \partial G / \partial N_i \) is then, for \( i = 1, \ldots, 6 \),

\[
\mu_i = e_i + p \Omega_i + T \ln x_i + \frac{\Omega_i}{\sum_{k=1}^{6} \Omega_k} \ln(1 - x_k)) + \lambda E_i.
\]

We now recognize that Voronoi cells of different values \( E_i \) but with the same size particle (small or large) are in equilibrium, each one being able to change to the other, but small particles cannot change to large particles, and therefore in equilibrium there exist only two independent values of \( \mu_i \), one for the small particles \( \mu_S \) and one for the large particles \( \mu_L \), and we have nine unknowns—six values of \( c_i \), two values of \( \mu \), and one Lagrange multiplier \( \lambda \). This is precisely balanced by the six equations (8), the Euler constraint, and the two constraints \( \sum_{i=1}^{6} \mu_i = \sum_{i=1}^{6} c_i = 1 / 2 \). These equations could be solved numerically using the precise values of \( \Omega_i(T) \) and \( \epsilon(T) \) as measured in the simulation. The approximate calculation of the entropy, however, does not warrant such a detailed calculation. In reality, on calculating the average areas of the cell types in the numerical simulations, we discover that to an excellent approximation these fall in two classes, smaller cells of area \( \Omega_S \) when small particles are in

FIG. 6. (Color online) Average energies of the Voronoi cells as a function of the temperature as measured in the simulations.

To get some systematics, we first define a “typical” energy for each type of topological cell, with an average taken over all cells with a given number of sides and a given particle type (big or small) in its center. The average is then \( E_i = (\sum_{k=1}^{N_i} E_k \left( \frac{2}{N_k} \right)^{1/2} ) \) where \( E_k \) is the number of edges associated with that cell type \( i \), \( r_{ik} \) is the distance to the particle in the adjacent Voronoi cell, and the average is over all particles of the same type \( i \). In Fig. 6 we present the values of these energies measured numerically as a function of the temperature, following a protocol of slow cooling. In the range of temperatures explored by simulations there are ten different cell types (large or small particle in squares, pentagons, hexagons, heptagons, and octagons), but octagons and squares are not shown since they already disappear at relatively high temperature, much above the glass transition.

We learn from these data graphs that the different cell types have clearly split energies throughout the interesting temperature range, and that these energies are only weakly dependent on the temperature. Within the temperature range of interest we can focus on the six types of cell; denote by \( \{N_i\}_i \) the number of cells of each type, with number of edges \( E_i \), ordering them by the mean energy \( e_i \) with \( i = 1 \) being the highest (large particle in a pentagon) and \( i = 6 \) being the lowest (small particle in a heptagon). Additional important properties of the cell types are their areas \( \Omega_i \) and their shapes; the first affects the enthalpy term and both affect the configurational entropy when we count the number of possible tilings of the plane.

With this in mind we can construct the statistical mechanics of this system by considering the free energy \( G = U + pV - TS \). We should stress at this point that one could aspire for more accurate statistical mechanics, considering, for example, not only the type of particle inside the Voronoi \( n \)-gon, but also who are the neighbors (small or large particles) (which resembles somewhat the plaquette expansions in statistical mechanics). Such a choice would have allowed a better treatment of the tendency of hexagons with large (or small) particles, for example, to crowd together to minimize the \( pV \) term in the free energy. The price is that the number
them, and larger cells of area $\Omega_L$ where large particles are enclosed. These areas again are only weakly dependent on the temperature. Then the whole system of equations simplifies to two analytically tractable sets of equations,

\[ \tilde{\mu}_L = \epsilon_i + T \ln c_i + \lambda E_i \quad \{i = 1, 2, 3\}, \]
\[ \tilde{\mu}_S = \epsilon_i + T \ln c_i + \lambda E_i \quad \{i = 4, 5, 6\}, \]

(9)

together with the above mentioned three constraints. In $\tilde{\mu}$ we have absorbed terms that added to $\mu$ in this special case.

In Fig. 7 we show the solutions of these equations when we use values of $\epsilon_i$ taken from Fig. 6 at $T=0$. We learn from these results that the statistical mechanics predicts that the first quasispecies to disappear are the large particles in pentagons and the small particles in heptagons. While the first is the highest in energy, the disappearance in tandem of the second is a result of the Euler constraint, and could not be guessed a priori. In previous work the first disappearing quasispecies were called liquidlike, since they are common in the liquid state and their concentration is exponentially small in the glass state. The region of temperature where their concentration falls off rapidly was identified with the region of slowing down. In fact, in [4] a quantitative relation between the concentration of these liquidlike quasispecies and the relaxation time was derived, explaining the slowing down as a result of an entropic squeeze. We will return to this issue in Sec. IV.

The statistical mechanics predicts a second transition (see Fig. 7). Below some temperature the concentration of hexagonal cells is predicted to be exponentially small, and the system retains only pentagons with small particles and heptagons with large particles. Indeed, in [4] it was found that a phase made of only pentagons with small particles and heptagons with large particles exists and is stable at low temperatures; see Fig. 8. We call such Voronoi quasispecies “glasslike” although clearly the phase shown in Fig. 8 is not a glass. The glassy phase is there as long as there is a competition between this type of phase and the phase-separated systems of small and large hexagons. Indeed, upon warming up such a phase, at a temperature roughly around $T_1$, a sizable number of hexagons appears to form the generic glassy state. Upon further warming, crossing a temperature roughly around $T_2$, a sizable number of liquidlike defects brings the system to a liquid state. The actual values of $T_1$ and $T_2$ can be understood from this model. Denote by $c_L$, $c_H$, and $c_G$ the concentrations of the liquidlike, hexagons, and glasslike quasispecies, and by $\epsilon_L=\epsilon_1+\epsilon_6=12.48$ the energy associated with the liquidlike quasispecies, $\epsilon_H=\epsilon_3+\epsilon_4=11.94$ the energy of the hexagons, and $\epsilon_G=\epsilon_5+\epsilon_7=11.76$ the energy of the glasslike defects. The theory predicts that the ratios $c_L/c_H$ and $c_H/c_G$ are of the order of $\exp[-(\epsilon_L-\epsilon_H)/T]$ and $\exp[-(\epsilon_H-\epsilon_G)/T]$, respectively. As an estimate of $T_2$ and $T_1$, take these ratios to be, say, of the order of $1\% \sim \exp(-5)$ and observe that such ratios are obtained for $T=T_2\approx0.11$ and $T=T_1\approx0.04$. It is important to notice that $\epsilon_H-\epsilon_G$ could be positive rather than negative, and then the system would crystallize on a hexagonal lattice. Such a lattice can exist in this system only when the particles phase-separate into two pure hexagonal lattices of small and large particles, respectively, with an interface in between. Such a phase may even be the ground state, but seems to be inaccessible in dynamical experiments starting from random organizations of small and large particles.

B. Nonexistence of Kauzmann temperature

Irrespective of whether the realized state at $T=0$ is the state shown in Fig. 8, or whether the ground state is made of hexagons, it is obvious now that any finite temperature will allow the appearance of other quasispecies (hexagons inside the phase in Fig. 8 or pairs of pentagons and heptagons in the hexagonal phase if the latter were the ground state). Focusing on the first situation, we understand that any pair of hexagons costs a given amount of energy $\epsilon_H$ which is a given
\(\Delta\) above the ground state \(\epsilon_G\). The calculation given by Eq. (7) is legitimate at any temperature \(T\), and the configuration entropy is approximately correct as stated there. One can improve the calculation of the entropy compared to the approximation employed above, but there is nothing extraordinary that is expected at any temperature. We thus state that the configurational entropy is expected to be an analytic function of \(T\) at any value of the temperature, and it can vanish only at \(T=0\). There is no Kauzmann temperature here or in any similar generic model.

C. The notion of fictive temperature

Notwithstanding all of the above, the system under study can slow down so much that upon reducing the temperature one has to wait for a very long time before equilibrium is reached. When the relaxation times are already very large, say at an initial temperature \(T_i\), any rapid decrease in the temperature of the heat bath to a final temperature \(T_f\) may result in a very lethargic response of the system, which may keep the concentrations of various quasispecies at values that are consistent with \(T_f\) rather than \(T_i\). It is then perfectly legitimate to introduce the notion of a fictive temperature \(T_{ic}\) \(\approx T_f\) as long as one is satisfied with short observation times. For longer and longer times the system will exhibit the process of aging, and in particular \(T_{ic}\) will converge to \(T_f\), reaching there with certainty if given enough time.

IV. SLOWING DOWN AND ENTROPIC SQUEEZE

The aim of this section is to explain the most important aspect of glass formation, i.e., the extreme slowing down in relaxation to equilibrium when the temperature is lowered. The riddle is as follows: the natural time scale is determined by the molecular jitter due to thermal motion. This time is typically of the order or \(10^{-12}\) s at room temperature. Glassy dynamics exhibits a relaxation time \(\tau_r\) of the order of seconds, or hours, sometimes years. How is it that such a huge gap in time scale is obtained without geometrical obstruction (as is the main theme of this paper)? We will explain that the issue is entropic squeeze, or the failure of entropy or "the number of available paths" to overcome the necessary energy climb required for relaxation.

In [3,32] it was argued that the relaxation time can be predicted if one knew the typical scale \(\xi\) that separates liquid-like quasispecies. In other words, having the concentration \(C_\ell\) of large particles in pentagons and small ones in heptagons one introduces a typical scale \(\xi\) by

\[
\xi \sim \frac{1}{\sqrt{C_\ell}},
\]

since the system has two dimensions. To connect between the relaxation time \(\tau_r\) and the length scale \(\xi\) it was asserted that for the viscous fluid there exists a free energy of activation \(\Delta G^*(T)\) associated with the relaxation event,

\[
\tau_r = \tau_0 \exp[\Delta G^*(T)/T],
\]

where \(\tau_0\) is a microscopic time scale of the order of a single-particle vibration time. The free energy of activation is estimated as the number of Voronoi cells \(N^*(T)\) involved in the relaxation event times the (temperature-independent) chemical potential per cell \(\Delta\mu\), \(\Delta G^*(T) = N^*(T)\Delta\mu\). The number \(N^*\) depends on whether the relaxation event is a one- [32] or two-dimensional event [3]. In the first case \(N^*(T) = \xi(T)/\bar{\Omega}\) while in the second \(N^*(T) = \pi \xi^2(T)/4\bar{\Omega}\), where \(\bar{\Omega}\) is the mean area of a Voronoi cell. We end up with the predictions

\[
\tau_r = \tau_0 \exp[\bar{\xi}(T)\Delta\mu/\bar{\Omega}T], \quad 1D \text{ event},
\]

\[
\tau_r = \tau_0 \exp[\pi\bar{\xi}^2(T)\Delta\mu/4\bar{\Omega}T], \quad 2D \text{ event}.
\]

These predictions were shown to fit the simulation data very well [3,32]. Here we want to derive the analytic form of these formulas from the identification of the dynamics on the graph.

The present language of geometrization as described above allows us to make these estimates more quantitative. We note that for every temperature \(T\) we expect an equilibrium concentration of quasispecies, i.e., colored pentagons, hexagons, and heptagons. We can therefore ask what happens if we perturb these concentrations a bit; say we increase the number of some quasispecies, how long will it take to decay back to equilibrium? Since we know that we can move by \(O(N^2)\) elementary steps of flips from any state to any state, all that remains is to determine the temperature dependence of this number for the relaxation process considered here. To achieve this calculation we note that the glass transition and the associated slowing down take place in the range of temperatures around \(T_1\) where the liquid-like quasispecies deplete rapidly. It is advantageous theoretically to focus on the range of temperatures around \(T_1\) where the hexagons deplete quickly (see Fig. 9), since then we have a smaller number of quasispecies to take into account, while the fundamental phenomenon of entropic squeeze is not very different. So think about a situation when the majority of
quasispecies are spatially disordered pairs of glasslike pentagons and heptagons, and set up the energy scale such that these pairs have energy zero. Next consider a temperature \( T \) where the equilibrium concentration \( c \) of hexagons is small, \( c \ll 1 \). Set up the energy units such that each pair of hexagons (one with a small and one with a large particle) has an energy \( \Delta = 1 \). Accordingly, the energy of this configuration is

\[
E = cN. \tag{14}
\]

On the other hand, the entropy of this configuration can be estimated as the logarithm of the number of ways the hexagons can be distributed, which is

\[
S \approx - N c \ln c = N c \ln(1/c). \tag{15}
\]

As a result we can compute

\[
\frac{\partial S}{\partial E} = \frac{1}{N c} \frac{\partial S}{\partial c} = - \ln c - 1 \approx - \ln c. \tag{16}
\]

Remembering the thermodynamic identity \( \partial E / \partial S = T \), we then conclude that the concentration of hexagons satisfies the relation

\[
c = \exp(-1/T). \tag{17}
\]

Accordingly we conclude that in two dimensions the average distance \( \xi \) between hexagons, which is \( \xi = 1/\sqrt{c} \), satisfies

\[
\xi \approx \exp(1/2T). \tag{18}
\]

Whenever a pair of hexagons is created, with high probability, this will be undone sometime in the future (by the inverse operation). Once they are separated in space by the distance \( \xi \), we need a number of flips which is at least of the order of \( \xi \), but maybe many more, in order to annihilate a pair. This is the fundamental process of relaxation at temperature \( T \) which we now proceed to estimate. In other words, we estimate how many flips are typically needed in order to get rid of one pair of hexagons, and we will show that the answer is super-Arrhenius.

We first note that, because the concentration \( c \) is small, any random flip will create a pair of hexagons and increase the energy by unity. Only flips that eliminate a pair of hexagons (to create a heptagon and pentagon) reduce the energy by unity. We denote the probability of such a rare event by \( \gamma \). It will be crucial that \( \gamma \) depends on the temperature as we will explain below. Out of all the other flips, assume that a fraction \( \alpha \) of flips does not change the energy (it has to involve a hexagon and one of a pentagon-heptagon pair). What is then the best way to move one hexagon a distance \( \xi \) until it can annihilate its counterpart? Assume that the path takes \( m \) flips where the energy increases by one unit (a typical flip), energy neutral over \( k \) flips, and then goes down in energy in \( m \) steps, with the constraint

\[
2m + k \geq \xi. \tag{19}
\]

Of course, these events can take place in any order. Clearly, we have a competition between the number of ways to arrange such a path and the energy barrier that needs to be surmounted, i.e., we need to sum up over all \( m \) and \( k \) the expression

\[
P(m, k) = \frac{(2m)!}{m! m! k!} (1 - \alpha - \gamma)^m \alpha^k \gamma^k e^{-mT}. \tag{20}
\]

Note that all the factors are smaller than 1 and therefore the sum is well approximated by the largest term, which occurs for \( m = \xi/2 \) and \( k = 0 \). We thus get

\[
P = \sum_{2m+k\geq \xi} P(m, k) \approx [(1 - \alpha - \gamma) \gamma e^{-1/T}]^{\xi/2}. \tag{21}
\]

The time we need to wait to see the event is proportional to the inverse of this probability, or

\[
\tau_e \approx \exp(\xi/T), \tag{22}
\]

which is precisely Eq. (12). This is the one appropriate for one-dimensional relaxation events as assumed here, and the generalization to other dimensions is obtained by modifying the relation between the density and \( \xi \). The non-Arrhenius nature of the relaxation time is due to the strong temperature dependence of \( \xi(T) \) (cf. \[32\]), which in turn is due to the fast disappearance of a class of quasispecies. This reduction of the number of quasispecies is responsible for the entropy squeeze.

We note in passing that the temperature dependence of \( \xi(T) \) determines the “fragility” of the glass under study. Fragility was introduced by Angell (see \[1\]) to characterize the degree of deviation from Arrhenius form of the relaxation time. In the present case it is obvious that the fragility is determined by the enthalpy difference between the excitations and the background, here hexagons vs pentagons and heptagons. We believe that this is quite general, but the elaboration of this idea is beyond the scope of this paper.

V. SUMMARY AND CONCLUSIONS

Glass-forming systems with hard cores can get jammed because of geometric constraints. In this paper we argued, on the basis of a generic example, that when the potentials are soft, the spectacular slowing down associated with the glass transition is in a sense more interesting, since it does not occur due to geometric jamming. Such systems never lose their ergodicity, and any configuration can be reached from any other in a polynomial (in \( N \)) number of steps, even though the number of configurations is exponential in the number of particles. We demonstrated explicitly that the configurational entropy in such systems is finite at any temperature, and thus neither the Kauzmann temperature nor the Vogel-Fulcher formula can be taken seriously. Both are the
results of an extrapolation that is not fundamental. Finally, we addressed the questions of the reason for slowing down and the mechanism of its super-Arrhenius temperature dependence. We showed by an explicit calculation for a generic relaxation step that near the glass transition, when the concentration of some quasispecies becomes very small, the entropic squeeze results in the inability of the entropic counting of paths to balance the energetic barriers, leading therefore to relaxation times that depend on the temperature much more sharply than expected from the Arrhenius form.

In summary, we focused on the topological properties of generic glass-forming systems, to clarify some fundamental issues that are not always clear in the literature. Needless to say, much of the interest in glass-forming systems, including their mechanical properties, calls for understanding further issues, including metric issues that are outside the scope of this paper. For some recent thoughts on these subjects we refer the reader to [19,31].

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[1] For a balanced review in which many of the common concepts and terminology are well spelled out, see M. D. Ediger, C. A. Angell, and S. R. Nagel, J. Phys. Chem. 100, 13200 (1996).
[2] W. Kauzmann, Chem. Rev. 43, 219 (1948).
[3] E. Aharonov, E. Bouchbinder, H. G. E. Hentschel, V. Ilyin, N. Makedonska, I. Procaccia, and N. Schupper, Europhys. Lett. 77, 56002 (2007).
[4] V. Ilyin, N. Makedonska, I. Procaccia, and N. Schupper, Phys. Rev. E 76, 052401 (2007).
[5] J.-P. Eckmann, J. Stat. Phys. 129, 289 (2007).
[6] See E. Donth, The Glass Transition, (Springer, Berlin, 2001), and references therein.
[7] D. Deng, A. S. Argon, and S. Yip, Philos. Trans. R. Soc. London, Ser. A 329, 549 (1989); 329, 575 (1989); 329, 595 (1989); 329, 613 (1989).
[8] D. N. Perera and P. Harrowell, Phys. Rev. E 59, 5721 (1999), and references therein.
[9] M. J. Demkowicz and A. S. Argon, Phys. Rev. Lett. 93, 025505 (2004).
[10] H. Shintani and H. Tanaka, Nat. Phys. 2, 200 (2006).
[11] For a particularly lucid and complete discussion of these issues, see A. Donev, Ph.D. dissertation, Princeton University, 2006, and references therein.
[12] J. H. Conway and N. J. A. Sloane, Sphere Packings, Lattices and Groups, Grundlehren der Mathematischen Wissenschaften (Springer, New York, 1999).
[13] G. H. Fredrickson and H. C. Andersen, Phys. Rev. Lett. 53, 1244 (1984).
[14] L. Davison and D. Sherrington, J. Phys. A 33, 8615 (2000).
[15] D. Sherrington, L. Davison, A. Buhot, and J. P. Garrahan, J. Phys.: Condens. Matter 14, 1673 (2002).
[16] G. Schliecker, Adv. Phys. 51, 1319 (2002).
[17] S. Léonard, P. Mayer, P. Sollich, L. Berthier, and J. P. Garrahan, J. Stat. Mech.: Theory Exp. (2007) P07017.
[18] G. Ben Arous and J. Černý, in Mathematical Statistical Physics, edited by A. Bovier, F. Dunlop, A. van Enter, and J. Dalibard, Proceedings of the Les Houches Summer School of Theoretical Physics, LXXIII (Elsevier, Amsterdam, 2006).
[19] V. Ilyin, N. Makedonska, I. Procaccia, and N. Schupper, Phys. Rev. E 76, 052401 (2007).
[20] V. Ilyin, I. Procaccia and N. Schupper, Weizmann Institute report, 2008 (unpublished).
[21] F. Santos, in Proceedings of the International Congress of Mathematicians, 2006, edited by M. Sanz-Sole, J. Soria, J. L. Varona, and J. Verdera, (European Mathematical Society, Zurich, 2006), Vol III, pp. 931–962.
[22] V. A. Malyshev, Usp. Mat. Nauk 54, 3 (1999).
[23] R. Mori, A. Nakamoto, and K. Ota, Graphs Combin. 19, 413 (2003).
[24] S. Negami, in Proceedings of the 10th Workshop on Topological Graph Theory, Yokohama, 1998, Yokohama Math Journal, Special Issue, p. 47.
[25] U. Pachner, Arch. Math., 30, 89 (1978).
[26] U. Pachner, Math. Z. 176, 565 (1981).
[27] K. Wagner, Jber. Deutsch. Math-Verein. 46, 26 (1936).
[28] P. Collet and J.-P. Eckmann, J. Stat. Phys. 121, 1073 (2005).
[29] D. J. Watts and S. H. Strogatz, Nature (London) 393, 440 (1998).
[30] W. T. Tutte, Can. J. Math. 14, 21 (1962).
[31] V. Ilyin, I. Procaccia, I. Regev, and N. Schupper, Phys. Rev. E 77, 061509 (2008).
[32] V. Ilyin, E. Lerner, T.-S. Lo, and I. Procaccia, Phys. Rev. Lett. 99, 135702 (2007).