Predictions of Dynamic Behavior under Pressure for Two Scenarios to Explain Water Anomalies

Pradeep Kumar,¹ Giancarlo Franzese,² and H. Eugene Stanley¹

¹Center for Polymer Studies and Department of Physics, Boston University, Boston, Massachusetts 02215, USA

²Departament de Física Fonamental, Universitat de Barcelona, Diagonal 647, Barcelona 08028, Spain

(Received 21 February 2007; revised manuscript received 8 November 2007; published 11 March 2008)

Using Monte Carlo simulations and mean field calculations for a cell model of water we find a dynamic crossover in the orientational correlation time τ from non-Arrhenius behavior at high temperatures to Arrhenius behavior at low temperatures. This dynamic crossover is independent of whether water at very low temperature is characterized by a "liquid-liquid critical point" or by the "singularity-free" scenario. We relate τ to fluctuations of hydrogen bond network and show that the crossover found for τ for both scenarios is a consequence of the sharp change in the average number of hydrogen bonds at the temperature of the specific heat maximum. We find that the effect of pressure on the dynamics is strikingly different in the two scenarios, offering means to distinguish between them.

DOI: 10.1103/PhysRevLett.100.105701

PACS numbers: 64.70.Ja, 05.40.-a, 64.70.qj

Two different scenarios are commonly used to interpret the anomalies of water [1,2]: (i) The *liquid-liquid critical point (LLCP)* scenario hypothesizes that supercooled water has a liquid-liquid phase transition line that separates a low-density liquid (LDL) at low temperature T and low pressure P and a high-density liquid (HDL) at high T and P and terminates at a critical point C' [3]. From C' emanates the Widom line $T_W(P)$, the line of maximum correlation length in the (T, P) plane. Response functions, such as the isobaric heat capacity C_P , the coefficient of thermal expansion α_P , and the isothermal compressibility K_T , have maxima along lines that converge toward $T_W(P)$ upon approaching C' [Figs. 1 and 2(a)]. (ii) The singularityfree (SF) scenario hypothesizes the presence of a line of temperatures of maximum density $T_{MD}(P)$ with negative slope in the (T, P) plane. As a consequence, K_T and $|\alpha_P|$ have maxima that increase upon increasing P, as shown using a cell model of water. The maxima in C_P do not increase with P, suggesting that there is no singularity [4] [Fig. 2(b)].

Above the homogeneous nucleation line $T_H(P)$ where data are available, the two scenarios predict roughly the same equilibrium phase diagram. Here we show that dvnamic measurements should reveal a striking difference between the two scenarios. Specifically, the low-T dynamics depends on local structural changes, quantified by the variation in the number of hydrogen bonds, that are affected by pressure differently for each scenario. We find this result by studying-using Monte Carlo (MC) simulations and mean field calculations -a cell model which has the property that by tuning a parameter its predictions conform to those of either the LLCP or the SF scenario [5]. This cell model is based on the experimental observations that on decreasing P at constant T, or on decreasing T at constant P, (i) water displays an increasing local tetrahedrality [6], (ii) the volume per molecule increases at sufficiently low P or T, and (iii) the O-O-O angular correlation increases [7].

The entire system is divided into cells $i \in [1, ..., N]$, each containing a molecule with volume $v \equiv V/N$, where $V \ge Nv_0$ is the total volume of the system, and v_0 is the hard-core volume of one molecule. The cell volume v is a continuous variable that gives, in d dimensions, the mean distance $r \equiv v^{1/d}$ between molecules. The van der Waals interaction is represented by a potential with attractive energy $\epsilon > 0$ between nearest-neighbor (NN) molecules and a hard-core repulsion at $R_0 \equiv v_0^{1/d}$.

For a regular square lattice, each molecule *i* has four bond indices $\sigma_{ij} \in [1, ..., q]$, corresponding to the four NN cells *j*, giving rise to q^4 different molecular orientations. Bonding and intramolecular (IM) interactions are accounted for by the two Hamiltonian terms

$$\mathcal{H}_{B} \equiv -J \sum_{\langle i,j \rangle} \delta_{\sigma_{ij}\sigma_{ji}}, \qquad (1)$$

where the sum is over NN cells, $0 < J < \epsilon$ is the bond energy, $\delta_{a,b} = 1$ if a = b and $\delta_{a,b} = 0$ otherwise, and

$$\mathcal{H}_{\mathrm{IM}} \equiv -J_{\sigma} \sum_{i} \sum_{(k,\ell)_{i}} \delta_{\sigma_{ik}\sigma_{i\ell}}, \qquad (2)$$

where $\sum_{(k,\ell)_i}$ denotes the sum over the IM bond indices (k, l) of the molecule *i* and $J_{\sigma} > 0$ is the IM interaction energy with $J_{\sigma} < J$, which models the angular correlation between the bonds on the same molecule. The total energy of the system is the sum of the van der Waals interaction and Eqs. (1) and (2).

At constant *P*, the density of water decreases for $T < T_{\text{MD}}(P)$ which the model takes into account by increasing the total volume by an amount $v_B > 0$ for each bond formed. Hence the total molar volume v of the system is

$$v = v_{\text{free}} + 2p_B v_B, \tag{3}$$

where v_{free} is a variable for the molar volume without taking into account the bonds, $p_B = N_B/(2N)$ is the fraction of bonds formed and N_B is the number of bonds [4,5].



FIG. 1 (color online). (a) The relevant part of the liquid phase diagram for the LLCP scenario. C' is the LDL-HDL critical point at the end of the first-order phase transition line (thick line) [5], $T_W(P)$ (thin line) is the Widom line, which we take to be the average between $T(|\alpha_P|^{\max})$ (\bigcirc) and $T(C_P^{\max})$ (\square) from panel (b), and Fig. 2(a), respectively. Upper and lower dashed line are quadratic fits of $|\alpha_P|^{\max}$ and C_P^{\max} , respectively, merging at the C'. $T(|\alpha_P|^{\max})$ and $T(C_P^{\max})$ overlap within the error bars. (b) $|\alpha_P|$ as a function of T for different P. (c) $|dp_B/dT|$, the temperature derivative of p_B , as a function of T for different P. In panel (a), $T(|dp_B/dT|^{\max})$ (\diamondsuit) overlaps $T_W(P)$.

We perform simulations in the *NPT* ensemble [5] for q = 6, $v_B/v_0 = 0.5$, $J/\epsilon = 0.5$, and for two different values of J_{σ}/ϵ : (i) $J_{\sigma}/\epsilon = 0.05$, which gives rise to a phase diagram with a LLCP [Fig. 1(a)], and (ii) $J_{\sigma} = 0$, which leads to the SF scenario [4]. We study two square lattices with 900 and 3600 cells, and find no appreciable size effects. We collect statistics over 10^6 MC steps after equilibrating the system for all *P* and *T*.

For $J_{\sigma}/\epsilon = 0.05$, $|\alpha_P|$ for $P < P_{C'}$ displays a maximum, α_P^{max} [Fig. 1(b)]. As *P* increases, α_P^{max} increases and shifts to lower *T*, converging toward $T_W(P)$ [Fig. 1(a)]. We find that the number of bonds, N_B , increases on decreasing *T*, and at constant *T* decreases for increasing *P*, and is almost constant at $T_W(P)$ [8]. This is consistent with trends seen both in experiments [6] and in simulations [9], suggesting that for $T > T_W(P)$ the liquid is less structured and more HDL-like, while for $T < T_W(P)$ it is more structured and more LDL-like.

We find that $|dp_B/dT|$ shows a clear maximum for all $P < P_{C'}$ which shifts to lower T upon increasing P [Fig. 1(c)]. Remarkably, we also find that the locus of $|dp_B/dT|^{\text{max}}$ coincides with the Widom line $T_W(P)$

[Fig. 1(a)] and that the value of $|dp_B/dT|^{\text{max}}$ increases on approaching $P_{C'}$. This is the same qualitative behavior as $|\alpha_P(T)|^{\text{max}}$ and $C_P(T)^{\text{max}}$, which are used to locate $T_W(P)$ [Figs. 1(b) and 2(a)]. The relation of $|dp_B/dT|$ with the fluctuations is revealed by its proportionality to $|\alpha_P(T)|$ and to the fluctuation of the number of bonds

$$\langle N_B^2 \rangle - \langle N_B \rangle^2 = \frac{2Nk_B T^2}{J - P \upsilon_B} \left| \frac{dp_B}{dT} \right|, \tag{4}$$

where k_B is the Boltzmann constant.

For $J_{\sigma} = 0$ (SF scenario) we observe no difference for the behavior of N_B and $|dp_B/dT|$. We further verify the prediction of the SF scenario [4] that C_P^{max} remains a constant upon increasing P [Fig. 2(b)].

Next, we study how this different behavior affects the dynamics. Previous simulations [10] found a crossover from non-Arrhenius to Arrhenius dynamics for the diffusion constant of models that display a LLCP, and showed that the temperature of this crossover coincided with $T_W(P)$. We calculate, for both scenarios, the relaxation time τ of $S_i \equiv \sum_i \sigma_{ii}/4$, which quantifies the degree of



FIG. 2 (color online). Temperature dependence of the specific heat C_P for both the LLCP and the SF scenarios, for seven values of P. (a) For the LLCP scenario, C_P has a maximum, the size of which increases with increasing pressure and diverges as $P \rightarrow P_{C'}$. (b) For the SF scenario, C_P also has a maximum, but its size does not increase with increasing pressure, consistent with the findings of the mean field calculations of Ref. [4].



FIG. 3 (color online). Dynamic crossover—large hatched circles of a radius approximately equal to the error bar—in the orientational relaxation time τ for a range of different pressures. (a) The LLCP scenario, with crossover temperature at $T_W(P)$. (b) The SF scenario, with crossover temperature at $T(C_P^{\text{max}})$. Solid and dashed lines represent Arrhenius and VFT fits, respectively. Notice that the dynamic crossover occurs at approximately the same value of τ for all seven values of pressure studied.

total bond ordering for site *i*. Specifically, we identify τ as the time for the spin autocorrelation function $C_{\sigma\sigma}(t) \equiv \langle S_i(t)S_i(0)\rangle/\langle S_i^2(0)\rangle$ to decay to the value 1/e.

For both scenarios we find a dynamic crossover (Fig. 3). At high T, we fit τ with the Vogel-Fulcher-Tamman (VFT) function

$$\tau^{\rm VFT} \equiv \tau_0^{\rm VFT} \exp\left[\frac{T_1}{T - T_0}\right],\tag{5}$$

where τ_0^{VFT} , T_1 , and T_0 are three fitting parameters. We find that τ has an Arrhenius T dependence at low T, $\tau = \tau_0 \exp[E_A/k_B T]$, where τ_0 is the relaxation time in the high-T limit, and E_A is a T-independent activation energy. We find that for $J_{\sigma}/\epsilon = 0.05$ the crossover occurs at $T_W(P)$ for $P < P_{C'}$ [Fig. 3(a)], and that for $J_{\sigma} = 0$ the crossover is at $T(C_P^{\text{max}})$, the temperature of C_P^{max} [Fig. 3(b)]. We note that for both scenarios the crossover is isochronic; i.e., the value of the crossover time τ_C is approximately independent of pressure. We find $\tau_C \simeq 10^{3/2}$ MC steps.

We next calculate the Arrhenius activation energy $E_A(P)$ from the low-*T* slope of log τ vs 1/*T* [Fig. 4(a)]. We extrapolate the temperature $T_A(P)$ at which τ reaches a fixed macroscopic time $\tau_A \ge \tau_C$. We choose $\tau_A = 10^{14}$ MC steps >100 sec [11] [Fig. 4(b)]. We find that $E_A(P)$ and $T_A(P)$ decrease upon increasing *P* in both scenarios, providing no distinction between the two interpretations. Instead, we find a dramatic difference in the *P* dependence of the quantity $E_A/(k_BT_A)$ in the two scenarios, increasing for the LLCP scenario and approximately constant for the SF scenario [Fig. 4(c)].

We can better understand our findings by developing an expression for τ in terms of thermodynamic quantities, which will then allow us to explicitly calculate $E_A/(k_BT_A)$ for both scenarios. For any activated process, in which the relaxation from an initial state to a final state passes through an excited transition state, $\ln(\tau/\tau_0) = \Delta(U + PV - TS)/(k_BT)$, where $\Delta(U + PV - TS)$ is the difference in free energy between the transition state and the initial state. Consistent with results from simulations and experiments [12,13], we propose that at low T the mechanism to relax from a less structured state (lower tetrahedral order) to a more structured state (higher tetrahedral order) corresponds to the breaking of a bond and the simultaneous molecular reorientation for the formation of a new bond. The transition state is represented by the molecule with a



FIG. 4 (color online). Effect of pressure on the activation energy E_A . (a) Demonstration that E_A decreases linearly for increasing P for both the LLCP and the SF scenarios. The lines are linear fits to the simulation results (symbols). (b) T_A , defined such that $\tau(T_A) = 10^{14}$ MC steps >100 sec [11], decreases linearly with P for both scenarios. (c) P dependence of the quantity $E_A/(k_BT_A)$ is different in the two scenarios. In the LLCP scenario, $E_A/(k_BT_A)$ increases with increasing P, and it is approximately constant in the SF scenario. The lines are guides to the eyes. (d) Demonstration that the same behavior is found using the mean field approximation. In all the panels, where not shown, the error bars are smaller than the symbol sizes.

broken bond and more tetrahedral IM order. Hence,

$$\Delta(U + PV - TS) = Jp_B - J_\sigma p_{\rm IM} - Pv_B - T\Delta S, \quad (6)$$

where p_B and p_{IM} , the probability of a satisfied IM interaction, can be directly calculated. To estimate ΔS , the increase of entropy due to the breaking of a bond, we use the mean field expression $\Delta S = k_B [\ln(2Np_B) - \ln(1 + 2N(1 - p_B))]\bar{p}_B$, where \bar{p}_B is the average value of p_B above and below $T_W(P)$.

We next test that the expression of $\ln(\tau/\tau_0)$, in terms of ΔS and Eq. (6),

$$\ln \frac{\tau}{\tau_0} = \frac{J p_B - J_\sigma p_{\rm IM} - P v_B}{k_B T} - \bar{p}_B \ln \frac{2N p_B}{1 + 2N(1 - p_B)}$$
(7)

describes the simulations well, with minor corrections at high *T*. Here $\tau_0 \equiv \tau_0(P)$ is a free fitting parameter equal to the relaxation time for $T \rightarrow \infty$. From Eq. (7) we find that the ratio $E_A/(k_BT_A)$ calculated at low *T* increases with *P* for $J_{\sigma}/\epsilon = 0.05$, while it is constant for $J_{\sigma} = 0$, as from our simulations [Fig. 4(d)].

In summary, we have seen that both the LLCP and SF scenarios exhibit a dynamic crossover at a temperature close to $T(C_P^{\max})$, which decreases for increasing P. We interpret the dynamic crossover as a consequence of a local breaking and reorientation of the bonds for the formation of new and more tetrahedrally oriented bonds. Above $T(C_P^{\max})$, when T decreases, the number of hydrogen bonds increases, giving rise to an increasing activation energy E_A and to a non-Arrhenius dynamics. As T decreases, entropy must decrease. A major contributor to entropy is the orientational disorder, that is a function of p_B , as described by the mean field expression for ΔS . We find that, as T decreases, p_B —hence the orientational order—increases. We find that the rate of increase of p_B has a maximum at $T(C_P^{\max})$, and as T continues to decrease this rate drops rapidly to zero—meaning that for $T < T(C_P^{\max})$, the local orientational order rapidly becomes temperature independent and the activation energy E_A also becomes approximately temperature independent, for the Eq. (6). Corresponding to this fact the dynamics becomes approximately Arrhenius.

We find that the crossover is approximately isochronic (correlation time independent of the pressure) consistent with our calculations of an almost constant number of bonds at $T(C_P^{\max})$. In both scenarios, E_A and T_A decrease upon increasing P, but the P dependence of the quantity $E_A/(k_BT_A)$ has a dramatically different behavior in the two scenarios. For the LLCP scenario it increases as $P \rightarrow P_{C'}$, while it is approximately constant in the SF scenario.

We interpret this difference as a consequence of the larger increase of the rate of change of p_B in the LLCP scenario, where $|dp_B/dT|$ diverges at finite $T_{C'}$, compared to the SF scenario, where $|dp_B/dT|$ can possibly diverge only at T = 0. Since experiments can detect local changes of water structure from HDL-like to LDL-like, (e.g., [14,15]), it is possible that our prediction on the dynamic consequences of this local change may be experimentally testable.

We thank C. A. Angell, M.-C. Bellissent-Funel, W. Kob, L. Liu, and S. Sastry for helpful discussions and NSF Grant No. CHE0616489 for support. G. F. also thanks the Spanish Ministerio de Educación y Ciencia (Programa Ramón y Cajal and Grant No. FIS2004-03454).

- [1] C.A. Angell, Annu. Rev. Phys. Chem. 55, 559 (2004).
- [2] P.G. Debenedetti, J. Phys. Condens. Matter 15, R1669 (2003); P.G. Debenedetti and H.E. Stanley, Phys. Today 56, No. 6, 40 (2003).
- [3] P.H. Poole et al., Nature (London) 360, 324 (1992).
- [4] S. Sastry *et al.*, Phys. Rev. E 53, 6144 (1996); L.P.N.
 Rebelo *et al.*, J. Chem. Phys. 109, 626 (1998).
- [5] G. Franzese and H.E. Stanley, Physica (Amsterdam) 314A, 508 (2002); J. Phys. Condens. Matter 14, 2201 (2002); 19, 205126 (2007); G. Franzese *et al.*, Phys. Rev. E 67, 011103 (2003).
- [6] G. D'Arrigo *et al.*, J. Chem. Phys. **75**, 4264 (1981); C. A. Angell and V. Rodgers *ibid.* **80**, 6245 (1984).
- [7] A.K. Soper and M.A. Ricci, Phys. Rev. Lett. 84, 2881 (2000) and references cited therein; For simulations E. Schwegler *et al.*, *ibid.* 84, 2429 (2000); P. Raiteri *et al.*, *ibid.* 93, 087801 (2004) and references cited therein.
- [8] We find p_B ≈ 0.8 at T_W(P), being somewhat smaller for increasing P, consistent with the value 0.795 used in [4] to trace the temperature of maximum K_T for J_σ = 0. We find that for J_σ/ε = 0.05 polynomial extrapolations of p_B at C' up to the fifth order lead to a value of p_B(C') = 0.55 ± 0.15. Mean field calculations of p_B compare well with simulations for T ≈ T_W(P) and for T ≪ T_W(P).
- [9] P. Kumar et al., Proc. Natl. Acad. Sci. U.S.A. 104, 9575 (2007).
- [10] L. Xu *et al.*, Proc. Natl. Acad. Sci. U.S.A. **102**, 16558 (2005); P. Kumar *et al.*, Phys. Rev. Lett. **97**, 177802 (2006).
- [11] Comparison with P. Kumar *et al.*, Phys. Rev. E **73**, 041505 (2006) shows that 1 MC step $> \tau_{\alpha} \sim 10$ ps, the α -relaxation time in supercooled water.
- [12] D. Laage and J. T. Hynes, Science 311, 832 (2006).
- [13] A. Tokmakoff, Science **317**, 54 (2007), and references cited therein.
- [14] M. Krisch et al., Phys. Rev. Lett. 89, 125502 (2002).
- [15] F.F. Li et al., J. Chem. Phys. 123, 174511 (2005).