Liquid-liquid equilibrium for monodisperse spherical particles

E. A. Jagla
Centro Atómico Bariloche and Instituto Balseiro, Comisión Nacional de Energía Atómica
(8400) S. C. de Bariloche, Argentina

A system of identical particles interacting through an isotropic potential that allows for two preferred interparticle distances is numerically studied. When the parameters of the interaction potential are adequately chosen, the system exhibits coexistence between two different liquid phases (in addition to the usual liquid-gas coexistence). It is shown that this coexistence can occur at equilibrium, namely, in the region where the liquid is thermodynamically stable.

I. INTRODUCTION

Traditionally, liquid-liquid separation for single component systems was not considered as a real possibility, because of the rapidly changing liquid structure, which smoothly varies on temperature and pressure. This view is changing. Experimentally, a whole family of network forming fluids, including some tetrahedrally coordinated materials, is suspected to have transitions between liquid states. Usually this transition occurs well inside the supercooled region, where the liquid is already in a glassy state, and then it is related to polyamorphism. The numerical modeling of these materials shows that interaction potentials can be constructed for which there are transitions between different liquid states. For instance, the potentials used to mimic the properties of water generate two liquid phases in the supercooled region. The main characteristic of these materials (and of the interaction potentials used to model them) which is responsible for their anomalous properties is the possibility of generating open, low coordinated network structures at low pressure, while collapsing to more compact structures at higher pressures.

This key ingredient is captured by a class of model spherical potentials, the so-called soft core Stell-Hemmer potentials. They include a strict hard core at some distance $r_0$ and a soft core at a larger distance $r_1$. In this way, two typical distances between particles occur, and there is a collapse from the largest to the smallest distance on applying pressure. The properties of these systems give a remarkably consistent description of most of the anomalous behavior of tetrahedrally coordinated materials.

The possibility of liquid-liquid separation has been extended in the last time by the finding that it can occur even at thermodynamic equilibrium. Katayama et al. report that phosphorus (which is a network forming fluid) shows liquid-liquid coexistence at equilibrium, providing the first experimental evidence of this phenomenon. In addition, Glosli and Ree find numerical evidence of liquid-liquid coexistence at equilibrium in carbon. This was the first direct numerical evidence of the phenomenon for a single component system. The interparticle potential used to model carbon-carbon interaction is non-isotropic, and incorporates information of the internal (electronic) degrees of freedom of the atoms. It is shown that one of the two coexisting liquids is mainly $sp^3$ coordinated, whereas the other shows a dominant $sp^2$ character.

On the basis of these results, it is natural to ask whether some kind of soft-core potentials (with no internal degrees of freedom for the particles) can produce an equilibrium liquid-liquid transition. We will show, by providing concrete examples, that soft core Stell-Hemmer potentials are able to do that, then showing that the phenomenon may occur also for systems that, in addition to being monodisperse, have particles which are spherical and ‘rigid’, i.e., in cases where internal degrees of freedom play no role. This is of basic interest since this model is much simpler, and then more transparent to analyze than real examples of material having liquid-liquid coexistence at equilibrium. It also provides a benchmark for the study of tetrahedrally coordinated materials with liquid-liquid critical points in the supercooled region. The relation between the existence of the liquid-liquid critical point and the anomalous thermodynamic properties of these materials can be studied here in a context of thermodynamic equilibrium. On a different context, our results make plausible the existence of liquid-liquid separation at equilibrium in particular cases of colloidal dispersions, which is of potential technological interest.

II. THE MODEL

We study a model of particles interacting through a potential which is the sum of attractive and repulsive parts $V(r) = V_A(r) + V_R(r)$. The repulsive part is given as

\[ V_R(r) = \begin{cases} \infty & \text{for } r < r_0 \\ \varepsilon_0 \frac{(r_1 - r)}{(r_1 - r_0)} & \text{for } r_0 < r < r_1 \\ 0 & \text{for } r > r_1, \end{cases} \quad (1) \]
nearly, a hard core at distance \( r_0 \), and a linear (repulsive) ramp between \( r_0 \) and \( r_1 \). The attractive part is taken also to be linear in \( r \), of the form

\[
V_A(r) = -\gamma (r_2 - r) / (r_2 - r_0) \quad \text{for} \quad r < r_2 \\
V_A(r) = 0 \quad \text{for} \quad r > r_2
\]

(\( r_2 > r_1, \gamma > 0 \)). The repulsive part of the potential favors one of the two distances \( r_0 \) or \( r_1 \) between neighbor particles depending on the value of external pressure and the attraction intensity \( \gamma \). Interparticle distance \( r_1(r_0) \) is favored for pressures lower (higher) than some crossover value \( P_0 \). The crossover pressure (for \( \gamma = 0 \)) is given roughly as \( P_0 = \varepsilon_0/(v_1 - v_0) \), where \( v_0 \) and \( v_1 \) are the specific volumes of the states with nearest particles at distances \( \sim r_0 \) or \( \sim r_1 \). For \( P \) near \( P_0 \) the system has an anomalously large compressibility. The attractive part of the potential may turn this anomaly into a first order transition through a van der Waals mechanism.

We will show results for two-dimensional (2D) and three-dimensional (3D) systems. The parameters we use are: \( r_1 = 1.72r_0, r_2 = 4.5r_0 \), and \( \gamma = 0.27\varepsilon_0 \) for 2D and \( r_1 = 1.72r_0, r_2 = 3.0r_0 \), and \( \gamma = 0.31\varepsilon_0 \) for 3D. Temperature will be measured in units of \( \varepsilon_0/k_B \), and pressure in units of \( \varepsilon_0/r_0^2 \) in 2D, and \( \varepsilon_0/r_0^3 \) in 3D. Calculations were done by standard Monte Carlo techniques, in systems with periodic boundary conditions. Some of the results to be presented were done in the NVT ensemble, where the volume of the system is kept fixed at each simulated value, and pressure is calculated through standard formulae. Other results correspond to the NPT ensemble, where external pressure is fixed, and the volume of the system is taken as an additional variable during the simulation.

**III. RESULTS**

**A. Two dimensions**

Fig. 3 shows the isotherms we get from the simulations for a system of 200 particles in the 2D case, near the position in which we find a liquid-liquid critical point for our parameters, namely \( T_{LL} \approx 0.10, P_{LL} \approx 0.07 \), and \( v_{LL} \approx 1.85r_0^2 \). We see in fact that when reducing \( T \) below \( \sim 0.1 \), the isotherms get a loop, typical of a first order transition. This transition separates two liquid phases that we call (following the notation in water) low density liquid (LDL) and high density liquid (HDL). We have checked that the position and characteristics of this transition do not change when going to a system of 1000 particles. As an example, we show in Fig. 3 the \( T = 0.105 \) and \( T = 0.0875 \) isotherms of systems of 200 and 1000 particles. Isotherms are seen to coincide with the 200 particles case, except within the coexistence region. Here the isotherm is more flat in the case of larger system. This is in fact what expected, since in an infinitely large system isotherms are strictly flat in the coexistence region. Fig. 3 shows snapshots of the system at \( T = 0.0875 \), within the LDL and HDL phases and in the coexistence region. The HDL (LDL) phase is characterized by a larger amount of particles at distance \( r_0 \) \( (r_1) \) from their neighbors. The snapshot at coexistence [Fig. 3(b)] shows indeed that different neighborhoods of high density and low density coexist in the system. The existence of a loop in the isotherms below the critical temperature, instead of a flat region, even when the system clearly separates in different regions corresponding to the two different phases [Fig. 3(b)] is due to the non-vanishing contribution to the total energy of the surface energy between the two phases. This contribution only vanishes in infinite systems.

From the isotherms of Fig. 3 we see that the LDL has a density anomaly, since there is a temperature range in which volume increases as temperature is reduced, at constant pressure. This density anomaly (which is reminiscent of the similar phenomenon in water) is known to occur in core-softened models even in the absence of an attractive part of the interparticle potential, namely, when \( U_A = 0 \).

We will now present the equilibrium \( P-T \) phase diagram of the system, close to the liquid-liquid critical point, to show clearly that the liquid-liquid coexistence we are observing occurs in thermodynamic equilibrium, and it is not preempted by the appearance of any other phase (gaseous or crystalline).

The \( P-T \) phase diagram is presented in Fig. 4, and the different borders between phases were obtained as explained in the following. The position of the equilibrium first order line between LDL and HDL phases in the \( P-T \) phase diagram was determined from the isotherms of Fig. 3 and from the values of the enthalpy \( h = e + P v \) (not shown) by thermodynamic integration, calculating differences in Gibbs free energy \( G \) from the relation

\[
\frac{G_2}{T_2} - \frac{G_1}{T_1} = \int_1^2 \frac{v}{T} dP - \frac{h}{T^2} dT
\]

where \( 1 \) and \( 2 \) stand for two set of values \( P_1, T_1 \) and \( P_2, T_2 \) and the integration is through an arbitrary (reversible) path in the \( P-T \) plane. In this way we can compare \( G \) on both sides of the first order line, performing an integration surrounding the critical point, and the first order line can be determined with good precision. This procedure is more reliable than the standard Maxwell construction, although the results we have obtained in this case are almost equivalent.

Concerning the crystalline structures, we know in fact that the present model shows a variety of different crystalline structures which are stable in different regions of the \( P-T \) plane. When testing for the stability of these structures, the fact that some of them take a very long time to appear spontaneously in simulations that start in the fluid phase is usually a problem, and then the
equilibrium coexistence line between these phases and the fluid has to be determined in some other way. We then applied the procedure already used in [11] consisting in the inclusion of an additional external potential, with an spatial periodicity corresponding to the crystalline lattice to be studied. The existence of this term (with a strength given by some parameter \( W \)) allows to construct a path in the \( P-T-W \) space that takes the system smoothly (i.e., without crossing any phase transition) from the fluid to the crystalline phase, and the difference in free energy \( \Delta G \) between the two phases at some fixed values \( P = P_0 \) and \( T = T_0 \) can be calculated by a generalization of expression (3), namely

\[
\Delta(G/T)|_{P_0,T_0} = \int_1^{T} \frac{v}{T} dP + \frac{e_W}{T W} dW - \frac{h + e_W}{T^2} dT, \quad (4)
\]

where \( e_W \) is the potential energy per particle in the external potential. In order to move the system from the crystalline to the fluid phase, the integration consists typically in the following close path: increasing \( W \) from zero to some large value; increasing \( T \) from \( T_0 \) to a large value; decreasing \( W \) down to 0; decreasing \( T \) down to \( T_0 \). In this way, the difference in free energy between the fluid phase and the possible crystalline phases was determined at fixed points in the \( P-T \) phase diagram. Then \( \Delta G \) can be calculated for the whole plane by thermodynamic integration of the pure phases. Using this procedure we have verified that the only crystalline phase that is thermodynamically stable within the metastability range is small. Typical isotherms around only close to the (liquid-gas) critical point, where the existence line can be numerically determined in a reliable way by the standard simulation methods we use. The liquid-gas critical point is estimated to be at \( P_{LG} \approx 0.02, T_{LG} \approx 0.37 \). Using the position of this critical point, we estimate the liquid-gas coexistence line by fitting to a van der Waals equation, chosen to give the correct values of \( T_{LG} \) and \( P_{LG} \). This is the line that is seen in the right part of Fig. 6 (note the change in the horizontal scale in this sector). At temperatures (\( \sim 0.1 \)) where we observe liquid-liquid coexistence, the transition pressure to the gas phase is of the order of \( 10^{-6} \). Then we are sure that the liquid-liquid transition occurs in a region in which liquid is thermodynamically stable with respect to gas.

The crystal-liquid-liquid triple point we obtained turns out into a gas-liquid-liquid triple point (still at equilibrium) by increasing the intensity of the attraction between particles. In fact, this moves all coexistence lines to lower pressures compared to the condensed-gas line, and at a certain point the crystal-liquid-liquid triple point is forced to be in the metastable region with respect to the gas phase. At this point, the gas-condensed line, together with the liquid-liquid line determine a liquid-liquid-gas triple point. For instance, increasing \( \gamma \) from 0.27 to 0.285 produces a liquid-liquid-gas triple point located at \( P \approx 10^{-6}, T \approx 0.10 \). The liquid-liquid critical point is in this case located at \( P \approx 0.03, T \approx 0.12 \). At still higher values of the attraction the whole liquid-liquid line moves into the metastable \( P < 0 \) region, and the equilibrium liquid-liquid transition is lost.

### B. Three dimensions

For the 3D case, we show in Fig. 7 the isotherms obtained for a system of 300 particles. They again show the existence of the liquid-liquid critical point. The curves are smooth for \( T > 0.08 \), whereas there is a jump in \( v \) at the coexistence pressure for \( T \) lower than this value. Also apparent from these curves is the existence (as in 2D) of density anomalies for the low density liquid.

The structure factor \( S(k) \) of the 3D liquid is shown for the two coexisting liquids at \( T = 0.075 \) in Fig. 8. The two main peaks corresponding to interparticle distance \( r_0 \) and \( r_1 \) are indicated. There is an abrupt change in the relative weight of these two peaks at the liquid-liquid transition, corresponding to an abrupt change in the mean distance between first neighbor particles.

For the simulations shown in Fig. 8 we did not find evidence of the appearance of any crystalline structure, suggesting that the liquid-liquid coexistence occurs at equilibrium. To be sure, we should investigate the stability of the different possible crystalline structures, as we did in the 2D case. For the 3D system however, this is a much harder work, as we actually do not know all the crystalline structures for this case (see [12]). Then we take another route, seeking for other indicators of stability of the liquid phase, to rule out the possibility that we are observing a liquid-liquid transition in a metastable region. So we will look for the fulfillment of the Hansen-Verlet and/or Lowen criteria for crystallization. Although they stand only as approximate, and mainly heuristic, they have been verified in a variety of cases. For simple (i.e. Lennard Jones) systems, the Hansen-Verlet criterion states that freezing occurs when the first peak of \( S(k) \) reaches a (quasi-)universal value
of $\sim 2.85$. A straightforward generalization of this criterion to ‘anomalous’ fluids tells that this relation should be checked for all peaks in $S(k)$. The results for $S(k)$ of Fig. 8 show that all peaks are lower than this value, suggesting liquid stability. The Lowen criteria for freezing indicates that this occurs when the relation between long and short times single particle diffusion coefficient drops below a value of $\sim 0.1$. This dynamical criterion is independent of the nature of the crystalline structure the system freezes to. Although it has been proposed and verified in systems with Brownian dynamics, we have verified in simple cases (i.e. hard spheres) that it is also fulfilled with the Monte Carlo dynamics, and so we apply it tentatively here. The relation between long and short time values of the diffusivity $D/D_0$ (calculated with the Monte Carlo dynamics) is shown in Fig. 8 along the $T = 0.075$ and $T = 0.09$ isotherms. The relation $D/D_0$ is in the whole range well above the value of 0.1 expected at freezing. Then all indicators coincide in telling that the liquid-liquid coexistence we are observing occurs in truly thermodynamic equilibrium. From Fig. 8 we also see that the evolution of $D$ on density shows a maximum which is well known to occur in tetrahedrally coordinates materials and it is also typical of core-softened potentials.

IV. CONCLUSIONS

In conclusion, we have shown that a system of identical, spherical, and rigid particles is able to display liquid-liquid coexistence at equilibrium (both in two and three dimensions). We showed explicitly this for an interaction potential that favors two different equilibrium distances between particles. The study of the present (or some related) model displaying liquid-liquid coexistence at equilibrium is likely to contribute to the understanding of the same phenomenon in cases where it occurs in the supercooled region, where it is much harder to study due to metastability against crystallization and long relaxation times. Our findings also open the possibility of observing liquid-liquid coexistence in monodisperse systems of rigid and spherical particles, i.e., colloidal systems, where interaction potentials with more than one equilibrium distance are easily obtained.

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FIG. 1. Isotherms from the simulation of a 2D system of 200 particles, at temperatures as indicated (symbols are the simulated points, lines are guides to the eye). There is a critical point at $T \approx 0.10, P \approx 0.07$, below which the isotherms get a typical van der Waals loop. Note the expansion upon decreasing $T$ at constant $P$ (a density anomaly) in the results for $v/r_0^2 > 2.1$. 

$T = 0.1050$

$\begin{array}{c}
0.1025 \\
0.1000 \\
0.0975 \\
0.0950 \\
0.0925 \\
0.0900 \\
0.0875 \\
\end{array}$

$P$

$\begin{array}{c}
0.15 \\
0.10 \\
0.05 \\
0.00 \\
\end{array}$

$\begin{array}{c}
1.4 \\
1.6 \\
1.8 \\
2.0 \\
2.2 \\
2.4 \\
\end{array}$

$v/r_0^2$
FIG. 2. Comparison of isotherm obtained in simulations of systems with 200 and 1000 particles. They are coincident, except within the first order loop, where the one corresponding to the larger system is more flat. This is the right tendency since the isotherm at coexistence is absolutely flat in the limit of systems of infinite size.

FIG. 3. Snapshots of the system of 1000 particles at the points indicated in Fig. 2. Black dots represent the strict hard core of the particles at $r_0$. In (a) and (c) the system is almost completely in one of the two liquid phases (HDL and LDL, respectively). In (b) there is a clear coexistence of the two phases.

FIG. 4. Pressure-temperature phase diagram of the two-dimensional model showing the existence of the liquid-liquid and liquid-gas critical points. The temperature of maximum density (TMD) is indicated.

FIG. 5. Isotherms near the liquid-gas transition for a two-dimensional system of 500 particles.
FIG. 6. Isotherms of a 3D system of 300 particles near the liquid-liquid critical point (located at $T \sim 0.08$, $P \sim 0.038$). Symbols are the simulated points, lines are guides to the eye.

FIG. 7. Structure factor of the two coexisting liquid phases of the 3D system at $T = 0.075$, $P \sim 0.032$.

FIG. 8. Relation between long and short time diffusivity $D/D_0$ along the $T = 0.075$ and $T = 0.09$ isotherms, for the 3D system.