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Variation of the dynamic susceptibility along an isochrone

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Koperwas et al. showed in a recent paper [Phys. Rev. Lett. 111, 125701 (2013)] that the dynamic susceptibility $\chi_4$, as estimated by dielectric measurements for certain glass-forming liquids decreases substantially with increasing pressure along a curve of constant relaxation time. This observation is at odds with other measures of dynamics being invariant and seems to pose a problem for theories of glass formation. We show that this variation is in fact consistent with predictions for liquids with hidden scale invariance: Measures of dynamics at constant volume are invariant along isochrones, called isomorphs in such liquids, but contributions to fluctuations from long-wavelength fluctuations can vary. This is related to the known noninvariance of the isothermal bulk modulus. Considering the version of $\chi_4$ defined for the $NVT$ ensemble, data from simulations of a binary Lennard-Jones liquid show in fact a slight increase with increasing density. This is a true departure from the formal invariance expected for this quantity.

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I. INTRODUCTION

A quantity of great interest in recent years in the context of supercooled, glass-forming liquids is the dynamic susceptibility $\chi_4$, associated with a four-point correlation function $S_4(k, t)$. This was originally introduced to understand spin-glass models [1], subsequently used to study dynamical heterogeneity in computer simulations of glass-forming liquids [2–7], and with the introduction of experimentally accessible approximations to $\chi_4$, in real liquids [8–11]. It quantifies the dynamical heterogeneities; in principle it can be interpreted in terms of a length scale $\xi_4$ characterizing dynamical fluctuations or the number of dynamically correlated particles during structural relaxation, but the precise relation is not trivial [12]; even assuming a scaling $\chi_4 \sim \xi_4^4$, the proportionality factor is nonuniversal [13]. The growth of $\chi_4$ (or $\xi_4$) is considered relevant for explaining the dramatic dynamical slow-down as the glass transition is approached, as well as other features of relaxation in viscous liquids [14]. In particular, it is believed that there is a unique relation between the relaxation time $\tau$ and the shape of the relaxation spectrum on the one hand [15], and the size of dynamical heterogeneities [16–21] on the other hand. As a recent example of such a claim, Flender et al. described a universal behavior in the relation between $\xi_4$ and $\tau$ rescaled by its value when violations of the Stokes-Einstein relation become apparent [22]. We note here that $\chi_4$ is not uniquely defined: It depends on the correlator of interest, and, as will be discussed in detail below, on the statistical ensemble (NVE, NVT, etc.).

Pressure has been increasingly exploited as an extra experimental parameter [23]. This has led to a focus on liquids obeying so-called power-law density scaling when temperature $T$ and pressure $p$ are varied; that is, liquids for which the relaxation time depends only on a scaled quantity $\Gamma \equiv p^\gamma / T$ where $p$ is the density and $\gamma$ a system-specific scaling exponent [23–28]. An experimentally observed feature of such liquids is so-called isochronal superposition according to which the relaxation spectra corresponding to the same relaxation time–but different densities and temperatures–superpose [15,29–31] (see Ref. [32] for an apparent exception). This suggests that the physics governing relaxation is the same at points in the phase diagram for which the relaxation time is the same.

Given the existence of such liquids, and the supposed link between $\tau$ and $\chi_4$, it is natural to investigate the behavior of $\chi_4$ along isochrones. This was recently done for the van der Waals glass-forming liquids o-terphenyl, glibenclamide, and phenylphthalein-dimethylether in Ref. [33]. Surprisingly, a significant variation was found, with the maximum $\chi_4$ decreasing as pressure (and therefore density) increased along an isochrone (see also Refs. [34,35]); in Ref. [36] the opposite behavior was reported for dibutyl-phthalate; in Ref. [37] no significant variation was found for four other liquids. Where a variation was seen, the interpretation was that the temperature- and density-related contributions to the dynamical heterogeneities are nonequivalent, since they contribute differently to different measures of dynamics. Thus the postulated unique relation between $\chi_4$ and the growth of $\tau$ must apparently be questioned; for instance, Alba-Simionesco et al. consider the variation along an isochrone of the number of correlated particles to contradict predictions from the random first-order transition (RFOT) theory [36]. On the other hand it is generally believed that, e.g., for van der Waals liquids, density and temperature changes affect the dynamics in the same way along an isochrone [15], raising the question: is this wrong or is the traditional $\chi_4$ quantity not the relevant measure of dynamic heterogeneities? We note that simulation results showing invariance of the $NVT$ quantity in Lennard-Jones systems were presented a few years ago [38,39]; the issue of ensemble dependence was not discussed though.

The purpose of this article is to throw light on this question using isomorph theory, which provides a theoretical framework for the density-scaling behavior mentioned above. Following extensive theoretical and simulation investigations [39–42] we have proposed the existence of a class of simple liquids [43]. We use the term Roskilde (R) liquids to distinguish from earlier senses of liquid simplicity [44]. Their key feature is the existence of isomorphs: curves in the phase diagram along which several properties are invariant to a good approximation [39], including all dynamical quantities, as long as volume is not allowed to fluctuate. One of these properties is
the relaxation time in appropriate reduced units; this suggests that liquids obeying density scaling (not necessarily power-law density scaling) can be identified as simple liquids in the R sense, and that isomorphs in experiments can be identified with isochrones.

II. ISOMORPHS

The theory of isomorphs—the formal theory underlying the concept of R liquids—takes as its starting point the following general definition of isomorphic state points: Two state points $(\rho_1, T_1)$ and $(\rho_2, T_2)$ are isomorphic if the Boltzmann factors of corresponding microstates are proportional:

$$\exp\left(-\frac{U(r_i^{(1)}, \ldots, r_N^{(1)})}{k_B T_1}\right) = C_{12} \exp\left(-\frac{U(r_i^{(2)}, \ldots, r_N^{(2)})}{k_B T_2}\right).$$

(1)

Here $U$ is the potential energy function and $C_{12}$ depends on the two state points, but not on which microstates are considered. Corresponding microstates means $\rho^{1/3} r^{(1)}_i = \rho^{1/3} r^{(2)}_i$, or $r^{(1)}_i = r^{(2)}_i$, where a tilde denotes so-called reduced units. Reduced units for lengths means multiplying by $\rho^{-1/3}$, for energies dividing by $k_B T$, and for times dividing by $(m/k_B T)^{1/2} \rho^{-1/3}$ (for Newtonian dynamics). An isomorph is a curve in the phase diagram consisting of points which are isomorphic to each other. From the definition it follows that all structural and dynamical correlation functions are invariant when expressed in reduced units. We speak of a quantity being formally isomorph invariant if its invariance follows from definition (1); because the proportionality of Boltzmann factors is typically only approximate, the actual extent to which a given quantity is invariant has to be checked empirically. Thermodynamic quantities which do not involve volume derivatives, such as the excess entropy $S_\text{ex}$ and specific heat at constant volume $C_V$, are also formally isomorph invariant. We define the density scaling exponent,

$$\gamma \equiv (\partial \ln T / \partial \ln \rho)_\text{ns},$$

(2)

as the slope of isomorphs in $(\ln \rho, \ln T)$ space [39]. To a good approximation this depends only on temperature; this is equivalent to approximating the potential energy of a Roskilde liquid with the following form [42]:

$$U(\vec{R}) \cong h(\rho) \vec{\Phi}(\vec{R}) + g(\rho).$$

(3)

Here $\vec{R}$ is the 3N-dimensional vector of positions in reduced units (i.e., multiplied by $\rho^{1/3}$), and $\vec{\Phi}(\vec{R})$ is a dimensionless function of the reduced coordinates. The scaling function $h(\rho)$ determines the shapes of isomorphs via

$$h(\rho)/T = \text{const};$$

(4)

its logarithmic derivative,

$$d \ln h/d \ln \rho,$$

(5)

is just $\gamma$. The term $g(\rho)$ depends on density (or volume) but not on the microscopic coordinates [42]: It contributes an extra noninvariant part to the free energy and its volume derivatives, including the bulk modulus. That this density-dependent term is actually nonlocal plays a role later on in the discussion. More recent developments of isomorph theory allow for the variation of $h(\rho)$ from one isomorph to another, or equivalently, that the exponent $\gamma$ can depend on temperature at fixed density [45,46]. A general method for identifying isomorphs is to consider the configurational adiabats, which are formally isomorphs (that is, the excess entropy is formally isomorph invariant); their slope in $(\ln \rho, \ln T)$ space is given by the fluctuation formula,

$$\gamma = \frac{\langle \Delta U \Delta W \rangle}{\langle (\Delta U)^2 \rangle},$$

(6)

which when combined with Eq. (2) allows the curves to be generated in a stepwise manner.

A crucial insight from this framework is that any quantity which is claimed to control, for example, the relaxation time must also be invariant on isomorphs. Thus it is vital to consider the isomorph invariance of different formulations of $\chi_4$—only one which is formally isomorph invariant can be relevant. We show below that the version of $\chi_4$ which has been estimated experimentally is not invariant on an isomorph, because the isothermal bulk modulus is not. We also show that an isomorph-invariant version of $\chi_4$ does exist, namely the version defined for the NVT ensemble.

III. ISOMORPH INVARIANCE OF $\chi_4$

We consider the different versions of $\chi_4$ and the approximate expressions used to determine it experimentally (see Ref. [47] for an experimental determination which does not use these approximations). For simplicity, in this section we consider a pure substance, so that concentration fluctuations need not be accounted for. Recall that $\chi_4$ can be defined as the variance of the correlator whose average is some correlation function of interest; thus it measures dynamical fluctuations.

If we consider a two-time equilibrium correlation function $C(t)$, we can write $C(t) = \langle C_2(t_0, t_0 + t) \rangle$ where $C_2(t_0, t_1)$ is the fluctuating two-time correlator. Here the average may be interpreted as over initial configurations keeping the initial time $t_0$ fixed and/or, as is done in practice with simulation data, over different initial times $t_0$ within the same trajectory. One now defines

$$\chi_4(t) \equiv N \sigma_\text{C}^2 = N \left( \langle C_2^2 \rangle - \langle C \rangle^2 \right),$$

(7)

where the total number of particles $N$ is included to give an intensive quantity.

Since $\chi_4$ measures fluctuations, it should not be too surprising that it depends on ensemble. Berthier et al. [9] have analyzed the ensemble dependence in detail. When going from an ensemble with a constrained global variable to one where it is free to fluctuate, $\chi_4(t)$ increases by a positive amount corresponding to the fluctuations induced by those of the unconstrained variable. For example, in going from NVE to NVT the energy is allowed to fluctuate; the additional contribution to $\chi_4$ involves the energy fluctuations themselves and the correlation between $C$ and $E$; this leads to a term involving the isochoric specific heat and the temperature derivative of $C(t)$:

$$\chi_4^{\text{NVT}}(t) = \chi_4^{\text{NVE}}(t) + \frac{1}{c_V/k_B} \left( \frac{\partial C(t)}{\partial \ln T} \right)_\rho^2.$$
Going further by allowing the volume to fluctuate leads to the $NPT$ ensemble and an additional term involving the isothermal bulk modulus and the density derivative of $C(t)$ [9]:

$$
\chi_4^{NPT}(t) = \chi_4^{NVT}(t) + \frac{\rho k_B T}{K_T} \left( \frac{\partial C(t)}{\partial \ln \rho} \right)_T^2 \tag{9}
$$

$$
\chi_4^{NVT}(t) = \chi_4^{NVE}(t) + \frac{1}{c_V/k_B} \left( \frac{\partial C(t)}{\partial \ln T} \right)_\rho^2 + \frac{\rho k_B T}{K_T} \left( \frac{\partial C(t)}{\partial \ln \rho} \right)_T^2 \tag{10}
$$

Here $c_V = C_V/N$ is the isochoric specific heat per particle, and $K_T$ the isothermal bulk modulus. For a pure substance as considered here this expression for $\chi_4^{NPT}$ is equivalent to the ensemble-independent quantity obtained by taking the $k \to 0$ limit of $\chi_4(k,t)$ after the thermodynamic limit (see Appendix B); for mixtures additional terms related to concentration fluctuations must be included. The above relation is based on the formalism for transforming between ensembles developed by Lebowitz et al. [48,49] but applied to the variance of a two-time dynamical quantity. In this formalism it must be realized that by the relation is based on the formalism for transforming between ensembles developed by Lebowitz et al. [48,49] but applied to the variance of a two-time dynamical quantity. In this formalism it must be realized that by the formalism for transforming between ensembles developed by Lebowitz et al. [48,49] but applied to the variance of a two-time dynamical quantity.

Thus the density derivative is proportional to the (isomorph-invariant) temperature derivative with a proportionality constant $-\gamma$. The latter is not isomorph invariant, but its variation over typically accessed densities is small and can often be neglected (see Refs. [36,50] for an exception). On the other hand the third term in Eq. (10) ($\rho$ term) includes also the factor $\rho k_B T/K_T$, which is the inverse reduced-unit isothermal bulk modulus. That it is not isomorph invariant follows from Eq. (3), whose term $g(\rho)$ contributes noninvariant parts to volume derivatives of the (free) energy, such as the bulk modulus.

The conclusion from the above analysis is that $\chi_4^{NPT}$ is not isomorph invariant, while $\chi_4^{NVE}$ is. This applies also to the estimators introduced by Berthier et al. which involve dropping the $NVE$ contribution:

$$
\chi_4^{NVT} \simeq \frac{1}{c_V/k_B} \left( \frac{\partial C(t)}{\partial \ln T} \right)_\rho^2 \tag{invariant}, \tag{14}
$$

$$
\chi_4^{NPT} \simeq \frac{1}{c_V/k_B} \left( \frac{\partial C(t)}{\partial \ln T} \right)_\rho^2 + \frac{\rho k_B T}{K_T} \left( \frac{\partial C(t)}{\partial \ln \rho} \right)_T^2 \tag{varies}, \tag{15}
$$

Thus isomorph theory predicts that there will be little variation along the isochrone when the $NVT$ version of $\chi_4$ is considered, so the latter is an allowed candidate for a quantity which controls the dynamics.

IV. EXPERIMENTAL TESTS

Data for the $NVT$ $\chi_4$ has already been published in Ref. [33], in the supplementary material. We reproduce their figure S2 in Fig. 1, which shows data from dielectric measurements made on the glass-forming liquid ortho-terphenyl over a range of temperatures and pressures. In the left panel different estimates of $\chi_4$ are plotted as a function of relaxation time while in the right panel data for relaxation time 100 s is plotted against pressure. Note that while it is in principle necessary to express the relaxation time in reduced units (rescaling by appropriate powers of temperature and density), at such viscous states the density and temperature change so little that the difference is negligible. In the right panel the lowest data set (circles) gives the estimation of the $\chi_4^{NVT}$ from Eq. (14), which is fairly constant as pressure is varied along the isochrone, while the stars represent the third term in Eq. (10) ($\rho$ term) which decreases as pressure (and density) increases, corresponding to the increase of the reduced bulk modulus. The top curve is the sum of the other two, representing the experimental estimate of $\chi_4^{NPT}$ via Eq. (15). It inherits the variation of the bulk modulus. Data for the latter (not shown) show a variation sufficient to

$$
\left( \frac{\partial \ln T}{\partial \ln \rho} \right)_X = -\frac{\rho}{\partial \ln \rho} T
$$

and note [Eq. (2)] that the left-hand side is just $\gamma$ (constant $X$ being the same as constant $s_{ex}$), thus

$$
\left( \frac{\partial X}{\partial \ln \rho} \right)_T = -\gamma \left( \frac{\partial X}{\partial \ln T} \right)_\rho
$$

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$$

$$
\chi_4^{NPT} \simeq \frac{1}{c_V/k_B} \left( \frac{\partial C(t)}{\partial \ln T} \right)_\rho^2 + \frac{\rho k_B T}{K_T} \left( \frac{\partial C(t)}{\partial \ln \rho} \right)_T^2 \tag{varies}, \tag{15}
$$

Thus isomorph theory predicts that there will be little variation along the isochrone when the $NVT$ version of $\chi_4$ is considered, so the latter is an allowed candidate for a quantity which controls the dynamics.
explain the variation here. The change of the estimate of $\chi_4^{\text{NPT}}$ is $-24\%$; the change of the estimate of $\chi_4^{\text{NVE}}$ is $-8\%$. Note that in both cases the term representing $\chi_0^{\text{NVE}}$ is missing, so the true percentage change for $\chi_4^{\text{NPT}}$ will be smaller. The $8\%$ change in the NVT estimate can be seen as a true deviation from isomorph invariance, since this quantity is formally isomorph invariant.

In contrast to Refs. [33–35], Alba-Simionesco et al. found an increase of the maximum of $\chi_4^{\text{NPT}}$ with increasing pressure for the liquid dibutyl-phthalate [36]. In this case the change in the reduced bulk modulus turns out to be relatively small, while there is an unusually large change in $\gamma$ (denoted $x$ in that work), from 2.5 to 4, as density increases, thus the increase in $\chi_4^{\text{NPT}}$. The authors presented these results as being in contradiction to the RFOF theory, but the implication that there is a problem with the theory is not necessarily valid, given the use of a non-isomorph-invariant definition of $\chi_4$.

V. SIMULATIONS

We have investigated the different contributions to $\chi_4$ in simulations. The important results are that the NVT estimate is invariant along an isochrone while the NPT estimator decreases, as expected. When the NVE contribution is included (which is not small), we find a slight increase in full NVT quantity. Thus the true deviation from (otherwise expected) isomorph invariance in this case has the opposite sign to what is seen in the NPT estimator.

Simulations were carried out on a binary Lennard-Jones system of 1024 particles using the Kob-Andersen potential parameters [51] and usual composition of 80\% large particles; technical details are given in Appendix A. A range of densities starting at the usual 1.2 and going up to 1.65 was simulated. Both NVE and NVT dynamics were used, but the thermostat relaxation time for the latter was chosen to be at each state point a few times the alpha relaxation time to give approximately constant energy during microscopic processes, to better correspond to the case of an ensemble of NVE trajectories as discussed in Sec. III. In addition it was held constant in reduced units along isochrones.

Since an isomorph is an idealized concept which formally is both an isochrone and a (configurational) adiabat, while in practice isochrones and adiabats do not exactly coincide, we have considered both an isochrone and an adiabat. To identify an isochrone we used the method of parametrizing the density scaling function $h(\rho)$ appearing in Eq. (4) that was used in Ref. [50]. It is formulated in terms of a reference density $\rho_*$, with respect to which the relative density $\tilde{\rho} \equiv \rho/\rho_*$ is defined:

$$h(\tilde{\rho}) = \rho^4 (\gamma_* (2 - 1) - \rho^2 (\gamma_* / 2 - 2)).$$

Here $\gamma_*$ is the scaling exponent at the reference density. For perfect isomorphs it should equal the fluctuation-based scaling exponent at $\rho_*$; in Ref. [50] a value was chosen which gave a good collapse of the relaxation time data over a broad range of density and temperature: For reference density $\rho_* = 1.6$, the value $\gamma_* = 4.59$ was used. This can be used to generate a set of $\rho$, $T$ values which constitutes an approximate isomorph, but it does not take into account the temperature dependence of $\gamma$. The value 4.59 was determined by a procedure which gives weight to more viscous state points; here we study less viscous state points due to the need for extra long runs for good $\chi_4$ statistics. The value of $\gamma$ observed from the fluctuations at density 1.6 is closer to 4.57; we find instead that we get a more exact isochrone if we generate the state points using this value. Note that this is a 0.5\% difference in $\gamma$; it is only relevant when trying to identify good isochrones on a linear (rather than logarithmic) time axis.

Figure 2 shows data for an isochrone containing the point $\rho = 1.2, T = 0.55$. In the main panel the self-part of the intermediate scattering function is plotted for densities between 1.2 and 1.65 [52]. The inset shows the relaxation time; it increases slightly, about 2\% over this range of densities. We have also simulated a configurational adiabat, identified as the curve whose logarithmic derivative ($\partial \ln T / \partial \ln \rho$)$_x$ is given by the fluctuation expression for $\gamma$ [Eq. (6)] at each state point. The procedure used was the same as in Ref. [45]: steps of 0.05 in density were taken, and Eq. (16) was re-parametrized at each step using the observed fluctuations [Eq. (6)]. For a system with perfect isomorphs these curves should be identical; for a real system they are close but not identical. A single parametrization of Eq. (16) using a relatively high reference density such as 1.6 (as opposed to re-parametrizing at each step in density) can generate a good isochrone over the whole simulated density range. But the observed $\gamma$ from fluctuations deviates from the that given by Eq. (16) as the density decreases (at density 1.20 the observed $\gamma$ is approximately 5.16 while the logarithmic derivative of $h(\tilde{\rho})$, parametrized by $\gamma_*(\rho = 1.60) = 4.57$, is closer to 5.30). This indicates a deviation of the adiabat from the isochrone. It is intriguing that a single parametrization of Eq. (16)—that is, determined by a
FIG. 2. (Color online) Self-intermediate scattering function for the Kob-Andersen system (large particles) for several state points along an isochrone with lowest density 1.20 (temperature 0.55), plotted against reduced time $\tilde{t}$. The inset shows the relaxation time $\tilde{\tau}$ defined as the time when the correlation function has fallen to $1/e$ along to curves: a isochrone (squares) determined by Eq. (16) parametrized by matching its logarithmic derivative to Eq. (6) for each step in density of size 0.05. The isochrone and adiabat share the state point ($\rho = 1.20, T = 0.55$). Lines are to guide the eyes.

A single value of $\chi_4$, at a given reference density, and used over the whole density range—generates a better isochrone than adiabat. On the latter the reduced relaxation time increases by about 10% over the range of densities considered here (inset to Fig. 2).

Figure 3 shows $\chi_4$ for the approximate isochrone in the NVE and NVT ensembles, the T term (or NVT estimator) and the $\rho$ term (which in combination with the T term gives the NPT estimator). Reduced units for time are used. For the first three there is an approximate collapse, while for the $\rho$ term there is clearly no collapse. This is in accordance with the expectation that the first three quantities are formally isomorph invariant while the fourth is not, while revealing that there is some variation even for the formally invariant quantities. A clearer idea of the trends as a function of density is given by considering the peak value of each $\chi_4(t)$ curve. These are shown in Fig. 4 for the isochrone and in Fig. 5 for the adiabat. Also shown are the maxima of the true NPT quantity obtained by adding the $\rho$ term to the NVT quantity. We have confidence that this is a good estimate; we have explicitly confirmed the decomposition of the NVT quantity, Eq. (8) (see Appendix A).

It can be seen that there appears, in fact, to be a slight increase in the NVE and NVT $\chi_4$. On the other hand, the estimator of the NVT value by the temperature derivative is quite flat, as invariant as the correlation function itself, while the estimator of the NPT value appears to decrease with increasing density, as expected from the behavior of...
the bulk modulus. Since the bulk modulus is not formally isomorph invariant, this decrease is not surprising. The deviation seen in the NVE or full NVT quantity is interesting because it represents a true deviation from isomorph invariance in a formally isomorph-invariant quantity. Unfortunately it does not appear in the experimentally accessible quantities. Ironically, when the full NPT quantity is considered, it is in practice perhaps the most invariant quantity due to a small percentage increase in the NVE and NVT quantities (a true deviation from isomorph invariance) and a large percentage decrease in the $\rho$ term (expected due to the bulk modulus). Since the latter term is relatively small the absolute changes tend to cancel, leaving the true NPT quantity quite flat.

VI. DISCUSSION

A. Interpreting the extra contributions (T term and $\rho$ term)

One must be careful not to over-interpret the different effects of pressure (or density) and temperature on the dynamics. The dynamics can only be said to depend on an isomorph invariant, for example, the excess entropy $s_{\text{ex}}$. To show explicitly how this can be represented in the contributions to $\chi_4$, we rewrite Eq. (10) in terms of derivatives with respect to the alternative variables $s_{\text{ex}}$ and $\rho$ using the chain rule and standard thermodynamic identities:

$$\chi_4^{\text{NPT}} - \chi_4^{\text{NVE}} = \left(\frac{c_{\text{v}}^4}{c_{\text{v}}}\right)^2 \left(1 + \frac{\rho T}{K_T} \frac{\partial \rho}{\partial s_{\text{ex}}} \right) \frac{\partial \rho}{\partial s_{\text{ex}}} + \frac{\rho T}{K_T} \frac{\partial \rho}{\partial s_{\text{ex}}}^{2} .$$

Note there is a crossterm involving both derivatives in this expression. For a general choice of independent variables there will be such a term; the choice of temperature and density (or volume) is special because these variables are statistically independent as is known from thermodynamic fluctuation theory [53]. We can interpret the coefficient of the first term by replacing $C$ with $s_{\text{ex}}$ (since the above expression is just a Lebowitz-type formula [48,49] with an extra factor $N$, we can certainly apply it to ordinary static quantities as well). Since the excess entropy is not a dynamical variable, its variance in the NVE ensemble is zero. Evaluating Eq. (17) gives the coefficient,

$$\left(\frac{c_{\text{v}}^4}{c_{\text{v}}}\right)^2 \left(1 + \frac{\rho T}{K_T} \frac{\partial \rho}{\partial s_{\text{ex}}} \right) \frac{\partial \rho}{\partial s_{\text{ex}}} \equiv \left(\frac{c_{\text{v}}^4}{c_{\text{v}}}\right)^2 \left(1 + R \right) \frac{\partial \rho}{\partial s_{\text{ex}}} \equiv N((\Delta s_{\text{ex}})^2)_{\text{NPT}} .$$

Thus this coefficient can be interpreted as the variance of excess entropy among the different NVE members of the NPT ensemble (note that it is not isomorph invariant due to the presence of $K_T$). The quantity $R \equiv \gamma^2 c_{\text{v}} / K_T \equiv \gamma^2 c_{\text{v}} \rho k_B T / K_T$ was also defined by Dalle-Ferrier et al. [11,36] to quantify the relative importance of density fluctuations compared to energy fluctuations on the dynamics. Here we see that more precisely it quantifies their relative contributions to fluctuations of excess entropy. Returning to $\chi_4$, in the case of an R liquid where $C(t)$ depends only on $s_{\text{ex}}$, we get

$$\chi_4^{\text{NPT}} - \chi_4^{\text{NVE}} = \left(\frac{c_{\text{v}}^4}{c_{\text{v}}}\right)^2 \left(1 + R \right) \frac{d \rho}{d s_{\text{ex}}} \left(\frac{d \rho}{d s_{\text{ex}}} \right)^2 \equiv N((\Delta s_{\text{ex}})^2)_{\text{NPT}} \frac{d \rho}{d s_{\text{ex}}} \left(\frac{d \rho}{d s_{\text{ex}}} \right)^2 .$$

Thus the variation of the dynamical quantity $\chi_4^{\text{NPT}}$ along an isomorph is, for an R liquid, due to the thermodynamic fact that a contour of $s_{\text{ex}}$ is not a contour of the NPT variance of $s_{\text{ex}}$. Another way to interpret this is to consider the NPT ensemble of NVE trajectories at each point along an isomorph. Members whose energy and volume are the mean energy and volume experience dynamics isomorphic with corresponding members elsewhere on the isomorph. Other members have energy and volume near the mean values and will statistically be very similar to those on the mean. But the spread of volume will vary along the isomorph: The ensemble will be narrower at high densities because the reduced bulk modulus typically increases with density. This is the sense in which the dynamical susceptibility is not invariant. Again we emphasize that the variation represented by the terms in a Lebowitz-type formula is variation of initial conditions, and not dynamical fluctuations.

B. Ensemble-independent dynamic susceptibility

We consider now an issue raised by the argument of Flenner and Szamel that the NPT version of $\chi_4$ is the correct one (for a pure substance; for a mixture extra terms accounting for concentration terms should be included) in the following sense [12,54]: Consider the four-correlation function $S_4(k,t)$ where $k$ should be distinguished from the wave number $q$ appearing in the self-intermediate scattering function. It denotes the wave vector associated with the Fourier transform of the correlator with respect to initial position (see Appendix B for details of our definition of $S_4$). At finite wave number $k$ the four-point correlation $S_4(k,t)$ or any wave-number-dependent response function is ensemble independent because the boundaries only couple to $k = 0$ behavior—alternatively measuring at finite wave numbers corresponds to sampling subsets of the system whose density, energy, and concentration can fluctuate while their mean values are fixed. Taking the $k \to 0$ limit after the thermodynamic limit gives therefore an ensemble-independent quantity. This raises the interesting question of the isomorph invariance of $S_4(k,t)$ at finite $k$. Its invariance in practice can be investigated in simulations, but what does the theory say—is $S_4(k,t)$ even formally isomorph invariant? Answering this is not straightforward, but consideration of it suggests how the theory of isomorphs can predict its own limit of validity. In particular, a reconciliation of the noninvariance of the bulk modulus and the idea of an ensemble-independent definition of $\chi_4$ suggests that isomorph invariance may somehow break down at long length scales.

Consideration of the term $g(\rho)$ in the approximate potential energy Eq. (3), which is responsible for the noninvariance of the bulk modulus, suggests how this might occur. This term is nonlocal, whereas the true potential energy is local. A way to restore locality to the approximation would be to replace $g(\rho)$ with a sum over particles $\sum_i g(\rho_{CG,i})$. Here $\rho_{CG,i}$ is a local density evaluated at the position of particle $i$, and
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Involving a coarse-graining length $l_{CG}$. For a sufficiently large $l_{CG}$ this would make no difference to the hitherto documented isomorph-invariant quantities, since these have all involved dynamics at short to moderate length scales. With such a local representation of potential energy for an R liquid we can predict that wave-number-dependent response functions (at least those with a longitudinal component) such as $s_4(k,t)$ are formally isomorph invariant only for $k$ larger than a crossover value of order $1/l_{CG}$, while for smaller $k$ we can expect deviations.

This line of reasoning has two consequences. First, it turns the discussion about which ensemble is relevant into one about which length scales are relevant. Second, the principle mentioned at the end of Sec. II states that only isomorph-invariant quantities can be relevant for determining isomorph-invariant measures of dynamics. This therefore suggests that density fluctuations and dynamical fluctuations at long wavelengths are not relevant for the usual measures of dynamics which involve relatively small wavelengths of order a few particle spacings. For a measure of dynamical heterogeneity satisfying the requirements of both ensemble independence and isomorph invariance one should consider $s_4(k,t)$ at some finite wave number $k$, which should be small enough to capture what correlated dynamics exists, but not arbitrarily small. This is tested in Fig. 6, which includes data for a system whose linear size is a factor of two larger ($N = 8192$). Little variation is evident: a slight decrease at the largest two wavelengths. If we assume that the terms accounting for concentration fluctuations are at least as invariant as the T term, then this is consistent with the fact that the $\chi_4^{NPT}$ maximum also varies little (Fig. 4). In that case the slope is positive, but the data are perhaps not precise enough to distinguish a change of +2.5% from one of −1.4%. Such little variation would normally be considered consistent with isomorph invariance, but the interpretation here becomes tricky: We already argued that the apparent invariance of the $NPT$ quantity is due to cancellation of a true violation of isomorph invariance in the $NVT$ quantity and the expected variation of the $\rho$ term. It is therefore not clear what can be concluded from these $s_4$ data. Investigation of a wider range of $k$ values (system sizes), temperatures (both more and less supercooled), and systems (including non-Roskilde liquids) is needed to resolve this issue.

C. Open questions

The analysis and discussion presented so far give rise to some open questions.

1. Is the cancellation of terms that leads to an almost invariant $\chi_4^{NPT}$ an accident or does it represent something deeper going on? At more viscous state points the T and $\rho$ terms are expected to account for a larger fraction of $\chi_4^{NPT}$, so the cancellation should be less effective. We see more or less the same behavior, however, at the lowest simulated temperatures (0.48 for the usual density 1.2; data not shown). As mentioned above, studies of less viscous state points and non-R systems are needed to clarify the general picture.

2. The line of reasoning about long-wavelength fluctuations suggests that the static structure factor $S(q)$ may show deviations from isomorph invariance at sufficiently small $q$ (this is also expected from the noninvariance of the bulk modulus). This should be investigated using large systems. Note that no deviations have been found in the (coherent) intermediate scattering function at small $q$: Veldhorst et al. studied the intermediate scattering function in a polymer model and found invariance at all wave numbers [55]. Extensive testing of this has yet to be done, however.

3. The definition of $s_4$ employed here (see Appendix B) includes contributions where the two wave vectors ($k$ and $q$) are parallel with each and ones where they are perpendicular. A decomposition into terms where the vectors $k$ and $q$ are parallel versus perpendicular, as done by Flenner et al. [22], might show interesting differences regarding isomorph invariance. In particular one could speculate, by analogy with the bulk versus shear modulus, that the longitudinal (parallel) case might show greater deviations from isomorph invariance than the transverse (perpendicular) case. Similarly the wave-number-dependent bulk and shear viscosities would be worth investigating at small $q$. Further theoretical work is required to elucidate the question of which wave-vector-dependent quantities, if any, are expected to be invariant at low $k$.

4. Another formally invariant quantity that increases slightly along an isomorph (adiabat) for Lennard-Jones liquids is $C_V$ [45]. In Ref. [45] it was shown that whether $C_V$ increases or decreases depends on a certain feature of the pair potential and one can speculate that $\chi_4^{NPT}$'s behavior depends similarly on the potential. While we have no theoretical argument for this, it is straightforward to check empirically.

VII. CONCLUSION

We have analyzed the different terms in the decomposition (10) which is the basis of experimental estimations of $\chi_4$
in pure substances, with a view to determining their formal isomorph invariance. We find that the \( \rho \) term, which accounts for volume fluctuations, spoils isomorph invariance due to its containing the bulk modulus, which is known to not be formally isomorph invariant. The fact that \( \chi^\text{NPT}_4 \) is not formally invariant has implications both for isomorph theory and for the physics of glass-forming liquids: Isomorph invariance cannot be expected to hold for dynamics at arbitrarily long wavelengths (take the bulk sound velocity, for example) while the usual measures of liquid dynamics can therefore not depend, or be controlled by, measures of long-wavelength fluctuations in the dynamics.

Data from simulations confirm the noninvariance of the \( \rho \) term, but also provide a means to check the \( \text{NVE} \) contribution which is not usually accessible experimentally. Interestingly, it, too, shows a variation, which tends to cancel that due to the \( \rho \) term. Thus the true \( \text{NPT} \) quantity turns out actually to be quite invariant, although theory does not predict it to be so.

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**APPENDIX A: TECHNICAL DETAILS OF SIMULATIONS**

A “shifted forces” cutoff of 2.5\( \sigma \) was used [56]. The software was RUMD [57] running on nVidia graphical processing units. The time step was chosen such that its reduced value was the same for all points along the isomorph; for \( \rho = 1.2, T = 0.55 \) the value was 0.005 in ordinary molecular dynamics units (based on the Lennard-Jones length and energy parameters for the large particles). For the runs where \( \chi_4 \) was calculated directly, the simulation run length corresponded to at least \( 10^4 \) relaxation times. For the runs at neighboring densities and temperatures, in order to numerically differentiate the correlation function, it was \( 10^5 \) relaxation times. The \( q \) value was chosen equal to that used by Kob and Andersen (7.25 for large particles) at density 1.20, but scaled proportional to \( \rho^{1/3} \) for other densities: \( q_\rho = q_{1.2}(\rho/1.2)^{1/3} \). Numerical differentiation was carried out at fixed \( q \) and \( t \) (i.e., in ordinary rather than reduced units). The relaxation time for the thermostat was chosen to be fixed in reduced units, and as mentioned above, at least a few times the relaxation time. Under these circumstances the energy is effectively constant on the time scale of relaxation processes, and the premise of Eq. (8), namely a canonical distribution of fixed-energy trajectories, is realized. See Fig. 7 for an illustration of the effect of thermostat relaxation time.

**APPENDIX B: DEFINITIONS OF 2- AND 4-POINT CORRELATION FUNCTIONS**

To give a precise definition of the two-point and four-point correlators we use, we start by defining a single-particle, two-time quantity \( f_{q,i}(t) \):

\[
f_{q,i}(t) = \frac{1}{2} \left( \cos(q \Delta x_i(t)) + \cos(q \Delta y_i(t)) + \cos(q \Delta z_i(t)) \right).
\]

(B1)

\[
F_{s}(q,t) = (F_{s}(q,t)) = \frac{1}{N_A} \left\langle \sum_{i} f_{q,i}(t) \right\rangle.
\]

(B3)

The variance of the correlator \( F_{s} \), multiplied by \( N \) (the total number of particles) is the dynamic susceptibility, as described in the main text. We note that at long times when particle positions have de-correlated from their initial values the variance is not zero but \( \frac{N_A^2}{N} \frac{1}{2} \langle \cos^2 \rangle \) which is 0.21 for the composition \( N_A/N = 0.8 \). This can be seen in Figs. 3(a) and 3(b), and differs from the case where the dynamical correlator is defined using an overlap function instead of a cosine.

In order to define the four-point function \( S_{4} \) we first define a quantity \( \rho(k,q,t) \) as the Fourier transform with respect to initial positions (x coordinate) of correlator,

\[
\rho(k,q,t) = \frac{1}{N_A} \sum_{i} f_{q,i}(t) \exp(i k x_i(0)).
\]

(B4)

(note that for \( t \to 0 \), any \( q \) this is just Fourier mode \( k \) of the density of A particles). Finally the four-point correlation function \( S_{4}(k,t) \) (suppressing the \( q \) dependence for more convenient notation) as

\[
S_{4}(k,t) \equiv N \langle \rho(k,q,t) \rho(-k,q,t) \rangle - \langle F_{s}(q,t) \rangle^2 \delta_{k,0}.
\]

(B5)
Note that $N$ is the total number of particles of all types. Setting $k = 0$ in this definition gives (the ensemble-dependent) $\chi_4$ as defined in the text; taking the $k \to 0$ limit after the thermodynamic limit gives an ensemble-independent quantity.