Dynamics and Thermodynamics beyond the critical point

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Sudden changes in the dynamical properties of a supercritical fluid model have been found as a function of pressure and temperature (T/Tc = 2–5 and P/Pc = 10–103), striving with the notion of a single phase beyond the critical point established by thermodynamics. The sound propagation in the Terahertz frequency region reveals a sharp dynamic crossover between the gas like and the liquid like regimes along several isotherms, which involves, at sufficiently low densities, the interplay between purely acoustic waves and heat waves. Such a crossover allows one to determine a dynamic line in the phase diagram which exhibits a very tight correlation with a number of thermodynamic observables, showing that the supercritical state is remarkably more complex than thought so far.

The supercritical fluid has since ever1, until recently, been considered as a unique state with intermediate properties of the liquid and gas phases. The thermodynamical observables show a continuous evolution along any P-T path in supercritical conditions. This fact is supported by the traditional view on liquid and gas as phases of the same symmetry, implying that these phases cannot be distinguished above the critical point where they do not coexist. On the other hand, this view assumes that physics of fluids is entirely determined by their structure. Modern approaches to liquid theory clearly show that this is not always the case. Dynamics plays often a crucial role. Early in 2005 L. Xu and collaborators have pointed out that the diffusion coefficient of water undergoes a non-Arrhenius (“fragile”) to Arrhenius (“strong”) crossover accompanying a thermodynamic liquid-liquid transition in the so called “no-man’s land” i.e. not accessible by current experimental techniques2.

More recently, it has been discovered that subtle but remarkable changes in supercritical fluids do occur in the positive sound dispersion (PSD)3–5. These changes posses the potential to part the phase diagram in a gas like and a liquid like region.

PSD is a phenomenon ascribed to the presence of one or more relaxation processes consisting in the increase of the sound velocity when the sound frequency is close to the relaxation frequency. In fact the relaxation processes time scale, τ, marks the border between the “viscous liquid-like” dynamic (for ω(Q)τ << 1, where ω and Q are the sound frequency and wave vector, respectively) to the “elastic solid-like” one (ω(Q)τ >> 1). Correspondingly, the longitudinal sound velocity undergoes a transition (PSD) from its “low” frequency adiabatic limit cs to its “infinite” frequency limit c∞ > cs. This effect has been observed in a wealth of liquids and glasses with typical relaxation frequency of the order of THz in simple liquids. On the other hand, PSD has never been observed so far in gases. Recently, PSD has been observed in supercritical fluids such as Argon (critical parameters Tc = 150.87 K and Pc = 4.898 MPa) and Oxygen (Tc = 154.6 K and Pc = 5.050 MPa) along isothermal lines, showing a cusp (discontinuity in the first derivative) as a function of pressure. This allows to define the high and low density supercritical fluids as liquid-like and gas-like supercritical fluids, respectively3–5. Based on such results, it has been proposed that the onset of PSD takes place in correspondence with the maxima of constant pressure specific heat cρ, defining the so called Widom line6, which establishes a tight link between dynamics (PSD) and thermodynamics (cρ). Unfortunately, in that case the study was performed along an isotherm at high temperature (T/Tc = 3.8), where the maximum of cρ is no longer visible and the Widom line could be only extrapolated.

A recent study indeed identified a dynamic crossover at the Widom line at lower temperatures where the maxima in cρ are still very well visible, based on anomalies in the self-diffusion coefficient and the excess two body entropy6. These results are at ease with previous findings of our group on the dynamic crossover related to collective modes. On the other hand, our view has been challenged by recent theoretical investigations. In one of these works6 it has been found that the extrema of different thermodynamic functions rapidly smear out above the critical point; this led the authors to question the potential link between dynamics and thermodynamics in the...
supercritical state, and to suggest the separation between liquid-like and gas-like regions to be rather determined by dynamics alone. The crossover in PSD of a model supercritical fluid was indeed assigned to a universal dynamic (Frenkel) line, where the equality between the relaxation time and the minimum period of transverse quasiharmonic waves holds\(^2\)–\(^5\). Unfortunately, PSD was not calculated, hence the advanced relationship between the potential dynamic crossover and the Frenkel line remains at the level of an interesting speculation calling for further investigations. Another possible way to identify two regions in the supercritical phase builds on the appearance of some elasticity upon crossing an aggregation line\(^6\). In addition, an alternative definition of the Widom line in terms of the loci of maxima of the volume of the correlation length was recently advanced\(^2\)–\(^5\). Interestingly, this line does not follow the slope of any response function extrema except in the close vicinity of the critical point, where it indeed follows the \(c_P\) maxima. This result adds a further call for a definitive clue on what is the right thermodynamic observable -if any- to correlate with the dynamic crossover.

**Results**

All this matter motivated us to deeply investigate what we think to be a milestone in the physics of the elusive supercritical state, i.e. the potential link between dynamics and thermodynamics. Specifically, based on molecular dynamics (MD) computer simulations, we investigated the onset of PSD along several isotherms down to temperatures where the line of maxima of \(c_P\) is well defined. In fact, we discovered that the onset of PSD occurs along this line up to temperatures where the maximum of \(c_P\) is still well defined and along the common minima of thermal diffusivity, \(D_T\), and kinematic shear viscosity, \(\nu\), at higher temperatures (note that kinematic longitudinal and shear viscosity \(D_L\) and \(\nu\) are related to the shear and bulk viscosity coefficients \(\eta_s\) and \(\eta_b\) as: \(D_L = (4/3)\eta_s + \eta_b/\rho\) and \(\nu = \eta_s/\rho\), where \(\rho\) is the density). The lines of extrema of these three observables are very close each other at sufficiently low temperatures, where maxima of \(c_P\) are well defined. Remarkably, we found that at sufficiently low densities in the gas-like regime, up to the dynamic crossover to the liquid-like regime, there is a strong interplay between purely acoustic waves and heat waves. These findings led us to the quantitative characterization of both types of wave properties, along with the quantitative determination of the crossover.

Some \(S(Q, \omega)\)s simulated on Ar at 700 K at four selected pressure and Q values are reported in figure 1, upper panel. The acoustic peak is increasingly more pronounced at high densities and small Q values. The corresponding sound dispersion \(c_L(Q)\) values have been divided by the adiabatic sound velocity value \(c_s\) (figure 1, lower panel). In case of no PSD this ratio must be equal to 1 at small Qs and to monotonically decrease at higher Qs on approaching the Q value of the first sharp diffraction peak (FSDP) of the static structure factor \(S(Q)\), resembling the bending of dispersion curves in crystals on approaching the Brillouin zone boundaries. On the other hand, when PSD is present, as it is the case here, the \(c_L(Q)/c_S\) ratio is expected to tend to 1 at very small Qs, to show a maximum upon increasing Q and then again to decrease at higher Qs on approaching the FSDP. PSD, i.e. the value of this maximum has been estimated by fitting a cubic polynomial function to the \(c_L(Q)/c_S\) ratio to account for finite Q sampling. The maximum takes place at around 3–5 nm\(^{-1}\), i.e. at the lengthscale \(~2\pi/Q^*\), where the inverse of structural relaxation time \(\tau(Q)\) equals the acoustic frequency \(\Omega(Q)\). This occurs at all the investigated pressures and the amount of PSD decreases on lowering density. Similar evolutions have been obtained along the other isotherms.

Density evolution of PSD is reported in Figure 2. Along all isotherms PSD shows a kink, being constant at low densities and increasing linearly at high densities. The extrapolation of the linear behaviour to low densities allows to estimate the density at which PSD is expected to vanish if coupling to the structural relaxation process was the only underlying, relevant mechanism; this density marks the onset of this kind of PSD or, in other words, the dynamic crossover between the gas-like and the liquid-like regime. On the other hand, the non-vanishing plateau in PSD at low densities indicates that for \(\rho \to 0\) some mechanism other than structural relaxation must exist. In order to shed light on potential mechanisms of PSD we used a thermo-viscoelastic model considering the possible propagation of heat and sound waves and the coupling between them\(^1\). A powerful method in the framework of generalized hydrodynamics is the approach of Generalized Collective Modes (GCM), representing acoustic excitations coupled to non-hydrodynamic processes as dynamic eigenmodes\(^1\)–\(^3\). The GCM approach explained correctly the “fast sound” phenomenon in disparate mass binary liquid alloys and predicted conditions of existence of non-hydrodynamic excitations like optic-like modes, heat waves etc\(^4\)–\(^7\). In Figure 3 (left frames) we show the frequency of acoustic modes and heat waves, as functions of Qs, at four selected densities at \(T = 363\) K. Heat waves were known to exist in solids as “second sound” excitations\(^8\)–\(^10\) while in liquids heat waves similarly to transverse sound waves cannot propagate on macroscopic distances\(^11\). This fact is reflected in the \(Q\) dependence of the frequency of heat waves vanishing at some finite \(Q_{h, \text{th}}\) value; heat waves can exist on spatial scales not larger than roughly \(2\pi/Q_{h, \text{th}}\). The dependence of sound dispersion on density shown in Fig. 3 gives evidence that below 5 nm\(^{-1}\) the deviation from the hydrodynamic dispersion law reduces with decreasing density. For low densities, however, there is also a strong increase of the deviation from the hydrodynamic dispersion around 10 nm\(^{-1}\), that is responsible of the overall PSD causing the plateau observed in Figure 2. We suggest that the origin of this increase at around 10 nm\(^{-1}\) for low-density states may be connected with heat waves because in this \(Q\) region at low density we observed the strong increase of their contributions to the \(S(Q, \omega)\) (Figure 3, right frames). These contributions (mode strengths) were obtained fit-free from the eigenvectors associated with acoustic and heat excitations. In the long-wavelength limit they tend to the values predicted by hydrodynamics\(^1\)–\(^2\). At high density the strength of the contribution of heat waves to the \(S(Q, \omega)\) is negligible, whereas it strongly increases upon decreasing density. This implies that at low densities, beyond the hydrodynamic region there exist two branches of strongly coupled acoustic-heat excitations, which in turn causes strong deviation from the hydrodynamic dispersion for the high-frequency branch.

**Discussion**

The reduced phase diagram (\(P/P_c-T/T_c\)) for Argon is reported in figure 4. It shows the simulation points (open dots) and the points where structural relaxation related PSD vanishes (full stars), according to the linear extrapolation of high density data as discussed above. Lines emanating from the critical point have been obtained from NIST data\(^12\): the red, blue and grey lines are obtained from extrema of \(c_P\), \(\nu\) and \(D_T\), respectively. These are among all possible lines emanating from the critical point and which can be considered as extensions of the liquid-vapour coexistence line into the supercritical fluid region. The line of maxima of \(c_P\) can be traced up to about 450 K (\(T/T_c = 2.98\)), as above this temperature these maxima severely smear out. On the other hand, the minima of \(D_T\) and \(\nu\) are still observable, even if broad, up to the highest temperature of this study (700 K, \(T/T_c = 4.64\)). The lines of extrema of \(c_P\), \(D_T\) and \(\nu\) are very close each other. Stars indicating the onset of PSD lie very close to these lines, putting the link between the dynamic crossover and thermodynamics on a very solid background. Indeed, the very existence of such a relationship has been the matter of what now appears to be a correct intuition in our previous works\(^3\)–\(^5\), where the relevant marker was thought to be only the line of \(c_P\) maxima. On the other hand, the fact that this line is not defined far from the critical point, where \(c_P\) maxima are not visible, made the connection between the onset of PSD and thermodynamics rather limited. Now, the connection between the dynamic crossover and the lines of \(D_T\) and \(\nu\) minima,
which are well defined also very far from the critical point, makes the link between dynamics and thermodynamics quite robust and general. Interestingly, the non monotonic density dependence of longitudinal kinematic viscosity, \(D_T\) and \(\nu\) has a remarkable consequence. These quantities define the hydrodynamic relaxation times for longitudinal and transverse dynamics of supercritical fluids\(^\text{22}\). The Q-width of the hydrodynamic regime, where collective excitations and relaxation processes have strictly hydrodynamic behaviour, depends on inverse of kinematic viscosities and \(D_T\)\(^\text{24,25}\). Recently the estimates for the hydrodynamic Q-range were proved by IXS experiments\(^\text{26}\). Hence, the minimum in the density dependence of these quantities corresponds to the widest Q-range of the hydrodynamic regime at given temperature, which in turn we found corresponding to the dynamic crossover observed in this work. Within this context, one may wonder how the crossover in the positive sound dispersion, established by this work to be the key parameter which allows to identify two different “phases”, one gas- and the other liquid-like, relates to the behaviour of diffusion coefficient, playing a role in distinguishing two different liquid phases in water\(^\text{2}\.\) In the water case, Kumar et al.\(^\text{27,28}\.\) identify the liquid-liquid transition reporting the breakdown of a linear dependence of the diffusion coefficient on density.

Figure 1 | Upper panel: \(S(Q, \omega)\) spectra obtained from MD simulations on supercritical Ar at four selected pressure (density) and Q values, at 700 K. Lower panel: \(c_L(Q)/c_S\) ratio at the same pressures. The dispersions (dots) have been fitted to a cubic polynom for a better estimation of the maximum. The \(c_S\) values have been obtained from NIST\(\text{23}\) up to the density corresponding to 10 Kbar, and extrapolated above this density using the formula: \(c_S(m/s) = 501.1744 + 0.178063(\rho)(kg/m^3) + 4.32538(10^{-4})(\rho)(kg/m^3)^2\). We propose this extrapolation as, on the basis of NIST\(\text{23}\) data for Ar at 700 K, it results that in the whole range of available data (from 50 to 10000 bar, (33.866 to 1506.5 kg/m\(^3\)) the sound speed shows a quadratic dependence on density with very good approximation (within about 0.1%).
fusivity as a function of the inverse structural relaxation time $1/\tau$. By virtue of the Maxwell relation, in turn, they eventually put forward the breakdown of the Stokes-Einstein relation. Although fascinating, this latter step relies on the assumption of a constant “infinite shear modulus”, the proportionality factor between diffusivity and $1/\tau$. The present work addresses the non constant behaviour of positive dispersion as function of the thermodynamic coordinates in the case of the gas-liquid transition in supercritical fluids, precisely the deviation of infinite frequency elastic properties from their hydrodynamic limit. It strongly suggests, therefore, that the amplitude of the structural relaxation was the dynamical observable related to the Widom line, rather than the relaxation time itself. This observation seems to indicate that the two phenomenon (liquid-liquid transition in water and gas-liquid transition in supercritical fluids), both related to the Widom line, may be not based on the same physics. In summary, we discovered two striking features in density dependence of PSD in supercritical fluids: i) linear behaviour of PSD at sufficiently high densities, that extrapolates right to the density of $C_P, D_T$ and $v$ extrema; ii) a plateau in the density behaviour of PSD for $p \to 0$, clearly indicating that this observable is originated by an additional collective process along with structural relaxation. The detailed analysis of dynamic eigenmodes and their contributions to $S(Q, \omega)$ revealed a novel effect of non-hydrodynamic heat waves on sound dispersion in low-density supercritical fluids. We quantitatively identified the dynamic crossover in a supercritical fluid model, at several temperatures, between the low density gas-like regime and the high density liquid-like regime, based on the analysis of collective modes with mesoscopic/microscopic wavelengths. In this framework, the unique role of heat thermal waves along with their interplay with sound waves have been unveiled. A remarkable outcome of this study is that the dynamic crossover shows a very tight correlation with the extrema of thermodynamic response functions and transport coefficients as $C_P, D_T$ and $v$, defining a narrow region emanating from the critical point, as emphasized in reference. This gives a new dimension to a recent work where a crossover in single particle dynamics was identified at the line of maxima of $C_P$ in a supercritical model fluid, in turns anticipated by similar results in the different context of the liquid-liquid transition in water. The deep link between dynamics and thermodynamics beyond a critical point is now rationalized on a very general background.

**Methods**

Molecular dynamics simulations were carried out on supercritical Argon at different densities at $T = 363, 453, 573$ and $700 \text{ K}$. The corresponding pressures were
obtained from NIST® up to 1 GPa and, above this pressure, as n(g/cm³) = A + B·log(P[GPa]), where A = -1.4186 g/cm³ and B = 9.42809·10⁻⁸ g/cm³ at 700 K, while the values at 573 K are reported elsewhere⁵. MD was done on the standard microcanonical ensemble for 2000 particles interacting through an ab initio potential with a 12 Å cutoff radius. We checked, at a few densities, that the same results are obtained with 4000 particles and 20 Å cutoff radius. The production runs were of 600000 time steps, for small densities they were extended up to 1.2 million time steps, each one of 2 fs. The energy drift over each production run was not higher than 0.2% that gave evidence of very good energy conservation in MD simulations. All of the calculated thermodynamic quantities agree with the values provided by the NIST® within 1%. Transport coefficients reported in Fig. 2, such as D_T and K_T, were used to calculate the Q-dependent contributions (mode strengths) to dynamic structure factors.

These densities represent fluctuations of conserved quantities and are the main dynamic variables for hydrodynamic description of liquid dynamics. The density-density time correlation functions F_n(Q,t) = <n'(Q)·n(Q+t)> , where the asterisk means complex conjugation, were calculated from the MD time evolution of n(Q,t), and additional averages were performed over the all possible directions of wave vectors corresponding to the same wave number Q. Numerical Fourier transform was used to obtain from the calculated density-density time correlation functions the dynamic structure factors S(Q,ω) and longitudinal current spectral functions J_0(Q,ω) = ε_0(ω)·S(Q,ω). The longitudinal sound velocity c_L(ω) is obtained as c_L(ω) = ω/√(k_0·ω·q), where ω_0(ω) is the sound frequency and it corresponds to the frequency of the maximum of J_0(Q,ω)². For analysis of contributions from collective modes to the maximum of J_L(Q,v), where L(Q)/ Q, was used to calculate the Q-dependent contributions (mode strengths) to dynamic structure factors.

Figure 4 Reduced phase diagram (P/P_c - T/T_c) for Argon. Open dots: P-T points used in MD simulations. Stars: onset of PSD at the four isotherms, based on linear extrapolation of the PSD data points to low densities (see figure 2). Lines (obtained from NIST®) loci of the extrema of c_p (red), D_T (grey) and v (blue). Dotted line: critical isochore.

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Author contributions
F.A.G., T.B., M.K., G.R., M.S. & T.S. designed research, performed research, wrote, reviewed and approved the manuscript. T.B. performed the numerical simulations.

Additional information
Competing financial interests: The authors declare no competing financial interests.

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