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Spatially Heterogeneous Dynamics and the Adam—Gibbs Relation in the Dzugutov Liquid

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We perform molecular dynamics simulations of a one-component glass-forming liquid and use the inherent structure formalism to test the predictions of the Adam—Gibbs (AG) theory and to explore the possible connection between these predictions and spatially heterogeneous dynamics. We calculate the temperature dependence of the average potential energy of the equilibrium liquid and show that it obeys the Rosenfeld—Tarazona law for low temperature, while the average inherent structure energy is found to be inversely proportional to temperature at low temperatures, consistent with a Gaussian distribution of potential energy minima. We investigate the shape of the basins around the local minima in configuration space via the average basin coordinates. This surface consists of a large number of local minima of varying depths, surrounded by potential energy barriers. At sufficiently low temperatures, the system resides near these local minima and the motion of the system can be described by vibrations around the minima, with infrequent transitions from one minimum to another.

The qualitative description of the potential energy landscape was formalized by Stillinger and Weber using the concept of the inherent structures (IS), defined as the configuration of the system at a local minimum of the potential energy hypersurface. Stillinger defined a basin of attraction as the set of points that map to the same local minimum upon minimization of the potential energy. On the basis of this operational definition, Stillinger and Weber proposed a mathematical formalism—referred to as the inherent structure formalism—to quantify the thermodynamic properties of the PEL, where the degeneracy of energy minima can be related to the entropy. Since then, this approach has become an essential tool for understanding the dynamic and thermodynamic properties of glass-forming liquids, proteins, and disordered spin systems.

The IS formalism facilitates the direct calculation of the configurational entropy by computer simulation through the enumeration of basins in the PEL that are accessible to the dynamic and thermodynamic properties of glass-forming liquids, proteins, and disordered spin systems.
system in equilibrium. Direct evaluation of $S_{\text{conf}}$ enabled testing of the phenomenological Adam–Gibbs (AG) theory that postulates a connection between thermodynamics and transport properties through a relation between $S_{\text{conf}}$ and viscosity $\eta$. This AG relation has been successful in rationalizing the slowing down of the transport properties of many glass-forming liquids approaching $T_g$. According to the theory, relaxation in supercooled liquids takes place through the activated motion of cooperatively rearranging regions (CRR), defined as the smallest region that can undergo a transition to a new configuration without a requisite simultaneous configurational change on and outside its boundary. The theory also postulates that the CRRs grow upon cooling, and thus, relaxation slows down because a concerted motion of larger and larger groups of particles is required. This idea has been further considered by a number of authors.

The notion that liquids approaching the glass transition should relax in dynamically correlated regions that have enhanced or diminished mobility relative to the average has received much attention over the past decade. This type of dynamics is commonly referred to as spatially heterogeneous dynamics. Numerous computer simulations and experimental works have clearly identified transient correlated regions of enhanced mobility. These regions form well-defined clusters, that can typically be decomposed further into groups of particles or molecules that follow each other in a stringlike fashion. Quantification of the size of these dynamically active regions clearly demonstrates that their size increases as a glass transition is approached. A recent theory of dynamic facilitation predicts that spatially heterogeneous dynamics is a necessary consequence of local dynamical facilitation and cooperative motion.

Despite the progress in quantifying the nature of spatially heterogeneous dynamics (SHD), a direct connection to the CRR that is central to the AG theory has been elusive, in part because the size of the CRR is not precisely defined from microscopic quantities. The AG approach does provide a prediction for the relationship between the size $z$ of a CRR and the configurational entropy, which provides a means to explore a possible relationship between the characteristic features of SHD and CRR. Recent work on a water model suggested that identifying $z$ as the average size of a cluster of mobile particles may satisfy the predicted AG relation.

Given that clusters are comprised of smaller groups of particles that move together in “strings”, it is also natural to ask whether the strings may be a possible, if not a better, candidate to represent the CRR envisaged by Adam and Gibbs. Hence, the primary focus of this work is to explore the connection between the dynamical clusters and strings that have been found in earlier works and the CRR of the AG theory. Our approach is as follows: We first evaluate the configurational entropy and dynamical heterogeneity of a one-component liquid described by the Dzugutov potential, a model for simple metallic liquids. The calculations of $S_{\text{conf}}$ are used to test the validity of the AG relation between $S_{\text{conf}}$ and the diffusion coefficient. We then investigate the extent to which mobile-particle strings or clusters may represent the CRR by comparing several different measures of the characteristic string and cluster size to the “minimum cooperatively rearranging region” of the AG theory.

The paper is organized as follows. In section 2, we present the mathematical formalism that describes the thermodynamics of the IS, which can also be found in refs 1, 7, 9, 48, and 49. We then briefly describe our model in section 3. In section 4, we discuss the IS properties of our system. Section 5 describes the evaluation of configurational entropy in detail. The validity of the AG theory and the connection of CRR with the dynamical clusters and strings of highly mobile particles are explored in sections 6 and 7. The conclusion is given in section 8.

II. Inherent Structure Thermodynamic Formalism

The central idea of the IS formalism is the notion that, at sufficiently low $T$, the dynamics of a liquid can be separated into vibrations within a single basin and infrequent transitions between basins. This partitioning is motivated by the fact that at $T$ approaching $T_g$, the time scales of intrabasin vibrational motion and interbasin diffusion differ by several orders of magnitude. Direct numerical evidence for the separation of the dynamics of a liquid into vibrations around and transitions between IS was provided by Schroder et al., where such separation becomes possible in the vicinity of the mode-coupling temperature $T_{\text{MCT}}$. Stillinger recognized the consequence of this partitioning is that the canonical partition function can be conveniently rewritten as a sum over all of the local potential energy minima and an integral around the surrounding basins of attraction.

The canonical partition function of a system of $N$ interacting particles is given by

$$Q_N(V,T) = \frac{\hbar^{3N}}{N!} \int \int \exp[-\beta H_N(r^N,p^N)] \, dr^N \, dp^N$$  (1)

where $H_N(r^N,p^N)$ is the Hamiltonian of the system that is expressed as

$$H_N(r^N,p^N) = \frac{1}{2m} \sum_{i=1}^{N} |p_i|^2 + U_N(r^N)$$  (2)

Here $r^N$ and $p^N$ are shorthand representations for the positions $\{r\}$ and momenta $\{p\}$ of the $N$ particles, and $U_N(r^N)$ is the total potential energy of the system. For the above Hamiltonian, the integration over the momenta can be carried out explicitly, and $Q_N(V,T)$ can be reduced to

$$Q_N(V,T) = \frac{\Lambda^{3N}}{N!} Z_N(V,T)$$  (3)

where $\Lambda = (2\pi\hbar^2/m)^{1/2}$ is the de Broglie thermal wavelength, $\beta = 1/k_BT$, and

$$Z_N(V,T) = \int \exp[-\beta U_N(r^N)] \, dr^N$$  (4)

is the configurational integral. In the IS formalism, the partitioning of the configuration space into nonoverlapping basins allows the partition function expressed above to be rewritten as

$$Q_N(V,T) = \sum_{\text{basins}} \exp(-\beta e_{\text{basin}}) \int_{R_{\text{basin}}} \exp[-\beta(U_N - e_{\text{basin}})] \, dr^N$$  (5)

where $R_{\text{basin}}$ is the set of points comprising a specific basin and $e_{\text{basin}}$ is the potential energy of the inherent structure corresponding to the basin. This last equation shows that the contributions to the partition function $Q_N(V,T)$ can be separated into two parts: the IS energy of all the distinct basins and the thermal excitation within the basins. If we introduce $\Omega(e_{\text{IS}}) \, d\epsilon_{\text{IS}}$ as the density of
states with IS energy between $e_{IS}$ and $e_{IS} + d e_{IS}$, then the above equation can be written as

$$Q_N(V, T) = \int d e_{IS} \Omega(e_{IS}) \exp[-\beta e_{IS} - \beta f_{basin}(\beta, e_{IS})] \tag{6}$$

where

$$-\beta f_{basin}(\beta, e_{IS}) = \ln \left( \int f_{basin} \exp[-\beta(U_N - e_{IS})] \frac{dN}{\Lambda} \right) \tag{7}$$

Here, $f_{basin}(\beta, e_{IS})$ is interpreted as the basin free energy with IS energy $e_{IS}$. Equation 6 is further simplified if we define the configurational entropy $S_{conf}(e_{IS})$ as

$$S_{conf}(e_{IS}) = k_B \ln \Omega(e_{IS}) \tag{8}$$

Then, eq 6 becomes

$$Q_N(V, T) = \int d e_{IS} \exp[-\beta(e_{IS} + f_{basin}(\beta, e_{IS}) - TS_{conf}(e_{IS}))] \tag{9}$$

In the thermodynamic limit, the free energy $A$ of the system can be obtained from this equation by employing a maximum integrand evaluation, which yields

$$A = \bar{e}_{IS} - TS_{conf}(\bar{e}_{IS}) + f_{basin}(\beta, \bar{e}_{IS}) \tag{10}$$

Here, $\bar{e}_{IS}$ is the average IS energy for a given T value that maximizes the integrand and thereby minimizes $A$.

The expression for the free energy eq 10 can be interpreted as follows. The first two terms on the right-hand side of eq 10 account for the average energy of the PEL minima visited and the degeneracy of the average IS energy $\bar{e}_{IS}$, respectively. The vibrational and the kinetic contributions are captured in the last term. Equation 10 thus provides a formal expression for the separation of configurational and vibrational contributions. Only an appropriate model for $f_{basin}(\beta, \bar{e}_{IS})$ is needed to fully evaluate $S_{conf}$, we shall see that the basin free energy is dominated by harmonic contributions, as seen in other model glass-forming liquids,\textsuperscript{5,6,49} with small corrections from anharmonicities that we can include in an ad hoc fashion.

### III. Model and Simulation

The model system we study is a one-component liquid using the Dzugutov interaction potential. The explicit form of the Dzugutov pair potential is\textsuperscript{51}

$$V = V_1 + V_2$$

$$V_1 = A(r^{-m} - B) \exp \left( \frac{c}{r - a} \right), \quad r < a$$

$$V_1 = 0, \quad r \geq a$$

$$V_2 = B \exp \left( \frac{d}{r - b} \right), \quad r < b$$

$$V_2 = 0, \quad r \geq b \tag{11}$$

where the parameters are compiled in Table 1. This pair potential is characterized by the presence of two repulsive regions and one attractive region. The potential is designed to favor local icosahedral ordering\textsuperscript{51,52} and is considered a generic model of simple metallic liquids, which have icosahedral coordination of the first neighbor shell. Further information about the model is found in refs 43 and 51.

### TABLE 1: Parameter Values for the Potential in Eq 11\textsuperscript{51}

<table>
<thead>
<tr>
<th>m</th>
<th>A</th>
<th>c</th>
<th>a</th>
<th>B</th>
<th>d</th>
<th>b</th>
</tr>
</thead>
<tbody>
<tr>
<td>16</td>
<td>5.82</td>
<td>1.1</td>
<td>1.87</td>
<td>1.28</td>
<td>0.27</td>
<td>1.94</td>
</tr>
</tbody>
</table>

Our molecular dynamics simulations are performed for a system of 2197 particles in a temperature range of $T = 0.44 - 1.0$. For all of the state points studied, the simulations are performed at isothermal conditions and at a constant density $\rho = 0.85$. Periodic boundary conditions are used in all three of the spatial directions. For each $T$ value studied, the liquid is cooled and equilibrated in a stepwise manner starting from $T = 1.6$. All of the units are quoted in Lennard-Jones reduced units: length in units of $\sigma$, temperature $T$ in units of $\varepsilon/k_B$, and time in units of $\sigma(m/\varepsilon)^{1/2}$. The mass $m$ and the distance $\sigma$ are set to unity.

To evaluate the entropy, we need to connect the range of densities and temperatures studied with the ideal gas. Hence, two additional sets of simulations are conducted. First, the system is simulated for temperatures ranging from $T = 2.0$ to $T = 5.0$ at constant density $\rho = 0.85$. Then, a second set of simulations at $T = 5.0$ is carried out in the density range of $0.85 - 0.001$.

The number of particles used in the present simulation is smaller than the $N = 17576$ used in ref 43. The calculation of the configurational entropy involves the diagonalization of a large Hessian matrix, to be described below. To accomplish this task in a reasonable amount of time, we have reduced the system size to $N = 2197$. Unfortunately, these smaller systems have a tendency to crystallize in simulations when supercooled for moderate times.\textsuperscript{53,54} To exclude crystallizing configurations from our analysis and to improve statistics, we simulate many independent samples at each $T$ and then remove those simulations that show a tendency toward crystal nucleation before beginning our analysis. The configurations that are crystallizing are identified by carefully inspecting the thermodynamic as well as the dynamic properties of each independent simulation. We use the behavior of the large system ($N = 17576$) as a benchmark and discard those configurations that have significantly different behavior. As a result, apart from its size, the properties of the system studied are within statistical error of that of the large system. Over all, our analysis is carried out by averaging over 23 independent simulations for the lowest $T$ value, while as many as 100 independent simulations are used for higher temperatures.

### IV. Inherent Structure Energy and Pair Distribution Function

As a first step toward characterizing the thermodynamics of the PEL, we sample the local potential energy minima nearest the system equilibrium trajectory on the PEL. This is accomplished by carrying out a conjugate gradient minimization\textsuperscript{55,56} of the potential energy starting from equilibrium liquid configurations, which are obtained from the molecular dynamics simulations. This process is often referred to as “quenching”. The associated quench rate is infinite, since $T$ is set to zero before energy minimization. For a given temperature of the starting equilibrium configurations, at least 115 equilibrium configurations are mapped onto the respective inherent structure configurations.

A basic property characterizing the minima on the landscape is the average potential energy $\bar{e}_{IS}$. The temperature dependence of $\bar{e}_{IS}$ has been studied by several authors.\textsuperscript{6,10,48,49,57–60} Figure 1 shows the plot of $\bar{e}_{IS}$ as a function of $T$. As in the previous
The histogram method is based on constructing a histogram or PEL method and the thermodynamic integration evaluated using two different methods, referred to as the TI method. The ti method is described in more detail elsewhere.7

Figure 1. Average, inherent structure energy per particle $\bar{E}_S/N$ as a function of $T$. Inset: $\bar{E}_S/N$ (circle) as a function of the inverse temperature for $T \leq 0.75$. The solid straight line is a guide for the eye.

Figure 2. Inherent structure pair correlation function $g(r)$ for $T = 0.44$ (solid line) and $T = 1.0$ (dashed line). Inset: pair correlation function $g(r)$ for the equilibrium liquid for $T = 0.44$ (solid line) and $T = 1.0$ (dashed line).

studies, $\bar{E}_S$ is nearly constant at high $T$. The IS average energy decreases upon cooling below the "onset of caging" indicating that the system populates deeper and deeper basins upon supercooling. Below this onset temperature, $S_{\text{IS}}$ is often found to follow a $1/T$ law for fragile liquids, which also appears true for this system (inset of Figure 1). This $1/T$ dependence has been mathematically shown to be a consequence of a Gaussian distribution of the probability density $P(\epsilon_{\text{IS}}/\beta)$. To gain insight into the spatial distribution of particles in a given potential energy minimum, we measure the pair correlation function $g(r)$. The results for two different temperatures, $T = 0.44$ and $T = 1.0$, are shown in Figure 2. As first recognized by Stillinger and Weber, the underlying structure obscured by the influence of vibrations in the equilibrium liquid becomes more apparent in the inherent structure. This is also the case for our system, where the splitting of the second peak in the pair correlation that arises from icosahehedral ordering becomes sharper as compared to the equilibrium liquid.

V. Evaluation of Entropy

In this section, we present the methods and the results for the entropy $S$, as well as the configurational $S_{\text{conf}}$ and vibrational $S_{\text{vib}}$ contributions. The configurational entropy can be evaluated using two different methods, referred to as the histogram or PEL method and the thermodynamic integration (TI) method. The histogram method is based on constructing the probability distribution $P(\epsilon_{\text{IS}},T)$ of the IS potential energy $\epsilon_{\text{IS}}$, which can be directly related to $S_{\text{conf}}$ with one unknown constant.

In this study, we will employ the TI method, which uses the fact that, upon deep supercooling, the motion of particles in configurational space is separable into vibrations within a basin and infrequent transitions between basins. As a result, the total entropy $S(T)$ can be expressed as the sum of the configurational entropy $S_{\text{conf}}(T)$ that results from the multiplicity of local potential energy minima sampled by the liquid and the vibrational entropy $S_{\text{vib}}(T)$ of typical basins sampled at $T$. Thus

$$S_{\text{conf}}(T) = S(T) - S_{\text{vib}}(T)$$

To obtain $S_{\text{conf}}(T)$, we need to evaluate both the overall $S(T)$, as well as the $S_{\text{vib}}(T)$. We will first focus on the evaluation of $S(T)$.

A. Total Entropy. To evaluate $S(T)$, we must connect our system with a reference system for which the exact value of $S$ is known. In this case, we will use thermodynamic integration to connect our system with an ideal gas state.

Generally speaking, thermodynamic properties can be split into a contribution from the ideal gas, and a contribution from interactions, typically referred to as the "excess" over the ideal gas value. For example, the total entropy $S$ can be written as $S = S_{\text{id}} + S_{\text{ex}}$, where $S_{\text{id}}$ and $S_{\text{ex}}$ are the ideal and excess terms of the entropy, respectively. Note that this should not be confused with the difference between the liquid and crystal entropies, sometimes also referred to as $S_{\text{cr}}$.

To determine the total entropy at a reference state point $R \equiv (T_R, V_R)$, where $T_R$ and $V_R$ are the reference temperature and volume, respectively, we perform thermodynamic integration at temperature $T_R = 5.0$ starting from a volume $V \rightarrow \infty$ (ideal gas limit) to the reference volume $V_R$ (corresponding to density 0.85). Following ref 64, we need to evaluate

$$S(R) = S(R') + \frac{U(R')}{T} + \int_{V_R}^{V} \frac{P_{\text{ex}}}{T} \, dV$$

where $S_{\text{id}} = -\frac{dA_{\text{id}}}{dT}$ is expressed as

$$S_{\text{id}} = N \left[ \ln \left( \frac{V}{N A^3} \right) + \frac{5}{2} \right]$$

To reduce the error in the numerical integration of $P_{\text{ex}}$ in eq 13, we use the fact that $P_{\text{ex}} = B_2(T)/k_B T/V^2$ as $V \rightarrow \infty$, where $B_2(T)$ is the first nontrivial virial coefficient. Therefore, we first subtract the virial term $B_2(T)/T/V^2$ from $P_{\text{ex}}$ and then integrate the difference over the system volume. The contribution arising from the first virial correction is then integrated analytically and added to the remaining calculation. The resulting expression is

$$\int_{V_R}^{V} \frac{P_{\text{ex}}}{T} \, dV = \int_{V_R}^{V} \frac{B_2(T)/k_BT}{T} \left[ \frac{N A^3}{V} \right] \, dV - B_2(T) V^2 / V_R$$

The values of the excess pressure $P_{\text{ex}} = P - P_{\text{id}}$ are plotted in Figure 3 as a function of $V$. From the behavior of $P_{\text{ex}}$ at large $V$ values, we find that $B_2(T) = 1.953$. Combining these results, we find that the total entropy at the reference state point $S(R)/k_B = 18.442$.

B. Entropy $S(T)$ at Different Temperatures. For a system with fixed $N$ and $V$ values, $dE = T \, dS$, where $E$ is the total energy of the system. We can use this relationship to evaluate the entropy at any $T$ along the isochoric path we study by integrating

$$S(T,V) = S(T_R,V_R) + \int_{T_R}^{T} \frac{dE}{T}$$
For hard spheres, Rosenfeld and Tarazona showed that the simulation. On the basis of the free energy functional calculation we fit the simulation data using the form

\[ U(T) = a + bT^{3/5} + cT \]  

As shown in Figure 4, this expression provides a good description of our numerical results for \( T \leq 0.75 \). To determine \( S(T) \) for \( T \leq 0.75 \), we first carry out an isochoric thermodynamic integration in the range of \( T_R \geq T \geq 0.75 \), using \( C_V \) obtained from the derivative of eq 20. This procedure links the entropy of the system \( S(T_R) \) at the reference temperature \( T_R = 5.0 \) to that at 0.75. Then, we use \( C_V \) obtained from the derivative of eq 19 to calculate \( S(T) \) at any \( T < 0.75 \). Figure 8 shows the plot of the total entropy \( S(T) \) as a function of \( T \) obtained in this way.

C. Vibrational Entropy \( S_{\text{vib}} \) and Basin Shape. We next turn our attention to the vibrational component of \( S \), which we will need to determine \( S_{\text{conf}} \). Previous studies have shown that the vibration around the basin minima can be described predominately by a harmonic expansion around the minimum, with minor corrections due to higher order anharmonic terms. Thus, \( S_{\text{vib}} \) can be expressed as the sum of harmonic \( S_{\text{harm}} \) and the anharmonic \( S_{\text{anh}} \) terms.

To determine if it is necessary to include anharmonic corrections to \( S_{\text{vib}} \), we first calculate the vibrational energy \( U_{\text{vib}} = U/N - 3/2 k_B T \); if the system is purely harmonic, \( U_{\text{vib}} = 3/2 k_B T \). Figure 5 shows that \( U_{\text{vib}} > 3/2 k_B T \), and thus there are higher order anharmonic terms that we must account for, as also found in the SPC/E model of water, the BKS model of silica, and the Lewis–Wahnström model of OTP.

1. Harmonic contribution to \( S_{\text{vib}} \). We first focus our attention on the harmonic contribution to the vibrational entropy resulting from the motion of the system within the basins. This can be calculated using the relation

\[ S_{\text{harm}} = \sum_{i=1}^{3N} \frac{3N-3}{k_B} \left( 1 - \ln \left( \frac{\hbar \omega_i}{k_B T} \right) \right) \]  

where \( \{ \omega_i \} \) values are the square roots of the eigenvalues of the Hessian matrix. The \( 3N \times 3N \) mass-weighted Hessian matrix elements \( H_{ij} \) are defined by

\[ H_{ij} = \frac{1}{\sqrt{m_m m_n}} \frac{\partial^2 U}{\partial r_i \partial r_j} |_{r_0} \]
where $m_i$ and $m_j$ are the masses of particle $i$ and $j$, respectively. These values are unity for our system. The Hessian matrix arises naturally by considering an expansion of $U$ near a local minimum $\epsilon_{15}$, that is,

$$U(r^N) = \epsilon_{15} + \sum_{i,j=1}^{N} (r_i - r_{15}) \frac{\partial^2 U}{\partial r_i \partial r_j} (r_j - r_{15}) + \text{higher order terms} \quad (23)$$

where $r^N$ is the $3N$-dimensional IS configuration at the local minimum $\alpha$. At the local minimum, the first derivative vanishes and $U(r^N) = \epsilon_{15}$. Hence, the square roots of the eigenvalues, or normal modes $\{\omega_i\}$, provide a description of the curvature of the basin near the basin minimum. It is this curvature that we refer to as “basin shape”. Note that all of the $\omega_i \geq 0$ since they are evaluated at a local minimum; there are three zero eigenvalues that account for the three independent translational directions of the entire system.

To quantify the basin shape, we evaluate the density of states $N(\omega)$, which is the histogram of $\{\omega_i\}$ (Figure 6). The spectrum of $\{\omega_i\}$ changes weakly with $T$. Only the position of the maximum slightly shifts to smaller values of $\omega$ with decreasing $T$. To more closely inspect this change, we calculate the ensemble average of the sum of the logarithms of the frequencies of normal modes

$$\langle \ln \omega \rangle = \left( \frac{1}{3N - 3} \sum_{i=1}^{3N-3} \ln \omega_i \right) \quad (24)$$

This quantity captures the average quadratic shape of a basin. The $T$ dependence of $\langle \ln \omega \rangle$ is shown in the inset of Figure 6. The $\langle \ln \omega \rangle$ value decreases weakly with decreasing $T$, indicating that basins become increasingly broad with increasing depth of the minima. Similar behavior has been found in the Lewis–Wahñström model of supercooled o-terphenyl (OTP). The magnitude of the changes on cooling are small compared to water, not unlike the binary LJ system. In contrast, in the SPC/E model of water the average basin frequency increases upon cooling, and hence the basins in that system become increasingly “sharp” with decreasing basin energy.

Using the calculated values of $\omega_i$ at each $T$, we plot $S_{\text{harm}} - (T/k_B)$ calculated using eq 21 in Figure 8.

2. Anharmonic Contribution to $S_{\text{th}}$. To evaluate the anharmonic contribution to $S_{\text{th}}$, we will exploit the fact that it is relatively easy to estimate the anharmonic contribution to the potential energy, $U_{\text{anh}}$, and then use the relation

$$S_{\text{anh}}(T) = \int_0^T \frac{1}{T} \frac{dU_{\text{anh}}(T')}{dT'} dT' \quad (25)$$

where we have assumed that $S_{\text{anh}}(T=0) = 0$, as well as $S(T=0) = 0$, in accordance with the third law of thermodynamics for a liquid.

Recalling that eq 23 describes the expansion of $U$ about a local minimum, the anharmonic contribution corresponds to the higher order terms in the expansion. Thus, $U_{\text{anh}}$ can be approximated as

$$U_{\text{anh}}(T) = U(T) - \bar{\epsilon}_{15}(T) - U_{\text{harm}}(T) \quad (26)$$

$U(T)$ and $\bar{\epsilon}_{15}(T)$ are found from the simulation, while $U_{\text{harm}}(T) = \frac{1}{2}(N-1)k_BT$. To perform the numerical integration of eq 25, the value of $U_{\text{anh}}(T)$ found from the simulation is further fitted to a polynomial in $T$ to the order $T^5$ using the expression

$$U_{\text{anh}}(T) = \sum_{k=2}^{5} c_k T^k \quad (27)$$

This form for $U_{\text{anh}}$ requires that $U_{\text{anh}}$ and its derivative with respect to $T$ (the anharmonic contribution to specific heat) vanish at $T = 0$, as we would expect.

In Figure 7, we show the plot of $U_{\text{anh}}$ and $S_{\text{anh}}$ obtained from these calculations. Additionally, $S_{\text{anh}}$ is plotted together with $S$ and $S_{\text{harm}}$ in Figure 8 to emphasize the extent of its contribution in the calculation of $S_{\text{conf}}$.

D. Configurational Entropy. Now that we have evaluated both $S(T)$ and $S_{\text{anh}}(T)$, we can evaluate $S_{\text{conf}} = S(T) - S_{\text{anh}}$. The plot of $S_{\text{conf}}$ is shown in Figure 8 together with the plots of $S$, $S_{\text{harm}}$, and $S_{\text{anh}}$. Extrapolation of $S_{\text{conf}}$ to lower $T$ values using the fitting forms presented indicates that $S_{\text{conf}} \rightarrow 0$ at $T \approx 0.33$. In the PEL approach to liquid dynamics, when the number of accessible basins becomes intensive in the number of particles, the system can no longer explore phase space and is said to have reached a vitrified state of thermodynamic origin. This state corresponds to $S_{\text{conf}} = 0$. The temperature where $S_{\text{conf}}$ vanishes is frequently referred to as the Kauzmann temperature.
observed for the BKS model of silica\(^0\) when \(\log D\) is plotted against \(1/TS_{conf}\) implying that the relation \(D \propto \exp(-A/TS_{conf})\), where \(A\) is a constant, may not be appropriate for silica. There, a better fit over a range of densities was obtained for the following relationship

\[
D/T \propto \exp(-A/TS_{conf})
\]  

(29)

Similarly, we find a better fit to a line when we plot \(\log(D/T)\) against \(1/TS_{conf}\) (Figure 9b). Note, however, the slight upward bend of the data at low \(T\) that remains. This behavior may indicate an enhanced value of the diffusion coefficient over that which would be predicted by the original Adam–Gibbs expression (see section VII.C) and the Stokes–Einstein relation\(^30,32,81\) and is not unexpected when substantial dynamical heterogeneity is present in the system\(^82\) as is the case here.

### VII. Dynamical Heterogeneity and Cooperatively Rearranging Regions

Central to the development of the AG approach is that motion of the system occurs through the independent relaxation of CRRs. The theory relates the minimum size of a cooperatively rearranging region \(z^*\) to the configurational entropy \(S_{conf}\) by

\[
z^* \propto \frac{1}{S_{conf}}
\]  

(30)

However, as mentioned earlier, a precise microscopic definition of the CRR is lacking. Here we are particularly interested in assessing a possible connection between the CRR envisaged by the AG theory and the dynamic strings and clusters of mobile particles that are the signatures of spatially heterogeneous dynamics.

#### A. Mobile Particle Clusters and Strings

One way to quantify the nature of spatially heterogeneous motion is to identify clusters of mobile particles. These clusters are typically defined as sets of highly mobile particles that are within the first neighboring distance of each other.\(^16,40\) These clusters can also be separated into subsets of particles that replace each other along one-dimensional stringlike paths. These subsets are referred to as strings. Typically, the mobile particles that make up strings and clusters are identified by selecting the particles with the \(\approx 5\%\) largest scalar displacement in a time interval \(t\).\(^15,16,39\) Both clusters and strings have been reported in a number of different systems and were found to be relevant to the process of cage rearrangement, bulk relaxation, and exploration of the potential energy landscape.\(^4,8,83\)

We first focus on the mobile particle clusters. We identify two mobile particles as belonging to the same cluster if they are separated by a distance no greater than that of 1.5, the position of the first minimum of \(g(r)\). By choosing many intervals \(t\) to define the mobility of particles, we can track the time evolution of these clusters. At any time, we can define a size of these clusters. There are two typical measures of the size: (i) the average size of a randomly chosen cluster, which we denote \(S_0\), and (ii) the average size of a cluster to which a randomly chosen molecule belongs, which we denote \(S_n\). The number-averaged cluster size is \(S_n = \langle n \rangle\), where \(n\) is the number of particles in the cluster. The \(S_0\) value is dominated by smaller clusters and is a number average. The weight-averaged cluster size \(S_w = \langle n^2 \rangle / \langle n \rangle\) gives more weight to larger clusters, since a randomly chosen particle is more likely in a large cluster.

A significant feature of the clusters is that, at any particular \(T\), the mean size of the clusters is time dependent, with a peak
Figure 9. Test of the validity of the Adam–Gibbs theory for the Dzugutov liquid: (a) the diffusion coefficient $D$ plotted against $1/T S_{\text{conf}}$ on a semilogarithmic axis and (b) $D/T$ plotted against $1/T S_{\text{conf}}$ on a semilogarithmic axis.

Figure 10. (a) Number averaged cluster size $S_n$ and (b) the number averaged string size $L_n$ plotted as a function of time. Note the strings are averaged over all of the strings including $l = 1$.

at intermediate time corresponding to the late-$\beta$-relaxation time of the MCT as shown in Figure 10a. For simplicity, we show only $S_n$, since the behavior of $S_w$ is qualitatively nearly identical (the numerical values are larger for $S_w$). The maximum values of both $S_n$ and $S_w$, which we denote $S^*_n$ and $S^*_w$, increase with decreasing $T$, consistent with the phenomenological argument of the AG theory that requires $z^*$ to increase as $T$ decreases. Hence, the cluster size is one possible candidate to describe CRR, as we will discuss.

We next identify strings that aggregate to form larger, highly ramified clusters. Strings of mobile particles are identified by taking snapshots of configurations that are separated by a time interval $t$ and then by finding highly mobile particles that replace each other within a radius $\delta = 0.6$ within the time interval $t$. Once the strings are found, we can define the mean size of the strings using either a number average $L_n$ or a weight average $L_w$ in analogy to the mobile particle clusters. Figure 10b shows that $L_n$ has nearly the same qualitative behavior as $S_n$. Since strings are the components that make up the clusters and also match the qualitative expectations of the CRR of the AG approach, these are also viable candidates for the CRR. It is important to note that strings are not simply small clusters. Strings are identified by verifying both neighborhood and the direction of the displacements of particles, while neighborhood is the only criterion for clusters. Such directional correlations may owe its origin to the local structure, where dynamic facilitation is channeled through the local excitations.

B. Potential Candidates for Describing CRR. To explore the possible candidates for the CRR of the AG theory, and possibly identify the best measure of the CRR, we examine the linearity of the relationship between the number-averaged and weight-averaged mean values of the clusters and strings and the inverse configurational entropy. In particular, we use the average values of the clusters and strings at the time when they are maximum to test the AG relation given in eq 30.

Strictly speaking, eq 30 requires that $z^* \to 0$ as $1/S_{\text{conf}} \to 0$. This requirement means that a linear fit between a measure, $X$, of the CRR and $1/S_{\text{conf}}$ must pass through the origin. With this in mind, we tested $X = S^*_n$, $S^*_w$, $L^*_n$, and $L^*_w$ defined in the previous section. Of these four candidates, we find that only $S^*_n$ and $L^*_w$ obey a linear relationship with $1/S_{\text{conf}}$ that passes through the origin (Figure 11). A similar result for $S^*_w$ was found in ref 42.; however, they found that $S^*_w - 1$ passes through the origin, and rationalized this by arguing that clusters of size one should not be considered CRR, and thus should be subtracted.

To better comprehend the variation in the intercept, we appeal to physical considerations that motivate a relaxation of the constraint that $z^*$ should be zero only when $1/S_{\text{conf}}$ is zero. The AG relation was originally proposed for liquids near the glass transition; the limit of $1/S_{\text{conf}} \to 0$ refers to the infinite temperature limit. At high $T$, cooperativity is not important for particle motion, and so it is reasonable to expect that $z^*$ should approach one at a finite value of $T$ and thus before $1/S_{\text{conf}} \to 0$. As a result, the extrapolated value of $z^*$ at the origin will be less than zero. Therefore, we relax the requirement that our fits should pass through the origin and test whether we obtain linear fits for $X - 1$ against $1/S_{\text{conf}}$, where we have subtracted one to eliminate those clusters or strings for which there is no cooperativity. We expect a fit of $X - 1$ vs $1/S_{\text{conf}}$ to pass through a positive, nonzero value of $1/S_{\text{conf}}$ at $X - 1 = 0$. We find a linear relationship for all of the candidates $X = S^*_n$, $S^*_w$, $L^*_n$, and $L^*_w$, as seen in Figures 12 and 13. Hence, all of these quantities are candidates for a measure of $z^*$.
At first glance, the plethora of possible measures for \( z^* \) is confusing. Which measure is most appropriate? To answer this question, we return to the formulation of the AG theory, where \( z^* \) in eq 30 refers to the minimum size that can give rise to a cooperative rearrangement. This naturally leads us to focus on the definition of \( z^* \) that is smallest and obeys eq 30. Therefore, we suggest that the string size is the more appropriate measure of the CRR. Why then should the cluster size follow eq 30, as found here and in ref 42? This is likely a result of the fact that the clusters consist of a collection of strings, which themselves obey the AG relation. But it is the strings, and not the clusters, that are the minimal units. Additionally, quantitative tests of the quality of the linear fit indicate that the linear fits to the string size are marginally better than those for the cluster size.

Figures 12 and 13 also include the weight-averaged and number-averaged mean cluster sizes and string sizes reported in ref 43 for a system containing 17,576 particles. Although we are unable to calculate \( S_{\text{conf}} \) in the large system, all of the other calculated bulk thermodynamic and dynamic quantities are indistinguishable within statistical error in the two systems, and thus we assume identical values for \( S_{\text{conf}} \). The figures show that the weight-averaged cluster sizes (which are larger than the number-averaged cluster sizes) are diminished in the small system at low \( T \), but the string sizes are essentially unchanged. This is expected since the largest clusters in the system exceed the smaller box size at low \( T \), but the largest strings do not exceed the box size. Thus, there are finite size effects in our measure of \( S_{\text{w}}^* \), and so it is not reliable to compare it with the AG expression. All of the other quantities should be unaffected.
It would be valuable to revisit the water system where the cluster size was found to describe the CRR,\textsuperscript{42} to test if strings provide a better candidate for CRR in that system, as well as extending this analysis to other systems. As a final remark, we point out that a test of the AG relation that employs “micro-strings”, groups of particles that move coherently (that is, simultaneously) within strings (ref 43), may constitute an alternative candidate for the CRR, but that lies outside the scope of this work.

C. Characteristic Time of the AG Theory and Relationship to Configurational Entropy. In Adam and Gibbs’ original paper,\textsuperscript{13} they proposed that the characteristic time required for the liquid to undergo a transition to a new configuration is proportional to the characteristic relaxation time of a cooperatively rearranging region. On this basis, they argued that

\[ t_{\text{transition}} \propto \exp\left(\frac{A}{TS_{\text{conf}}}\right) \]  

(55)

Since we have shown the strings (and clusters of strings) of highly mobile particles to be likely candidates for the CRRs, it is reasonable to propose that this characteristic time is given by the characteristic time of a string or cluster (which are essentially the same). We thus propose the following

\[ t_{\text{str}}^{\max} \propto \exp\left(\frac{A}{TS_{\text{conf}}}\right) \]  

(56)

This expression is tested in Figure 14, where we see that a good linear fit is obtained when \( \ln t_{\text{str}}^{\max} \) is plotted vs \( 1/TS_{\text{conf}} \) over the admittedly limited decades of available data to us. The data for \( t_{\text{str}}^{\max} \) is included in the figure for comparison. Unfortunately, the crystallization tendencies of the Dzugutov liquid prevent us from obtaining equilibrium data at lower \( T \) values. It would be useful to test this expression in other liquids where additional decades in characteristic time scales are more easily obtainable. We note that a plot of either \( D/r_{\text{str}}^{\max} \) or \( (DIT)/r_{\text{str}}^{\max} \) vs \( T \) for \( r_{\text{str}}^{\max} \) corresponding to the time when the weight or number-averaged string or cluster size is maximum, is not constant for all \( T \) values, demonstrating that the constant \( A \) in the numerator of the exponential of the AG expressions in eq 28 (or eq 29) and 31 is different in the two expressions. This is consistent with the derivation of the two expressions, which require several assumptions pertinent to the particular quantity under consideration.

VIII. Conclusion

One of the overarching goals of studying the correlated dynamics in supercooled liquids is to incorporate information about dynamic heterogeneities into theoretical approaches to the glass transition. To make progress in this direction, we need to identify connections between bulk relaxation phenomena and the strings and clusters of mobile particles. The approach presented by Adam and Gibbs proposes relations for the entropy with the bulk dynamics and the size of clusters. Combining these relations gives a possible direct theoretical connection between heterogeneity and relaxation.

By relating the average size \( S_{\text{n}} \) of mobile clusters to the diffusion constant \( D \) and the configurational entropy \( S_{\text{conf}} \), Giovambattista et al. argued that \( S_{\text{n}} \) can be interpreted as the size of the CRR of the AG theory. Noting that the description of the CRR in the AG theory emerges from a heuristic argument, this connection sheds light into the possible origins of the AG theory, which has proved to be extremely useful over the last several decades to explain the nature of slowing dynamics in glass-forming liquids. Part of the motivation of this work was to explore the findings for mobile-molecule clusters in ref 42 for a different class of material; in so doing, we also investigated several predictions of the inherent structure formalism for the Dzugutov liquid. An additional motivation for this work was to explore to what extent strings, which include directional correlations not used to define mobile-particle clusters, may provide a better measure of the CRR of the AG theory.

We find that the temperature dependence of the average potential energy of the equilibrium liquid obeys the Rosenfeld—Tarazona \( T^{3/5} \) law for low temperatures. Furthermore, the temperature dependence of the average inherent structure energy \( \tilde{e}_{\text{IS}} \) is found to follow a \( 1/T \) law in the deeply supercooled regime, while \( \tilde{e}_{\text{IS}} \) is essentially constant at high \( T \). The shape of the basins around the local minima in configuration space, as detected by the average basin vibrational frequency, is observed to become slightly broader upon cooling similar to observations in the Lewis—Wahnström model of supercooled \( \alpha \)-terphenyl (OTP) and in contrast to that observed for the SPC/E model of water in which the basins become increasingly sharp upon cooling.

We evaluated the configurational entropy \( S_{\text{conf}} \) of the Dzugutov liquid, and we investigated the validity of the Adam—Gibbs theory in that system. We find that the Adam—Gibbs relation between bulk dynamics and \( S_{\text{conf}} \) holds for the Dzugutov liquid at the studied \( T \), as was observed in other systems. We also explored the connection between the dynamical clusters or strings that are the signatures of spatially heterogeneous dynamics, and the CRRs of the Adam—Gibbs theory. Similar to the observation for the SPC/E model of water, we find a linear relationship between the size of mobile-particle clusters and \( 1/S_{\text{conf}} \), which in turn provides a possible connection between clusters and the CRR. We further tested the possible relationship between the size of mobile-particle strings and \( S_{\text{conf}} \) and found a marginally better linear relationship than that found for the cluster size. This marginal difference between the strings and the clusters in better describing the CRR is likely a consequence of the fact that clusters are themselves composed of strings. Nonetheless, the fact that strings show a slightly better linear relationship, combined with the fact that these are the more elementary units of cooperativity, suggests that strings are better candidates for the CRR within the language proposed by the AG theory. We performed tests of the characteristic time of the string and cluster size as the characteristic relaxation time of the AG theory, which further support this proposition. Simulations that are able to probe lower \( T \) values will be
valuable, as the range over which we have been able to test this relation is relatively limited because of the crystallization tendencies of the Dzugutov liquid at lower T values.

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References and Notes

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The constants in the polynomial fit to $P_e$ are $a_2 = 4.71 \times 10^7$, $a_3 = 1.42 \times 10^{11}$, $a_4 = 3.37 \times 10^{14}$, and $a_5 = 1.47 \times 10^{18}$. The $a_2$ value is directly related to $B_2$.

To evaluate eq 14, we use $p = 0.063507$ kJ ps mol$^{-1}$ and assume the unit of energy to be $\epsilon = 1$ kJ mol$^{-1}$, the unit of length to be $\sigma = 1$ nm, and the unit of time to be $\tau = 1$ ps. We obtain $S_0(R)/k_B = 23265.5$ using these values. Note that the choice of constants will play no role when we return to reduced units.


(72) The best fit of $U$ using eq 19 yields $a = -7221.98$ and $b = 6292.87$, in the temperature range $T \leq 0.75$.

(73) The best fit of $U$ using eq 20 yields $a' = -6771.84$, $b' = 6492.36$, and $c' = -785.32$. It is interesting to note that with a different set of fitting parameters, this functional form can describe the entire $T$ range of the simulation data.

(74) The coefficients $c_i$ we find for eq 27 are $c_2 = -783.2$, $c_3 = 4500.3$, $c_4 = -6241.1$, and $c_5 = 2917.6$.


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