Pressure-energy correlations in liquids. V. Isomorphs in generalized Lennard-Jones systems

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(Received 14 October 2010; accepted 6 April 2011; published online 28 April 2011)

This series of papers is devoted to identifying and explaining the properties of strongly correlating liquids, i.e., liquids with more than 90% correlation between their virial W and potential energy U fluctuations in the NVT ensemble. Paper IV [N. Gnan et al., J. Chem. Phys. 131, 234504 (2009)] showed that strongly correlating liquids have “isomorphs,” which are curves in the phase diagram along which structure, dynamics, and some thermodynamic properties are invariant in reduced units. In the present paper, using the fact that reduced-unit radial distribution functions are isomorph invariant, we derive an expression for the shapes of isomorphs in the WU phase diagram of generalized Lennard-Jones systems of one or more types of particles. The isomorph shape depends only on the Lennard-Jones exponents; thus all isomorphs of standard Lennard-Jones systems (with exponents 12 and 6) can be scaled onto a single curve. Two applications are given. One tests the prediction that the solid-liquid coexistence curve follows an isomorph by comparing to recent simulations by Ahmed and Sadus [J. Chem. Phys. 131, 174504 (2009)]. Excellent agreement is found on the liquid side of the coexistence curve, whereas the agreement is less convincing on the solid side. A second application is the derivation of an approximate equation of state for generalized Lennard-Jones systems by combining the isomorph theory with the Rosenfeld-Tarazona expression for the temperature dependence of the potential energy on isochores. It is shown that the new equation of state agrees well with simulations. © 2011 American Institute of Physics. [doi:10.1063/1.3582900]

I. INTRODUCTION

This is the final paper in a series1–4 investigating the properties of strongly correlating liquids,5 i.e., liquids that have strong correlations between their constant-volume equilibrium fluctuations of potential energy, \( U(t) \), and virial6,7 \( W(t) \equiv -1/3 \sum_i \mathbf{r}_i \cdot \nabla_i U(\mathbf{r}_1,...,\mathbf{r}_N) \) where \( \mathbf{r}_i \) is the position of particle \( i \) at time \( t \). As is well known, the average virial \( W \) gives the configurational contribution to the pressure:

\[
pV = Nk_BT + W. \tag{1}
\]

Letting \( \Delta \) denote instantaneous deviations from equilibrium mean values, the \( WU \) correlation is quantified by the correlation coefficient \( R \) (with \( \langle \ldots \rangle \) denoting equilibrium average):

\[
R = \frac{\langle \Delta W \Delta U \rangle}{\sqrt{\langle \Delta W^2 \rangle \langle \Delta U^2 \rangle}}. \tag{2}
\]

Perfect correlation gives \( R = 1 \). As a pragmatic definition we have chosen “strongly correlating liquids” to designate liquids that have \( R \geq 0.9 \) in the \( NVT \) ensemble (constant volume, \( V \), and temperature, \( T \)).

Strongly correlating liquids have simpler physics than liquids in general, an observation that has particular significance for the highly viscous phase.8–18 Thus it has been shown that strongly correlating viscous liquids to a good approximation have all eight frequency-dependent thermoviscoelastic response functions19–21 given in terms of just one22 (i.e., are single-parameter liquids in the sense of having dynamic Prigogine-Defay ratio19 close to unity,2,20,22). Strongly correlating viscous liquids moreover obey density scaling23–27 to a good approximation, i.e., their dimensionless relaxation time \( \tilde{\tau} \equiv \tau \rho^{1/3} \sqrt{k_BT/m} \) (where \( m \) is the average particle mass) depends on density \( \rho = N/V \) and temperature as \( \tilde{\tau} = F(\rho^\gamma/T)^{26-30} \).

Paper I3 presented computer simulations of 13 different systems, showing that van der Waals type liquids are strongly correlating, whereas hydrogen-bonding liquids like methanol or water are not. Strongly correlating liquids include1,2,5,22,29 for instance, the standard Lennard-Jones (LJ) liquid, the Kob-Andersen binary LJ (KABLJ) mixture, an asymmetric rigid-bond dumbbell model, a seven-site united-atom toluene model, and the Lewis-Wahnström OTP model. Paper II2 analyzed the cause of \( WU \) correlations with a focus on the LJ potential. The strong correlations were related to the well-known fact that an inverse power-law (IPL) pair potential, \( v(r) \propto r^{-n} \) where \( r \) is the distance between two particles,31–42 implies perfect \( WU \) correlation,

\[
\Delta W(t) = \gamma \Delta U(t) \tag{3}
\]
with \( \gamma = n/3 \). Around the potential energy minimum, the LJ potential is well described by an “extended” inverse power-law potential (eIPL), \( v_{\text{IPL}}(r) \equiv Ar^{-\gamma} + B + Cr. \) At constant volume the linear term contributes little to the virial and potential-energy fluctuations: when one nearest-neighbor distance increases, others decrease in such a way that the sum is almost constant. Thus systems interacting via the LJ potential inherit strong WU correlations from an underlying inverse power-law—they have a “hidden scale invariance.”3,28

Paper III4 gave further numerical evidence for the explanation for strong WU correlations presented in Paper II, and theoretical results were given on the statistical mechanics and thermodynamics of the hidden scale invariance that characterizes strongly correlating liquids. It was also shown that strong virial potential-energy correlations are present even in out-of-equilibrium situations—the hidden scale invariance is a property of the potential energy surface, not just of the equilibrium states.

Paper IV4 introduced the concept of “isomorphs” in the phase diagram of a strongly correlating liquid. Starting from a single assumption a number of isomorph invariants were derived. In particular, structure and dynamics were shown to be invariant on isomorphs when reduced units are used.

In the present paper further simulation results supporting the isomorph predictions are presented for systems interacting via a multicomponent generalized LJ potential:

\[
v_{ij}(r_{ij}) = v^{(m)}_{ij}(r_{ij}) + v^{(n)}_{ij}(r_{ij}),
\]

where \( v^{(k)}_{ij}(r_{ij}) \) is an IPL potential involving the two particles \( i \) and \( j \):

\[
v^{(k)}_{ij}(r_{ij}) \equiv \varepsilon_{ij}(\sigma_{ij}/r_{ij})^k.
\]

Section II briefly reviews the isomorph concept. For systems interacting via a generalized LJ potential, a prediction for the shape of the isomorphs in the WU phase diagram is derived in Sec. III and demonstrated to fit well to simulation results.43 Interestingly, the isomorph shape depends only on the exponents \( m \) and \( n \). Thus, e.g., all 12-6 LJ systems have the same isomorphs in the WU phase diagram. In Sec. IV we present two applications of the theory: One tests the Paper IV prediction that solid-liquid coexistence curves are isomorphs. The second application gives an approximate equation of state for systems interacting via generalized LJ potentials; this is arrived at by combining the present theory with Rosenfeld and Tarazona’s expression for the isochoric temperature dependence of the potential energy.

II. ISOMORPHS

A. Definition

We term a microscopic configuration “physically relevant” if its influence on the thermodynamics and dynamics of the system is not a priori negligible (Paper IV). For instance, any configuration with very strong particle overlap is physically irrelevant. Note, however, that some unlikely configurations like transition states are physically relevant.

Two state points (1) and (2) with temperatures \( T_1 \) and \( T_2 \) and densities \( \rho_1 \) and \( \rho_2 \), respectively, are defined to be isomorphic (Paper IV) if they obey the following: any two physically relevant configurations of state points (1) and (2), \( (\mathbf{r}_1^{(1)}, ..., \mathbf{r}_N^{(1)}) \) and \( (\mathbf{r}_1^{(2)}, ..., \mathbf{r}_N^{(2)}) \), which trivially scale into one another,

\[
\rho_1^{1/3} \mathbf{r}_i^{(1)} = \rho_2^{1/3} \mathbf{r}_i^{(2)} (i = 1, ..., N),
\]

have proportional configurational Boltzmann statistical weights:

\[
e^{-U(\mathbf{r}_1^{(1)}..., \mathbf{r}_N^{(1)})/k_B T_1} = C_{12} e^{-U(\mathbf{r}_1^{(2)}..., \mathbf{r}_N^{(2)})/k_B T_2}.
\]

Here \( U(\mathbf{r}_1, ..., \mathbf{r}_N) \) is the potential energy function and it is understood that the constant \( C_{12} \) depends only on the state points (1) and (2).

The property of being isomorphic defines a mathematical equivalence relation on the set of state points. The corresponding equivalence classes are smooth curves in the phase diagram termed isomorphs or isomorphic curves.

B. The approximate nature of isomorphs

Equation (7) implies identity of the normalized Boltzmann probabilities for scaled physically relevant configurations of isomorphic state points. As detailed in Paper IV this identity implies that several quantities are invariant along an isomorph. Examples include the configurational entropy, the isochoric specific heat, all multiparticle distribution functions in reduced units (in particular, the radial distribution function(s)), reduced-unit dynamics (both for Newtonian and stochastic dynamics), normalized reduced-time autocorrelation functions, reduced-unit transport coefficients, etc. It was further shown in Paper IV that a jump from equilibrium at one state point to an isomorphic state point brings the system instantaneously to equilibrium (see also Ref. 44).

IPL potentials are Euler homogeneous functions, i.e., obey \( U(\lambda \mathbf{r}_1, ..., \lambda \mathbf{r}_N) = \lambda^{-\gamma} U(\mathbf{r}_1, ..., \mathbf{r}_N) \). Such purely repulsive potentials – which do not describe real intermolecular interactions – are the only potentials that have 100% correlation between virial and potential-energy fluctuations; this reflects the well-known IPL virial-theorem identity \( W = (n/3)U \). Likewise, only IPL liquids have exact isomorphs; they satisfy Eq. (7) with \( C_{12} = 1 \) for all configurations of two state points obeying \( \rho_1^2/T_1 = \rho_2^2/T_2 \) where \( \gamma = n/3 \).

Appendix A of Paper IV showed that strongly correlating liquids generally have isomorphs to a good approximation. Consequently, to a good approximation these liquids inherit a number of the exact invariants that IPL liquids have along their curves given by \( \rho^\gamma/T = \text{Const.} \), namely, all IPL invariants that do not rely on the IPL identity \( C_{12} = 1 \). Examples include the above mentioned thermodynamic, static, and dynamic isomorph invariants. IPL invariants that are not inherited by strongly correlating liquids in general include, e.g., the Helmholtz free energy over temperature, the potential energy over temperature, and the reduced-unit compressibility.

The fact that strongly correlating liquids have isomorphs to a good approximation should be understood as follows.
Any liquid has curves in its phase diagram of constant excess entropy, curves of constant isochoric specific heat, curves of constant reduced relaxation time, curves of constant reduced-unit transport coefficients, etc. For a strongly correlating liquid these curves are almost identical, and they define its (approximate) isomorphs. Points on these isomorphs have approximately identical structure and approximately identical dynamics as probed, e.g., by normalized reduced-time-autocorrelation functions.

For clarity of presentation below we shall not repeatedly mention that the existence of isomorphs is not exact for generalized LJ systems. Accordingly, we shall speak of isomorphs as a unique, well-defined concept. In other words: all properties of isomorphs derived below in generalized LJ systems are in principle – as well as practice – approximate.

C. Generating isomorphs in simulations

The structure in reduced units ($\bar{r} \equiv \rho^{1/3} r$) is predicted to be invariant along an isomorph (Paper IV). The excess entropy, $S_{\text{ex}} = S - S_{\text{ideal}}$, depends only on structure and was shown in Paper IV also to be invariant along an isomorph ("excess" refers to the quantity in question in excess to that of an ideal gas with same density and temperature). In the below simulations we generate an isomorph as a set of state points in the phase diagram with constant $S_{\text{ex}}$. To change density and temperature keeping $S_{\text{ex}}$ constant, the following identity is used (Paper IV):

$$\gamma \equiv - \left( \frac{\partial \ln T}{\partial \ln V} \right)_{S_{\text{ex}}} = \frac{V}{T} \left( \frac{\partial S_{\text{ex}}}{\partial V} \right)_T = \frac{V \beta_{\text{ex}}^T}{C_{\text{ex}}^V},$$

where $\beta_{\text{ex}}^T \equiv (\partial p_{\text{ex}}/\partial T)_V = \frac{1}{\rho} (\partial W/\partial T)_V$ and a Maxwell relation was applied. Utilizing the standard fluctuation formulæ this leads (Paper IV) to:

$$\gamma = \left( \frac{\partial \ln T}{\partial \ln \rho} \right)_{S_{\text{ex}}} = \frac{\langle \Delta U \Delta W \rangle}{\langle \Delta U \rangle^2}.$$  

The procedure applied to generate isomorphs can be summarized as: (1) an equilibrium $NVT$ simulation was performed at a given state point; (2) $\gamma$ was calculated from the fluctuations using Eq. (9); (3) the density $\rho$ was changed slightly (of order 1%), and $\gamma$ was used to calculate the corresponding change in temperature in order to keep $S_{\text{ex}}$ constant (Eq. (9)); 4) a simulation at the new state point was performed, and the procedure was repeated.
KABLJ mixture. For comparison, Fig. 1(b) shows the same radial distribution function for the large (A) particles of the energy surface in reduced units (\(\tilde{\tau}_a\)) for four different binary generalized LJ mixtures.

The insignificant change of \(\gamma\) along isochors (black circles in Fig. 3) means that these to a good approximation are straight lines in the \(WU\) phase diagram. This is illustrated in Fig. 4 for four different binary generalized LJ mixtures.
This implies $U_m > 0$ for all configurations, and therefore also for all thermodynamic averages. Combining this with Eq. (14) implies $W > \frac{\pi}{4} U$. If all interactions have an attractive part ($\epsilon_{ij}^{(n)} < 0$) we have $U_n < 0$, which in combination with Eq. (15) implies $W > \frac{m}{3} U$. As an example, the region of the $WU$ phase diagram accessible to the standard 12-6 LJ potential is given by $W > 4U$ and $W > 2U$ (the later inequality being relevant when $U \leq 0$). See Fig. 5 for an illustration.

A. Parametric description of isomorphs

Along an isomorph the structure is invariant in reduced units (Fig. 1). From this it follows that $U_k / \rho^{k/3}$ is invariant along an isomorph ($\tilde{\rho} \equiv \rho / \rho_*$):

$$\frac{U_k}{\rho^{k/3}} = \left( \sum_{i,j} \frac{\epsilon_{ij}^{(k)}}{\epsilon_{ij}} \right)^k \tilde{\rho}^k = \left( \sum_{i,j} \tilde{\epsilon}_{ij}^{(k)} \right)^k \tilde{\rho}^k,$$

(16)

$$= \sum_{i,j} \epsilon_{ij}^{(k)} \left( \frac{\rho^{k/3}}{\rho_*} \right)^k \tilde{\rho}^{k/3} U_k^*.\quad (17)$$

If we let “$\ast$” denote a reference point, $(W_*, U_*)$, we can use Eqs. (14) and (15) to get $(U_m^*, U_n^*)$ and find for other state points on the same isomorph (where $\tilde{\rho} \equiv \rho / \rho_*$):

$$U_k = \left( \frac{\rho}{\rho_*} \right)^{k/3} U_k^* = \tilde{\rho}^{k/3} U_k^*.\quad (18)$$

Combining Eq. (18) with Eqs. (11) and (13) we obtain a parametric description of an isomorph in the $WU$ phase diagram:

$$U = \tilde{\rho}^{m/3} U_m^* + \tilde{\rho}^{n/3} U_n^*, \quad (19)$$

Once the numbers $U_m^*$ and $U_n^*$ have been determined at the reference state point using Eqs. (16) and (17), the entire isomorph to which this state point belongs is thus traced out in the $WU$ phase diagram using Eqs. (18) and (19). Alternatively, given a collection of isomorphic state point generated, e.g., as described in Sec. II C, the predicted relation between $U$, $W$, and $\tilde{\rho}$ can be tested by linear regression:

$$U = \tilde{\rho}^{m/3} U_m^* + \frac{n}{3} \tilde{\rho}^{n/3} U_n^*.\quad (20)$$

This is done in Fig. 6 for the 12-6 KABLJ isomorph presented in Figs. 1 and 2. The potential energies were fitted by linear regression by means of Eq. (21). Subsequently, the virials were compared to Eq. (22) using the parameters estimated from the potential energies (Eq. (21)), i.e., without performing a new fit.

We note here two important consequences of the above. (1) If the reference state point $(U_*, W_*)$ generates the isomorph $(U(\tilde{\rho}), W(\tilde{\rho}))$, then the reference state point $(aU_*, aW_*)$ generates the isomorph $(aU(\tilde{\rho}), aW(\tilde{\rho}))$. This means that all isomorphs for a given system have the same shape in the $UW$ diagram and can be scaled onto one another. (2) The shape of the isomorphs depends only on the exponents, $m$ and $n$. This means that all 12-6 LJ systems have the same isomorphs in the $WU$ phase diagram – they may be scaled onto a single “master isomorph” (see also Eq. (27) below).
Two isomorphs belong to the KABLJ system, the third results from simulations of the Wahnström binary LJ liquid \(^{46}\) (WBLJ), and the fourth (SCLJ) points give simulation results, the dashed curve the prediction of Eq. \((28)\). Figure 7 shows four isomorphs in the \(WU\) phase diagram for three different 12-6 LJ systems.\(^{47}\) Figure 8 shows the same isomorphs scaled onto the 12-6 master isomorph, Figure 9 shows an isomorph for the 18-6 KABLJ system and compares it to a 12-6 LJ isomorph. The agreement between simulation results and predicted shapes is quite good.

**B. Eliminating the density parameter: The “master isomorph” equation**

Combining Eq. \((18)\) with Eqs. \((14)\) and \((15)\) we get

\[
\bar{\rho}_m^{m/3} = \frac{U_m}{U_n} = \frac{3W - nU}{3W - mU}.
\]

Eliminating density and rearranging, we find an invariant for the isomorph:

\[
\left(\frac{W - \frac{n}{3}U}{W - \frac{n}{3}U^*}\right)^m = \left(\frac{W - \frac{m}{3}U}{W - \frac{m}{3}U^*}\right)^n.
\]

Choosing the reference point with zero potential energy \((U_0, W_0) = (0, W_0^*)\) (this reference point is guaranteed to exist if all interactions have attraction), we get:

\[
\frac{(W - \frac{n}{3}U)^m}{(W - \frac{n}{3}U^*)^m} = \frac{(W - \frac{m}{3}U^*)^n}{(W - \frac{m}{3}U^*)^n}.
\]

\(W_0^*\) is a unique number identifying the isomorph. Equation \((26)\) implies that the isomorph is the solution to

\[
\left(\frac{W - \frac{m}{3}U}{W_0^* - \frac{m}{3}W_0^*}\right)^{m/n} = \left(\frac{W - \frac{n}{3}U}{W_0^* - \frac{n}{3}W_0^*}\right)^{n/m}.
\]

Here it can be seen directly that – as argued above – for a given set of exponents \((m, n)\) all isomorphs have the same shape, suggesting the name “master isomorph”; \(W_0^*\) is the parameter that determines the scale of each isomorph. For \(m = 2n\) (e.g., the standard 12-6 LJ potential) the solution to this (quadratic) equation is:

\[
2 \frac{W}{W_0^*} = 1 + 4 \frac{n}{3} \frac{U}{W_0^*} \pm \sqrt{1 + 4 \frac{n}{3} \frac{U}{W_0^*}}.
\]

This has real solution(s) whenever \(U/W_0^* \geq -\frac{3}{4n}\), where equality corresponds to \(W = 0\). Figure 5 plots Eq. \((28)\) with \(n = 6\) for two values of \(W_0^*\).

**IV. APPLICATIONS**

**A. Solid-liquid coexistence**

In Paper IV it was argued briefly that Eq. \((7)\) predicts that solid-liquid coexistence curves are isomorphs. In more detail the argument goes as follows. First one notes that an isomorph cannot cross a solid-liquid coexistence curve: recall that the isomorph concept relates to equilibrium ensemble probabilities. For two isomorphic state points all pairs of microscopic configurations related by Eq. \((6)\) have identical Boltzmann probabilities according to Eq. \((7)\). Consequently, the two isomorphic state points must also have the same macroscopic phase behavior. In other words, if an isomorph were to cross the coexistence curve, part of the isomorph would have liquid
microstates as the most likely and part of it would have mixed crystalline-liquid microstates as the most likely (having in mind a density-temperature phase diagram). This would violate the proportionality of Boltzmann statistical weights for scaled states, the property that defines an isomorph.

Given that an isomorph cannot cross the solid-liquid coexistence curve, the next point is to note that—because all state points belong to some isomorph—a curve infinitesimally close to the coexistence curve must be an isomorph. This argument applies in the $W$, $U$ phase diagram, as well as in the density-temperature or the pressure-temperature phase diagrams. In the first two cases there is a multitude of isomorphs in the coexistence region; for instance, one particular isomorph will be characterized by 53% liquid and 47% crystal, etc. In the pressure-temperature phase diagram these “coexistence” isomorphs collapse into one, of course.

The prediction that the melting curve, both slightly to the liquid side and slightly to the solid side, are isomorphs has a number of consequences, and it sheds new light on some well-known phenomenological melting rules (as well as exceptions to these rules for non-strongly correlating liquids, see Paper IV). Thus, for instance, the observation that the Lindemann melting criterion is pressure independent for a given liquid follows from the isomorph property of the melting line (whereas the existence of a universal Lindemann criterion does not follow). Indeed, as is easy to show from paper IV the isomorph theory implies Ross’ “generalized Lindemann melting law” from 1969,\(^\text{48}\) according to which the reduced configurational partition function of the crystalline phase $Q^* = \int d\mathbf{R} \exp[-\beta(U(\mathbf{R}) - U(0))]$ is invariant along the melting curve (here, $U(0)$ is the potential energy of the crystal with all atoms located at their lattice sites, and the tilde denotes reduced coordinates). Further well-known melting rules, which the isomorph theory explains, include (Paper IV): the reduced viscosity, the reduced surface tension, the reduced diffusion constant, and the reduced heat conductivity are all invariant in the liquid phase along the melting curve. Likewise, the reduced-unit static structure factor is invariant along the melting curve. These rules have been confirmed for several liquids in both experiment and simulation (see the references of Paper IV).

![FIG. 10. Test of isomorph predictions for the liquid side of the solid-liquid coexistence curve of the single component generalized LJ potential ($n = 6$). Left panel: Eq. (21), middle panel: Eq. (22); data points are simulation results from Ahmed and Sadus.\(^\text{49}\) straight lines are linear regression fits. The right panel plots the fit parameters from Eqs. (21) and (22) against each other, demonstrating that the fits to energies and virials, respectively, are consistent.](image1)

![FIG. 11. Same data as in Fig. 10, plotting potential energy versus density (left panel) and virial versus density (right panel). Dashed curves: isomorph predictions (Eqs. (19) and (20)), using the parameters determined in Fig. 10.](image2)

![FIG. 12. Solid-liquid phase behavior for the single-component 12-6 LJ system. Main figure: filled symbols mark the coexistence region in the WU diagram. Blue circles are from Ref. \(^\text{49}\) and red squares are from Ref. \(^\text{50}\). Full curves are the isomorph prediction (Eq. (28)) with $W_m^*/N = 49.8$ and 56.8, respectively. Open circles indicate results from simulations in the solid and liquid phase, respectively. Both phases are strongly correlating with $R \geq 0.99$. The inset shows the solid-liquid phase behavior in a $\rho$, $T$-diagram with logarithmic axes. Dashed red curves are the isomorph prediction $\rho^*/T = \text{const}$ using $\gamma$ found using Eq. (9) from the simulations of the two phases (open circles) giving $\gamma = 5.24$ and 5.18, respectively. Full black curves are free fits giving $\gamma = 5.13$ and 6.49, respectively.](image3)
Like other isomorph predictions following from Eq. (7), the prediction that solid-liquid coexistence curves are isomorphs is only exactly obeyed for systems that obey Eq. (7) exactly, i.e., perfectly correlating systems. Indeed, it is straightforward to show that in IPL systems the phase behavior depends only on $\rho^{n/3}/T$, and thus the co-existence curve is an exact isomorph. The interesting question now is: how well do strongly correlating systems, e.g., generalized LJ systems, follow the prediction? Note that we do not expect the liquid-gas co-existence curves to be isomorphs (and indeed they are not); the gas-phase is not strongly correlating and thus Eq. (7) cannot be fulfilled in the relevant part of configuration space. In contrast, at solid-liquid coexistence both phases involved are strongly correlating for LJ systems (Paper I). Below we briefly turn to which degree the isomorph prediction is fulfilled.

Ahmed and Sadus used in a recent paper a novel numerical method for determining solid-liquid coexistence. They reported results for the generalized LJ potential with the repulsive exponent $m$ varying from 12 to 7 and fixed attractive exponent $n = 6$ (Ahmed and Sadus termed the repulsive exponent $n$, not $m$; we keep, however, the above notation). Figure 10 shows the state points reported by Ahmed and Sadus for the liquid side of the solid-liquid coexistence curve compared to the isomorph predictions as expressed in Eqs. (21) and (22). The right panel shows that the parameters found by fitting to Eqs. (21) and (22), respectively, are consistent. Figure 11 shows the same data plotting potential energy (left panel) and virial (right panel) versus density. Figure 12 shows the 12-6 LJ solid-liquid phase behavior in the $WU$-diagram and in the $\rhoT$-diagram (inset), including here also data from Mastny and de Pablo.50

The prediction that solid-liquid coexistence curves are isomorphs agrees well on the liquid side with the simulation results of Ahmed and Sadus as well as with the results for 12-6 LJ from Mastny and de Pablo.50 On the solid side the agreement is less convincing (see Fig. 12); in particular the logarithmic derivative in the $\rhoT$ diagram (slope in the inset) differs by 25%. At present we do not have an explanation for this. It might be related to inter-particle distances longer than the first peak of the radial distribution function playing a more important role for the phase behavior on the solid side than on the liquid side.

### B. An approximate equation of state for generalized LJ systems

Equations (19) and (20) give the potential energy and the virial as functions of density along an isomorph. The temperature variation along the isomorph can be found by integrating Eq. (9). Since $\gamma$ generally changes only little (Fig. 3), in many situations it is a good approximation to assume it constant, leading to the relation $T = \hat{\rho}^{1/\gamma}T_s$ along an isomorph, where $T_s$ is the temperature at the reference point. Taken together, Eqs. (19), (20), and (9) thus imply that from a single reference point $(\rho_s, T_s, U_s, W_s)$ we have a prediction for $(\rho, T, U, W)$ on the isomorph to which the reference point belongs.

Figure 4 demonstrated that in the $WU$ phase diagram, isochors to a good approximation are straight lines with slope $\gamma$, i.e., that one can write $W(\rho, T) = W_0(\rho) + \gamma(\rho)U(\rho, T)$. The only ingredient missing to generate a full equation of state expressed as $U = U(\rho, T)$ and $W = W(\rho, T)$ is a relation between temperature and potential energy or virial on an isochore.

Rosenfeld and Tarazona derived from density functional theory an expression for the potential energy on an isochore:

$$ U(\rho, T) = U_0(\rho) + \alpha(\rho)T^{1/5}. $$

Rosenfeld and Tarazona noted that the expression “...provides a good estimate of the thermal energy (...) near freezing densities only for predominant repulsive interactions,” and confirmed their expression by simulations of IPL potentials with different exponents. Equation (29) has since been shown, however, to be an excellent approximation also for several models with attraction, including the KABLJ mixture.53–55 This was regarded as a bit of a mystery, but can now be understood as a consequence of the fact that dynamics and heat capacity of strongly correlating liquids are well reproduced by a corresponding IPL system – a consequence of the hidden scale invariance of strongly correlating liquids discussed briefly in the introduction (see Papers I–III).

Figure 13 tests the Rosenfeld-Tarazona prediction for the systems investigated in Fig. 4. The prediction works very well, but less so for the supercooled Wahnström BLJ system, for which the formation of extended “Frank-Kasper” clusters at low temperatures affects the temperature dependence of the potential energy.56

Combining Eq. (29) with our results on isomorphs and isochors we can construct an approximate equation of state for generalized LJ systems. The idea is to map any state point $(\rho, T, U, W)$ to the corresponding isomorphic state point $(\rho_*, T_*, U_*, W_*)$ on a reference isochore where it is implicitly understood that the parameters $(W_0, \gamma, U_0, \alpha)$ are evaluated at the reference isochore $\rho_*$:

$$ T_s(\rho, T) = T_0^{1/\gamma}, \quad \hat{\rho} \equiv \rho/\rho_* $$

FIG. 13. Test of Rosenfeld-Tarazona scaling (Eq. (29)) plotting potential energy versus $T^{1/5}$ on isochors for the systems investigated in Fig. 4.
Applying the fitting procedure to the simulated state points on the five parameters found by fitting to the reference density in Figs. 4 and 13 are (38.3, 5.16, −8.63, 2.61), illustrating the robustness of the fitted equation of state. Applying the fitting procedure to the simulated state points on the $T = 0.550$ isotherm gives the parameters (38.5, 5.19, −8.63, 2.61). Right panel: same data as above in a $pT$ diagram.

\[ U_*(\rho, T) = U_0 + \alpha (T \bar{\rho}^{-\gamma})^{3/5}, \] (31)

\[ W_*(\rho, T) = W_0 + \gamma U_*(\rho, T). \] (32)

From $(U_*, W_*)$ we can calculate the contribution to the potential energy from the two IPL terms of the potential (compare Eqs. (14) and (15)):

\[ U_m^*(\rho, T) = \frac{3W_0 - (n - 3\gamma)(U_0 + \alpha (T \bar{\rho}^{-\gamma})^{3/5})}{m - n}, \] (33)

\[ U_n^*(\rho, T) = -\frac{3W_0 + (n - 3\gamma)(U_0 + \alpha (T \bar{\rho}^{-\gamma})^{3/5})}{m - n}. \] (34)

Finally, we use isomorphic scaling (Eqs. (19) and (20)) to go back to the $(\rho, T, U, W)$ state point:

\[ U(\rho, T) = \bar{\rho}^{m/3}U_m^*(\rho, T) + \bar{\rho}^{n/3}U_n^*(\rho, T), \] (35)

\[ W(\rho, T) = \frac{m}{3} \bar{\rho}^{m/3} U_m^*(\rho, T) + \frac{n}{3} \bar{\rho}^{n/3} U_n^*(\rho, T). \] (36)

The equation of state given by Eqs. (33) to (36) contains four parameters $(W_0, \gamma, U_0, \alpha)$ for the 12-6 KABLJ system using merely two state points. The resulting equation of state is shown as full curves in Fig. 14, where it is compared to simulation results shown as data points. The agreement is good. The largest deviations are seen when changing density away from the reference density, most evident for the isotherm in the $WU$ diagram and the isomorph in the $pT$ diagram. Better agreement is expected if the density dependence of $\gamma$ is taken into account, but such a correction will not be attempted here. To summarize, combining the isomorph theory with two further approximations, a state-point independent exponent $\gamma$ and the Rosenfeld-Tarazona expression, a realistic equation of state with four parameters is arrived at for generalized LJ systems.

V. CONCLUSIONS

Paper IV demonstrated that strongly correlating liquids have isomorphs, i.e., curves in the phase diagram along which structure and dynamics to a good approximation are invariant in reduced units. The main new results in this paper are predictions specific to generalized Lennard-Jones systems and supporting numerical evidence.

Starting from the invariance of structure along isomorphs, a prediction for the shape of these in the $WU$ phase diagram was presented and shown to agree well with simulation results. It was shown that for a given system, the isomorphs all have the same shape in the $WU$ phase diagram; they are simply scaled versions of each other. Furthermore, the isomorph shape depends only on the exponents $m$ and $n$, i.e., all systems with, say, 12-6 LJ interactions have the same isomorphs in the $WU$ phase diagram. What differs from system to system is the values of the density and temperature along an isomorph.

Throughout Papers I-V the $WU$ phase diagram was referred to extensively. This particular phase diagram has not been used much before, but it is evident that for strongly correlating liquids several features stand out when presented
in the $W$ phase diagram. Prominent examples are that isochors are straight lines (with slope $\gamma$) and the existence of the “master” isomorph discussed above for generalized LJ systems.

The class of strongly correlating liquids includes, we believe, all van der Waals liquids and metallic liquids (but excludes hydrogen-bonding, covalent, as well as strongly ionic liquids). More simulations, as well as experiments, are needed to substantiate that the class of strongly correlating liquids is this large, but preliminary simulations of simple molecular models like $^{28}$ the asymmetric dumbbell and the Lewis-Wahnstrom OTP three-site model are encouraging. In particular, the isomorph properties of more complex molecule models need to be simulated, as well as theoretically contemplated; the scaling properties of molecules with fixed bond lengths are not trivial. It should be noted that even a quite complex model system like $^{28}$ the asymmetric dumbbell and the molecular models like $^{28}$ the asymmetric dumbbell and the

33. Unless otherwise stated, simulations were performed using a newly developed molecular dynamics code optimized for NVIDIA graphics cards. The code is available as open source at http://rumd.org. Potentials were cut and shifted at 2.5 $\sigma_{NG}$. Reported potentials and virials do not include contributions beyond the cut-off.