Structure and Correlations for Harmonically Confined Charges

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(Dated: April 5, 2022)

Abstract

Coulomb charges confined by a harmonic potential display a rich structure at strong coupling, both classical and quantum. A simple density functional theory is reviewed showing the essential role of correlations in forming shell structure and order within the shells. An overview of previous comparisons with molecular dynamics and Monte Carlo simulations is summarized and extended.

PACS numbers:
I. INTRODUCTION

The problem considered here is the structure and correlation among $N$ equal Coulomb charges confined by an external harmonic potential. It is a generalization of the Thomson problem [1] for charges confined to the surface of a sphere, posed 117 years ago, to three dimensions and finite temperatures. The ground state for the harmonic confinement is well-studied, exposing a rich shell structure (distribution of particles localized about well-defined radii with localization on each radius similar to those of the single sphere Thomson problem). Within classical mechanics, these ground state results have been quantified in detail via shell models, molecular dynamics simulation (MD), and Monte Carlo simulation (MC) [2–7]. They are also realized experimentally for dusty plasmas. The corresponding results for confined charges at finite temperatures is the extension described here. We provide here a summary of our theoretical work in collaboration with the Bonitz group at the Institut fur Theoretische Physik und Astrophysik, Christian-Albrechts Universitat, Kiel [8–11]. In addition we describe the method for extension to quantum theory and list some remaining outstanding problems.

The primary effect of temperature on the classical ground state shell models is to broaden the sharp shell structure and smooth their angular distribution due to thermal motion. The governing parameters are the Coulomb coupling constant $\Gamma$ (ratio of Coulomb to thermal energies of a pair) and the average number of charges $\overline{N}$. The number of shells is fixed by $\overline{N}$ while their relative resolution (sharpness) is determined by $\Gamma$. The zero temperature ground state results from shell models are recovered in the limit of large $\Gamma$. A simple approximate density functional theory described below is able to capture these results quantitatively, in comparison with those from Monte Carlo simulations. It is based on approximating correlations among the charges in the trap by those in the uniform one component plasma. The close relationship of correlations in these two quite different systems has been confirmed by MD simulation [10, 11].

At still larger $\Gamma$, corresponding to lower temperatures, the rotational invariance of the fluid phase is broken and the particles within each shell become localized about sites close to those of the Thomson problem for a single shell. Those localized domains are approximated here by Gaussian distributions centered at these sites, and the correlations among them are calculated showing good agreement with results from MC simulation.
The effects studied here result from strong coupling conditions for which there are relatively few theoretical methods available. In the classical case the density functional model described below is confirmed by MD and Monte Carlo simulations. The latter simulations are not available for the quantum case, and the quantum density functional model has problems at finite temperatures. However, it has been shown that the quantum system can be mapped onto a corresponding classical system with quantum effects embedded in effective Coulomb and trap potentials [12]. Applications to the one component plasma (jellium) show good agreement with quantum Monte Carlo results [13, 14]. This approach has been applied subsequently to the case of quantum charges in a harmonic trap [15, 16] as described below. In this way a broad scope of confined Coulomb systems of interest can be addressed. Figure 1 gives a simple overview of the parameter space.

It is a pleasure to dedicate this contribution to our friend and colleague of many years, Professor John (Jack) Sabin. Jack has been an inspiration for all that is expected of those with an academic career, exemplifying the best in teaching, research, and administration. His cheerful nature and good will have brightened the lives of all who knew him well.

II. DENSITY FUNCTIONAL THEORY

Consider $N$ particles of charge $q$ at equilibrium in a harmonic trap at inverse temperature $\beta$. The free energy is a function of $\beta$ and a functional of the non-uniform density $n(r)$, $F(\beta \mid n)$. The equilibrium density profile is determined from

$$\frac{\delta F(\beta \mid n)}{\delta n(r)} = \mu - V(r), \quad V(r) = \frac{1}{2}m\omega^2r^2. \quad (1)$$

The potential $V(r)$ is the confining harmonic trap. The free energy functional can be separated into that for a system without interactions, $F_0(\beta \mid n)$, and a remainder $F_{ex}(\beta \mid n)$ containing all effects of Coulomb interactions among the particles

$$F(\beta \mid n) = F_0(\beta \mid n) + F_{ex}(\beta \mid n). \quad (2)$$

A formal representation for the excess free energy in terms of pair correlations can also be written exactly

$$F_{ex}(\beta \mid n) = -\int_0^1 dy (1-y) \int d\mathbf{r}d\mathbf{r}' n(\mathbf{r}) n(\mathbf{r'}) \beta^{-1} c^{(2)}(\mathbf{r}, \mathbf{r}' \mid yn), \quad (3)$$
FIG. 1: Overview of the relevant parameter space. Here \( r_s \) is the Wigner-Seitz radius in terms of the Bohr radius of the confined particles (\( r_s = r_0/a_b \)), and \( t \) is the temperature relative to the ideal gas Fermi temperature per particles (\( t = k_B T/\varepsilon_F \)) [16].

where \( c^{(2)}(\mathbf{r}, \mathbf{r}' \mid n) \) is the direct pair correlation function

\[
\beta^{-1} c^{(2)}(\mathbf{r}, \mathbf{r}' \mid n) \equiv -\frac{\delta F_{\text{ext}}(\beta \mid n)}{\delta n(\mathbf{r}) \delta n(\mathbf{r}')} .
\] (4)

In this way Eq.(1) is an equation for the local density in terms of the pair correlations of the direct correlation function [9]

\[
\frac{\delta F_0(\beta \mid n)}{\delta n(\mathbf{r})} = \mu - V(\mathbf{r}) + \int_0^1 dy \int d\mathbf{r}' n(\mathbf{r}') \beta^{-1} c^{(2)}(\mathbf{r}, \mathbf{r}' \mid y n) .
\] (5)

The average density \( \bar{n} \) is defined by

\[
\bar{n} = \frac{N}{V} , \quad \bar{N} = \int d\mathbf{r} n(\mathbf{r}) .
\] (6)
The system is self-confined with spherical symmetry. The maximum radius $R$ is the point at which Coulomb repulsion force on a particle is balanced by the harmonic trap confinement

$$\mathcal{N} \frac{q^2}{R^2} = m\omega^2 R, \quad V = \frac{4}{3}\pi R^3.$$  \hspace{1cm} (7)

The mean distance between particles $r_0$ is introduced by

$$\frac{4}{3}\pi r_0^3 = 1.$$ \hspace{1cm} (8)

Scaling the coordinates with respect to $r_0$ in the above equations gives the dimensionless form

$$\frac{\delta F^*_0(n^*)}{\delta n^*(r^*)} = \beta \mu - \frac{1}{2} \Gamma r^* + \int_0^1 dy \int d(r^* n^*(r'^*)) e^{(2)}(r^*, r'^*; y n^*),$$ \hspace{1cm} (9)

where

$$F^*_0(n^*) = \beta F_0(\beta | n); \quad n^*(r^*) = n(r) r_0^3, \quad \Gamma = \beta m\omega^2 r_0^2 = \beta \frac{q^2}{r_0}.$$ \hspace{1cm} (10)

The parameter $\Gamma$ is the Coulomb coupling constant (Coulomb energy of a pair at the average distance relative to the thermal energy $\beta^{-1}$). The constant $\beta \mu$ can be eliminated in terms of $\mathcal{N}$.

Up to this point the results apply for both quantum and classical mechanics. The classical case is considered more explicitly next.

III. CLASSICAL MECHANICS

Within classical statistical mechanics $F_0(\beta | n)$ can be written exactly as a functional of the density

$$F^{(0)}(\beta | n) = -\frac{1}{\beta} \int d\mathbf{r} n(r) \left( 1 - \ln (n(r)\lambda^3) \right).$$ \hspace{1cm} (11)

Here $\lambda = (\hbar^2 \beta / 2\pi m)^{1/2}$. Then eq.(9) becomes [9]

$$\ln (n^*(r^*)) = \ln \left( \left( \frac{\lambda}{r_0} \right)^3 e^{\beta \mu} \right) - \frac{1}{2} \Gamma r^* + \int_0^1 dy \int d(r^* n^*(r'^*)) e^{(2)}(r^*, r'^*; y n^*),$$ \hspace{1cm} (12)

The constant first term on the right side can be eliminated in terms of $\mathcal{N}$. Consequently the dimensional density profile and associated free energy density profile depend only on these two parameters.

The solutions to (12) are expected to confirm the following qualitative behavior observed from ground state energy functions [8], and Monte Carlo and MD simulations [7]. For given
N the profiles have a strong dependence on the coupling strength $\Gamma$. At very small values the density profile is rotationally symmetric and monotonically decreasing to zero from a maximum at $r^* = 0$. At increasing values the radial dependence develops local maxima, referred to as shells. This is illustrated in Figure 2. The number of shells increases with $N$ and their width sharpens with increasing $\Gamma$. The shell populations are greater for increasing radius and grow linearly with $N$.

Eventually, at sufficiently large $\Gamma$ rotational symmetry is broken. The uniform distribution of particles within each shell distorts to local domains for the associated population. Their locations are close to those of the Thomson problem - the ground state configuration for a given number of charges confined to a sphere. In the following the extent to which approximations to the density functional theory (12) captures these features is described.

A. Fluid phase

The approximate determination of the pair correlations described by the direct correlation functional in (12) is motivated as follows. First, it is shown elsewhere that the functional
FIG. 3: Comparison of the pair distribution of particles in a one-component plasma (OCP) and harmonic trap. MD calculations for the OCP were performed using Sarkas.[20, 21]

\( F_{ex}(\beta | n) \) is exactly equal to the corresponding system with a uniform neutralizing background (inhomogeneous jellium), i.e. charges in a uniform neutralizing background with the same harmonic potential [17]

\[
F_{ex}(\beta | n) = F_{jex}(\beta | n). \tag{13}
\]

The advantage of this is that the OCP has a finite uniform limit in the absence of the harmonic confinement, which is the uniform one component plasma OCP.

\[
c^{(2)}(\mathbf{r}, \mathbf{r}'; \lambda n) \rightarrow c^{(2)}(\mathbf{r}, \mathbf{r}'; \mathbf{\mu}) = c^{(2)}_{OCP}(\|\mathbf{r} - \mathbf{r}'\|; \mathbf{\mu}). \tag{14}
\]

It has been observed elsewhere [11] that the distribution of pairs within the trap without reference to their center of mass position are almost identical to those of the OCP, see Figure 3. Therefore, as an approximation for the fluid phase (14) OCP correlations have been used. Evaluation of the direct correlation function for the OCP still poses a formidable many body problem at strong coupling. However, it is a well-studied problem and an excellent approximation, the adjusted hypernetted chain approximation (AHNC), is known [18, 19]. The solutions to (12) with these two approximations for \( c^{(2)}(\mathbf{r}, \mathbf{r}'; n) \) give all of the above
expected properties for the density profile quantitatively in comparison to MD and MC results, across the entire domain of $\Gamma$ and $N$. An example is illustrated in Figure 4.

**B. Ordered phase**

In the fluid phase the particles are uniformly distributed throughout each shell. As the coupling increases eventually the particles enter an ordered state where rotational symmetry is broken within the shell. Figure 5 shows the angular correlations within the outer shell from a molecular dynamics simulation, for three values of the coupling constant corresponding to fluid and ordered phases. The system consisted of $N = 38$ charges, with 32 in the outer shell. The pair correlation function $g(\theta)$ is the probability to find a particle displaced on the shell by an angle $\theta$ from an arbitrary reference particle. As in a uniform fluid the peaks represent nearest neighbor, next nearest neighbor, etc. At the lower values of $\Gamma$ there is not much qualitative difference in the angular correlations, but the correlation peaks are narrowing and some structure is starting to develop in the form of shoulders in the middle peaks. For significantly larger values of $\Gamma$ however, a definite structure appears within each peak of the correlation graph. In particular, the two broad peaks at $\Gamma = 100$ that appeared at 70 and 110 degrees have condensed into triplets at $\Gamma = 1000$. In addition, the two peaks
FIG. 5: Angular correlations within a single shell from MD simulation. The harmonic trap contained 38 particles, with 32 in the outer shell. MD simulations were performed with LAMMPS.[22, 23]

at 40 and 140 degrees are showing the beginning formation of a doublet structure with the presence of shoulders at $\Gamma = 1000$. This set of doublet and triplet features persist at much higher $\Gamma$, without the appearance of any more peaks.

These features can be modeled using a thin-shell model where the particle configuration results from the Thomson problem (minimum energy configuration for charges constrained to a sphere). The specific ordering depends on the number of charges because of the spherical geometry. In Figure 6 the Thomson configuration for a randomly chosen particle from a system containing $N = 32$ charges are shown along with constant-angle planes showing the angular displacement of the other particles. To compare more directly, the angular correlations of all 32 charges in the Thomson problem were calculated. A plot of the angular displacement of all pairs is shown in Figure 7. The charges occur in two different angular correlation structures. These are shown in Figure 8. The combination of these two populations account for the specific angular correlation structure for the case of $N = 32$ charges.

To account for thermal effects the charges were modeled using a Gaussian function along
FIG. 6: A system of 32 charges in the Thomson problem. Horizontal circles connect those charges that are at constant angle $\theta$ from the chosen particle at the top of the figure.

The sphere of the form

$$f(\theta) = \sqrt{\frac{\alpha}{\pi}} \exp\left(-\alpha(\theta - \theta_0)^2\right)$$

(15)

where $\theta_0$ is the angle corresponding to the Thomson site and $\alpha$ is a parameter that increases with decreasing temperature. Fig 9 shows how this model reproduces the correct splitting, from the four broad peaks at small $\Gamma$ which condense to the characteristic doublets and triplets at higher $\Gamma$. Figure 10 shows the effect of the fitting parameter $\alpha$ showing how it models the effect of thermal motion. This supports the idea of considering the Thomson sites on a sphere to be analogous to a fundamental lattice for the ordered state, to the extent that the shell can be approximated as thin.

In principle, the value of $\alpha$ in the ordered phase, for given $\Gamma, \overline{N}$ should be obtained from minimizing the above free energy functional using the assumed Gaussian density profile. In the fluid phase the value of $\alpha$ would be large, representing a uniform profile. At very large $\Gamma$, approaching the ground state, the value of $\alpha$ would approach zero. However, the assumption of the Thomson sites should first come from solutions of (12). This could be quite difficult since ground state studies from simulations suggest there are many metastable
FIG. 7: Number of each angle present between each pair of particles in the Thomson system for $N = 32$ charges.

configurations as well.

IV. QUANTUM MECHANICS

The above classical description of confined charges at strong coupling has exploited the methods of density functional theory, liquid state theory, MD, and MC. At low temperatures, at or below the Fermi temperature, quantum effects become important and many of these classical methods do not apply directly. An accurate quantum theory at finite temperatures, strong coupling, and confinement is still a challenging problem. Numerical methods such as quantum Monte Carlo are applicable at zero temperature but become less controlled as the temperature increases, particularly for fermions. A quite different approach is to develop an exact mapping of the quantum equilibrium structure to an effective classical problem. This has been done recently, allowing application of the above classical methods to quantum systems [12]. Structure and correlation calculated in this way for the OCP have shown good accuracy in comparison with recent quantum Monte Carlo simulations [13, 14].

More recently the effective classical representation of a system of quantum charges in a
FIG. 8: Angular distribution for the two types of angular configuration for $N = 32$ charges in the Thomson problem. The first type of angular configuration has particles at eight specific angles between $\theta = 0^\circ$ and $\theta = 180^\circ$. There are 20 particles in this configuration. The remaining 12 particles are in a configuration with six specific angles to the other particles. Plots show the total number of bonds that each population contributes for the entire system; their sum gives the results in Figure 7.

The harmonic trap has been explored [15, 16]. The primary differences from the classical description above are modifications of the Coulomb potential and the trap potential to accommodate quantum effects of diffraction and exchange symmetry. A new parameter appears, in addition to $\Gamma$ and $\overline{N}$, the temperature relative to the Fermi temperature ($t = k_B T/e_F$ where $e_F$ is the ideal gas Fermi energy per particle). For $t \gg 1$ the above description of
FIG. 9: Solid lines: evolution of the thermally-broadened angular correlations from the Thomson problem as the parameter $\alpha$ increases. Peak height increases with larger $\alpha$. Crosses: comparative results from MD at $\Gamma = 250$.

FIG. 10: Angular correlations from the thermally-broadened Thomson system for two values of the fitting parameter $\alpha$. Here there are $N = 32$ charges in the system. The structure follows the same behavior as MD simulations for 32 particles in a shell.
shell structure and correlations is recovered for strong coupling. At smaller $t$ the effects of exchange degeneracy are incorporated by imposing the exact noninteracting density profile for the ideal gas. This is non-trivial since the classical representation of the quantum ideal gas has effective interactions. The additional effects of exchange and diffraction are included via the direct correlation function with modified Coulomb interactions. Figure 11 shows a self-similar change in the classical two shell structure being compressed due to quantum effects on the harmonic potential. There is much more to be done with this classical description of a quantum system in the low $t$ domain. At much smaller $N$ connection to other studies of quantum dots and ultracold gases should be useful. Other properties such as spin polarization, coherent control of trap properties, charge dependence and others acceptable to direct observation can be addressed. A different direction for application of the results here is obtained by the replacement of the harmonic trap with a Coulomb potential to calculate the electron distribution about an ion. This is a solved problem of quantum chemistry, but its extension to a random configuration of ions is of intense current interest for warm, dense-matter applications, e.g. the electron density in the presence an ion configuration. Such densities are required to compute the forces in quantum molecular dynamics simulations for the ions in warm, dense matter at finite temperatures where traditional density functional methods fail (e.g. the traditional Kohn-Sham self-consistent equations for temperatures near
the Fermi temperature). Here those self-consistent equations are replaced with the classical integral equations of AHNC. This advantage has been stressed by Dharma-wardana [24].

V. DISCUSSION

The extreme conditions of long range Coulomb charges, confined at strong coupling and finite temperatures lead to complex structures: radial shell structure and broken symmetry angular ordering within the shells. At the classical conditions a density functional representation with strong coupling correlations from the OCP is able to capture quantitatively the transition from simple uniform filling at weak coupling to the formation of "atomic" shell structure. At infinite coupling the ground state ordered state is closely related to the Thomson problem (sharp shell radii, no interactions between shells), extended here to finite temperature (e.g., like Debye-Waller broadening). The classical fluid phase with uniform angular distribution is now well studied by MC, MD, and theory - the features discussed in reference [9] are given quantitatively as a function of $\Gamma, \bar{N}$, e.g. number of shells, occupancy, amplitude, location. The classical ordered phase also is well studied in the ground state, but less so at finite temperatures. The onset of localization within shells seems not to be sharp but rather gradual, as is the formation of shells, as a function of $\Gamma$. The Thomson sites associated with the occupation number for a given shell provide a good reference for this localization, as confirmed by ground state minimum energy models. The latter models, and simulation, indicated that there are metastable configurations with similar energy so the minimization requires care. It is possible that a more controlled limit is obtained from the limit of finite temperature studies as described here.

The quantum case is well-studied at the ground state in the context of quantum dots, nanodevices, and related systems. The case of higher temperatures and transition to classical behavior is more limited, both from theory and simulation. The classical map method described here is particularly well-suited for this domain, but has not been explored very much. Some of the advantages for states of warm, dense matter have been outlined in reference [24]. Another interesting question is the line for Wigner crystallization (see Figure 1), whose location is known only at $T = 0$ and in the high temperature classical limit.
VI. ACKNOWLEDGEMENT

Much of the work summarized here was done in collaboration with the Bonitz group at Christian Albrechts University. The work of JW and JD was supported by US DOE Grant DE-SC0002139.

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