Variational average atom in quantum plasmas (VAAQP)—first numerical results

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Abstract

A new code called VAAQP (variational average atom in quantum plasmas) is briefly described and its first results in the case of aluminium at solid density and temperatures between 0.05 and 12 eV are reported. The code is based on a new fully variational approach to plasmas at local equilibrium with both bound and free electrons treated quantum mechanically. This model which is derived from two first terms of the cluster expansion appears to be the quantum extension of the well-known atom-in-cell model based on the Thomas–Fermi theory (Thomas–Fermi average atom) that has been proposed in 1949 by Feynman, Metropolis and Teller. Similar to the case of Feynman et al.

\textsuperscript{1}’s model the VAAQP approach, due to its fully variational character, respects the virial theorem and uses a simple formula for the electronic pressure. Comparisons to results obtained using other approaches are also shown and discussed in the aluminium case. The results confirm the feasibility of the quantum variational model in the warm dense matter regime. Effects of the variational treatment can lead in this regime to significant differences with respect to existing non-variational models.

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1. Introduction

We present here a code called VAAQP which is a numerical application of a new fully variational plasma model with all quantum electrons. Models of plasmas at local thermodynamic equilibrium with quantum bound and free electrons are necessary in order to calculate the photo-absorption cross sections and the equation of state in the modelling of inertial fusion devices, in astrophysics, in laboratory plasma studies and in warm dense matter experiments. Up to now the existing average-atom models in which all electrons
are treated as quantum [1–3] have not been fully variational in the sense that they were not respecting the virial theorem. Recently, a fully variational model with all quantum electrons respecting the virial theorem has been proposed [4, 5]. This variational model is in our opinion a first relatively simple and coherent model of plasma matter with quantum electrons. Atomic structure calculations and electronic pressure obtained from this model are in principle thermodynamically coherent. The structure elements can be used further to calculate plasma radiative and transport properties consistent with the equation of state.

One may expect our model to work well in the case of strongly coupled plasmas in which the ion–ion correlation function has roughly the form of a cavity (as it will be explicitly stated in section 2). We focus our attention here on the electronic structure. This model may be supplemented by different approximations to ion free energy and pressure contributions.

Among the most interesting applications of plasma models are cases corresponding to the warm dense matter (WDM) regime. Special attention is thus paid to matter at high density (i.e. solid density) and relatively low temperature (around a few eV or below). However this regime is also the most difficult to account for in the VAAQP code. The main reason for this difficulty is the presence of the long-range Friedel oscillations. For this reason in this preliminary study we focus our attention on a WDM case for aluminium.

This paper is organized as follows. Section 2 contains a short review of the model. Section 3 is devoted to a brief description of the VAAQP code. Some results obtained with the code for aluminium (Al) in the WDM regime are presented in section 4. Differences between models requiring the neutrality of the Wigner-Seitz (WS) sphere (like in the INFERNO model [1, 3]) and the variational model are discussed in this section. We recall that the WS sphere radius $R_{WS}$ is simply related to the ion density $n_i$ by the formula: $R_{WS} = (3/(4\pi n_i))^{1/3}$.

Comparisons with experiments and state-of-the-art simulations are presented in section 5.

2. Variational theory of the average atom in jellium

The VAAQP code uses a new model [4, 5] of plasma at thermodynamical equilibrium with bound and continuum electrons treated within the same formalism. In this model, the plasma equilibrium is determined by three parameters: atomic number $Z$, temperature $T$ and ion density $n_i$. All other quantities, including mean ionization $Z^*$, can be determined from variational equations using for instance the density functional theory (DFT). As shown in [4, 5] in the Thomas–Fermi (TF) approximation to the electron density the present model leads naturally to the TF ion-in-cell approach proposed in 1949 by Feynman et al [6].

The starting point of the present model is the cluster expansion [7–9] of the free energy per unit volume from which the two first orders are retained:

$$f(n_i, Z, T) = f_0 + \langle f \rangle_1 + \cdots,$$

where $f_0$ is the free energy per unit volume of a uniform electron gas with an unknown density $n_0$.

$$n_0 = n_i Z^* \quad (2)$$

In the zeroth order, electron charge density is neutralized by a uniform ion background. $\langle f \rangle_1$ is the average contribution to the free energy density of one ion immersed in that jellium. In the first order the system is no more homogeneous. The cluster expansion in the first order leads to

$$\langle f \rangle_1 = n_i \int d^3r \left( f_{\text{ion+jellium}}(X; n_i, Z, T; \mathbf{r}) - f_0(n_0, T) \right). \quad (3)$$

The subtraction of the zeroth-order term assures the convergence of the above integral. In what follows all structure variables, including $n_0$, will be denoted by $X$. 

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The free energies per ion are thus defined as follows:

\[ F(X; n_i, Z, T) = F_0 + \Delta F_1 = \frac{f_0(n_0, T)}{n_i} + \langle f \rangle_1 n_i. \]  

(4)

All structure variables \(X\) are determined from the minimization of the free energy \(F(X; n_i, Z, T)\) with additional conditions:

- In the first order, we assume that the non-central ions are absent inside a cavity with a radius \(R\)

\[ n_0 \theta(r - R). \]  

(5)

- The neutrality of all charges, including the cavity, leads to the equation

\[ Z + \int d^3r \{n(\vec{r}) - n_0 \theta(r - R)\} = 0. \]  

(6)

- The total electron number per unit volume has the following cluster expansion:

\[ Z n_i = n_0 + n_i \int d^3r \{n(\vec{r}) - n_0\} + \cdots. \]  

(7)

Substituting equation (7) into equation (6) leads to the identity: \(R = R_{WS}\)

The minimization leads to self-consistent-field (SCF) equations similar to those obtained in [1–3] plus a new one which allows us to determine \(n_0\) or \(Z^*\):

\[ \int d^3r \{V_{el}(\vec{r}) \theta(r - R)\} = 0, \]  

(8)

where \(V_{el}(r)\) is the electrostatic part of the SCF potential [4, 5].

It is interesting to note that equation (8) means that in the model, the electronic SCF potential does not interact with the non-central ions charge distribution. There is a possibility to extend this model to other non-central ions charge distributions. This subject will not be considered in the present paper.

The main difference between INFERNO-type models [1, 3] and the present one is the determination of the mean ionization \(Z^*\). In the INFERNO type models, the mean ionization is obtained from the neutrality of the WS sphere whereas in the present one it is obtained from the variational principle (8). In the present model, the WS sphere is in general non-neutral.

### 3. The VAAQP code

The VAAQP code finds the solution to the variational equations described above (see equations (1)–(8)). First, a family of solutions to the SCF equations in the four-parameter space \((Z, T, n_i, Z^*)\) is calculated. Among them is found such that fulfils equation (8) (see figure 1). The code has three options to calculate the electron density: semiclassical (TF), quantum-non-relativistic (Schrödinger) or quantum-relativistic (Dirac) formalism.

In the case of calculations using quantum density models, continuum state contributions to observables are integrated over energy using a resonance-catching adaptive-mesh-refinement method. The SCF solutions in the four-parameter space \((Z, T, n_i, Z^*)\) are obtained by iteration using the method described in [10]. The electron density and total potential are calculated using finite difference numerical methods inside the region extending from the radius equal to zero up to an asymptotic radius. At this radius the electron density is matched to its asymptotic form which is calculated using the linear response theory of a homogeneous dense plasma [11, 12]. Solutions are considered valid if they are insensitive to the choice of the matching
Figure 1. Left-hand term of equation (8) versus $Z^*$. Example of solution in the case of Al plasma at 5 eV temperature and 2.7 g cm$^{-3}$ matter density. Each point on the figure corresponds to a SCF solution in the four-parameter space $(Z, T, n_i, Z^*)$.

Figure 2. Plot of the electron density versus radius for Al at 0.25 eV temperature and 2.7 g cm$^{-3}$ matter density with an enlarged view of the Friedel oscillations.

radius and are not mesh dependent. The matching radius depends mostly on the range of the Friedel oscillations and becomes larger as temperature is lower (see figure 2).

Instead of looking for solution to equation (8), the code has other options allowing one to fulfil other conditions as for instance the neutrality of the WS sphere.

4. Model comparison in the case of the warm dense aluminium

The code has been applied in the case of Al at typical condition of WDM, i.e. solid density (2.7 g cm$^{-3}$) and temperatures between 0.05 and 12 eV. In this regime, Friedel oscillations decay on a scale much larger than the WS radius (decay length $(2b_0^F)^{-1} = 55.2 a_0$ at 0.05 eV temperature, WS radius $R_{WS} = 2.99 a_0$; $b_0^F = \mu^{1/2}(-1 + \sqrt{1 + (\pi T/\mu)^2})^{1/2}$ is the imaginary part of the zeroth-order pole of the Fermi–Dirac function). In such cases, the region of numerical calculations should extend over several tens of $R_{WS}$. It is important to recall that the present approach is based on the assumption that the SCF potential decays exponentially due to screening and decay of the Friedel oscillations which is only valid at finite temperatures [11, 12].

The exchange-correlation term is taken in the local density approximation (LDA). For this exploratory study, in order to have direct comparisons with the Thomas–Fermi–Dirac (TFD) model, Dirac exchange term [13, 14] was used. Iyetomi and Ichimaru finite-temperature exchange-correlation term [15] has also been tested. In the temperature regime of interest in
Figure 3. Mean ionization $Z^*$ as defined in equation (2) versus temperature in the case of Al at 2.7 g cm$^{-3}$ matter density. Solid line corresponds to the variational model with TF density. Plus symbols denote the neutral WS sphere model with TF density. Squares correspond to the variational model with non-relativistic quantum density. Circles denote the neutral WS sphere model with non-relativistic quantum density.

In this study, the differences between results obtained using these two approximations appeared to be relatively small.

Figure 3 displays the behaviour of the mean ionization $Z^*$ that was defined in equation (2). This definition is the only one justified in the framework of the TF and quantum cases [5] and can be also applied to any model as it corresponds to the asymptotic value $n_0$ of the electron density, which is related to the chemical potential. Other definitions related to the total electron density are sometimes used. One of those is the value of the electron density at the WS radius. This definition is motivated by the fact that it is identical to the one related to $n_0$ in the TF case. Another possible definition could be the difference between the atomic number $Z$ and the sum of the bound levels occupation numbers. However, definitions of the number of bound (or free) electrons per atom relying on the summation of the occupations of bound (or localized free) states encounter well-known difficulties due to the fact that they are not related to any well-defined quantum operator. Definitions other than equation (2) will not be considered in this paper.

Results of four approaches are presented in figure 3. The first one is the result of the variational model with calculated non-relativistic quantum density. The second one is the result of the model with electron density calculated using the non-relativistic quantum density but in which equation (8) is replaced by the condition of the neutrality of the WS sphere (we call this approach NWS after ‘neutral WS sphere’):

$$ Z = \int_{|\vec{r}| \leq R_{WS}} d^3r n(\vec{r}) \quad (9) $$

The condition based on equation (9) is used in the INFERNO model. However, calculations reported here with equation (9) do not correspond exactly to the original INFERNO model as in our case the potential outside the WS sphere is not equal to zero which is the case in that model. Two calculations using the TF formalism are also reported, one with equation (8),
another with equation (9). According to [4, 5], in the TF case, these two approaches are strictly equivalent which is confirmed by figures 3 and 4.

As it can be observed in figure 3, results from quantum calculations with equation (8) can differ significantly from those obtained by using equation (9). Differences are especially pronounced at low temperature and tend to vanish as the temperature increases. It appears also that in the case of Al at solid density and at temperature below 2.5 eV, the atomic structure is different in the two quantum models. The 3s shell is found to exist within the variational model whereas it is absent when using the NWS model. These differences are mainly related to the quantum behaviour of the density outside the WS sphere, namely to the Friedel oscillations which are present in this regime.

Electronic pressure is calculated from the formula

\[ P = -f_0 + n_0(\mu + V_{xc}(n_0)) + V_{el}(R_{WS}). \]  

(10)

\[ f_0 = f_0^0 + f_{xc}^0 \] is the free energy per unit volume of a uniform electron gas including exchange correlation. \( f_0^0 \) is the ideal gas free energy for the grand-canonical ensemble, namely in atomic units

\[ f_0^0 = -\frac{2\sqrt{3}}{3\pi^2} T^{5/2} J_{3/2} \left( \frac{\mu}{T} \right) + n_0\mu. \]  

(11)

In the present study \( f_{xc}^0 \) was set to \( \epsilon_{xc} \) from [14]. \( V_{xc} \) is the exchange-correlation part of the SCF potential [4, 5]

\[ V_{xc}(n) = \frac{\partial f_{xc}^0(n)}{\partial n} = -\left( \frac{3n}{\pi} \right)^{1/3}. \]  

(12)

Equation (10) is obtained from the variational theory and applies only to the case where equation (8) is fulfilled. The quantum NWS model is not variational and the use of the above pressure formula is only given for comparison purposes. In the case of models that are not fully variational the electronic pressure is often calculated by numerical differentiation of the free energy with respect to the ion density.

As can be seen in figure 4, relative differences in calculated pressures from the two quantum models can be about 40% in the low temperature region and decrease as temperature increases. Pressures obtained using the quantum NWS model at low temperatures seem to be relatively close to those obtained from the TFD model.

Figures 3 and 4 both display curves that are continuous with respect to temperature. In the case of the quantum variational model, it is worth stressing that continuity is preserved despite the fact that the 3s shell is disappearing between 2 and 2.5 eV. This is due to a careful treatment of continuum resonance in the VAAQP code.

5. Comparison with experiments and simulations

VAAQP provides only electronic contribution to the pressure. In order to compare our results to experiments and simulations in the regime of interest in this paper, we need to add an ion contribution to the pressure. We have chosen to use the simple formula given in [16, 17] which is based on Hansen’s one-component plasma (OCP) simulations. Here, this contribution is of the order of about 20% or less. Following Mazevet and Zerah, we have calculated several mass density–temperature points corresponding to the SESAME 3700 principal Hugoniot curve (see [18] and references therein). In figure 5 are displayed results from those variational quantum calculations as well as available experimental data from [19–42]. Mazevet and Zerah simulation results are also shown for comparison.
Figure 4. Electronic pressure $P$ versus temperature in the case of Al at 2.7 g cm$^{-3}$ matter density. Solid line corresponds to the variational model with TF density. Plus symbols denote the neutral WS sphere model with TF density. Squares correspond to the variational model with non-relativistic quantum density. Circles denote the neutral WS sphere model with non-relativistic quantum density.

Figure 5. Comparison along the principal Hugoniot of Al between VAAQP variational quantum calculations, ab initio simulations from [18] and experiments from [19–42]. Total pressure $P$ versus mass density $\rho$ is presented.

This first comparison indicates that the pressure results compare rather well with experiments and state-of-the-art ab initio simulations in this region which is especially difficult from the numerical point of view in our case.

6. Conclusion

In this paper, a fully variational model of the average atom in quantum plasmas is applied for the first time in a numerical code called VAAQP. It is shown on some chosen examples that
the code is able to calculate variational self-consistent equilibrium at high matter density and relatively low temperatures. In this regime, Friedel oscillations of the electron density and SCF potential present a real challenge from the numerical point of view.

Effects of the variational treatment are studied by comparison to results from a model requiring the neutrality of the WS sphere. The variational model can lead to electronic pressures, mean ionizations and atomic structures that differ significantly from those obtained using existing models especially in the low temperature region.

Comparisons with experiments and ab initio simulations are also shown. Let us recall however that the main objectives of this work is to open the way to atomic physics and radiative properties calculations which could be coherent with the variational equation of state and the virial theorem. Such calculations can be based on methods similar to that used in the VAAQP code.

The work on the present model is in progress. Among objectives is the application of the method presented in this paper to the superconfigurations as indicated in [5].

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