Entropy-driven liquid–liquid separation in supercooled water

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Twenty years ago Poole et al. (Nature 360, 324, 1992) suggested that the anomalous properties of supercooled water may be caused by a critical point that terminates a line of liquid–liquid separation of lower-density and higher-density water. Here we present an explicit thermodynamic model based on this hypothesis, which describes all available experimental data for supercooled water with better quality and with fewer adjustable parameters than any other model suggested so far. Liquid water at low temperatures is viewed as an ‘athermal solution’ of two molecular structures with different entropies and densities. Alternatively to popular models for water, in which the liquid–liquid separation is driven by energy, the phase separation in the athermal two-state water is driven by entropy upon increasing the pressure, while the critical temperature is defined by the ‘reaction’ equilibrium constant. In particular, the model predicts the location of density maxima at the locus of a near-constant fraction (about 0.12) of the lower-density structure.

Cold and supercooled water have been the subject of intensive experimental, theoretical and computational studies for the last several decades. Still, the famous statement of 1972 by Franks ‘of all known liquids, water is probably the most studied and least understood’ remains topical. This is especially true for metastable supercooled water, which is now a focal point of debates. On the other hand, there is a growing interest in the prediction of properties of supercooled water. In particular, in applied atmospheric science it is commonly accepted that the uncertainties in numerical weather prediction and climate models are mainly caused by poor understanding of properties of water in tropospheric and stratospheric clouds, where liquid water can exist in a deeply supercooled state.

FIG. 1. Hypothetical phase diagram of cold water. Supercooled water exists between the melting temperature $T_M$ and the homogeneous ice nucleation temperature $T_H$. Below $T_H$, there may exist a liquid–liquid critical point, marked by ‘C’, which terminates a liquid–liquid coexistence curve. The location of this critical point is shown as predicted by our model. The adopted location of the liquid–liquid coexistence curve is close to similar suggestions of Mishima, Kanno and Angell. The extension of the coexistence curve into the one-phase region is the line of maximum fluctuations of the order parameter, the Widom line. Thin solid lines represent phase boundaries between the different ices.

Intriguing liquid–liquid phase separation and the existence of the second critical point in water still remain a plausible hypothesis which needs further verification. In view of the inaccessibility of the LLCP to direct experiments, development of an equation of state, based on a solid physical concept and able to accurately describe all available experimental data, might help in resolving the supercooled-water dilemma.

In this paper, we offer an approach to thermodynamics of phase separation in supercooled water alternative to common views. Following Bertrand and Anisimov, we assume that liquid water is a non-ideal athermal ‘solution’ of two supramolecular arrangements, which undergoes phase separation driven by non-ideal entropy upon increase of the pressure. In the athermal two-state model, the non-ideality driven by entropy determines the critical pressure, whereas the critical temperature follows from the condition of ‘chemical reaction’ equilibrium. We have developed an explicit equation of state, which is based on the athermal two-state model and which describes all available experimental data for supercooled water, both H$_2$O and D$_2$O, with better quality and with fewer adjustable parameters than any other model suggested so far. We have also shown that the popular two-state ‘regular-solution’ scenario, in which phase separation is energy-driven, fails to describe the experimental data on water.

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I. HOW A PURE LIQUID CAN UNMIX

The only possible fluid–fluid phase transition in simple one-component substances, such as argon or methane, is separation into liquid and gas. This transition is driven by attraction forces between molecules. The critical temperature of the liquid–gas separation is uniquely proportional to the depth of the intermolecular interaction potential, which is a superposition of the attraction and repulsion potentials. The simplest model of the liquid–gas transition is the lattice gas, which, despite its simplicity, reflects the most important features of fluid phase behaviour. A central concept in physics of phase transitions is the notion of the order parameter. The order parameter in fluids is the difference between the densities of liquid and vapour. Thus, in simple one-component fluids only one liquid phase and only one critical point are possible. To observe a liquid–liquid separation one needs a mixture of two or more species. Remarkably, the same lattice-gas model that describes the liquid–gas transition in pure substances can also explain liquid–liquid demixing in weakly-compressible binary solutions. In binary liquid solutions the order parameter is the difference in concentrations of two liquid phases, while the critical temperature is proportional to the difference in the depths of the interaction energies between like and unlike molecules.

However, such a simple fluid-phase behaviour is not a law of nature. As explained by Mishima and Stanley, if the intermolecular potential of a pure fluid could exhibit two minima, the interplay between these minima may define the critical temperature and pressure of liquid–liquid separation. Application of this picture to water implies that the second, liquid–liquid, critical point is driven by molecular interaction energy just like the liquid–gas critical point.

The hypothesized existence of two liquid states in pure water can be globally viewed in the context of polyamorphism, a phenomenon that has been experimentally observed or theoretically suggested in silicon, liquid phosphorus, triphenyl phosphate, and in some other molecular-network-forming substances. Commonly, polyamorphism in such systems is described as energy-driven. However, there is an ambiguity in terminology adopted in Refs. and , where the term ‘density, entropy-driven’ is used for an energy-driven phase separation. There is an example of entropy-driven liquid polyamorphism, intermediate between two limiting cases: mostly entropy-driven phase separation (lattice liquid-like) near the critical point and mostly energy-driven (lattice gas-like) separation into two amorphous states observed in water at about K. We show in this paper that the actual behaviour of supercooled water appears to be much closer to the lattice-liquid behaviour than to the lattice-gas behaviour.

The entropy-driven separation of a lattice liquid can be further specified in terms of a two-state model. The pure liquid is assumed to be a mixture of two interconvertible states or structures of the same molecules, whose ratio is controlled by thermodynamic equilibrium. The existence of two states does not necessarily mean that they can separate. If these states form an ideal ‘solution’, the liquid will remain homogeneous at any temperature or pressure. However, if the solution is non-ideal, a positive excess Gibbs energy, , would cause phase separation. If the excess Gibbs energy is associated with a heat of mixing , the separation is energy-driven. If the excess Gibbs energy is associated with an excess entropy , the separation is entropy-driven. The entropy-driven nature of such a separation means that the two states would allow more possible statistical configurations, and thus higher entropy, if they are unmixed.

Two-state models for liquid water have a long history, dating back to the 19th century. More recently, two-state models have become popular to explain liquid polyamorphism. Ponyatovsky et al. and Moynihan assumed that water could be considered as a ‘regular binary
solution’ of two states, which implies that the phase separation is driven by energy. None of these two-state models for water has been used for a quantitative description of available experimental data.

II. VIRTUAL LIQUID–LIQUID CRITICALITY IN SUPERCOOLED WATER

There is no direct experimental evidence of the LLCP in real water, but it is indirectly supported by thermodynamic arguments based on density measurements\(^\text{13}\) and by critical-like anomalies of thermodynamic response functions\(^\text{6–8}\). The known existence of two states of glassy water\(^\text{28}\), as well as experiments on nano-confined water, which does not crystallize\(^\text{34,35}\), are also consistent with the possibility of a ‘virtual’ liquid–liquid separation. This possibility is also supported by simulations of some water-like models, such as ST2\(^\text{36}\). The exact location of the liquid–liquid critical point in these models is uncertain\(^\text{27}\). Moreover, for the mW model\(^\text{37}\) it has been recently shown that spontaneous crystallization occurs before a possible liquid–liquid separation could equilibrate\(^\text{38,39}\). However, the anomalies observed in the metastable region of the mW model\(^\text{37–40}\) might still be associated with the existence of a virtual LLCP in the unstable region.

The first attempt to develop an equation of state for supercooled water, based on the assumption that the LLCP exists, and on the asymptotic theory of critical phenomena\(^\text{41,42}\), was made by Fuentevilla and Anisimov\(^\text{43}\) and further elaborated and clarified by Bertrand and Anisimov\(^\text{40}\). In particular, both works estimated the LLCP critical pressure below 30 MPa, much lower than most of simulated water-like models predicted. Holten et al.\(^\text{44}\) used the same asymptotic equation of state, also in a mean-field approximation\(^\text{45}\), but introduced the noncritical backgrounds of thermodynamic properties in a thermodynamically consistent way. The resulting correlation represents all available experimental data for supercooled water, H\(_2\)O and D\(_2\)O, within experimental accuracy, thus establishing a benchmark for further developments in this field. However, there is a concern regarding the application of the asymptotic theory to a broad range of temperatures and pressures including the region far away from the assumed critical point. Such an extension makes the description of experimental data inevitably semi-empirical since all non-asymptotic physical features are absorbed by the adjustable backgrounds of thermodynamic properties. This fact underlines the need to develop a closed-form theoretically-based equation of state which would satisfy the asymptotic critical anomalies and, at the same time, describe regular behaviour far away from the critical region.

III. WATER AS AN AETHERMAL SOLUTION OF TWO STATES

We assume pure liquid water to be a mixture of two interconvertible states or structures A (HDL) and B (LDL). The fraction of molecules in state B is denoted by \(x\), and is controlled by the ‘reaction’

\[ A \rightleftharpoons B. \tag{1} \]

The states A and B could correspond to different arrangements of the hydrogen-bonded network.\(^\text{36}\) The Gibbs energy per molecule \(G\) is the sum of the contributions from both states,

\[ G = (1 - x)\mu^A + x\mu^B = \mu^A + x\mu^{BA}. \]

where \(\mu^A\) and \(\mu^B\) are the chemical potentials of A and B in the mixture. The variable \(x\) is conjugate to \(\mu^{BA} = \mu^B - \mu^A\). If A and B form an athermal non-ideal solution, the Gibbs energy of the mixture is

\[ \frac{G}{k_B T} = \frac{G^A}{k_B T} + x\frac{G^{BA}}{k_B T} + x \ln x + (1 - x) \ln(1 - x) + \omega x(1 - x), \]

and the chemical-potential difference is then

\[ \mu^{BA} = G^{BA} + k_B T \left[ \frac{x}{1 - x} + \omega(1 - 2x) \right], \]

where \(G^{BA} = G^B - G^A\) is the difference in Gibbs energy per molecule between pure configurations A and B, \(k_B\) is Boltzmann’s constant, and \(\omega = \omega(P)\) is the interaction parameter, which depends on pressure but not on temperature.

Considering \(x\), the fraction of B, as the ‘reaction coordinate’ or ‘extent of reaction’\(^\text{47}\), the condition of chemical reaction equilibrium,

\[ \left( \frac{\partial G}{\partial x} \right)_{T,P} = \mu^{BA} = 0, \]

yields the equilibrium constant \(K = K(T, P)\) of reaction (1) as

\[ \ln K = \frac{G^{BA}}{k_B T}, \]

and defines the equilibrium fraction \(x_e\) through

\[ \ln K + \ln \frac{x_e}{1 - x_e} + \omega(1 - 2x_e) = 0. \tag{3} \]

The expression (2) combined with the equilibrium condition \(x = x_e\) is our equation of state for the two-state supercooled water. The non-ideality of the two-state mixture is entirely associated with the excess entropy of mixing \(S^E = -\omega x(1 - x)\), while the heat (enthalpy) of mixing is zero. The parameter \(\omega\) determines the critical pressure through its pressure dependence, which we approximate as

\[ \omega = 2 + \omega_0 \Delta \tilde{P}, \]

where \(\Delta \tilde{P} = (P - P_c)/\rho_c k_B T_c\) with a subscript ‘c’ denoting critical parameters. An alternative, regular-solution model would have a factor \(w/k_B T\) in place of \(\omega\) in equation (2), where the interaction parameter \(w\) would determine the critical temperature. For a regular solution, the excess entropy is zero, while the heat of mixing is \(H^E = wx(1 - x)\), thus making phase separation purely energy driven.
The conditions for the critical point of liquid–liquid equilibrium,
\[
\frac{\partial^2 G}{\partial x^2} \bigg|_{T,P} = 0, \quad \frac{\partial^3 G}{\partial x^3} \bigg|_{T,P} = 0,
\]
yield in the case of the athermal-solution model the critical parameters \( x_c = 1/2 \) and \( \omega(P_c) = 2 \). At pressures below the critical pressure, \( \omega < 2 \) and the Gibbs energy versus fraction \( x = x_c \) shows a single minimum. Above the critical pressure, \( \omega > 2 \) and there are one or two minima in the Gibbs energy, depending on the value of the equilibrium constant \( K(T,P) \).

If there are two minima, the minimum with the lowest Gibbs energy represents stable equilibrium, and the other minimum corresponds to a metastable state. When \( \ln K(T,P) = 0 \), both minima have the same Gibbs energy, representing two-phase equilibrium. The critical temperature \( T_c \) and the location of the liquid–liquid transition (LLT) curve in the \( P-T \) plane are thus determined by the dependence of the equilibrium constant \( K \) on temperature and pressure. Since the LLT curve is defined as an analytical function of temperature and pressure \( \ln K(T,P) = 0 \), this function is to be obtained from the experimental data. In Supplementary Section 1 we explain how we match the LLT curve of the two-state model to the experimentally expected shape and location. The resulting expression for \( \ln K \) is
\[
\ln K = \lambda(\Delta \hat{T} + a \Delta \hat{P} + b \Delta \hat{T} \Delta \hat{P}),
\]
where \( \Delta \hat{T} = (T - T_c)/T_c \). The parameter \( a = -\rho_c k_B dT/dP \) is the slope of the LLT curve at the critical point, and \( b \) determines the curvature. The parameter \( \lambda \) is proportional to the heat of reaction (1), while the product \( \nu = \lambda a \) is proportional to the volume change of the reaction. Since \( \lambda / \nu = \Delta S / \rho_c k_B \Delta V = Q/T \Delta V \), the heat of reaction is asymptotically related to the latent heat of phase separation through \( \lambda \Delta x = Q/k_B T_c \), while \( \Delta x = \Delta S / k_B \lambda = Q / \lambda k_B T_c \) serves as the order parameter along the LLT.

The Gibbs energy \( G^A \) of the pure structure A defines the background properties and is approximated as
\[
G^A = \sum_{m,n} c_{mn}(\Delta \hat{T})^m(\Delta \hat{P})^n, \tag{4}
\]
where \( m (0 \to 3) \) and \( n (0 \to 5) \) are integers and \( c_{mn} \) are adjustable coefficients. Expressions for thermodynamic properties which follow from our model are given in Supplementary Section 2.

IV. EFFECTS OF CRITICAL FLUCTUATIONS

Thermodynamics predicts the divergence of fluctuations at the critical point. Entropy fluctuations are proportional to the isobaric heat capacity \( C_p \), volume/density fluctuations are proportional to the isothermal compressibility \( k_T \), and cross entropy–volume fluctuations are proportional to the thermal expansion coefficient \( \omega_T \). The two-state model described above is essentially mean-field, not affected by fluctuations. Being expanded near the critical point in powers of \( x - x_c \), equation (2) takes the form of a Landau expansion\(^2\). The procedure to include the effects of critical fluctuations is well developed and known as crossover theory\(^4\). This procedure is fully described in Supplementary Section 3. The variables \( x - x_c \) and \( P - P_c \) are renormalized such that the behaviour close to the critical point agrees with the asymptotic critical behaviour\(^4\), and crosses over to mean-field behaviour given by the equation of state, (2) and (3), far away from the critical point.

V. DESCRIPTION OF EXPERIMENTAL DATA AND DISCUSSION

We have fitted both the mean-field and the crossover formulation of our two-state model to experimental data, as described in Supplementary Section 4. We adopt the location of the liquid–liquid coexistence parallel to the homogeneous nucleation curve, which appears to be close to a suggestion of Mishima\(^7\) based on the shape of metastable melting curves of different ices, and a ‘singularity’ line of Kanno and Angell\(^7\) based on the extrapolation of the compressibility anomalies. The optimum locations of the critical point form a narrow band in the \( P-T \) diagram. The best fit for the critical point is obtained at about 227 K and 13 MPa, with \( \lambda = 2.3 \) and \( \omega_T = 0.35 \). The LLT curve was chosen to intersect the band of LLCP locations at the optimal value of the critical pressure (Fig. 2) where it has a dimensionless slope of \( 1/\alpha = 15.3 \). This location of the critical point is about 10% of \( \rho_c k_B T_c \) higher than that optimized by the mean-field approximation (see Supplementary Section 4A). This shift is induced by critical fluctuations, as follows from crossover theory.

The description of the density and the response functions
FIG. 3. Density of cold and supercooled water as a function of temperature and pressure. Black curves are the predictions of the crossover two-state model. $T_m$ (thin black) indicates the melting temperature and $T_H$ indicates the homogeneous nucleation temperature. The thin blue line is the liquid–liquid equilibrium curve, with the critical point C. The red line is the line of maximum density, and the green line is the line of a constant LDL fraction of about 0.12. Symbols represent experimental data. Mishima’s data have been shifted by at most 0.3% to bring them into agreement with data for stable water, as described in Ref. 44.

are shown in Figs. 3 and 4. A significant improvement compared to the previous works is that our equation of state does not show any sign of additional thermodynamic instability beyond the liquid–liquid separation. We now believe that the mechanical and thermal instability below the LLT, reported previously, were associated with the asymptotic nature of the equation of state, which caused negative backgrounds of the compressibility. The number of adjustable background coefficients in equation (4) is now 14, to be compared with $16^{44,45}$ (see Supplementary Table S1). Another improvement is the agreement of the coexistence densities, shown in Fig. 5, with the experimental densities of low-density and high-density amorphous water. We also fitted our model to all available experimental data for supercooled $D_2O$ with the same quality as for $H_2O$ (see Supplementary Section 4B). With the same number of adjustable parameters, a regular-solution two-state model fails (about ten times higher root-mean-square error) to describe experimental data on supercooled water.

In Fig. 6, we compare the LDL fraction $x$ predicted by our equation of state and that obtained in simulations of the mW model by Moore and Molinero. Remarkably, both results show a similar temperature dependence. In contrast, this fraction for an ideal LDL/HDL solution would be almost a linear function of temperature. The experimental maximum density line is located approximately along the line of constant LDL fraction $x = 0.12$ as shown in Fig. 3.

We confirm the previous finding of a critical pressure that is much lower than found in simulations. We believe that the low critical pressure reflects the entropy-driven nature of liquid–liquid criticality in supercooled water. As shown by Stokely et al., the LLCP critical pressure is determined by the ratio of hydrogen-bond cooperativity and hydrogen-bond covalent strength. The higher the ratio, the lower the critical pressure.

The necessity to develop a microscopic model for water, which would be consistent with its athermal two-state character, and which would clarify the microscopic nature of the order parameter, $(x - x_c)/x_c$, is evident.

Finally, the problem of the influence of the hypothesized liquid–liquid separation on homogeneous ice nucleation remains unresolved. The fact that the LLT curve is located just below the homogeneous nucleation curve and imitates its shape suggests that homogeneous ice nucleation may be caused by the entropy/structure fluctuations associated with the liquid–liquid transition. A connection between the change in the structure of liquid water and the crystallization rate of ice was shown for the mW model. Whether the LLCP of simulated water-like models is in the metastable region or just projected to be in the unstable region, the liquid–liquid criticality still may be responsible for the observed anomalies in the accessible domain.

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FIG. 4. Response functions as a function of temperature. a, Isothermal compressibility. Pressures, from top to bottom: 0.1 MPa, 10 MPa, 50 MPa, 100 MPa, 150 MPa, 190 MPa, 300 MPa, 400 MPa. b, Heat capacity at constant pressure $C_P$ (open circles: data from Archer and Carter) and at constant volume $C_V$ (calculated) at 0.1 MPa. See Supplementary Section 5 for more details. c, Thermal expansivity. Pressures, from top to bottom: 380 MPa, 200 MPa, 100 MPa, 0.1 MPa. In a, b, and c, the curves are the prediction of the crossover two-state model, and the symbols represent experimental data.

FIG. 5. Density along the liquid–liquid transition curve. The dashed line represents the mean-field two-state model; the solid line represents the crossover two-state model. The open circles are the densities of the low-density amorphous (LDA) and high-density amorphous (HDA) phases of water at 200 MPa. One can notice that the crossover LLT curve is flatter than the LLT curve in the mean-field approximation and that the actual position of the critical point is shifted to a lower temperature by critical fluctuations.

FIG. 6. Fraction of molecules in the low-density state. The fraction $x$ is shown for the two-state model at 0.1 MPa, in the case of an athermal solution (solid) and an ideal solution (dotted). The dashed curve is the fraction of four-coordinated water molecules, i.e., the fraction of water molecules with four neighbours, for mW water simulations performed by Moore and Molinero (see Supplementary Section 6). The temperature is scaled by the temperature of maximum density $T_{MD}$ (250 K for the mW model and 277 K for real water).

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