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Hidden scale invariance of metals

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Density functional theory (DFT) calculations of 58 liquid elements at their triple point show that most metals exhibit near proportionality between the thermal fluctuations of the virial and the potential energy in the isochoric ensemble. This demonstrates a general "hidden" scale invariance of metals making the condensed part of the thermodynamic phase diagram effectively one dimensional with respect to structure and dynamics. DFT computed density scaling exponents, related to the Grüneisen parameter, are in good agreement with experimental values for the 16 elements where reliable data were available. Hidden scale invariance is demonstrated in detail for magnesium by showing invariance of structure and dynamics. Computed melting curves of period three metals follow curves with invariance (isomorphs). The experimental structure factor of magnesium is predicted by assuming scale invariant inverse power-law (IPL) pair interactions. However, crystal packings of several transition metals (V, Cr, Mn, Fe, Nb, Mo, Ta, W, and Hg), most post-transition metals (Ga, In, Sn, and Tl), and the metalloids Si and Ge cannot be explained by the IPL assumption. The virial-energy correlation coefficients of iron and phosphorous are shown to increase at elevated pressures. Finally, we discuss how scale invariance explains the Grüneisen equation of state and a number of well-known empirical melting and freezing rules.

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I. INTRODUCTION

Scale invariance plays an important role in many branches of science. It greatly simplifies a given phenomenon by reducing the parameter space and introducing universalities over length or time scales. Some examples of this are the size distribution of earthquakes [1], Brownian motions of microscopic particles [2], cosmic microwave background radiation [3,4], and biological fractal structures [5] such as those of lung tissue or Romanesco broccoli. In the physics of matter, scale invariance controls the properties of a fluid near the gas-liquid critical point [6]. This paper establishes an approximate "hidden" scale invariance in the dense liquid part of the thermodynamic phase diagram of certain elements from *ab initio* computations. While the above classic examples of scale invariance cover several decades of variations, the scale invariance of condensed matter covers much smaller length-scale differences. Nonetheless, hidden scale invariance implies a great simplification, namely that the thermodynamic phase diagram becomes effectively one dimensional.

Specifically, we have performed *ab initio* density functional theory (DFT) computations on 58 liquid elements at their triple point. We infer hidden scale invariance of metals from strong correlations in thermal fluctuations of virial W (the potential part of the pressure) and potential energy U. We find that metals in general possess hidden scale invariance. These results give the first *ab initio* quantum-mechanical confirmation of the picture proposed in Refs. [7–13], according

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to which some systems have strong virial potential-energy correlations in the condensed phases, whereas this is not the case for other systems—typically those with strong directional bonding. Liquids belonging to the former class of systems were originally referred to as "strongly correlating," but to avoid confusion with strongly correlated quantum systems the term Roskilde (R) simple system is now sometimes used [14–19]. Although the focus here is on monatomic liquids, experimental and molecular-dynamics simulation results have shown that (R) simple systems include some molecular van der Waals bonded systems [20], polymers [21,22], and crystalline solids [23]. Returning to the elements, molecular liquids like N₂ [9] and the noble gases [7] are also expected to be simple [8].

What is the microscopic origin of hidden scale invariance? To answer this, we start by noting that at high pressure the dominant interatomic forces are harshly repulsive [24–32]. These forces can be modeled approximately by scale-invariant inverse-power-law (IPL) pair potentials $\propto r^{-n}$ [33–48] in which r is the interparticle distance, plus a density-dependent constant $g(\rho)$ taking long-ranged attractive interactions into account [24,27]. By Euler's theorem for homogeneous functions, there is exact correlation between the fluctuations of the virial and the potential energy in the IPL case. Strong correlations between virial and potential energy are, however, not necessarily a consequence of approximately scale-invariant pair interactions, but rather a property of the multidimensional energy function $U(\mathbf{R}, V)$ per se. This fact motivates the present paper in which we show from first-principles computations that many metals possess hidden scale invariance also at *low* pressure, i.e., when virtually uncompressed compared to the zero-pressure state.

For metals, the outer electrons result in complex many-body interactions of the atoms, which this study takes explicitly into account. At low pressure hidden scale invariance is nontrivial since the forces on the atoms are not at all dominated by

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pairwise additive repulsive IPL-type forces [49]. Hidden scale invariances have been found to be absent if the multidimensional energy function $U(\mathbf{R}, V)$ has prominent contributions characterized by two or more length scales [8,9]. Examples of this are water with hydrogen bonds and core repulsions and the Dzugutov model with core repulsions supplemented by repulsive forces on second-nearest neighbors [8,9]. However, it is possible to have multibody interactions and hidden scale invariance. Below we do not make any assumptions on the potential energy landscape and verify hidden scale invariance for the majority of metals directly from a quantum-mechanical description. Density functional theory (DFT) currently provides the best means to do so [50-53], and although DFT can only approximate the true electronic behavior, it provides accurate predictions on a broad scale, such as crystal structures, densities, and melting points.

The remainder of the paper is organized as follows: Sec. II gives the theoretical background of hidden scale invariance, and Sec. III gives details on the *ab initio* DFT method. Results are presented in Sec. IV, and in Sec. V we relate our findings to the Grüneisen equation of state and well-known empirical melting and freezing rules.

II. THEORY: HIDDEN SCALE INVARIANCE OF CONDENSED MATTER

Certain condensed-matter systems are characterized by a "hidden scale invariance" as reflected in the following approximate representation of the potential energy function at density $\rho = N/V$ [11–13]:

$$U(\mathbf{R}) \cong h(\rho)\tilde{\Phi}(\tilde{\mathbf{R}}) + Ng(\rho),$$
 (1)

where the coordinates of the system's N particles have been merged into a single vector, $\mathbf{R} \equiv (\mathbf{r}_1, \mathbf{r}_2, \dots, \mathbf{r}_N)$, and the reduced coordinate vector is $\tilde{\mathbf{R}} \equiv \rho^{1/3} \mathbf{R}$. The intensive functions $h(\rho)$ and $g(\rho)$ both have dimension energy, and $\tilde{\Phi}$ is a dimensionless, state-point-independent function of the dimensionless variable $\hat{\mathbf{R}}$, i.e., a function that involves no lengths or energies. Physically, Eq. (1) means that a change of density to a good approximation leads to a linear affine transformation of the high-dimensional potential energy surface. Thus, if temperature is adjusted in proportion to $h(\rho)$, state points in the thermodynamic phase diagram are reached where the particles move in the same way according to Newton's laws, except for a uniform scaling of space and time. Sets of such thermodynamic state points are referred to as isomorphs, and along the isomorphs structure and dynamics are identical in properly reduced units to a good approximation [10]. The local slope $d \ln(T)/d \ln(\rho)$ of the isomorph is given by

$$\gamma(\rho) = d \ln(h(\rho)) / d \ln(\rho) \tag{2}$$

and is referred to as the density scaling exponent [11–13,21,22,54–57]. Thus, the phase diagram becomes effectively one dimensional, and density and temperature merge into a single parameter.

Hidden scale invariance is revealed in the thermal fluctuations at a single state point. There are two contributions to the pressure: the ideal-gas pressure $p_{\rm id}$ —a term that is always present and which only depends on the velocities of the

particles (atoms)—plus a term deriving from the interaction between the particles. The latter is the so-called virial W, and it only depends on the coordinates of the particles. W is an extensive quantity of dimension energy, and the general pressure relation is $p = p_{id} + W/V$. Hidden scale invariance dictates that fluctuations of virial and potential energy are strongly correlated in the NVT ensemble [7–9,11]:

$$W(\mathbf{R}, V) \cong \gamma(\rho)U(\mathbf{R}, V) + \text{const.},$$
 (3)

where the virial is defined by $W(\mathbf{R}, V) \equiv -V \partial U(\mathbf{R}, V)/\partial V$ [58,59]. We omit the explicit volume dependence of the potential energy in the following for brevity. Below we use the Person correlation coefficient

$$R = \langle \Delta W(\mathbf{R}) \Delta U(\mathbf{R}) \rangle / \sqrt{\langle [\Delta W(\mathbf{R})]^2 \rangle \langle [\Delta U(\mathbf{R})]^2 \rangle}$$
 (4)

of virial and potential energy fluctuations to determine to which degree the multibody energy function $U(\mathbf{R})$ has hidden scale invariance. Here, $\langle \ldots \rangle$ indicates a thermodynamic average in the NVT ensemble and Δ denotes the difference to the average value. R is an intensive quantity with a value between -1 and 1. A value close to 1 indicates hidden scale invariance. The density scaling exponent is given by [10]

$$\gamma = \langle \Delta W(\mathbf{R}) \Delta U(\mathbf{R}) \rangle / \langle [\Delta U(\mathbf{R})]^2 \rangle. \tag{5}$$

It was previously conjectured that metals possess a hidden scale invariance, but this was based on assuming Lennard-Jones-type pair potentials [7]. Reference [8] showed via the effective medium theory that pure copper, as well as a magnesium alloy, exhibit hidden scale invariance. We do not make such assumptions in this paper, as detailed in the following section.

Before proceeding we note that the isomorph theory was recently generalized by defining hidden scale invariance via the property that the order of potential energies is maintained for uniform scaling of configurations, i.e., $U(\mathbf{R}_a) < U(\mathbf{R}_b) \Rightarrow U(\lambda \mathbf{R}_a) < U(\lambda \mathbf{R}_b)$ [60].

III. METHOD: DENSITY FUNCTIONAL THEORY

DFT molecular dynamics is a computationally efficient method with a quantum-mechanical treatment of the electron-density field. Examples of recent successes of DFT are computation of anharmonic contribution due to phonon-phonon interactions for fcc crystals of metals [61], dynamics of water dissociative chemisorption on a Ni surface [62], and accurate computations of the melting points of period-three metals [63].

In this paper we present calculations of 58 elements using the Vienna *ab initio* simulation package (VASP) [64] employing the projector augmented wave (PAW) method [65] with the frozen-core assumption and the Perdew-Burke-Ernzerhof (PBE) exchange-correlation functional [66]. Dispersion corrections are not included, but they are not expected to be important for metals near the melting point [63]. We use periodic simulation cells containing 125 atoms for most elements except for the elements Li, Na, Mg, and Al where we use 256, 250, 384, and 256 atoms, respectively. Initial equilibration trajectories cover between 9 and 24 ps corresponding to several structural relaxation times. Constant temperature is obtained with a Langevin thermostat with a coupling time of 1 ps. Table I contains the electronic configurations of the calculated

TABLE I. Correlation coefficients R [Eq. (4)] and density scaling exponents γ [Eq. (5)] of the elements at the estimated DFT triple points T, ρ . Values in round parenthesis are the statistical uncertainties. N is the number of atoms in the periodic simulation cell. The electronic configurations of the calculated electrons as well as the plane wave cutoff energies E_{max} are listed in the fourth and fifth columns. The last two columns list the σ parameter of the IPL approximation [Eq. (8)] and the correlation between IPL and DFT energy fluctuations. Experimental values of T and ρ at the triple points are given in square brackets. Note that here ρ denotes the mass density rather than the number density [40,67–69].

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Elei	nent	N	Electrons	$E_{\rm max}$ [eV]	<i>T</i>	[exp.] [K]	ρ [g/c	[exp.] cm ³]	p [GPa]	R	γ	σ	$R_{ m IPL}$
3	Li	256	$2s^1$	140.0	470	[454]	0.534	[0.512]	0.29	0.72(0.04)	1.0(0.1)	2.03(0.27)	0.75(0.04)
4	Be	125	$2s^2$	308.8	1850	[1560]	1.570	[1.690]	-0.27	0.75(0.03)	1.8(0.3)	1.16(0.09)	0.50(0.13)
5	В	125	$2s^22p^1$	318.6	2400	[2349]	2.245	[2.08]	-1.69	0.10(0.12)			
6	C	125	$2s^22p^2$	273.9	6000	[4800]	1.514	[1.37]	-1.04	-0.04(0.18)			
11	Na	250	$3s^1$	62.1	370	[371]	0.941	[0.927]	-0.02	0.84(0.02)	1.9(0.1)	1.25(0.05)	0.89(0.01)
12	Mg	384	$3s^2$	98.5	900	[923]	1.581	[1.584]	0.07	0.90(0.01)	2.6(0.2)	1.13(0.03)	0.83(0.08)
13	Al	256	$3s^23p^1$	116.4	1000	[933]	2.352	[2.375]	-0.27	0.88(0.05)	4.0(0.2)	1.02(0.01)	0.91(0.02)
14	Si	125	$3s^23p^2$	245.3	1700	[1687]	2.827	[2.57]	0.51	0.79(0.03)	4.1(0.2)	0.94(0.02)	0.79(0.04)
15	P	125	$3s^23p^3$	255.0	650	[317]	1.850	[1.74]	-0.43	0.01(0.17)	(0.2)	015 1(0102)	0.77(0.0.1)
16	S	125	$3s^23p^4$	258.7	550	[388]	1.784	[1.819]	0.07	0.13(0.12)			
19	K	125	$3p^{6}4s^{1}$	116.7	350	[337]	0.815	[0.828]	0.07	0.86(0.11)	1.6(0.4)	1.40(0.21)	0.87(0.20)
20	Ca	125	$3p^64s^2$	266.6	1200	[1115]	1.378	[1.378]	-0.44	0.80(0.11)	1.9(0.1)	1.40(0.21)	0.87(0.20)
21	Sc	125	$4s^23d^1$	154.8	1900	[1814]	2.800	[2.80]	-0.38	0.63(0.24)	1.4(0.5)	1.43(0.56)	0.83(0.04)
22	Ti	125	$4s^23d^2$	178.3	3900	[1941]	4.161	[4.11]	-0.38 -0.48	0.03(0.24)	2.0(0.2)	0.96(0.04)	0.81(0.19)
23	V	125	$3p^64s^2d^3$	263.7	2500	[2183]	5.773	[5.5]	-0.13	0.81(0.07)	2.6(0.6)	1.02(0.12)	0.81(0.05)
24	Cr	125	$3p^{6}4s^{1}d^{5}$	265.7	2600	[2180]	6.735	[6.3]	-1.55	0.90(0.04)	3.3(0.9)	0.98(0.07)	0.83(0.07)
25	Mn	125	$4s^23d^5$	269.9	2400	[1519]	7.973	[5.95]	0.66	0.93(0.02)	3.6(0.2)	0.95(0.03)	0.72(0.08)
26	Fe	125	$3p^64s^23d^6$	267.9	2400	[1811]	8.338	[6.98]	0.03	0.95(0.02)	3.6(0.1)	1.00(0.01)	0.90(0.04)
27	Co	125	$4s^23d^7$	268.0	1870	[1768]	8.924	[7.75]	0.03	0.93(0.01)	3.5(0.1)	1.06(0.01)	0.94(0.01)
28	Ni	125	$4s^23d^8$	269.5	2000	[1728]	8.189	[7.81]	-0.17	0.92(0.03)	3.5(0.1)	1.03(0.02)	0.96(0.02)
29	Cu	125	$4s^{1}3d^{10}$	295.4	1480	[1358]	8.020	[8.02]	-0.72	0.90(0.02)	4.1(0.2)	1.01(0.01)	0.94(0.01)
30	Zn	125	$4s^23d^{10}$	276.7	760	[693]	6.570	[6.57]	-0.61	0.53(0.12)	3.3(0.9)	1.03(0.04)	0.43(0.17)
31	Ga	125	$4s^24p^1$	134.7	500	[303]	5.967	[6.095]	-0.41	0.74(0.09)	3.3(0.5)	1.09(0.04)	0.65(0.09)
32	Ge	125	$4s^23d^{10}4p^2$	310.3	1250	[1211]	5.600	[5.60]	-1.46	0.82(0.15)	4.8(1.1)	0.91(0.02)	0.80(0.15)
33	As	125	$4s^24p^3$	208.7	1300	[1090]	5.220	[5.22]	0.58	-0.04(0.08)	1.0(1.1)	0.51(0.02)	0.00(0.12)
34	Se	125	$4s^24p^4$	211.6	550	[494]	3.838	[3.99]	0.16	-0.02(0.21)			
37	Rb	125	$4p^65s^1$	121.9	340	[312]	1.460	[1.46]	0.01	0.80(0.03)	2.3(0.6)	1.12(0.16)	0.91(0.02)
38	Sr	125	$4s^24p^65s^2$	229.4	1100	[1050]	2.375	[2.375]	-0.24	0.88(0.14)	1.9(0.7)	1.19(0.22)	0.89(0.14)
39	Y	125	$4s^24p^65s^24d^1$	202.6	1850	[1799]	4.359	[4.24]	-0.47	0.60(0.09)	1.3(0.2)	1.47(0.25)	0.82(0.07)
40	Zr	125	$4s^24p^65s^24d^2$	229.9	2500	[2128]	6.305	[5.8]	0.01	0.82(0.07)	2.2(0.5)	1.13(0.11)	0.93(0.03)
41	Nb	125	$4p^65s^14d^4$	208.6	2900	[2750]	7.669		0.96	0.89(0.04)	3.2(0.2)	1.01(0.02)	0.91(0.01)
42	Mo	125	$5s^14d^5$	224.6	3000	[2896]	9.330	[9.33]	0.42	0.89(0.03)	3.2(0.4)	0.98(0.04)	0.77(0.10)
43	Tc	125	$5s^24d^5$	228.7	2500	[2430]	10.606		-0.43	0.87(0.09)	4.0(0.4)	0.94(0.02)	0.65(0.18)
44	Ru	125	$5s^14d^7$	213.3	2800	[2607]	11.200	[10.65]	1.18	0.94(0.05)	4.6(0.5)	0.97(0.02)	0.85(0.07)
45	Rh	125	$5s^{1}4d^{8}$	229.0	2400	[2237]	10.807	[10.7]	-1.86	0.88(0.07)	5.2(0.4)	0.96(0.01)	0.89(0.03)
46	Pd	125	$5s^14d^9$	250.9	1900	[1828]	10.380	[10.38]	-1.32	0.92(0.04)	4.9(0.5)	0.98(0.01)	0.94(0.01)
47	Ag	125	$5s^{1}4d^{10}$	249.8	1350	[1235]	9.320	[9.320]	0.98	0.90(0.03)		1.00(0.01)	
48	Cd	125	$5s^24d^{10}$	274.3	650	[594]	7.996	[7.996]	0.61	0.78(0.07)	5.5(0.6)	0.99(0.02)	0.84(0.04)
49	In	125	$5s^25p^1$	95.9	600	[430]	6.859	[7.02]	0.28	0.90(0.05)	4.2(0.5)	1.02(0.04)	0.88(0.08)
50	Sn	125	$5s^25p^2$	103.2	900	[505]	6.685	[6.99]	-0.89	0.88(0.06)	4.7(0.9)	0.99(0.03)	0.83(0.04)
51	Sb	125	$5s^25p^3$	172.1	950	[904]	6.530	[6.53]	0.68	0.40(0.15)	3.0(1.7)	1.03(0.33)	0.41(0.09)
52	Te	125	$5s^25p^4$	175.0	750	[723]	5.700	[5.70]	-0.64	0.03(0.22)			
55	Cs	125	$5s^25p^66s^1$	220.3	330	[302]	1.826	[1.843]	0.04	0.90(0.22)	1.7(0.6)	1.33(0.77)	0.96(0.45)
56	Ba	125	$5s^25p^66s^2$	187.2	1050	[1000]	3.338	[3.338]	-0.41	0.64(0.24)	1.0(0.4)	1.89(1.15)	0.69(0.29)
57	La	125	$5s^25p^66s^25d^1$	219.3	1280	[1193]	5.940	[5.94]	-1.86	0.72(0.23)	1.7(0.6)	1.25(0.30)	0.86(0.16)
72	Hf	125	$5p^{6}6s^{2}5d^{2}$	220.3	2600	[2506]	12.349	[12]	-1.92	0.83(0.03)	2.2(0.3)	1.16(0.07)	0.91(0.02)
73	Ta	125	$5p^{6}6s^{2}5d^{3}$	223.7	3450	[3290]	15.000	[15]	0.45	0.90(0.03)	3.3(0.3)	1.02(0.02)	0.91(0.06)
74	W	125	$5p^{6}6s^{2}5d^{4}$	223.1	3900	[3695]	16.803	[17.6]	-0.78	0.86(0.08)	3.7(0.6)	0.99(0.03)	0.88(0.06)
75	Re	125	$6s^25d^5$	226.2	3650	[3459]	18.787	[18.9]	0.98	0.82(0.09)	4.5(0.7)	0.97(0.02)	0.81(0.10)
76	Os	125	$6s^25d^6$	228.0	3450	[3306]	19.741	[20]	0.46	0.86(0.06)	5.1(0.4)	0.97(0.01)	0.85(0.04)
77	Ir	125	6s ¹ 5d ⁸	210.9	2900	[2719]	19.275	[19]	- 0.91	0.71(0.07)	5.1(0.4)	0.96(0.01)	0.83(0.05)

TABLE I.	(Continued.)
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Eler	nent	N	Electrons	E _{max} [eV]	<i>T</i>	[exp.]	ρ [g/o	[exp.]	p [GPa]	R	γ	σ	$R_{ m IPL}$
78	Pt	125	6s ¹ 5d ⁹	230.3	2200	[2041]	18.532	[19.77]	- 1.00	0.87(0.06)	6.0(1.4)	0.97(0.02)	0.94(0.03)
79	Au	125	$6s^15d^{10}$	229.9	1470	[1337]	16.690	[17.31]	1.06	0.86(0.14)	7.9(1.6)	0.95(0.02)	0.92(0.06)
80	Hg	125	$6s^25d^{10}$	233.2	470	[234]	11.917		0.48	0.84(0.13)	4.4(1.9)	1.00(0.07)	0.67(0.19)
81	Tl	125	$6s^26p^1$	90.1	600	[577]	11.220	[11.22]	0.53	0.90(0.09)	3.7(0.2)	1.06(0.02)	0.86(0.12)
82	Pb	125	$6s^26p^2$	98.0	900	[601]	10.671	[10.66]	0.48	0.90(0.10)	4.1(0.7)	1.03(0.03)	0.94(0.09)
83	Bi	125	$6s^26p^3$	105.0	580	[545]	10.050	[10.05]	-0.27	0.10(0.19)			
84	Po	125	$6s^26p^4$	159.7	850	[527]	9.039		-0.01	0.36(0.21)	2.5(1.7)	1.13(0.40)	0.35(0.23)

electrons as well as the cutoff energies of the plane wave basis set. All calculations were done non-spin-polarized, and the Brillouin zone was sampled at its center, the Γ point. Statistical uncertainties are estimated by dividing MD trajectories into statistically independent blocks. Details on estimating DFT triple points are given in the Appendix.

IV. RESULTS

A. Correlated virial and potential energy fluctuations

Figure 1 shows the results of DFT computations on the first six period-three elements. Each subfigure gives a scatter plot of virial versus potential energy of configurations taken from NVT equilibrium simulations of the liquid phase at the triple points. For the metals (Na, Mg, and Al) and the metalloid (Si) the scatter plots show strong correlations, implying that Eq. (3) is obeyed to a good approximation. The value of the Pearson correlation coefficient R [Eq. (4)] quantifies how well

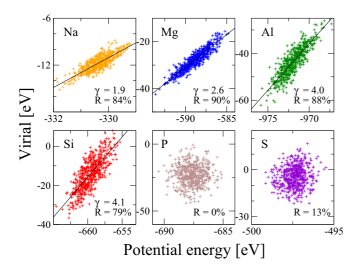


FIG. 1. (Color online) Results from *ab initio* quantum-mechanical calculations of the first six period-three elements in the liquid phase at the triple point. Each subfigure shows a scatter plot of the potential energy $U(\mathbf{R})$ and the corresponding virial $W(\mathbf{R})$ for configurations of the NVT ensemble. Fluctuations of virial and potential energy are strongly correlated for the metals Na, Mg, and Al and the metalloid Si, but not for the covalently bonded nonmetals P and S. The strong correlation validates hidden scale invariance for metals. Each subfigure reports the Pearson correlation coefficient R [Eq. (4)] and the density-scaling exponent γ of Eq. (3) determined as the linear-regression slope [Eq. (5)].

the approximation is obeyed [7–11]. The nonmetals P and S do not exhibit strong WU correlations.

B. Invariant structure and dynamics

To demonstrate hidden scale invariance directly, we investigated liquid magnesium in more detail. The results are summarized in Fig. 2, which studies isomorphic as well as isochoric state points for temperatures between 900 and 1600 K. Panels (a) and (b) give the radial distribution functions for the isomorphic and isochoric state points, respectively, while (c) and (d) give the translational order parameter Q suggested by Truskett, Torquato, and Debenedetti [70] and the reduced diffusion constant. We see that structure and dynamics are almost invariant along the isomorph, which confirms magnesium's hidden scale invariance.

The isomorph in Fig. 2 is determined as follows: First we choose to identify state points with density increases of 9%, 20%, and 30% relative to that of the triple point. Then

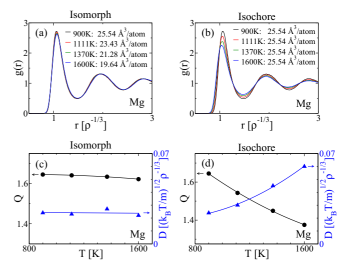


FIG. 2. (Color online) Predicted isomorph invariants of magnesium along an isomorph and an isochore for the same temperature variation. Panels (a) and (b) show results for the radial distribution function g(r) in reduced units; panels (c) and (d) show the translational order parameter Q of Debenedetti $et\ al.$ [70] and the reduced diffusion constant D. The reduced unit structural and dynamical quantities g(r), Q, and D all vary much less along the isomorph [(a) and (c)] than along the isochore [(b) and (d)]. m is the atomic mass.

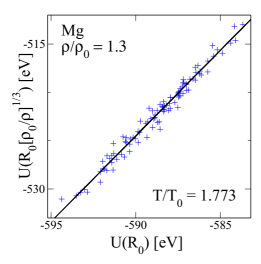


FIG. 3. (Color online) Determination of the isomorphic temperature at $\rho/\rho_0=1.3$ for Mg using Eq. (6) ($\rho_0=1.581$ g/cm³ is the estimated triple point density listed in Table I). The ratio T/T_0 is given as the ratio of the fluctuations of the potential energy of the scaled and unscaled configurations.

temperatures along the isomorph are determined by relating the Boltzmann factors of scaled configurations [10]:

$$\exp(-U(\mathbf{R}_0)/k_BT_0) \propto \exp(-U(\mathbf{R})/k_BT).$$
 (6)

100 representative configurations \mathbf{R}_0 at the triple point (T_0, ρ_0) are rescaled to the new density ρ according to $\mathbf{R} = \mathbf{R}_0[\rho_0/\rho]^{1/3}$. By taking the logarithm of Eq. (6), we see that T/T_0 is the ratio of the fluctuations of the energy at the scaled configurations $U(\mathbf{R})$ and the energy of the reference configurations $U(\mathbf{R}_0)$. Figure 3 relates the energies of these configurations at the initial density ρ_0 and at the target density ρ . The computed temperatures of the isomorphic states are 1110, 1370, and 1600 K, respectively.

C. The periodic table of hidden scale invariance

The results of Fig. 1 inspired us to study the elements in general to investigate whether all liquid metals exhibit hidden scale invariance. The results for 58 elements are summarized in Fig. 4 and Table I. We excluded some nonmetallic elements (gray in Fig. 4) for which standard semilocal density functionals are inaccurate. Metallic liquid elements all have strong or fairly strong virial potential-energy correlations at the triple point as quantified in the virial potential-energy correlation coefficient *R*. Most of the metals have *R* larger than 80%, however, a few metallic elements show correlation coefficients in the range of 50–80%. Scale invariance is expected to be worse for these elements.

As mentioned, all systems have the hidden-scale-invariance property at high pressure where repulsive pair interactions dominate (see Sec. IV G). Moreover, crystals generally have stronger virial potential-energy correlations than liquids [23]. We therefore conclude that metallic elements are (R) simple in the entire condensed-phase part of the phase diagram, i.e., exhibit hidden scale invariance to a good approximation. This excludes state points close to the critical point, as well as those of the gas phase far from the melting line. Table I also reports

the computed DFT scaling exponents γ , which we discuss in detail in the following section, Sec. IV D.

D. Density scaling exponents

For a monatomic liquid of N classical particles (above the Debye temperature) the density scaling exponent γ can be determined purely from thermodynamic responses [10]:

$$\gamma = [\gamma_G - k_B/c_v]/[1 - 3k_B/2c_v], \tag{7}$$

where $\gamma_G = \alpha_p K_T/\rho c_v$ is the thermodynamic Grüneisen parameter. Here, $\alpha_p = [\partial V/\partial T]_p/V$ is the isobaric thermal expansion coefficient, $K_T = -V[\partial p/\partial V]_T$ is the isothermal bulk modulus, $c_v = C_V/N$ is the isochoric heat capacity per atom, and k_B is the Boltzmann constant. The density scaling exponent is proportional to the thermodynamic Grüneisen parameter up to the accuracy of the Dulong-Petit approximation $c_v \simeq 3k_B$: $\gamma \simeq 2\gamma_G - 2/3$ (see inset of Fig. 5). In Fig. 5 we compare 16 experimentally determined density scaling exponents [71] to the values computed with DFT for the liquid state (see Table I). Incidentally, our computations show that the Lennard-Jones model does not properly reflect the physics of most metals because the DFT values of the density-scaling exponents are generally significantly smaller than $\gamma \simeq 6$ of the Lennard-Jones model [7].

Interestingly, with the exception of Fe and Tl, the elements Na, K, Rb, and Cs forming the more open body centered cubic (bcc) crystal structure have lower scaling exponents than the elements Mg, Al, Co, Ni, Cu, Zn, Ag, Cd, Au, and Pb that form close packed (cp) structures—face centered cubic (fcc) or hexagonal close packed (hcp). We explore this further in the following Sec. IV E.

E. Revisiting the inverse power-law model

As mentioned in the introduction, hidden scale invariance may be explained if the interactions in a metal can be approximated with the IPL model [33–48]: $U(\mathbf{R}) \simeq U_{\rm IPL}(\mathbf{R})$ where

$$U_{\text{IPL}}(\mathbf{R}) = Ng(\rho) + A \sum_{i < j}^{N} |\mathbf{r}_j - \mathbf{r}_i|^{-3\gamma}, \tag{8}$$

where $A = k_B T \sigma^{3\gamma} \rho^{-\gamma}$ is a material dependent constant. The mean field term $Ng(\rho)$ takes long-ranged attractive interactions into account [24,27]. Scale invariance is not "hidden" for the IPL model but a trivial result: Temperature and density merge into the dimensionless parameter $\sigma =$ $[A\rho^{\gamma}/k_BT]^{1/3\gamma}$ —i.e., σ is the single parameter of the phase diagram and isomorphs are given by $T \propto \rho^{\gamma}$. To quantify the accuracy of this approximation we list the correlation coefficients between the IPL energy and the DFT energy, $R_{\rm IPL} = \langle \Delta U_{\rm IPL}(\mathbf{R}) \Delta U(\mathbf{R}) \rangle / \sqrt{\langle [\Delta U_{\rm IPL}(\mathbf{R})]^2 \rangle \langle [\Delta U(\mathbf{R})]^2 \rangle},$ in Table I and color the symbols indicating the crystal structures in Fig. 4 accordingly. Elements with strong correlations between W and U fluctuations also have strong correlations between $U_{\rm IPL}$ and U. Figure 6 compares the experimentally determined structure factor S(q) of Mg at the triple point [72–74] with that of the IPL model (without any free parameters). The agreement is excellent. The deviation at small q vectors, related to the bulk modulus, is due

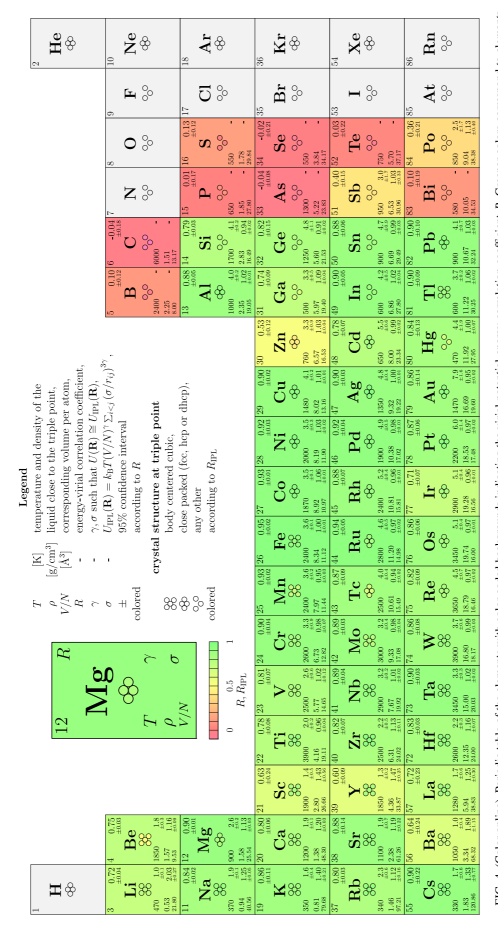


FIG. 4. (Color online) Periodic table of the elements with color coded backgrounds indicating the virial potential-energy correlation coefficient R. Greener colors correspond to elements with strong correlations between fluctuations of virial and potential energy in the NVT ensemble of the liquid state at the triple point. These elements exhibit hidden scale invariance to a good approximation. Most metalloids and all nonmetals investigated have weak to vanishing correlations. Gray elements were not considered since standard semilocal density functionals are naccurate for these. See Legend and Table I for more details. Note that here ρ denotes the mass density rather than the number density.

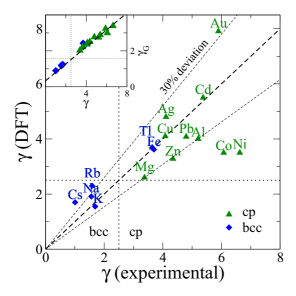


FIG. 5. (Color online) Density scaling exponents from experimental thermodynamic data collected in Ref. [71] compared to *ab initio* DFT computations for liquids. Elements forming bcc structures are marked with blue diamonds, while elements forming one of the close-packed structures (fcc or hcp) are marked with green triangles. The inset compares the experimental values of γ to those of the Grüneisen parameter γ_G . The dashed line indicates the prediction based on the Dulong-Petit approximation ($c_v \simeq 3k_B$).

to the mean-field $g(\rho)$ treatment of long-ranged attractive interactions [75].

Following the seminal 1972 paper by Hoover, Young, and Grover [34], we look for a connection between the density scaling exponents and crystal packings. An IPL liquid

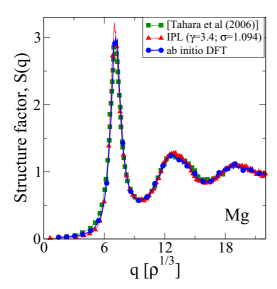


FIG. 6. (Color online) Experimental structure factor of Mg at the triple point [73] (green squares), compared to the IPL prediction (red triangles). The prediction has no free parameters: The exponent $\gamma=3.4$ of the IPL potential [Eq. (8)] is given by experimental values [71] of c_v , α_p , and K_T [using Eq. (7)], and $\sigma=1.094$ was chosen to match that of the solid-liquid coexistence shown in Fig. 7. The blue dots are the results of an *ab initio* DFT calculation.

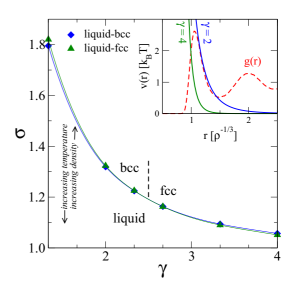


FIG. 7. (Color online) Phase diagram of the IPL model [Eq. (8) with $k_BT = \rho = 1$ and $g(\rho) = 0$] computed with the interface pinning method [80]. Green triangles are the liquid sides of the liquid-fcc transitions, blue diamonds are liquid-bcc transitions. Full lines are guides to the eye. The vertical dashed line indicates the bcc-fcc transition at $\gamma = 2.5$ [34,77]. The inset shows the pair energies and a representative liquid radial distribution at coexistence.

is known [33,34,40,41,47,76,77] to crystallize into a close packed (cp) fcc crystal structure when the IPL γ is above 2.5, while the more open body centered cubic (bcc) crystal is stable at lower values of γ (Fig. 7). The IPL model exhibits a polymorphic cp-bcc transition when $\gamma < 2.5$ also seen for many metals [40,78,79]. Thus, at the triple point elements with $\gamma < 2.5$ are expected [34] to form bcc crystals while elements with $\gamma > 2.5$ are expected to form cp crystals (fcc or hcp). Figure 8 shows the agreement of this prediction with the experimental crystal structure formed at the triple point (using the DFT values of γ given in Table I).

The prediction holds for many elements despite the small free energy differences between the respective crystal structures. However, it fails for several transition metals (V, Cr, Mn, Fe, Nb, Mo, Ta, W, and Hg), most post-transition metals (Ga, In, Sn, and Tl), and the metalloids Si and Ge. Moreover, the IPL model predicts that metals that form bcc structures at

Н														Не			
Li	Ве	Ве										В	С	N	О	F	Ne
Na	Mg	Mg										Al	Si	P	S	Cl	Ar
K	Ca	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kr
Rb	Sr	Y	Zr	Nb	Мо	Тс	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Те	I	Xe
Cs	Ba	La	Hf	Та	W	Re	Os	Ir	Pt	Au	Hg	T1	Pb	Bi	Po	At	Rn

FIG. 8. (Color online) Periodic table where green elements have a crystal structure at the triple point [40,78,79] predicted by assuming IPL interactions (see Sec. IV E). The IPL prediction fails for the red elements. Zn and Cd have been colored yellow since their distorted hcp crystals makes the IPL prediction ambiguous. All colored elements have hidden scale invariance judged from their highly correlated WU fluctuations (see Fig. 4 and Table I).

TABLE II. Correlation coefficients and scaling exponents for the period-three metals in the liquid phase at the triple point. Comparing the last two columns shows that the melting-line scaling exponent γ_m agrees with the density-scaling exponent γ . Numbers in parenthesis indicate the statistical uncertainties.

Element	R	γ	$\gamma_{ m m}$
Na	0.84(0.02)	1.9(0.1)	1.7(0.5)
Mg	0.90(0.01)	2.6(0.2)	2.6(0.5)
Al	0.88(0.05)	4.0(0.2)	4.6(0.8)

the triple point also have a low temperature cp phase; this is not always the case. For all of these elements we find significant WU correlations (see Fig. 4). Thus, we conclude that scale invariance can be present even when the element's crystal structure cannot be accurately predicted by IPL-like interactions. This also illustrates the importance of choosing an ab initio method for investigating hidden scale invariance.

F. The melting lines follow isomorphs

The melting line follows an isomorph to a good approximation [10,11,63]. In Table II this prediction is validated for the period-three metals by showing that the melting-line scaling exponent $\gamma_{\rm m} \equiv {\rm d} \ln T_{\rm m}/{\rm d} \ln \rho$ [23,63,80,81], calculated from the slope of the melting line, agrees with that of the isomorph, the density-scaling exponent γ .

G. Elevated pressure behavior of Fe and P

Fe shows a fairly good correlation between the virial and the potential energy. Interestingly, the melting line $T_m(\rho)$ reported in Refs. [82,83] coincides to a good degree with an extrapolation along the isomorph starting from the triple point, using $T_m \propto \rho^{\gamma}$ with $\gamma = 3.6$ (see Table I).

To exemplify that the behavior of elements becomes simpler at high pressure we compare in Fig. 9 the WU scatter plot of Fe at the triple point [Fig. 9(a)] with simulation results at the pressure 310 GPa [Fig. 9(b)] corresponding to a pressure in Earth's liquid outer core and near the melting line of pure iron [82,83]. The correlation coefficient increases from R=0.95 to R=0.98. Figure 9(c) shows that WU fluctuations are uncorrelated in the gas-liquid coexistence

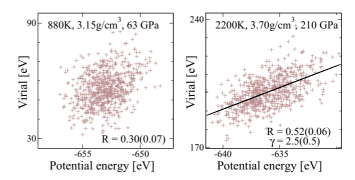


FIG. 10. (Color online) High-pressure calculations of phosphorus (P) show that the correlation coefficient increases with increasing pressure.

regime, which is presumably associated with the formation of a gas-liquid interface in the simulation cell and the onset of critical fluctuations. Likewise, the WU correlation coefficient of the nonmetal phosphorus (P) increases at high pressures from nearly uncorrelated at the triple point to R=0.30 and R=0.52 at (p=63 GPa, T=1880 K) and (p=210 GPa, T=2200 K), respectively (Fig. 10).

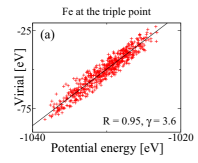
V. CONCLUDING REMARKS

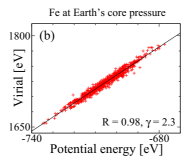
A. The Grüneisen equation of state

It has been known for a long time that at pressures high enough to result in non-negligible compression, solids and liquids generally obey the Grüneisen equation of state from 1912 (also referred to as the Mie-Grüneisen equation) [26,84]. This expresses proportionality between pressure and energy E per volume V as follows:

$$p = \gamma_{\rm G}(\rho)E/V + C(\rho) \tag{9}$$

in which $\gamma_G(\rho)$ is the Grüneisen parameter (Sec. IV D) and $C(\rho)$ the "cold pressure," both of which are functions only of the density. The Grüneisen equation has been applied for describing condensed matter in a wide variety of high-pressure situations, ranging from the core of Earth [82] to various forms of explosions [84,85]. The well-proven Hugoniot shock-adiabatic method is available for determining $\gamma_G(\rho)$ experimentally [84,86,87]. At high pressure in the dense fluid phase not too far from the melting line [88] the virial





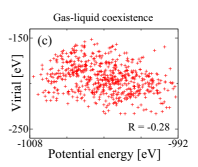


FIG. 9. (Color online) Virial potential-energy scatter plots for iron. Panel (a) shows a scatter plot of potential energy and virial of 125 iron atoms at the DFT triple point. Panel (b) shows that the correlation increases at elevated pressure, here chosen corresponding to a pressure at Earth's core and near the melting line of pure iron (310 GPa, 9000 K) [82,83]. Panel (c) shows that the correlation is low in the gas-liquid coexistence region.

term dominates. For instance, the ideal-gas contribution to the pressure is below five percent throughout the liquid outer core of Earth. The configuration-space analog of the Grüneisen equation is the relation

$$W \cong \gamma(\rho)U + C(\rho)V. \tag{10}$$

Equation (10) follows from the hidden scale invariance [10]. The scaling exponent $\gamma(\rho)$ has a simple relation to $\gamma_G(\rho)$ given in Eq. (7) [10].

B. Empirical melting and freezing rules

As a consequence of our finding, many empirical melting and freezing rules now find a concise explanation. Specifically, a number of invariants along the melting line of metals (and model systems) have been known for years with no good explanations. These rules follow from hidden scale invariance, because the melting and freezing lines are both isomorphs [11] and the rules all involve isomorph invariants. A famous melting rule is the Lindemann criterion, according to which a crystal melts when the thermal vibrational atomic displacement is about 10% of the crystal's interatomic distance [36,89–91]. While our work does not imply a universal value of 10%, it does imply invariance of the Lindemann quantity along the melting line. There are also other empirically wellestablished freezing rules of invariance, for instance the Hansen-Verlet rule that a liquid crystallizes when the first peak of the structure factor reaches the value 2.85 [92], the Andrade equation predicting constant reduced-unit viscosity along the freezing line [93,94], the Raveche-Mountain-Streett criterion [95] of a quasiuniversal ratio between maximum and minimum of the radial distribution function at freezing, Lyapunov-exponent based criteria [96], or the criterion of zero higher-than-second-order liquid configurational entropy at crystallization [97]. Connecting the melting and freezing lines is the rule of invariant constant-volume melting entropy [98,99].

ACKNOWLEDGMENTS

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APPENDIX: ESTIMATIONS OF DFT TRIPLE POINTS

The main focus of the paper is to investigate the possibility of hidden scale invariance of elements in the low-pressure part of the liquid phase. An unbiased choice is to use state points

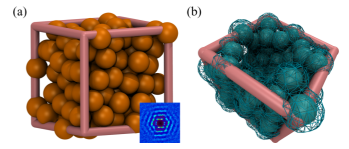


FIG. 11. (Color online) Panel (a) shows an example of a crystal-lized configuration (Ga). The inset shows the computed diffraction pattern. In such cases the computations were redone at higher temperature to avoid crystallization. Panel (b) shows a DFT configuration of Fe at the experimental triple point. DFT overestimates the triple point density by 16%; thus cavities are formed when iron is simulated at the experimental density.

near the DFT triple points. However, an accurate determination of the triple point is beyond the scope of the present paper and also not relevant for the present work. The DFT triple point is estimated by performing an NpT computation at a state point close to, or slightly above, the experimental melting temperature and at ambient pressure (within an accuracy of 2 GPa or less, see Table I). The NpT ensemble is realized using the Parinello-Rahman method [100] with a fictitious mass of 100 atomic units and a thermal coupling time of 1/3 ps. The finite cutoff of the plane wave basis set requires the exertion of an additional pressure to compensate for the missing basis set functions. This pressure is referred to as Pulay stress, and it was estimated in a separate calculation. If the system froze at the experimental temperature, for instance due to finite size effects, the temperature was increased until the system stayed liquid throughout the entire trajectory. An example of a crystallized configuration is shown in Fig. 11(a). For some systems, such as Ti, Ga, Sn, and Hg, the temperature had to be increased significantly compared to the experimental melting temperature. However, we only found small variations of γ and R upon increasing the temperature by a factor of 2. The final state points used to compute the UW correlations in the NVT ensemble are listed in Table I.

For Mn, Fe, and Co the computed liquid densities are 25%, 16%, and 15% higher than the respective experimental densities. Simulations of these liquids at the experimental density using standard density functional theory led to internal surfaces and cavities between the atoms as shown in Fig. 11(b). The reason for this sizable deviation of the DFT liquid densities from experiment needs to be investigated in the future. A likely explanation is strong paramagnetic fluctuations at the atomic sites that would require a treatment beyond standard density functional theory, for instance using dynamical mean field theory. Such fluctuations can increase the typical bond length by some 5%.

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