First-Principles Equation of State Database for Warm Dense Matter Computation

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We put together a first-principles equation of state (FPEOS) database for matter at extreme conditions by combining results from path integral Monte Carlo and density functional molecular dynamics simulations of the elements H, He, B, C, N, O, Ne, Na, Mg, Al and Si as well as the compounds LiF, B4C, BN, CH4, CH2, C2H3, CH, C2H, MgO, and MgSiO3. For all these materials, we provide the pressure and internal energy over a density-temperature range from ~0.5 to 50 g cm⁻³ and from ~10⁴ to 10⁷ K, which are based on ~5000 different first-principles simulations. We compute isobars, adiabats and shock Hugoniot curves in the regime of L and K shell ionization. Invoking the linear mixing approximation, we study the properties of mixtures at high density and temperature. We derive the Hugoniot curves for water and alumina as well as for carbon-oxygen, helium-neon, and CH-silicon mixtures. We predict the maximal shock compression ratios of H₂O, H₂O₂, Al₂O₃, CO, and CO₂ to be 4.61, 4.64, 4.64, 4.89, and 4.83, respectively. Finally we use the FPEOS database to determine the points of maximum shock compression for all available binary mixtures. We identify mixtures that reach higher shock compression ratios than their end-members. We discuss trends common to all mixtures in pressure-temperature and particle-shock velocity spaces. In the supplementary material, we provide all FPEOS tables as well as computer codes for interpolation, Hugoniot calculations, and plots of various thermodynamic functions.

I. INTRODUCTION

A rigorous and consistent theoretical description of warm dense matter (WDM) has been identified [1–3] as a central goal to the development of key plasma technologies, such as inertial and magnetic confinement fusion [4–9], shock physics [10–12], and high energy astrophysics [13, 14]. WDM represents materials at solidification [4–9], shock physics [10–12], and high energy astrophysics, such as inertial and magnetic confinement fusion [4–9], shock physics [10–12], and high energy astrophysics [13, 14]. WDM represents materials at solidification [4–9], shock physics [10–12], and high energy astrophysics [13, 14]. WDM represents.

Despite these challenges, the development of a rigorous and consistent theoretical framework to describe WDM remains to be of high importance because it represents states of matter on the pathway to reaching fusion conditions. Predicting with high accuracy the equation of state (EOS) as well as transport and optical properties at extreme pressure and temperature conditions is the primary motivation for developing new methods. Significant progress towards this goal has been made with laboratory experiments [11, 23–26] and first-principles (FP) computer simulations [27]. Since hydrodynamic simulations typically guide the design of dynamic compression experiments and they rely on accurate EOS tables to be predictive, computer simulations of any material and thermodynamic condition that can be probed with laboratory experiments are of high interest. FP computer simulations, that are based on the fundamental laws of quantum mechanics, enabled us to compute the EOS of materials over a wide range of conditions that also include planetary and stellar interiors. In giant planets [28], not only hydrogen-helium mixtures [29–34] but also rocky materials [35–41] are exposed to pressures of tens of megabars and temperature of ~10⁴ K. Accurate EOSs are needed to complete the spacecraft measurements of giant planets in our solar system to better characterize their interior structure and evolution [42–44]. The discovery of thousands of exoplanets with ground-based observations and space missions [45–47] has considerably broadened the range of conditions and materials of interest [48–50].

Stellar interiors encompass a wide range of temperatures from 10⁴–10⁹ K. The most detailed information came from observing the normal mode oscillations of our Sun [51–53]. Such astero-seismological observations now...
improve our understanding of distant stars [54]. For the first time, the frequencies of a number of normal modes in a giant planet have been determined with high precision through the detection of spiral density waves in Saturn’s ring by the Cassini spacecraft [55].

In this article, we build a FPEOS database for WDM computation by combining the results of two computer simulation methods, path integral Monte Carlo (PIMC) calculations and density functional theory molecular dynamics (DFT-MD) simulations. Alternative methods to perform these calculations include orbital-free density functional theory [56–58], Thomas-Fermi molecular dynamics [59], or average atom models [60–63] or their combination [64, 65]. With their approximations, all these methods enable one to compute the properties of WDM independently for one set of temperature-density condition at a time. In this regard, they differ from conventional EOS models that start from a cold curve and then introduce nuclear and electronic excitations by constructing elaborate free energy models. Multi-material databases like the Quotidian EOS [66] and many Sesame models [67] rely on that approach.

Here we instead rely exclusively on predictions from FP computer simulations in order to build an FPEOS database to characterize 11 elements and 10 compounds over a wide range of temperature and density conditions. We exclude nuclear reactions from consideration even though they occur at the highest temperatures that we study. We predict the shock Hugoniot curves and study a variety of binary mixtures by invoking the ideal mixing approximation at constant pressure, $P$, and temperature, $T$. In Ref. [68], this approximation has been shown to work remarkably well for WDM computations for temperatures above $2 \times 10^5 \text{K}$ and the shock compression ratio exceeding $\sim 3.2$. With the goal of making WDM computations more reliable and efficient, we make available as supplemental material all EOS tables as well as the C++ computer codes for their interpolation. Python code is provided to generate graphs of shock Hugoniot curve, isentropes, isobars, and isothersms for compounds and user-defined mixtures [69].

II. METHODS

A. PIMC Simulations

The equations of state in Table I were assembled by combining published results from PIMC simulations at high temperature and from Kohn-Sham density DFT-MD simulations at lower temperature. The PIMC simulation method is based on the early work on superfluid $^4\text{He}$ that introduced the permutation sampling to the path integral computations [98–100]. The algorithm was subsequently extended to fermionic systems by introducing the restricted paths approach [101–103]. The first results of this simulation method were reported in the seminal works on liquid $^3\text{He}$ [102] and dense hydrogen [104, 105]. Simulations of one-component plasmas [106–108] and of hydrogen-helium mixtures [109] followed. In Ref. [79], it was demonstrated that the free-particle nodal approximation worked sufficiently well to study hot, dense carbon and water, which paved the way for performing the PIMC simulations of elements from hydrogen to neon (Z=10), as shown in Tab. I. In Ref. [88], Hartree-Fock orbitals were used to efficiently incorporate localized electronic states into the nodal structure, which extended the applicability of fermionic PIMC simulations to heavier elements up to silicon (Z=14).

The PIMC method is based on the thermal density matrix of a quantum system, $\hat{\rho} = e^{-\beta \hat{H}}$, that is expressed as a product of higher-temperature matrices, $e^{-\beta \hat{H}} = (e^{-\tau \hat{R}})^M$. The integer $M$ represents the number of steps along the path in imaginary time. $\tau = \beta / M$ is the corresponding time step. The path integral emerges when the operator $\hat{\rho}$ is evaluated in real space,

$$\langle R|\hat{\rho}|R'\rangle = \frac{1}{N!} \sum_P (-1)^P \int_{R\rightarrow R'} dR_t e^{-S[R_t]}.$$  \hspace{1cm} (1)

The sum over $P$ represents all permutations of identical fermions that in combination with the $(-1)^P$ factor project out the antisymmetric states. For sufficiently small time steps, $\tau$, all many-body correlation effects vanish and the action, $S[R_t]$, can be computed by solving a series of two-particle problems [98, 110, 111]. The advantage of this approach is that all many-body quantum correlations are recovered through the integration over paths. The integration also enables one to compute quantum mechanical expectation values of thermodynamic observables, such as the kinetic and potential energies, pressure, pair correlation functions and the momentum distribution [100, 112]. Most practical many-body implementations of the path integral method rely on Monte Carlo sampling techniques because the integral has $D \times N \times M$ dimensions in addition to sum over permutations. ($D$ is the number of spatial dimensions; $N$ is the number of particles.) The method becomes increasingly efficient at high temperature because the length of the paths scales like $1/T$. In the limit of low temperature, where few electronic excitations are present, the PIMC method becomes computationally demanding and the Monte Carlo sampling can become inefficient. Still, the PIMC method avoids any exchange-correlation approximation and the calculation of single-particle eigenstates, which are embedded in all standard Kohn-Sham DFT calculations.

The PIMC simulations were performed with the CUPID code [113] using periodic boundary conditions. The necessary computer time foremost depends on the number of electrons and the number of path integral time steps. Earlier simulations of hydrogen were performed with 32 electrons [70] while later calculations [73] used between 64 and 256 electrons depending on density. The helium calculations in Ref. [77] employed 64 and 114 electrons. For the simulations of boron [78] and B$_4$C [91],
slightly larger cells with 30 nuclei and 150 (B) or 156 (B$_4$C) electrons were used. For the simulations of elemental nitrogen, oxygen, magnesium, silicon, we employed cells with 8 nuclei and 56, 64, 96, and 112 electrons, respectively. For BN and MgSiO$_3$, PIMC simulations with 144 electrons were performed. A detailed finite-size study is provided in the supplementary material of Ref. [83].

### B. DFT-MD Simulations

All DFT-MD simulations were performed with the Vienna Ab initio Simulation Package (VASP) [114]. We used the hardest projector augmented wave [115] pseudopotentials that were available for that code. The Perdew-Burke-Ernzerhof (PBE) [116] functional or the local density approximation [117, 118] were employed to incorporate exchange-correlation effects. We used cubic simulation cells with periodic boundary conditions and, to improve efficiency, we used a smaller number of atoms at the highest temperatures than we employ at lower temperature. As shown in our previous work [82, 87, 93, 95], this is not detrimental to the accuracy of the EOS data at high temperatures.

The Mermin functional [119] was used throughout to incorporate the effects of excited electronic states at elevated temperatures. The temperature condition where we switched from DFT-MD to PIMC depends on the material. For low-Z materials like helium, we already switched to PIMC at 10$^8$ K while for elements from Na through Si, we performed DFT-MD simulation for temperatures as high as 2×10$^9$ K. The agreement between the EOSs derived with PIMC and DFT-MD methods is fairly good. The deviations in pressure were found to be 2% or less while the internal energies typically deviated by ∼5 Ha/nucleus or less. This means that fundamental approximations, like the nodal structure in PIMC and the choice an exchange-correlation functional in DFT methods, do not prevent us from constructing consistent EOS tables for all 21 materials under consideration. This also suggests that the most fundamental electronic properties were accurately described with both method and that the numerical approximations were reasonably well controlled. Moreover, in Ref. [91, 92, 120] it was shown that alternate DFT methods such Fermi operator expansion and spectral quadrature as well as different pseudopotentials and exchange-correlation functionals gave consistent results.

With the described approximations, one typically finds that the predictions from FP simulations agree with results from laboratory experiments at extreme temperatures and pressures because at the present time, these measurements have error bars that are larger those of high-pressure experiments at room temperature, which have enabled us to benchmark the accuracy of different FP methods [121]. Still as experiments at high temperature and pressure become more precise in the future, one will need to revisit the fundamental approximations (Fermion nodes in PIMC and exchange-correlation description in DFT) and the controllable approximations (finite size effects in both methods, convergence with respect to simulation duration in DFT-MD and PIMC).

| Material     | Number of isochores | Minimum density [g cm$^{-3}$] | Maximum density [g cm$^{-3}$] | Minimum temperature [K] | Maximum temperature [K] | Number of EOS points | References |
|--------------|---------------------|-------------------------------|-------------------------------|--------------------------|--------------------------|-----------------------|------------|
| Hydrogen     | 33                  | 0.001                         | 798.913                      | 15625                    | 6.400×10$^8$             | 401                   | [70–75]    |
| Helium       | 9                   | 0.387                         | 10.457                        | 500                      | 2.048×10$^8$             | 228                   | [76, 77]   |
| Boron        | 16                  | 0.247                         | 49.303                        | 2000                     | 5.174×10$^8$             | 314                   | [78]       |
| Carbon       | 9                   | 0.100                         | 25.832                        | 5000                     | 1.035×10$^8$             | 162                   | [79, 80]   |
| Nitrogen     | 17                  | 1.500                         | 13.946                        | 1000                     | 1.035×10$^8$             | 234                   | [81]       |
| Oxygen       | 6                   | 2.486                         | 100.019                       | 10000                    | 1.035×10$^8$             | 76                    | [82]       |
| Neon         | 4                   | 0.895                         | 15.026                        | 1000                     | 1.035×10$^8$             | 67                    | [83]       |
| Sodium       | 9                   | 1.933                         | 11.600                        | 10000                    | 1.293×10$^8$             | 193                   | [84, 85]   |
| Magnesium    | 23                  | 0.431                         | 86.110                        | 20000                    | 5.174×10$^8$             | 371                   | [86]       |
| Aluminum     | 15                  | 0.270                         | 32.383                        | 10000                    | 2.156×10$^8$             | 240                   | [87]       |
| Silicon      | 7                   | 2.329                         | 18.632                        | 50000                    | 1.293×10$^8$             | 85                    | [88, 89]   |
| LiF          | 8                   | 2.082                         | 15.701                        | 10000                    | 1.035×10$^8$             | 91                    | [90]       |
| B$_4$C       | 16                  | 0.251                         | 50.174                        | 20000                    | 5.174×10$^8$             | 291                   | [91]       |
| BN           | 16                  | 0.226                         | 45.161                        | 20000                    | 5.174×10$^8$             | 311                   | [92]       |
| CH$_4$       | 16                  | 0.072                         | 14.376                        | 6736                     | 1.293×10$^8$             | 247                   | [93, 94]   |
| CH$_2$       | 16                  | 0.088                         | 17.598                        | 6736                     | 1.293×10$^8$             | 248                   | [93, 94]   |
| C$_2$H$_3$   | 16                  | 0.097                         | 19.389                        | 6736                     | 1.293×10$^8$             | 247                   | [93, 94]   |
| CH           | 16                  | 0.105                         | 21.000                        | 6736                     | 1.293×10$^8$             | 248                   | [93, 94]   |
| C$_2$H       | 16                  | 0.112                         | 22.430                        | 6736                     | 1.293×10$^8$             | 245                   | [93, 94]   |
| MgO          | 19                  | 0.357                         | 71.397                        | 20000                    | 5.174×10$^8$             | 286                   | [95]       |
| MgSiO$_3$    | 16                  | 0.321                         | 64.158                        | 6736                     | 5.174×10$^8$             | 284                   | [96, 97]   |

TABLE I. Parameters of the 21 EOS tables in this database. A total of 4869 first-principles calculations were combined.
IV. RESULTS FOR SINGLE COMPOUNDS

![Graph](image)

**FIG. 1.** Density-temperature-pressure conditions of our first-principles simulations of helium [77]. Isobars, isentropes, and shock Hugoniot curves are shown. In the upper panel, we include the interior conditions of Jupiter [44, 122] and main-sequence stars of different masses [13] as well as the highest-pressure conditions reached in recent shock wave experiments on CH [123], B$_4$C [91], MgO [124], BN [92], and MgSiO$_3$ [125], and CO$_2$ [126]. The corresponding temperatures were not measured but derived from simulations.

In this section, we outline the basic functions of our FPEOS database for single compounds. In the following two sections, we will discuss the properties of specific mixtures and then query the database to compute properties of all binary mixtures. In Table 1, we provide the density, $\rho$, and temperature ranges of the EOS tables of eleven elements and ten compounds in our database. We chose helium as an example to illustrate the calculations and plots that our database provides for all these 21 materials. In Fig. 1, we directly plot the EOS points from the first-principles simulations in $T$-$P$ and $T$-$\rho$ spaces. We added isobars that we obtained via a 2D spline interpolation of $P(\rho, T)$ that we also employ to interpolate the internal energy, $E(\rho, T)$. As a guide for future ramp compression experiments, we also plotted a collection of isentropes that we derived from the relationship $\frac{\partial P}{\partial T}_S = -T \frac{\partial^2 P}{\partial \rho^2} V / \partial E V$.

Then we added different shock Hugoniot curves that predict the states generated in dynamic compression experiments. By only measuring the shock and particle velocities, they provide a direct way to determine the EOS of materials at extreme conditions. The sample material initially has the internal energy, pressure, and volume, $\{E_0, P_0, V_0\}$. Under shock compression, the material reaches a final state denoted by $\{E(\rho, T), P(\rho, T), V\}$. The conservation of mass, momentum, and energy across the shock front leads to the Rankine-Hugoniot relation [10, 127, 128],

$$[E(\rho, T) - E_0] + \frac{1}{2} [P(\rho, T) + P_0] [V - V_0] = 0. \quad (2)$$

The volume, $V$, follows from the density, $\rho = m/V$. For helium, we set $\rho_0 = 0.1235 \text{ g cm}^{-3}$. In Fig. 2, we show the resulting shock Hugoniot curve that has a pronounced compression maximum of $\rho/\rho_0 = 5.32$ at $T = 151,000 \text{ K}$ and $P = 370 \text{ GPa}$. If internal degrees of freedom are excited at high $T$ and $P$, the typical compression ratio of an ideal gas ($\rho/\rho_0 = 4$) can be exceeded because these excitations increase the internal energy $E$, which is then compensated by a decrease in volume to satisfy Eq. (2). For conditions under consideration in this article, it is the excitation of K and L shell electrons that introduce one or two compression maxima into the Hugoniot curves that we compute here.

In Fig. 2, we also show the effect of relativistic electrons that increases the shock compression for temperatures above $10^8 \text{ K}$. Since relativistic effects are not included in our PIMC computations, we derived them for an ideal electron gas assuming complete ionization. We also show a Hugoniot curve with radiation effects. Assuming an ideal black body behavior, we very approximately derived the photon contribution to the EOS using $P_{\text{rad}} = (4\sigma/3c)T^3$ and $E_{\text{rad}} = 3P_{\text{rad}}V$, where $\sigma$ is the Stefan-Boltzmann constant and $c$ is the speed of light in vacuum. We find that radiative effects are important for temperatures above $5 \times 10^6 \text{ K}$, which are well above the temperature necessary to completely ionize the K shell electrons of the helium nuclei.

For every material, our PIMC and DFT-MD results can be combined into a single EOS table that can be smoothly interpolated. In Fig. 3, we plot the internal energy of helium for a collection of isochores. To reduce the range of the Y axis, we removed the contribution...
from ideal Fermi gas of electrons and classical nuclei. In this figure, we also show that our results converge to the predictions of the Debye plasma model [21] in the limit of very high temperature. At lower temperatures, this model quickly fails because it does not include any bound states.

IV. RESULTS FOR MIXTURES

In Ref. [68], we demonstrated that ideal mixing approximation works well for temperatures above \(2 \times 10^5\) K and shock compression ratios greater than \(\sim 3.2\). The magnitude of nonideal mixing effects were found to be small and the shock Hugoniot curve of BN, B\(_4\)C, MgO, and MgSiO\(_3\) could all be reproduced with high precision by mixing the EOSs of the elemental substances at constant pressure and temperature. The good agreement included the regimes of K and L shell ionization that lead to compression maxima on Hugoniot curves. This remarkable agreement is the basis for the mixture calculations that we implemented into our FPEOS database.

Neglecting all inter-species interactions, the linear mixing approximation assumes all extensive properties of the mixtures can be derived by adding the contributions from the components 1 and 2 at the \(P\) and \(T\) conditions of interest as follows:

\[
V_{\text{mix}} = N_1 V_1 + N_2 V_2 ,
\]

\[
m_{\text{mix}} = N_1 m_1 + N_2 m_2 ,
\]

\[
E_{\text{mix}} = N_1 E_1 + N_2 E_2 ,
\]

where all variables have been normalized per formula unit. \(N_1\) and \(N_2\) specify how many formula units of species 1 and 2 are contained in one unit of the mixture. The mass density of the mixture is given by \(\rho_{\text{mix}} = m_{\text{mix}}/V_{\text{mix}}\).

In Fig. 4, we compare the resulting shock Hugoniot curve of various mixtures of silicon and CH plastic. Plastics are typical coating materials of ICF capsules that may be doped with heavier elements to modify their behavior to absorb radiation. Elemental silicon was predicted to have two compression maxima that correspond
We find excellent agreement with the revent shock wave to be sufficient to eliminate the lower L shell maximum. While the oxygen curve shows two compression maxima, already a carbon content of C:O=1:2 appears not to respectively obtain $E_0=188.158$ and $188.1574$ Ha/CO. As expected, the resulting Hugoniot curve falls in between those of elemental carbon and oxygen. While the oxygen curve shows two compression maxima, already a carbon content of C:O=1:2 appears to be sufficient to eliminate the lower L shell maximum. We find excellent agreement with the revent shock wave experiments on CO$_2$ by Crandall et al. [126]. All measurements agree with our predictions within error bars.

Neon is the most difficult material to transform into a metal, followed by helium [130]. In Fig. 6, we study the shock properties of mixtures of the two inert gases. Neon exhibits two compression maxima while helium shows one at much lower pressure. Consequently, shock compression maxima shift down in pressure with increasing helium contents. A small neon contents as low as He:Ne=4:1 appears to be sufficient to cause two compression maxima.

Finally, in Fig. 7 we compare the shock properties of water [59, 131, 134], hydrogen peroxide, and alumina with those of their elemental constituents. We predict water to exhibit two compression maxima that, despite a shift to higher pressures, are similar to the K and L shell ionization maxima of the oxygen. For compression ratios between 3.4 and 3.7, the Hugoniot curve that we derived with the linear mixing approximation is in very good agreement with fully interacting DFT-MD results of Ref. [131]. This adds support to the prediction in Ref. [68] that the linear mixing approximation works very well for compression ratios of 3.2 and larger. Both theoretical Hugoniot curves are in agreement with the re-analysis in Ref. [132] that shifted the experimental data point obtained by Podurets et al. [133] to slightly lower densities.

We predict the shock Hugoniot curve of H$_2$O$_2$ to exhibit only a single compression maximum at $T=$...
3.784×10^6 K, \( P = 141 \, 600 \, \text{GPa} \) and \( \rho / \rho_0 = 4.639 \). Despite having a higher atomic oxygen fraction than \( \text{H}_2\text{O} \), the lower L shell ionization appears only as a shoulder in the Hugoniot curve of \( \text{H}_2\text{O}_2 \), which is a consequence of its higher initial density. 1.713 g cm\(^{-3}\). A density increase reduces the compression maxima along the Hugoniot curve because particles interact more strongly, which increases the pressure and thus reduces the compression ratio (see Figs. 2 and 5 as well as Ref. [77]). If the initial density would be lowered to 1.35 g cm\(^{-3}\) or less, the ionization of L shell electrons would again lead to a separate compression maximum. We set \( E_0 = -151.48932 \, \text{Ha per formula unit (FU)} \) in all Hugoniot calculations of \( \text{H}_2\text{O}_2 \).

For the computation of shock Hugoniot curve of alumina (\( \text{Al}_2\text{O}_3 \)) were assumed a corundum crystal structure and used \( E_0 = -708.807 \, \text{Ha/FU} \) and \( \rho_0 = 3.9929 \, \text{g cm}^{-3} \) for the initial conditions. The resulting Hugoniot curve only exhibits a single maximum, which is a surprise because oxygen and aluminum both show separate K and L shell maxima. However, these maxima are offset in pressure from one another and since both nuclei are present in this compound, their combined effects remove the L shell maximum. Furthermore the initial density of alumina is rather high, which reduces the magnitude of any compression maximum.

### V. RESULTS FROM DATABASE APPLICATIONS

In Ref. [97], it was shown that the regimes of pressure and thermal ionization can be distinguished from the slope, \( \frac{\partial E}{\partial \rho} \)\_\( T \). At low density and high temperature, this slope is negative because with decreasing density, more and more free-particle states become available, more electrons become ionized, and as a result, the internal energy increases. This is called the thermal ionization regime, which is often described by the Saha ionization equilibrium [15]. Conversely, at high density the slope \( \frac{\partial E}{\partial \rho} \)\_\( T \) becomes positive for two reasons. First, there is the confinement effect, which increases the kinetic energy of the free electrons and, second, the orbitals of the bound electrons hybridize and may even be pushed into the continuum of free-particle states, which is commonly referred to as pressure ionization. In Fig. 8, we employ the condition, \( \frac{\partial E}{\partial \rho} \)\_\( T \) = 0, to distinguish between these two ionization regimes for six materials selected from our database. As expected, one finds low-Z materials like helium and \( \text{CH}_4 \) to switch from thermal to pressure ionization at a lower density compared to BN, nitrogen, carbon, and oxygen. Still, if one plots these transition lines in temperature-pressure space, they move much closer together (see second panel of Fig. 8).

Furthermore, our FPEOS database enables us to efficiently compute the shock Hugoniot curves of all 21 compounds and 194 meaningful mixtures. In Fig. 9, we compare the conditions of shock compression maxima on all resulting Hugoniot curves for a 1:1 mixing ratio of
FIG. 8. The dividing lines between the regimes of thermal and pressure ionization are shown in temperature-density and temperature-pressure spaces.

FIG. 9. Conditions of the compression maximum on the shock Hugoniot curves of 21 elements and compounds as well as their mixtures. All materials show compression maxima larger than 4.3 that reflect the effects from electronic excitation. Low-Z materials like helium and helium-rich mixtures already reach their compression maxima for lower temperatures while silicon-rich mixtures require much higher temperatures to excite their K shell electrons.

FIG. 10. Conditions of the compression maximum on the shock Hugoniot curves of 21 elements and compounds as well as their mixtures. All materials show compression maxima larger than 4.3 that reflect the effects from electronic excitation. Low-Z materials like helium and helium-rich mixtures already reach their compression maxima for lower temperatures while silicon-rich mixtures require much higher temperatures to excite their K shell electrons.

We can identify a number of trends in Fig. 9 but in general predicting the compression maxima of a specific mixture is not trivial [63, 135]. We find the mixtures of silicon exhibit a compression maximum at higher temperature, which is consistent with the ionization of the K shell electrons. Mixtures of helium tend to exhibit a compression maximum at lower temperature. However, mixtures with hydrogen do not follow this trend because it may be the other element in the mixture with hydrogen that is responsible for introducing the compression maximum.

To study this trend, we study how strongly the temperature of shock compression maximum correlates with the average nuclei charge \( \langle Z \rangle \) of the mixture in Fig. 10. As expected, one finds some support for the trend of \( T_{\text{max}} \) to increase with \( \langle Z \rangle \) but the correlation is not very strong. There are many mixtures with silicon that have \( T_{\text{max}} \sim 2 \times 10^6 \) K but there are also several mixture with helium that have a similar \( T_{\text{max}} \).

In Fig. 11, we converted the conditions of maximal shock compression into a \( u_p-u_s \) plot. The shock and particle velocities were derived from \( u_p = \sqrt{\xi \eta/m} \) and \( u_s = \sqrt{\xi/(\eta m)} \) where \( \xi = (P_1 - P_0)V_0, \eta = 1 - V_1/V_0, \) and \( m \) is the mass of one formula unit. We find that shock and particle velocities at maximum compression very closely follow the linear relationship, \( u_s^{\text{max}} = 1.2727 \times u_p^{\text{max}} - 0.8588 \) km/s, over a wide \( u_p \) range from 50 to 350 km/s. The largest deviations from this trend are only +1.2% and −1.9%. This relationship, that we derived for the different compression maxima, shares similarities with the linear \( u_p-u_s \) relationships that have been constructed for individual materials [84, 96] or groups of materials like fluid metals [136]. For a very high particle velocities of \( \sim 400 \) km/s, the shock velocity has been found to rise faster than linear [84, 96] but the corresponding pressures and temperatures (\( \sim 10^6 \) GPa and \( \sim 10^7 \) K) cannot yet be reached in present-day planar shock experiments.

When we plot the temperature-pressure conditions of all computed Hugoniot maxima in log-log space in
FIG. 10. Temperature condition of the compression maximum on the shock Hugoniot curves of 21 elements and compounds as well as their mixtures is shown as a function of the average ionic charge $\langle Z \rangle$ of these materials. While generally the compression maximum shifts towards higher temperature with increasing $Z$, the correlation is found to be not very strong.

FIG. 11. The particle and shock velocities are shown for the points of maximum compression along shock Hugoniot curves of 21 elements and compounds as well as their mixtures. With good accuracy, the dataset can be represented by the linear relationship of $u_s = 1.2727 \times u_p - 0.8588 \text{ km/s}$ (dashed orange line). The largest positive deviation is seen for a Si:Al mixture, in which case the fit underpredicts $u_s$ by 5.2 km/s, or 1.2%. The largest negative deviation is found for a H:CH$_4$ mixture, in which case the fit overpredicts $u_s$ by 5.4 km/s, or 1.9%.

FIG. 12. Pressure-temperature conditions for the points of maximum compression along shock Hugoniot curves of 21 elements and compounds as well as their mixtures. As expected, pressure and temperature are highly correlated, which can be represented by the fit $\log(T/\text{K}) = 0.5074 + 3.7469 \log(P/\text{GPa})$. However, specific materials deviate substantially from this fit. The temperature of the maximum compression point of a H:Ne Hugoniot curve is 38% lower than this fit would imply. Conversely, the temperature of a B:Al mixture is 49% higher than predicted by the fit.

VI. CONCLUSIONS

By assembling results from $\sim 5000$ FP computer simulations of 21 elements and compounds, we have constructed a general-purpose FPEOS database for computation of matter at extreme conditions. It is our goal
FIG. 13. Conditions of the compression maximum on the shock Hugoniot curves of eight cases for which the mixtures exhibit a higher compression ratio than its endmembers. The lines emerge because a range of mixing ratios were considered.

to make the calculations of shock Hugoniot curves and a ramp compression paths so efficient – without compromising the precision of PIMC and DFT-MD methods – that they become routine in the design and the analysis of WDM experiments. We thus provided our EOS tables as well the C++ and Python codes for the interpolation and the generation of various thermodynamic functions as supplementary material [69].

By invoking the linear mixing approximation at constant pressure and temperature, we first studied a selected number of binary mixtures, computed their shock Hugoniot curves, and related the resulting compression maxima to the ionization of L and K shell electrons. Then we applied our database to study the behavior of these maxima in 194 mixtures and identified trends in pressure, temperature, particle and shock velocity. Finally we identified eight unusual mixtures that should exhibit a higher shock compression ratio than their respective endmembers.

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