Which mechanism underlies the water-like anomalies in core-softened potentials?

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ermington and Debenedetti [5] and Netz et al. [4] found, in SPC/E water, that there exists a hierarchy between the density and diffusion anomalies as follows. The diffusion anomaly region, inside which the mobility of particles grow as the density is increased, englobes the density anomaly region, inside which the system expands upon cooling at constant pressure. Even thought these observations are supported by experimental results [3], there is no clear explanation why the thermodynamic and dynamic anomalies should be found in the same locus of the pressure temperature phase-diagram.

Realistic simulations of water [7–9] have achieved a good accuracy in describing the thermodynamic and dynamic anomalies of water. However, due to the high number of microscopic details taken into account in these models, it becomes difficult to discriminate what is essential to explain the anomalies. On the other extreme, a number of isotropic models were proposed as the simplest framework to understand the physics of the liquid state anomalies. From the desire of constructing a simple two-body isotropic potential capable of describing the complicated behavior present in water-like molecules, a number of models in which single component systems of particles interact via core-softened (CS) potentials have been proposed. They possess a repulsive core that exhibits a region of softening where the slope changes dramatically. This region can be a shoulder or a ramp [10–22]. Ramp and continuous shoulder-like potentials exhibit thermodynamic, dynamic, and structural anomalies [20–22]. However the square discontinuous shoulder shows no thermodynamic

Abstract. Using molecular dynamics simulations we investigate the thermodynamics of particles interacting with continuous and discrete versions of a core-softened (CS) intermolecular potential composed by a repulsive shoulder. Dynamical and structural properties are also analyzed by the simulations. We show that in the continuous version of the CS potential the density at constant pressure has a maximum for a certain temperature. Similarly the diffusion constant, $D$, at a constant temperature has a maximum at a density $\rho_{\text{max}}$ and a minimum at a density $\rho_{\text{Dmin}} < \rho_{\text{Dmax}}$, and structural properties are also anomalous. For the discrete CS potential none of these anomalies are observed. The absence of anomalies in the discrete case and its presence in the continuous CS potential are discussed in the framework of the excess entropy.

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1 Introduction

While the majority of fluids contract upon cooling, water expands when cooled below $T = 4^\circ\text{C}$ at atmospheric pressure [1]. This effect is called density anomaly. Besides the density anomaly, there are more than sixty other anomalies known for water [2]. The diffusivity is one of them. For normal liquids the diffusion coefficient, $D$, decreases under compression. However, experimental results have shown that for water at temperatures approximately below 10°C the diffusion coefficient increases under compression and has a maximum [3]. The temperature of maximum density (TMD) line inside which the density anomaly occurs, and the line of maximum in diffusivity are located in the same region of the pressure-temperature (P-T) phase diagram of water. Simulations for water also show thermodynamic and dynamic anomalies. The simple point charged/extended (SPC/E) model for water exhibits a TMD line in the P-T phase diagram. The diffusion coefficient has a maximum and a minimum that define two lines at the P-T phase diagram, the lines of maximum and minimum in the diffusivity coefficient [4–6]. Similarly to the experimental results, the TMD and the lines of maximum and minimum in the diffusion are located at the same region at the P-T phase diagram for the SPC/E model. Errington and Dehenedetti [5] and Netz et al. [4] found, in SPC/E water, that there exists a hierarchy between

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anomaly in three dimensions [11]. All these potentials have in common the presence of two representative repulsive scales in the potential, \( \sigma \) and \( \sigma_1 \), where the closest scale, \( \sigma < \sigma_1 \), has the higher potential energy \( U(\sigma) > U(\sigma_1) \).

One question that arises in this context is why a continuous shoulder potential like the one described by de Oliveira et al. [20,21] has density and diffusion anomalies while the square shoulder described by Franzese et al. [11] has no anomaly? Moreover, why are the thermodynamic and the dynamic anomalies linked? In order to shade some light in the reasons for the presence of the density and the diffusion anomalies in CS potentials, both the discontinuous shoulder potential and the continuous version are analyzed in the framework of the excess-entropy-based formalism [23] recently applied to similar systems by Errington et al. [24] and Chakraborty and Chakravarty [25]. Within this approach the presence of the density and the diffusion anomalies are related to the density dependence of the excess entropy, \( s_{\text{ex}} \). We will follow this surmise and compute the excess entropy for both the discontinuous and the continuous shoulder potentials. From the analysis of \( s_{\text{ex}} \) for the discontinuous model we are able to derive a simple argument for the presence or absence of anomalies in CS potentials.

The outline of the paper is as follows. We present the details of the two models in Section 2, details of the simulations, and the P-T phase-diagram for both models are presented in Section 3. In Section 4 comparisons between the behavior of the excess entropy of the continuous and discontinuous models are made and from that a condition for the presence or absence of anomalies in CS potentials is proposed. Conclusions end these sections.

2 The models

The first model we study here consists of a system of \( N \) particles of diameter \( \sigma \) inside a cubic box whose volume is \( V \), resulting in a density number \( \rho = N/V \) [20] interacting with a continuous shoulder potential given by

\[
U^*(r) = 4 \left[ \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^{6} \right] + a \exp \left[ - \frac{1}{c^2} \left( \frac{r - r_0}{\sigma} \right)^2 \right],
\]

where \( U^*(r) = U(r)/\epsilon \). The first term of equation (1) is a Lennard-Jones potential of well depth \( \epsilon \) and the second term is a Gaussian centered on radius \( r = r_0 \) with height \( a \) and width \( \epsilon \). In previous publications we have studied this model with setting \( a = 5, r_0/\sigma = 0.7 \) and \( c = 1 \) (see Fig. 1) [20,21,26]. This potential has two natural length scales: one close to the hard core, \( \sigma \), and another at a further distance where the potential has its lower value. This last length we call \( \sigma_1 \).

The second model we study here is a system of \( N \) particles of diameter \( \sigma \) inside a cubic box whose volume is \( V \), resulting in a density number \( \rho = N/V \) but interacting with a discontinuous shoulder potential given by

\[
U^*(r) = \begin{cases} 
\infty & r < \sigma \\
1 & \sigma_1 > r > \sigma \\
0 & r > \sigma_1.
\end{cases}
\]

where \( U^*(r) = U(r)/\epsilon \). This potential has two natural length scales: the hard core distance, \( \sigma \), and the outer core, \( \sigma_1 \). Here we analyze the case \( \sigma_1/\sigma = 1.75 \) illustrated in Figure 2. Here we use dimensionless pressure, \( P^* \), temperature, \( T^* \), and density, \( \rho^* \), that are given in units of \( \sigma^3/\epsilon, k_B/\epsilon \), and \( \sigma^4 \), respectively and \( k_B \) stands for the Boltzmann constant.

3 Details of simulations

For the continuous shoulder potential we performed molecular dynamics simulations in the canonical ensemble using 500 particles. Detail of the simulations can be

![Fig. 1](image1.png)

**Fig. 1.** Interaction potential from equation (1) with parameters \( a = 5, r_0/\sigma = 0.7 \) and \( c = 1 \), in reduced units.

![Fig. 2](image2.png)

**Fig. 2.** Interaction potential from equation (2) with parameter \( \sigma_1/\sigma = 1.75 \) in reduced units.
found in reference [20]. Figure 3 shows the P-T phase-diagram we have obtained in a previous publication [20]. The isotherms have minima that define the temperature of maximum density. The TMD line encloses the region of density anomaly. We also have studied the mobility associated with the potential described in equation (1) [20]. The diffusion was also calculated using the mean-square displacement averaged over different initial times. The behavior of $D$ as a function of $\rho$ goes as follows. At low temperatures, the behavior is similar to the behavior found in SPC/E supercooled water [4]. The diffusivity increases as the density is lowered, reaches a maximum at $\rho_{\text{Dmax}}$ (and $D_{\text{Dmax}}$) and decreases until it reaches a minimum at $\rho_{\text{Dmin}}$ (and $D_{\text{Dmin}}$). The region in the P-T plane where there is an anomalous behavior in the diffusion is bounded by ($T_{\text{Dmin}}, P_{\text{Dmin}}$) and ($T_{\text{Dmax}}, P_{\text{Dmax}}$) and their location is shown in Figure 3. The region of diffusion anomalies ($T_{\text{Dmax}}, P_{\text{Dmax}}$) and ($T_{\text{Dmin}}, P_{\text{Dmin}}$) lies outside the region of density anomalies like in SPC/E water [4].

In order to simulate the discrete potential shown in Figure 2 we used collision driven molecular dynamics techniques [27]. 500 particles were put into a simulation box with periodic boundary conditions and the rescaling velocities scheme was used for every 1000 collisions until equilibration time to achieve the desired temperature. After thermalization particles were allowed to move under microcanonical ensemble. In the collision driven molecular dynamics simulation, kinetic energy has to be rigorously conserved. Hence, no special mechanism is necessary in order to simulate a desired temperature and the NVE ensemble becomes the natural choice. The equilibration and production times in reduced units were 350 and 650 respectively. Figure 4 shows the P-T phase-diagram for the discontinuous shoulder potential. The isochors for different temperature and pressures show no minima so no density anomaly is present.

**4 Excess entropy and anomalies**

Why the discontinuous shoulder potential has no density or diffusion anomalies and its continuous counterpart has both of them? We can gain some understanding about the presence or absence of these anomalies and the shape of the potential by analyzing the density dependence of the excess entropy [24]. The excess entropy is defined as the difference between the entropy of the real fluid and that of an ideal gas at the same temperature and density. Errington et al. have shown that the density anomaly is given by the condition $\Sigma_{\text{ex}} = (\partial s_{\text{ex}}/\partial \ln \rho)_T > 1$ [24]. They have also shown that the two body contribution of $s_{\text{ex}}$,

$$s_{\text{ex}} \approx s_2 = -2\pi \rho \int [g(r) \ln g(r) - g(r) + 1] r^2 dr,$$

(2)

gives a good approximation of $s_{\text{ex}}$. Here we will show that the use of this approximation also shade some light in the link between the thermodynamic and dynamic anomalies. The radial distribution function, $g(r)$, is proportional to the probability to find a particle at a distance $r$ to another particle placed at the origin. Errington et al. [24] have also suggested that the diffusion anomaly can be predicted by using the empirical Rosenfeld's parameterization [28]. They found the condition $\Sigma_2 > 0.42$ for a diffusion anomalous behaviour. They also claim that $\Sigma_2 > 0$ is a good estimate for determining the region where structural anomaly occurs [24].

In order to understand the differences between the continuous and the discontinuous shoulder potentials we test the excess entropy criteria described above in both potentials. The radial distribution functions for different temperatures and densities for both potentials were then obtained by the molecular dynamic simulation method described in Section 3.

Figure 5 illustrates the two-body contribution of excess entropy for the continuous potential given by equation (1). $s_2$ is negative and its slope changes from negative to positive what indicates the presence of structural
anomaly. Figure 6 shows the behavior of $\Sigma_2$ with density for a fixed temperature for the continuous model. The horizontal lines at $\Sigma_2^i = 0, 0.42,$ and $1$ indicate the threshold beyond which there are structural, diffusion, and density anomalies respectively. In accordance with Figure 3 the density anomalous region shown in Figure 6 occurs in an interval of density smaller than the interval where the diffusion is anomalous. The two-body excess entropy also show the presence of structural anomaly what corroborates results of simulations of structural parameters [21].

Figure 7 illustrates the two-body excess entropy of the discontinuous potential given by equation (2). $s_2$ is negative and its slope negative is for almost all temperatures and densities. The filled circles show the region where the system crystallization occur. Figure 8 shows the behavior of $\Sigma_2$ with density for fixed temperatures for the discontinuous model. The line $\Sigma_2^1 = 1$ is never crossed so no density anomaly should be expected what is in good agreement with the Figure 4. The line $\Sigma_2^i = 0.42$ is also never crossed what indicates that diffusion anomaly is not expected in the discontinuous model. This is also in agreement with results for similar step potentials where no diffusion anomaly is found for large steps [29]. The line $\Sigma_2 = 0$ is crossed for temperatures $T^* = 0.2$ what would suggest the presence of structural anomaly. However this has to be taken with a grain of salt since Errington et al. [24] demonstrated that $s_2$ overestimates the region of anomalies. In this sense, a detailed study of translational and orientational order parameters [21] is necessary prior to any affirmative on the presence of structural anomaly for this discontinuous shoulder.
Even thought the excess entropy criteria is quite useful for predicting if the anomalies would be present for a certain potential it does not provide an easy and intuitive method for understanding why the continuous shoulder has anomalies and the discontinuous one does not have. In principle, both potential exhibit similar characteristics. Both potentials have two repulsive scales and consequently the radial distribution function in both cases has two peaks, one close to the hard core and another close to the distance \(r = \sigma_1\) as illustrated in Figure 9 for the continuous potential and in Figure 10 for the discontinuous potential. A closer look at the radial distribution reveals an important difference between the two cases. For the continuous potential for densities and temperatures if \(T^* \approx \rho^* \approx \epsilon/\sigma^2\) has two repulsive scales and consequently the radial distribution function in both cases \(g(r)\) increases with density both at the hard core and at \(r = \sigma_1\) for any temperature and density (see Fig. 10 for example). Notice that the radial distribution function in both cases has significative changes with the change in density at the two natural scales.

Finally we propose that a two scale potential has thermodynamic, dynamic and structural anomalies for some temperature and densities if \(\partial g(r)/\partial \rho > 0\) for \(r \approx \sigma\) and \(\partial g(r)/\partial \rho < 0\) for \(r \approx \sigma_1\). If this requirements would not be fulfilled for any temperature and density no anomaly would be present. This proposition is based on the physical picture that within the anomalous region for a fixed temperature an increase in density implies an increase in the number of particles close to the hard core. This particles move from the distance \(\sigma_1\) to \(\sigma\). For a discontinuous potential this requires an activation energy of \(U = \epsilon\) while for the continuous case it can be done continuously. As it was shown by Netz et al. [29] for the steps potential, anomalies would only be observed if the discontinuity in \(U^*\) would be below a certain threshold.

Now we shall test if this simple hypothesis is in agreement with Errington et al. criteria. First we compute \(\Sigma_2\) as

\[
\Sigma_2 = \left( \frac{\partial s_2}{\partial \ln \rho} \right)_T = s_2 - 2\pi \rho^2 \int g(r) \frac{\partial g(r)}{\partial \rho} r^2 dr
\]

In this expression, the first term is always negative for all two scale potentials. The integral in the second term, as we have observed above, is dominated by the the values at the two scales, \(\sigma\) and \(\sigma_1\).

For the continuous potential in the region where the anomaly is present \(\ln g(r \approx \sigma) < 0\) while \(\ln g(r \approx \sigma_1) > 0\) (since \(g(r \approx \sigma) < 1\) and \(g(r \approx \sigma_1) > 1\) respectively). Also \(\partial g(\sigma)/\partial \rho > 0\) while \(\partial g(\sigma_1)/\partial \rho < 0\). Consequently the second term in equation (3) is positive. This allows for a zero or positive values of \(\Sigma_2\) for appropriated densities and temperatures. Therefore our criteria is in accordance with Errington et al. criteria.

In resume, in this paper we have calculated the excess entropy and its derivative for both continuous and discontinuous shoulder potential. For the continuous case, using the Errington et al. criteria indicates that this potential has density, diffusion and structural anomalies as we have shown in previous publications. For the discontinuous potential the criteria indicates that no thermodynamic and dynamic anomalies are present and its not conclusive for structural anomalies. Direct calculations of the P-T temperature phase-diagram confirms the excess of entropy prediction. On basis of these results we propose a criteria for predicting if a two scale potential has or not anomalies. Our criteria provides a simple picture for the anomalies not being observed in the discontinuous shoulder potential.

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