Restricted path integral Monte Carlo simulations are used to calculate the equilibrium properties of hydrogen in the density and temperature range of \(9.83 \times 10^{-4} \leq \rho \leq 0.153\) g cm\(^{-3}\) and \(5000 \leq T \leq 250,000\) K. We test the accuracy of the pair density matrix and analyze the dependence on the system size, on the time step of the path integral, and on the type of nodal surface. We calculate the equation of state and compare with other models for hydrogen valid in this regime. Further, we characterize the state of hydrogen and describe the changes from a plasma to an atomic and molecular liquid by analyzing the pair correlation functions and estimating the number of atoms and molecules present.

I. INTRODUCTION

In spite of the simple composition, hydrogen exhibits a surprisingly complex phase diagram, which is the subject of numerous experimental and theoretical approaches. In this work, we study the high temperature regime of \(5000 \leq T \leq 250,000\) K where hydrogen undergoes a smooth transition with increasing temperature from a molecular fluid through an atomic regime and finally to a two component plasma of electrons and protons (see Fig. 1). The properties of hydrogen in this regime are crucial for the evolution of stars and the characteristics of the Jovian planets.

A variety of simulation techniques and analytical models have been developed to describe hydrogen at low density. This regime has been studied with chemical models \(^1\)\(^–\)\(^3\) that describe hydrogen as a mixture of interacting molecules, atoms, free protons, and electrons. The chemical composition is determined by minimizing an approximate free energy function constructed from known theoretical limits. In this paper, we focus on low and intermediate densities \(9.83 \times 10^{-4} \leq \rho \leq 0.153\) g cm\(^{-3}\) corresponding to \(14 \geq r_c \geq 2.6\), where one expects the chemical models to work well although the properties of hydrogen are determined by the complex interplay of long-range Coulomb forces leading to strong coupling and bound states as well as degeneracy effects.

All these effects can also be described from first principles simulation. There are \textit{ab initio} methods such as restricted path integral Monte Carlo simulations (PIMC) \(^4\)\(^–\)\(^6\), density functional theory molecular dynamics (DFT-MD) \(^7\)\(^,\)\(^8\) and wave-packet molecular dynamics \(^9\),\(^10\). The focus of the work is to test the equation of state (EOS) derived from chemical models and the actual density-temperature limits of the validity of the chemical picture. Additionally, we provide data to determine the parameters of the free energy models. Chemical models are expected to become inaccurate in regions of high density where the lifetime of molecules reduces to a few molecular vibrations \(^8\).

We present results from more accurate PIMC simulations. First, we analyze the accuracy of the pair density matrix and the error of the ‘‘time’’ discretization. Then we analyze the finite size dependence of the derived EOS and discuss the fixed node errors by comparing results from simulations using free particle (FP) nodes and variational nodes. Furthermore, we calculate pair correlation functions that we use in conjunction with a cluster analysis to characterize the state of hydrogen at different temperatures and densities. We use atomic units (lengths in Bohr radii and energies in Hartrees) throughout this work except where indicated otherwise.

II. PATH INTEGRAL MONTE CARLO METHOD

A. Restricted path integral

The density matrix (DM) of a fermion system at temperature \(k_B T = 1/\beta\) can be written as an integral over all paths \(R_t\),

\[
\rho(R_0, R_\beta; \beta) = \frac{1}{N!} \sum_{\pi} (-1)^\pi \oint_{R_0 \rightarrow \beta R_\beta} dR_t e^{-S[R_t]},
\]

(2.1)

\(R_t\) stands for the entire paths of \(N\) particles in three-dimensional space \(R_t = (r_{1t}, \ldots, r_{Nt})\) beginning at \(R_0\) and ending at \(\beta R_\beta\). \(\pi\) labels the permutation of the particles and \((-1)^\pi\) to its signature. For nonrelativistic particles interacting with a potential \(V(R_t)\), the action of the path \(S[R_t]\) is given by,

\[
S[R_t] = \int_0^\beta dt \left[ \frac{m}{2 \hbar} \frac{dR(t)}{dt}^2 + V(R(t)) \right] + \text{const.}
\]

(2.2)
In practice one discretizes [11] the path into a finite number of imaginary time slices $M$ corresponding to a time step $\tau = \beta / M$.

For fermionic systems the integration is complicated due to the cancellation of positive and negative contributions to the integral (the fermion sign problem). It has been shown [12,13] that one can evaluate the path integral by restricting the path to only specific positive contributions. One introduces a reference point $R^*$ on the path that specifies the nodes of the DM, $\rho(R, R^*, t) = 0$. A node-avoiding path for $0 < t \leq \beta$ neither touches nor crosses a node: $\rho(R(t), R^*, t) \neq 0$. By restricting the integral to node-avoiding paths,

$$\rho_f(R_0, R^*, \beta) = \int dR_0 \rho_f(R_0, R^*; 0) \times \oint_{R_0 - R_0 = Y(R^*)} dR e^{-S[R]}, \quad (2.3)$$

[$Y(R^*)$ denotes the restriction] the contributions are positive and therefore PIMC represents, in principle, a solution to the sign problem. The method is exact if the exact fermionic DM is used for the restriction. However, the exact DM is only known in a few cases. Most applications have approximated the fermionic DM by a determinant of single particle DMs,

$$\rho(R, R^*; \beta) = \begin{pmatrix} \rho_1(r_1, r'_1; \beta) & \cdots & \rho_1(r_N, r'_1; \beta) \\ \cdots & \cdots & \cdots \\ \rho_1(r_1, r'_N; \beta) & \cdots & \rho_1(r_N, r'_N; \beta) \end{pmatrix}. \quad (2.4)$$

This approach has been extensively applied using the FP nodes including applications to dense hydrogen [4–6]. It can be shown that for temperatures larger than the Fermi energy, the interacting nodal surface approaches the FP nodal surface. In addition, in the limit of low density, exchange effects are negligible: the nodal constraint has a small effect on the path and therefore its precise shape is not important.

To gain some quantitative estimate of the possible effect of the nodal restriction on the thermodynamic properties, it is necessary to try an alternative. In addition to FP nodes, we used nodal surface of a variational density matrix (VDM) [14] derived from a variational principle that includes interactions and atomic and molecular bound states. This method was first combined with PIMC in [15]. One assumes the DM is a Slater determinant of single particle Gaussian functions,

$$\rho_1(r, r'; \beta) = (\pi w)^{-3/2} \exp\{-((r - m)^2 / w + d)\}, \quad (2.5)$$

where the variational parameters are the mean $m$, squared width $w$, and amplitude $d$. These parameters are determined as a function of imaginary time using a system of coupled differential equations given in Ref. [14]. The initial conditions at $\beta \to 0$ are $w = 2\beta$, $m = r'$, and $d = 0$ in order to regain the correct FP limit. It follows from the variational principle that at low temperature, the VDM goes to the lowest energy wave function within the variational basis. For an isolated atom or molecule this will be a bound state, in contrast to the delocalized state of the FP DM. A further discussion of the VDM properties is given in Ref. [14]. Note that these nodes are only used to determine the nodal restriction in Eq. (2.3). The complete potential is taken into account in the path integral action as discussed in detail in Ref. [11].

B. Accuracy of the method

The numerical implementation of the PIMC method requires one to make several approximations. Inaccuracies can be caused by statistical errors from the Monte Carlo (MC) integration, inaccuracies in the numerically determined pair density matrices, a dependence on the time step of the path integral because of $N$-body ($N \geq 3$) correlations, finite size effects, and nodal errors from approximations in the trial density matrix. Their effects on the accuracy of the computed thermodynamic averages are quantitatively estimated in this section.

Statistical errors in the estimators for the thermodynamic quantities are calculated from the block averages generated by the MC simulations. The correlations between blocks are taken into account by performing a blocking analysis [16]. The resulting error bars (one standard deviation) are given for all observables throughout this paper as a number in parenthesis referring to the least significant digit.

In the following discussion, we compare the internal energy and the pressure calculated using the virial theorem,

$$3Pv = 2K + V, \quad (2.6)$$

where $v$ is the volume of the simulation cell, $K$ the kinetic energy, and $V$ the potential energy. Accurate estimation of the pressure requires a high accuracy in the kinetic and the potential energies because they tend to cancel out. In the molecular regime at low density, both terms, dominated by intramolecular contributions, cancel to a large extent leaving behind the molecular gas pressure. As a result, the pressure
FIG. 2. Accuracy study of PIMC simulations of the isolated hydrogen atom using different orders $n_A$ in the expansion formula Eq. (2.7) for the action and energy. The calculated $2K + V$ (exact value equals zero) and the deviation from the exact internal energy of $-13.6$ eV are shown for different orders from simulations at $T=10,000$ K using $\tau^{-1}=10^6$ K.

is, in general, significantly more sensitive to approximations than other quantities such as the internal energy.

1. Pair density matrix

If one only used the bare potential as in Eq. (2.2) (the primitive approximation for the action), the convergence would be very slow [17] and would result in extremely inefficient many-particle simulation. Instead, we numerically solve the two-particle problem with the matrix squaring technique [18]. Numerical representations of the exact pair density matrices are stored in tables used by the PIMC simulation program by expanding in the small variables $s$ and $z$

$$u(r, r'; \tau) = \frac{1}{2} [u_0(r; \tau) + u_0(r'; \tau)]$$

$$+ \sum_{k=1}^{n_A} \sum_{j=0}^{k} u_k(q; \tau) z^{2j} s^{2(k-j)},$$  \quad (2.7)

where

$$q = \frac{1}{2} (|r| + |r'|), \quad s = |r - r'|, \quad z = |r| - |r'|,$$  \quad (2.8)

and $r$ and $r'$ denote the separation of the two particles at adjacent time slices. The accuracy of these tables is crucial for all computed results. Using the precomputed pair density matrices allows one to employ a much larger time step because one starts with a solution of the two-particle problem. Figure 2 shows how accurate this method is. The internal energy of an isolated hydrogen atom at sufficiently low temperature ($T=10,000$ K) in a large box ($L=26$) is compared with the exact ground state energy of $-13.6$ eV. The temperature was chosen low enough so that excited states can be neglected; the contribution to the energy from the occupation of the first excited state is $7 \times 10^{-5}$ eV at this temperature. Also shown is how well the kinetic energy $K$ and the potential energy $V$ satisfy the virial theorem $2K + V = 0$, thus determining the accuracy with which the pressure can be determined. For a time step of $\tau^{-1}=10^6$ K, the analysis shows a quick convergence with the order of terms considered in the action expansion Eq. (2.7). Using terms up $n_A=3$ reduces the error to 0.033(3) eV in energy and to 0.039(8) eV for $2K + V$. This corresponds to an inaccuracy in the pressure equivalent to a noninteracting molecular gas at $T = 260(30)$ K.

2. Time step dependence

Employing the pair approximation of the density matrix does not include correlation effects for three or more particles (for example, between one electron and two protons). We now estimate how small the time step must be to obtain a given accuracy for an isolated molecule. Figure 3 shows results for different time steps and temperatures with the nuclei kept fixed at the equilibrium position of $R=1.4008$. From the virial theorem, it follows that $2K + V = 0$ at a sufficiently low temperature. The exact energy per atom is $-15.973$ eV [19]. The $T$ dependence is small suggesting that the electrons are in the ground state. However, one finds a significant dependence on the time step. Using $\tau^{-1}=2 \times 10^6$ K reduces the error in the energy per atom to 0.036(3) eV and in $2K + V$ to 0.090(16) eV. The time step error is larger than the errors of the inaccuracies in the pair density matrices discussed above. The error in $2K + V$ corresponds to a pressure of a noninteracting molecular gas at $T=700(100)$ K, which provides us with an approximate limit of accuracy in the equation of state calculated from many-particle simulations with $\tau^{-1}=2 \times 10^6$ K.

3. Finite size dependence

The estimation of the finite size errors is more difficult to assess because the needed PIMC simulations are computationally much more demanding. The required computer time increases rapidly with the number of particles making it challenging to obtain converged results for paths corresponding to large systems.
Most results from many-body simulations reported in this work were calculated with \( N = 32 \) pairs of electrons and protons in a periodically repeated simulation cell. To study the effect on \( N \), we performed simulations for \( N = 16 \) and 64 pairs of protons and electrons for a density of \( r_s = 2.6 \) and \( T > 10000 \) K. We chose the highest density under consideration because one expects the finite size dependence to be largest there due to the stronger interaction between the atoms.

The finite size dependence of the pressure, shown in Fig. 4, is small at high temperatures but grows to approximately \( 4(2)\% \) near \( T = 30,000 \) K. In this regime, the hydrogen undergoes structural changes involving the formation of atoms, which affect the pressure. This study provides us only with an estimate of the finite size dependence. An extrapolation to \( N \to \infty \) would require significantly larger systems, not currently feasible at low temperatures. One should note the increased size of the error bars for \( T \approx 30,000 \) K, which indicate that the two curves in Figs. 4 and 5 are still in agreement within the error bars.

Figure 5 shows the finite size error of the internal energy. The smaller systems are more strongly bound by approximately 0.2 eV per atom. For lower densities, we expect this value to be smaller because, at high temperature, the finite size errors are primarily caused by the interaction of the charges with their own images in periodic boundary conditions. This introduces artificial correlations with a too negative potential energy. The correction is inversely proportional to system size \( (1/N) \). For lower temperatures, this interaction is reduced due to screening effects and the formation of bound states. This explains why the finite size error of the internal energy increases with temperature and leads to lower values for small systems.

4. Nodal Approximation

In the above, we have studied controlled approximations. The only uncontrolled approximation in the restricted PIMC method is the use of trial density matrix to constrain the paths. The nodal surfaces are important only if the electrons are degenerate: at low temperatures or at high densities. Recall that in this work we focus on hydrogen only at low density, where the electrons are bound in atoms and molecules and have a low or moderate degeneracy. Even at low density, one still needs a nodal surface in order to prevent the
TABLE I. Pressure and internal energy per atom from PIMC simulations with 32 pairs of particles and $r^{-1} = 2 \times 10^6$ K using free particle nodes except for * where VDM nodes were employed instead. The probabilities $x$ for finding a proton in a given state for the three dominant species are derived from a cluster analysis. $P_{\text{perm}}$ is the permutation probability for the electrons.

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<th>$r_s$</th>
<th>$T$(K)</th>
<th>$P$(GPa)</th>
<th>$E$(eV)</th>
<th>$x_{H^+}$</th>
<th>$x_H$</th>
<th>$x_{H_2}$</th>
<th>$P_{\text{perm}}$</th>
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</table>
formation of unphysical clusters like H$_3$ and H$_4$ or even the collapse of the entire system, but the precise shape of the nodes is not important at low density as shown in Fig. 6. Comparing FP and VDM nodes for $r_s = 2.6$, one only finds differences in the pressure for $T < \approx 15625$ K, which is approximately where, at this density, the system shows a significant molecular signature (see Sec. III B and Fig. 16). In this regime, FP nodes systematically lead to a too high pressure, while simulations with VDM nodes stay closer to the prediction of a semiempirical chemical model [1]. At a lower density $r_s = 4$ results from FP and VDM nodes agree within the error bars. The differences in the internal energy, as shown in Fig. 7, using FP and VDM nodes are significantly smaller than the pressure deviations. One either finds agreement within the error bars or that VDM nodes predict lower internal energies, which was used in [15] to show that VDM nodes are the more accurate nodal surface. The observed energy differences did not exceed 0.1 eV per atom.

For even lower density, the nodes are less relevant because they become only important in a collision of two molecules, which occur less frequently at lower density. This trend can also be understood in terms of the degeneracy of the electrons. The degree of degeneracy manifests itself in the path integral formalism by the probability for the electrons to be involved in a permutation. At high temperature, the paths are very short and permutations are rare. At low temperature and high density, the paths are long and can form long permutation cycles. However in hydrogen at low density, the paths are localized due to the attraction in atoms and molecules and permutations are rare. Figure 1 shows that the permutation probability never reaches 1% for $r_s = 4$ (see Table I). For higher densities, the permutation probability is increased as indicated by the contour lines. This is consistent with the temperature and density dependence of the nodal error in pressure and internal energy discussed above.

III. RESULTS

A. Equation of state

Table I gives the complete set of energies and pressures at six densities and eight temperatures. We now compare these results with several models for hydrogen. We begin our discussion by studying the internal energy per atom as a function of temperature shown in Figs. 8–11 for two selected densities and temperatures.

- Fig. 8: Internal energy per atom vs temperature for a density of $r_s = 10$ comparing the SC-EOS [1] with PIMC calculations.
- Fig. 9: Internal energy per atom vs temperature for a density of $r_s = 10$ as shown in Fig. 8 but here for lower temperatures also including results from the activity expansion (ActEx) [20] and the fluid variational theory (FVT) [21].
- Fig. 10: Internal energy per atom vs temperature as shown in Fig. 8 but here for a density of $r_s = 2.6$.
- Fig. 11: Internal energy per atom vs temperature for a density of $r_s = 2.6$ as shown in Fig. 10 but here for lower temperatures also including results from the activity expansion (ActEx) [20] and the FVT [21].
densities corresponding to \( r_s = 10.0 \) and 2.6. Generally, we find a fairly good overall agreement with the EOS by Sau-\( \text{man} \) and Chabrier\(^{[1]} \) over the entire temperature and density range discussed in this work. The agreement is particularly good in the molecular and atomic regime for \( r_s = 10.0 \), as shown in Fig. 9. There the SC energies are within the error bars of the PIMC results. At higher temperature shown in Fig. 8, we find systematic deviations of up to 5 eV per atom at \( T = 250 \,000 \) K. They indicate that the SC energies are too low at high temperatures and too high at intermediate temperatures (see Fig. 8). One possible explanation for the deviations at high temperature is that the SC model underestimates the degree of ionization (see discussion in \( \text{[22]} \)).

We also studied these deviations as a function of density. The cross-over temperature, above which the SC-EOS underestimates the energy, increases with density. At \( r_s = 10.0 \), the crossover is near \( 70 \,000 \) K compared to \( 130 \,000 \) K (Fig. 10) at \( r_s = 2.6 \). At temperatures below \( 20 \,000 \) K for \( r_s = 2.6 \), one also finds some small deviations up to 0.5 eV per atom (Fig. 11).

Figure 12 shows a comparison of energy versus density for several temperatures. It shows that the SC-EOS overestimates the energy for \( T = 5000 \) K and \( 31 \,250 \) K and underestimates it for \( 125 \,000 \) K for densities higher than those corresponding to \( r_s = 2.6 \).

Now let us compare the pressure from the SC-EOS with that from the PIMC simulation using Eq. (2.6) in Table I. We find remarkably good agreement of the entire range of temperature and density under consideration. For a low density such as \( r_s = 10.0 \), this is shown in Fig. 13. As expected, one finds that both methods interpolate between the limit of an ideal Fermi gas at high temperatures and noninteracting molecular gas at low temperatures. Figure 14 confirms the good agreement at a higher density of \( r_s = 2.6 \). As a result of the strong interactions at this density, one finds that the pressure at low temperatures is significantly above the non-interacting molecular gas limit. We find that the SC-EOS underestimates the pressure by about 3% for \( T > 62 \,500 \) K. This difference is outside the error bar from the approximations in PIMC discussed in Sec. II B and could be interpreted as a further indication, in addition to the observed energy deviations, that the SC model underestimates the degree of ionization at high temperatures. For intermediate temperatures \( 62 \,500 \leq T \leq 15 \,625 \) K, one finds pressure differences, which are of the same magnitude as the finite size effects in PIMC. For temperatures below 15 \,625 K, the increased statistical errors in the PIMC pressure are of the same size as the observed deviations.

In Fig. 15, we show the pressure as a function of density, which confirms the good agreement. The figure also indicates that, at 5000 K and \( r_s > 4 \), the pressure is close to the pressure of a noninteracting molecular gas.

In our comparison, we also included results from the activity expansion by Rogers \( \text{[20]} \), which shows very good agreement in pressure and internal energy (see Figs. 9, 11, and 12). The differences are small but increase with density. In the molecular and atomic regime, one also finds good agreement with the FVT by Juranek and Redmer \( \text{[21]} \) as shown in Figs. 9, 11, 13, and 14. For higher temperatures, the FVT model is not applicable since it does not include ionization of atoms.
B. Pair correlation functions

There are four different pair correlation functions that can be directly obtained from many-body simulations and provide direct information about the state of the system. Shown in the following figures is an extensive set of pair correlations that allow one to estimate the microscopic structure of the system and allow a direct comparison with other simulations. The proton-proton pair correlation functions from PIMC simulations with free particle nodes are shown in Fig. 16. For $T \approx 20,000$ K a peak at the bond length of 1.4 emerges, which clearly demonstrates the formation of molecules. We found it useful to multiply the pair correlation function by an extra density factor $n = N/v$ so that the area under the peak is proportional to the molecular fraction. The peak height gets smaller with decreasing density as a result of entropic dissociation of the molecules, driven by the number of unbound states at low density. Thermal dissociation also reduces the number of molecules with increasing temperature. For $r_s \approx 2$, we expect that pressure dissociation diminishes the number of molecules with increasing density [17] but this density range is beyond the scope of this paper.

The proton-electron pair correlation function multiplied by the density is shown in Fig. 17. The peak near the origin shows the increased probability of finding an electron near a proton due to the Coulomb attraction. The peak height decreases with temperature and increases with density because of thermal ionization and entropy ionization, respectively. At low temperature, the peak can be interpreted as occupation of bound states although (unbound) scattering states can also contribute. From proton-electron pair correlation alone, one cannot distinguish between an atomic and a molecular state.

Figures 18 and 19 show the electron-electron pair correlation functions for pairs with anti-parallel spin. The peak at small separations comes from the formation of the molecular bond. For pairs of electrons with parallel spin, one always finds a strong repulsion due to the Pauli exclusion principle and to a lesser extent to the Coulomb repulsion. This is shown in Fig. 19.

C. State of hydrogen

In this section, we discuss the phase diagram of hydrogen as shown in Fig. 1. The diagram shows the approximate location of the molecular, the atomic, and the plasma regimes. The PIMC simulations, since they are based on the basic description in terms of electrons and protons, do not directly lend themselves to determining the number of compound particles such as molecules and atoms. (Methods for determining this from PIMC simulations will be discussed in a future publication.) In order to obtain an estimate of the atomic and molecular fraction, we employed a cluster analysis of the PIMC path configurations. As described in [23], we consider two protons as belonging to one cluster if they are less than 1.9$\alpha_B$ apart. An electron belongs to one particular
cluster if it is less than $1.4a_B$ away from any proton in the cluster. The two cut-off radii were chosen from the molecular and atomic ground state distribution. This analysis gives reasonable estimates for the molecular and atomic fractions at low temperatures. At high temperature, it overestimates the number of bound states because even in an (unbound) collision, two particles are counted falsely as being part of the cluster. We corrected for this by applying an additional criterion: a particle can only be considered as bound if the difference in action to remove it from the cluster is positive. This method leads to the expected corrections at high temperature. (The regime boundaries in Fig. 1 discussed below are hardly affected by the additional correction.)

The lower dashed line represents the region where 60% of the protons are bound in molecules. When the number of protons bound in atoms (i.e., with an electron) drop below 40% we labeled this state as plasma as shown in the upper dashed-dotted line. It should be emphasized that the location of these lines depends on the choice of these limits as well as on the cut-off radii used to determine the clusters in this place. Figure 1 also shows the location of isobars, which appear as almost straight lines in this double logarithmic graph. The slope is different from the ideal gas because the pressure depends on ionization and dissociation.

Table I shows the fraction of the three most frequently found species: molecules, atoms, and free protons where $x$ is defined as the probability of finding a proton in a certain compound particle. It should be noted that the sum: $x_{H^+} + x_{H_2^+}x_{H_2}$ is less than 1 since other clusters have a nonzero probability. The largest contributions besides those listed are $H_2$ with a maximum of 0.06 for $r_s=2.6$ and $T=15625$ K followed by $H_3$ with $x=0.03$ and $H^-$ with $x=0.02$. Even larger clusters occur very infrequently. The cluster analysis also gives an estimate for the fraction of free electrons, which agrees well with the number of ionized protons: $x_{H^+}$.

Figure 20 shows a comparison of the fraction of molecules and ionized atoms for $r_s=10$ comparing the cluster analysis of the PIMC results with the SC model [1] and the FVT [21].

![Figure 20](image20.png)

FIG. 20. Fraction of molecules and free protons as a function of temperature for $r_s=10$ comparing the cluster analysis of the PIMC results with the SC model [1] and the FVT [21].
analysis leads to smaller number of molecules than predicted by the SC model.

Summarizing, one can say that the cluster analysis provides us with reasonable estimates for the number of atoms and molecules in the considered density range. We caution the reader that other definitions of atoms and molecules, possible in a many-body hydrogen, while giving qualitatively similar results, may differ quantitatively. Our computed numerical values must be used with caution. A rigorous, many-body definition of a bound state remains to be applied. Several ideas are discussed in [24].

IV. CONCLUSIONS

In this work, we studied the high-temperature equation of state of hydrogen at low and intermediate densities and find a remarkably good agreement with the SC-EOS. Generally, one finds that the deviations in the energy are more pronounced than the differences in the pressure. We find significant deviations in the EOS of temperatures \( \approx 100,000 \) K, most likely caused by an underestimate of the degree of ionization at those temperatures. In future work, we will extend this comparison to higher densities. There one expects to find substantial differences between the SC model and PIMC, which manifest themselves in a different shock Hugoniot curve [15].

ACKNOWLEDGMENTS

We would like to thank E.L. Pollock for useful discussions as well as D. Saumon, F. Rogers, and H. Juranek for providing us with their EOS tables. This research was funded by the U.S. Department of Energy through the University of California under Subcontract No. B341494 and in part by the University of California/Lawrence Livermore National Laboratory (LLNL) under Contract No. W-7405-Eng-48. We used the computational facilities at the LLNL and at the National Center for Supercomputing Applications.