

Optical properties of highly compressed polystyrene: An *ab initio* studyS. X. Hu (胡素兴),^{1,*} L. A. Collins,² J. P. Colgan,² V. N. Goncharov,¹ and D. P. Kilcrease²¹Laboratory for Laser Energetics, University of Rochester, 250 East River Road, Rochester, New York 14623, USA²Theoretical Division, Los Alamos National Laboratory, Los Alamos, New Mexico 87545, USA

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Using all-electron density functional theory, we have performed an *ab initio* study on x-ray absorption spectra of highly compressed polystyrene (CH). We found that the *K*-edge shifts in strongly coupled, degenerate polystyrene cannot be explained by existing continuum-lowering models adopted in traditional plasma physics. To gain insights into the *K*-edge shift in warm, dense CH, we have developed a model designated as “single mixture in a box” (SMIAB), which incorporates both the lowering of the continuum and the rising of the Fermi surface resulting from high compression. This simple SMIAB model correctly predicts the *K*-edge shift of carbon in highly compressed CH in good agreement with results from quantum molecular dynamics (QMD) calculations. Traditional opacity models failed to give the proper *K*-edge shifts as the CH density increased. Based on QMD calculations, we have established a first-principles opacity table (FPOT) for CH in a wide range of densities and temperatures [$\rho = 0.1\text{--}100\text{ g/cm}^3$ and $T = 2000\text{--}1\,000\,000\text{ K}$]. The FPOT gives much higher Rosseland mean opacity compared to the cold-opacity-patched astrophysics opacity table for warm, dense CH and favorably compares to the newly improved Los Alamos ATOMIC model for moderately compressed CH ($\rho_{\text{CH}} \leq 10\text{ g/cm}^3$), but remains a factor of 2 to 3 higher at extremely high densities ($\rho_{\text{CH}} \geq 50\text{ g/cm}^3$). We anticipate the established FPOT of CH will find important applications to reliable designs of high-energy-density experiments. Moreover, the understanding of *K*-edge shifting revealed in this study could provide guides for improving the traditional opacity models to properly handle the strongly coupled and degenerate conditions.

DOI: [10.1103/PhysRevB.96.144203](https://doi.org/10.1103/PhysRevB.96.144203)**I. INTRODUCTION**

Optical properties characterize the response of materials to external electromagnetic radiation ranging from microwaves to x rays. For materials at ambient conditions, optical spectroscopic probes often reveal the intricacies of the electronic structure. Heating and compression can cause variations in the spectroscopic features, reflecting the changing environment experienced by the atoms in the material [1–6]. In turn, such spectroscopic measurements can be used to accurately infer information about the electron density and temperature in materials under extreme conditions, provided that reliable theoretical models can be applied. Traditional atomic models [7–11] exist for plasma spectroscopy under classical plasma conditions. These models are employed to incorporate density and temperature effects, including line broadening and shifting, into the calculations of plasma spectroscopy [12–14] by atomic physics codes and generally provide a reliable guide to understanding the spectral features in diluted and hot plasma conditions. These traditional models can break down, however, for dense plasmas. For example, recent experimental studies [15–19] have revealed a lack of understanding of line shifting in hot but dense plasmas and have stimulated renewed theoretical interest [20–23] in the important physical concept of *continuum lowering* or *ionization potential depression* (IPD). So far, these studies have been limited to a strongly coupled but nondegenerate plasma regime (i.e., solid density or moderately compressed hot plasma conditions). How atomic physics can be modified further for extremely compressed materials at relatively low temperatures remains to be answered. Namely, the optical

properties of strongly coupled, fully degenerate materials under high compression might be completely different from our understandings gained from existing studies.

Matter at extreme conditions of density and temperature exists in many natural environments such as in astrophysical objects like brown dwarfs and planetary interiors. Recently, various high-energy-density (HED) facilities have achieved such exotic material conditions. For example, the imploding shell of an inertial confinement fusion (ICF) target [24–28] can reach compressions of hundreds or even thousands times ambient conditions with its temperature still remaining below its Fermi temperature [29,30]. In such HED systems, the optical properties (opacity/emissivity) determine the radiation energy transport. Therefore, a theoretical understanding of the optical properties of extremely dense and warm matter becomes essential in order to complement the experimental campaigns. This endeavor will not only provide reliable opacity/emissivity tables for ICF/HED experimental designs but will also make it possible to examine extreme HED systems using traditional spectroscopic tools.

Polystyrene (CH) is one of the materials often chosen for high-energy-density-physics (HEDP) and ICF target designs. Its optical properties, such as x-ray absorption under extreme conditions, are crucial for understanding the response of dense material mixtures to electromagnetic radiations. In this paper, we present a comprehensive *ab initio* study of the opacity of polystyrene over a wide range of densities and temperatures. Using the quantum molecular dynamics (QMD) method, based on an all-electron density functional theory (DFT), we thoroughly investigated the behavior of x-ray absorption in response to changes in various conditions. Our *ab initio* results revealed certain mechanisms behind the shift in the carbon *K* edge in very dense polystyrene that cannot be

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explained by existing continuum-lowering models. Instead, the behavior of the K -edge shift can only be understood through combining the lowering of the continuum and the rising of the Fermi surface. Performing such *ab initio* calculations over the various densities ($\rho_{\text{CH}} = 0.1\text{--}100\text{ g/cm}^3$) and temperatures ($T = 2000\text{--}1\,000\,000\text{ K}$), we produced a first-principles opacity table (FPOT) of CH for ICF/HEDP applications. We find that our QMD-based FPOT gives an-order-of-magnitude higher opacity than the widely used astrophysics opacity table (AOT) [31] for soft x rays below the carbon K edge. On the other hand, comparisons with modern opacity libraries such as the newly revised ATOMIC model [32] agree favorably with FPOT for a wide range of CH conditions (up to $\rho_{\text{CH}} \approx 10\text{ g/cm}^3$). However, ATOMIC underestimates the CH opacity by a factor of 2–3 for extremely high densities ($\rho_{\text{CH}} \geq 50\text{ g/cm}^3$) at which the ATOMIC model gives much larger K -edge upshifting than QMD predictions.

The paper is arranged as follows: The DFT-based QMD method is briefly described in Sec. II, which is followed in Sec. III by a detailed study of K -edge shifting in dense CH plasmas. The established FPOT is compared with the widely used AOT and ATOMIC in Sec. III C, including the total Rosseland mean opacity and the grouped opacity tables. Our conclusions are presented in Sec. IV.

II. AB INITIO METHOD FOR X-RAY ABSORPTION IN HED SYSTEMS

Our all-electron QMD calculations were performed with the Vienna *ab initio* simulation package (VASP) [33–35] with electrons treated quantum mechanically by a plane-wave finite-temperature DFT description. In contrast to most QMD calculations that employ frozen-core pseudopotentials for warm, dense CH, [36–40] we represent the electron-ion interaction by the Coulombic potential, $-Z/r$ with $Z = 1$ or $Z = 6$, respectively, for hydrogen and carbon ions. The electron exchange correlation is given by the Perdew-Burke-Ernzerhof (PBE) functional [41] in the generalized-gradient approximation. Under the Born-Oppenheimer approximation, the self-consistent electron density is first determined for a randomly chosen ion configuration by solving the Kohn-Sham DFT equation. [42] Next, the classical ions are moved by the combined electronic and ionic forces, using Newton's equation. This molecular dynamics procedure is repeated for thousands of time steps from which both static and optical properties can be directly evaluated. Finally, the electrons and ions are in thermodynamic equilibrium with equal temperatures ($T_e = T_i$).

Since the pure Coulomb potential describes the electron-ion interactions, we must take high-energy (large-momentum) plane waves in order to sample the tightly bound $1s$ -core electron of carbon. Therefore, we first test the energy cutoff (E_{max}) for the plane-wave expansion in solving the Kohn-Sham equation. Figure 1 presents the convergence results for CH at a density of $\rho = 4.0\text{ g/cm}^3$ with a temperature of $T = 125\,000\text{ K}$, in which the (a) pressure and (b) internal energy are plotted as a function of the maximum E_{max} . In these Γ -point calculations, we have also examined the supercell size effect by varying the number of atoms from $N = 8$ to $N = 54$. Figure 1 indicates that converged static properties of the system

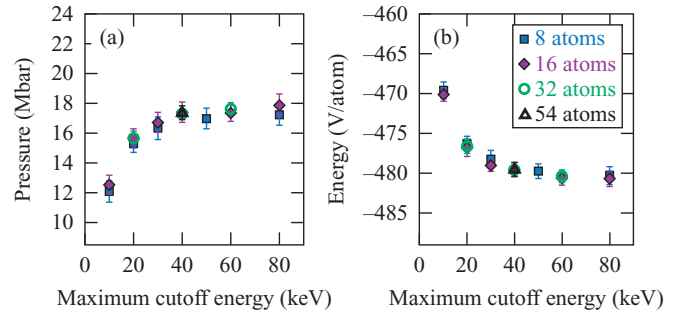


FIG. 1. Results of convergence tests for polystyrene (CH) at $\rho = 4.0\text{ g/cm}^3$ and $T = 125\,000\text{ K}$: (a) pressure and (b) internal energy as a function of the energy cutoff (E_{max}) and the number of atoms (N) in the supercell. The pure Coulombic potentials are used in our *all-electron* quantum molecular dynamics (QMD) calculations. The convergence of static properties is generally reached for $E_{\text{max}} \geq 40\text{ keV}$ and $N \geq 16$.

can be obtained for $E_{\text{max}} \geq 40\text{ keV}$ and $N \geq 16$ for which the variation of P and E is within $\sim 1\%$. This larger energy cutoff is required to accurately sample the deeply bound $1s$ orbital. We used the energy cutoff of $E_{\text{max}} \geq 40\text{ keV}$ and the number of atoms $N \geq 16$ to perform all of the all-electron QMD calculations presented in this paper.

After we ran the calculations for thousands of QMD steps, we obtained a sufficient trajectory of ionic configurations. We then chose five to ten uncorrelated snapshots from these ionic configurations to calculate the x-ray absorption spectra of dense CH by computing the velocity dipole matrix elements D_{mn} using the DFT wave functions from VASP. The quantity D_{mn} then determines the frequency-dependent Onsager coefficients via the Kubo-Greenwood formula as [43]

$$L_{ij}(\omega) = \frac{2\pi(-e)^{4-i-j}}{3Vm_e^2\omega} \sum_{mn} F_{mn} |D_{mn}|^2 \times \left(\frac{E_m + E_n}{2} - H \right)^{i+j-2} \delta(E_m - E_n - \hbar\omega), \quad (1)$$

where $V = 1/\rho$ is the atomic specific volume, E_m (E_n) is the energy of the m th (n th) state, and H is the enthalpy (per atom) of the system. The quantity F_{mn} is the difference between the Fermi-Dirac distributions for states m and n at temperature T . To obtain the frequency-dependent absorption coefficient $\alpha_K(\omega)$, we need the electric conductivity and the index of refraction of the system. From the real part of the electric conductivity, $\sigma_1(\omega) = L_{11}(\omega)$, the imaginary part of the electric conductivity can be evaluated from the principal value integral:

$$\sigma_2(\omega) = -\frac{2}{\pi} P \left(\int \frac{\omega' \sigma_1(\omega')}{\omega'^2 - \omega^2} d\omega' \right). \quad (2)$$

The dielectric function $\varepsilon(\omega) = \varepsilon_1(\omega) + i\varepsilon_2(\omega)$ can then be calculated by

$$\begin{aligned} \varepsilon_1(\omega) &= 1 - \frac{4\pi}{\omega} \sigma_2(\omega), \\ \varepsilon_2(\omega) &= \frac{4\pi}{\omega} \sigma_1(\omega). \end{aligned} \quad (3)$$

Using the dielectric function, we obtain the real part of the refraction index:

$$n(\omega) = \sqrt{\frac{|\varepsilon(\omega)| + \varepsilon_1(\omega)}{2}}. \quad (4)$$

Finally, the mass-absorption coefficient (α_m) is equal to the absorption coefficient (α_K) divided by the mass density [44], i.e.,

$$\alpha_m(\omega) = \frac{\alpha_K(\omega)}{\rho} = \frac{4\pi \bar{\sigma}_1(\omega)}{c \times \bar{n}(\omega) \rho}, \quad (5)$$

with the speed of light c . The “bar” over σ_1 and n stands for averaging over a sample of uncorrelated snapshots. Such *ab initio* calculations of optical properties [45,46] have been used to build a first-principles opacity table (FPOT) of deuterium for ICF applications [47]. Since the QMD calculations are *limited to certain maximum photon energy* (due to the number of bands used), we extrapolate the monotonically decreasing high-energy tail of the electrical conductivity to higher energies, by using a *Drude-type fitting formula* with an additional ω^3 term described in Ref. [47].

To test the sensitivity of the mass-absorption coefficient to the number of atoms and the k -point sampling, we performed a series of tests for dense polystyrene at a density of $\rho = 4.0 \text{ g/cm}^3$ and a temperature of $T = 125 \text{ 000 K}$, which are shown in Fig. 2 for the absorption coefficient as a function of photon energy $h\nu$. For the $N = 16$ and Γ -point calculations [Fig. 2(a)], we varied the energy cutoff (E_{max}) from 20 to 60 keV, demonstrating that the overall difference is negligible between $E_{\text{max}} = 40 \text{ keV}$ and $E_{\text{max}} = 60 \text{ keV}$, while the $E_{\text{max}} = 20 \text{ keV}$ calculation gives a slight energy shift near the K edge of carbon, $h\nu \approx 280 \text{ eV}$. Therefore, setting

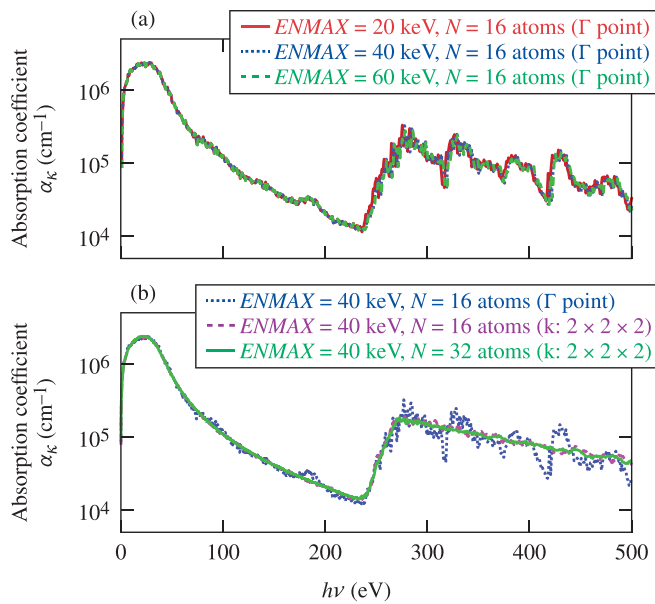


FIG. 2. Convergence testing results of x-ray absorption spectra for CH at $\rho = 4.0 \text{ g/cm}^3$ and $T = 125 \text{ 000 K}$: (a) E_{max} varying from 20 to 60 keV with 16-atom and Γ -point calculations; (b) changing the number of atoms and k point with $= 40 \text{ keV}$. Convergence of x-ray absorption calculations is generally reached for $E_{\text{max}} \geq 40 \text{ keV}$ and $N \geq 16$.

$E_{\text{max}} \geq 40 \text{ keV}$ should guarantee convergent results. Finally, we plot the k -point convergence results in Fig. 2(b) along with a variation in N from $N = 16$ to $N = 32$. Compared to the Γ -point-only calculation, a $2 \times 2 \times 2$ Monkhorst-Pack grid gave much smoother results, especially above the carbon K edge. Moreover, the $N = 16$ calculation (dashed red line) is almost identical to the $N = 32$ case (solid green line) shown in Fig. 2(b). Therefore, all of our results are from convergent QMD calculations, using $E_{\text{max}} \geq 40 \text{ keV}$, $N \geq 16$, and a k -point grid of $2 \times 2 \times 2$.

Under the multigroup diffusion approximation, the Rosseland mean opacity K_R is often used for the radiation transport in hydrodynamics simulations. In general, the grouped Rosseland mean opacity is calculated by

$$K_R(\omega_1:\omega_2) = \frac{\int_{\omega_1}^{\omega_2} n(\omega)^2 \frac{\partial B(\omega, T)}{\partial T} d\omega}{\int_{\omega_1}^{\omega_2} n(\omega)^2 \frac{1}{\alpha_m(\omega)} \frac{\partial B(\omega, T)}{\partial T} d\omega}, \quad (6)$$

for a group of photon energies between $\hbar\omega_1$ and $\hbar\omega_2$. The Planck function,

$$B(\omega, T) = \frac{\hbar\omega^3}{4\pi^3 c^2} \frac{1}{e^{\hbar\omega/k_B T} - 1},$$

depends on the emitting photon energy and the material temperature. We obtain the total/gray Rosseland mean opacity for $\omega_1 = 0$ and $\omega_2 = \infty$.

III. RESULTS AND DISCUSSION

A. Shifting of carbon K edge in highly compressed polystyrene

This subsection presents our *ab initio* results for x-ray absorption spectra of CH compressed to densities from $\rho = 4 \text{ g/cm}^3$ up to $\rho = 100 \text{ g/cm}^3$. We will focus on understanding the shift of the carbon K edge in highly compressed polystyrene. For the case shown in Figs. 1 and 2, polystyrene was compressed to $\rho = 4 \text{ g/cm}^3$ with a temperature of $T = 125 \text{ 000 K}$. According to Eq. (5), the x-ray absorption coefficient depends on the real parts of the electrical conductivity $\sigma_1(\omega)$ and the refractive index $n(\omega)$. Figure 3 shows

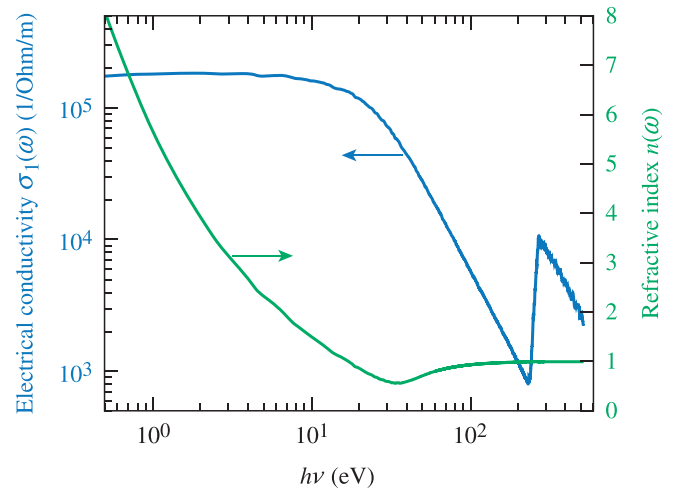


FIG. 3. The electrical conductivity and the refractive index are plotted as functions of the photon energy (frequency) for the case of polystyrene at $\rho = 4.0 \text{ g/cm}^3$ and $T = 125 \text{ 000 K}$.

the variation of these two quantities with respect to the electromagnetic wave frequency (photon energy). For CH at this condition, the plasma frequency ω_p is so high that the energy associated with plasma oscillation is $\hbar\omega_p \approx 35$ eV. We see from Fig. 3 that the refractive index has a minimum at $\omega = \omega_p$ (strong absorption); $n(\omega)$ increases dramatically for $\omega \ll \omega_p$ and approaches $n(\omega) = 1$ for $\omega \gg \omega_p$, as expected. The electrical conductivity has a relatively small variation at low frequencies, dropping exponentially for $\omega > \omega_p$ until the $1s$ -core electrons become accessible for conduction. The sharp jump of $\sigma_1(\omega)$ at $h\nu = E_K \approx 272$ eV by an order of magnitude indicates the carbon $1s$ -core electrons are involved in conduction (absorption). For $h\nu > E_K$ the electrical conductivity exhibits a monotonic decrease as does the absorption coefficient. The calculated carbon K edge of $E_K \approx 272.3$ eV is about $\Delta E \approx 11.7$ eV smaller than the experimental value in ambient CH. This is a well-known problem in DFT for the PBE exchange-correlation functional, which somewhat underestimates the band gap in general. We have scanned the PBE-DFT calculations from a low CH density of $\rho = 0.1$ g/cm³ (approaching isolated-atom limit) to $\rho = 1.0$ g/cm³ and $\rho = 10$ g/cm³. When the resulting K edges are compared with experimental K -edge measurements for an isolated carbon atom and solid CH, the observed deficit of $\Delta E \approx 11.7$ eV is *not significantly changed*. Thus we have added this energy deficit for the K edges calculated from our PBE calculations (discussed below). We have found that the more accurate hybrid Heyd-Scuseria-Ernzerhof (HSE) exchange-correlation functional [48] can provide a much better band gap (ΔE reducing to ~ 2 eV) but is computationally more time consuming than PBE. In addition, the self-energy calculation through the Green's function G and the screened Coulomb interaction W (GW) in the “quasiparticle” picture should give more accurate energy levels than the PBE-DFT calculations presented here. However, for our bare Coulomb potentials used here these GW calculations are found to be extremely expensive and memory demanding, if not impossible, due to the very high cutoff energy ($E_{\text{cut}} \geq 40$ keV) required. Even for the very large computational resources currently available to us (maximum memory of 512 GB per node), we could not obtain reliable GW results at this time.

In Fig. 4, we examine the x-ray absorption spectra of CH compressed to a higher density of $\rho = 10$ g/cm³ at three temperatures: $T = 31\,250$ K, $125\,000$ K, and $500\,000$ K. Figure 4 indicates that the peak absorption moves to a slightly higher frequency as ω_p shifts up because of the increasing electron density in comparison with the case of $\rho = 4$ g/cm³ (Fig. 2). For the three temperatures, Fig. 4 shows that the carbon K edge shifts down as the temperature increases from $T = 31\,250$ K to $T = 500\,000$ K; the K edge also becomes less steep at higher temperatures (see the inset of Fig. 4). To understand these temperature-induced features of the carbon K edge, we plotted the density of states (DOS's) as a function of energy in Fig. 5 for the three corresponding cases. For the lowest temperature of $T = 31\,250$ K, the narrow peak at $E \approx -220$ eV is the $1s$ -core electron energy band of carbon ions for $10\times$ -compressed CH. Compared with the $1s$ -core electron binding energy (-282 eV) of an isolated carbon atom, the observed upshift of core electron energy is caused both by the screening of ionized free electrons to the C^{6+} core and

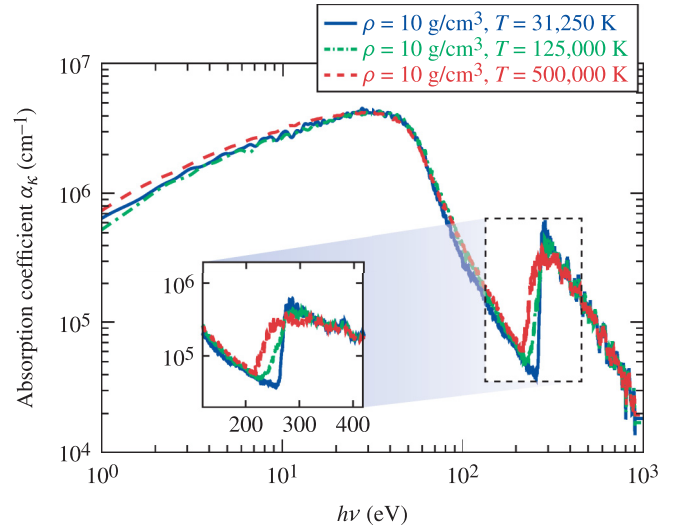


FIG. 4. The x-ray absorption coefficient α_K is plotted as a function of the photon energy (frequency) for CH at a higher density of $\rho = 10$ g/cm³ with temperatures varying from $T = 31\,250$ K to $T = 500\,000$ K. The small K -edge movement is seen for the highest-temperature case of $T = 500\,000$ K (dashed red line).

by the microfield influences from surrounding ions. Since we always set the continuum level at $E_C = 0$ in these calculations, the energy upshifting of $1s$ -core electrons corresponds exactly to the traditional view of “continuum lowering” (or “ionization potential depression”). Namely, the energy gap between the $1s$ -core energy level and the continuum is decreased. At such relatively low temperatures ($T = 31\,250$ K and $T = 125\,000$ K), the Fermi surface is located at around $E_F \approx 50$ eV (marked by vertical lines in Fig. 5), which means the energy bands below E_F are fully occupied. Therefore, the $1s$ -core electrons can only make a transition to the unoccupied bands above E_F . This determines the K -edge location at $E_K = E_F - E_{1s}$,

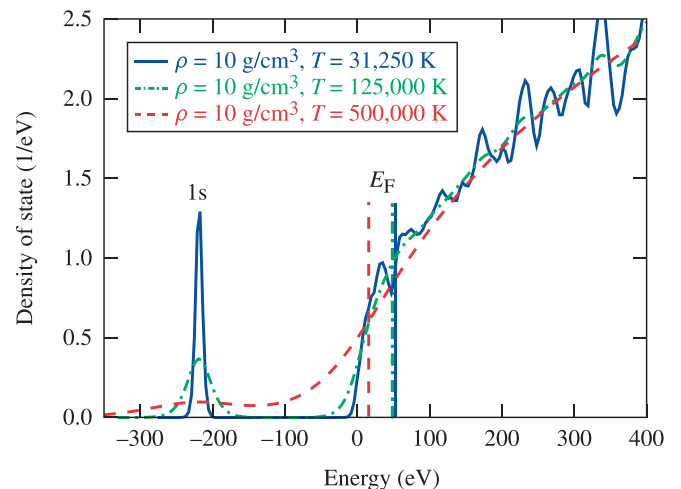


FIG. 5. The density of state (DOS) is plotted for each corresponding case shown in Fig. 4. The Fermi-energy levels are marked by the vertical lines for each case. The downshift of the Fermi surface at $T = 500\,000$ K explains the K -edge movement seen in Fig. 4 (dashed red line) at this higher temperature.

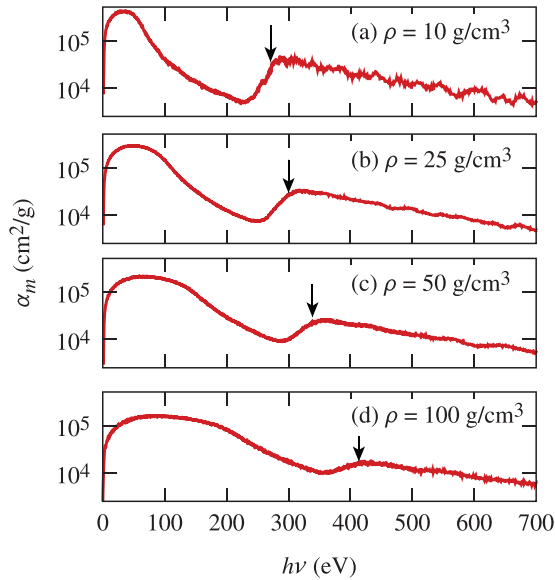


FIG. 6. The x-ray absorption spectra are plotted as a function of the photon energy (frequency) for CH at different densities varying from (a) $\rho = 10 \text{ g/cm}^3$ to (d) $\rho = 100 \text{ g/cm}^3$ with a fixed temperature of $T = 125\,000 \text{ K}$. The K -edge upshifting with CH density increasing is clearly seen.

which accounts exactly for the K edge seen in Fig. 4 (blue and green curves). Because the temperature is well below the Fermi temperature ($T < E_F/k$), the Fermi distribution has a sharp cutoff around $E \sim E_F$ so that the lower the temperature, the sharper the K edge (illustrated by the inset in Fig. 4). When the temperature increases to $T = 500\,000 \text{ K}$, Fig. 4 shows that the K edge shifts down by $\sim 30 \text{ eV}$ (solid red line) with respect to the low- T cases. This is caused by the downshift of the Fermi surface because of thermal excitation of electrons to high-energy bands, as indicated by the dashed red lines in Fig. 5. The $1s$ band is also significantly broadened as T increases, where the Fermi smearing is used to determine the partial occupancies on each orbital. For a fixed density, the K -edge shifting of tens of eV can be used as a measure of material temperature (see the inset in Fig. 4).

Next, we examine the x-ray absorption spectra at even higher densities. The results are plotted in Fig. 6 for CH densities of 10, 25, 50, and 100 g/cm^3 at the same temperature of $T = 125\,000 \text{ K}$. The black arrow in each panel of Fig. 6 marks the location of the carbon K edge for each situation. One sees that the K edge continuously upshifts as the density increases, while the contrast of the K edge is gradually reduced. To understand why the K edge shifts in the way shown in Fig. 6, we plot the DOS as a function of band energy in Fig. 7 for the corresponding cases with the vertical dashed line indicating the location of the Fermi surface. Figure 7 shows that (1) the $1s$ band of the carbon ion moves up and broadens in response to increasing ρ ; and (2) the Fermi surface continuously elevates with increasing density (leading to the reduction of absorption caused by Pauli blocking, discussed in the next section). As a consequence, the K edge is again determined by $E_K = E_F + E_C - E_{1s}$ (note that we set the “continuum” at $E_C = 0$). The reduced K -edge contrast can be attributed to the density-induced broadening seen in Fig. 7. For

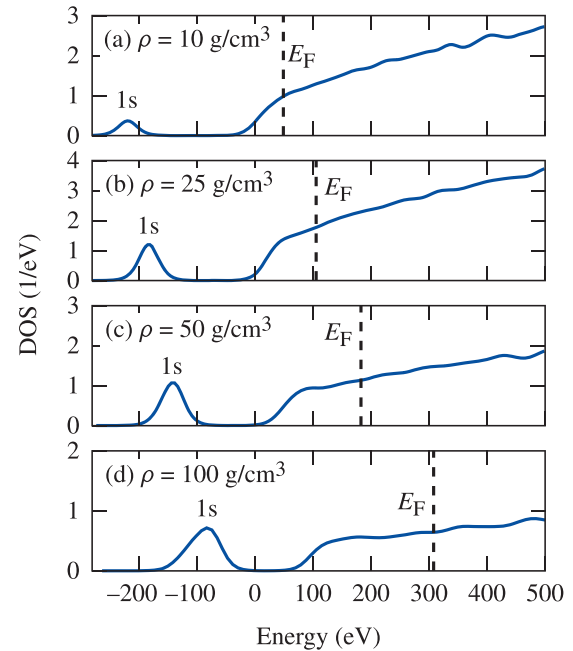


FIG. 7. The DOS for each corresponding case shown in Fig. 6. Again, the vertical dashed line in each panel marks the Fermi surface location for each case.

example, the width of the $1s$ band varies from $\Delta E_{1s} \approx 38 \text{ eV}$ to $\Delta E_{1s} \approx 64 \text{ eV}$, respectively, for density increasing from $\rho = 25 \text{ g/cm}^3$ to $\rho = 100 \text{ g/cm}^3$.

B. Single mixture in a box (SMIAB) model

Figure 8 plots the carbon K -edge locations from our QMD calculations as a function of the CH density for the same temperature of $T = 125\,000 \text{ K}$. Our QMD results are represented by the blue diamonds, in which an error bar of $\Delta E \sim 11.7 \text{ eV}$ is assigned to account for the gap inaccuracy of the PBE exchange-correlation functional (discussed above). To test if the traditional density-dependent continuum-lowering models can explain these observations, we compare our QMD results with the different predictions of the carbon K edge in Fig. 8. Four IPD models, including Ecker-Kröll [8], Stewart-Pyatt [7], modified ion sphere [9], and Crowley [11], are used to calculate the carbon K edges with the corresponding mass densities obtained for different ion charge states $Z = 1$ to 5 by the Fermi-Dirac average-atom model. These IPD models, except the Crowley, did not include the Pauli blocking (degeneracy) effect. We also compare with the ATOMIC model [49–52]. ATOMIC accounts for IPD by an occupation probability formalism through use of explicitly computed plasma microfields [10] that gradually destroy atomic bound states through field ionization as the density increases. ATOMIC also includes an explicit factor that accounts for the rising of the Fermi surface (also known as Pauli blocking [53]) that in this case, causes the carbon K edge to increase. These model predictions are represented by the dashed colored lines with symbols in Fig. 8. Once again, the K edge for an ion charge state is calculated in these models by $E_K = E_C - E_{1s}$ (isolated), where E_C is the model-predicted IPD and E_{1s} (isolated) is the $1s$ -electron ionization potential

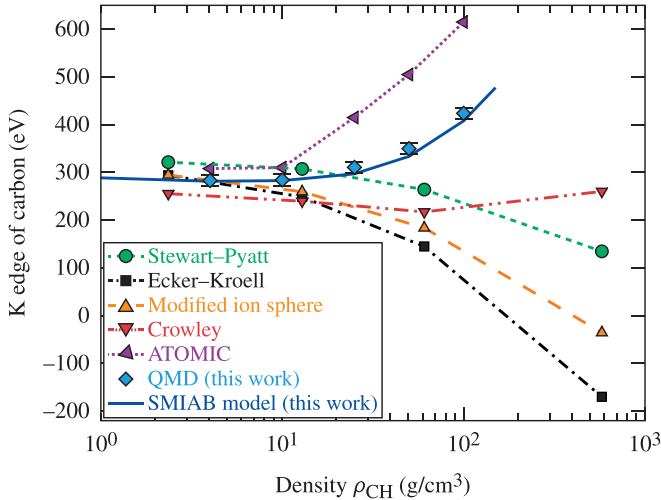


FIG. 8. The QMD-predicted K edges of carbon are plotted as a function of CH density. Predictions from five different continuum-lowering models, including the Stewart-Pyatt [7], Ecker-Kröll [8], modified ion-sphere [9], and Crowley models [11], and the newly improved ATOMIC code (Los Alamos) using the Hummer-Mihalas continuum-lowering model [10] with the consideration of Pauli blocking, are also drawn in the same figure with open symbols. The “single mixture in a box” (SMIAB), developed in this work, is indicated by the blue solid line.

of an isolated carbon ion at a charge state of Z . As can be seen from Fig. 8, the first three models predict the K -edge downshifting as ρ_{CH} (Z) increasing, while the Crowley model, taking the electron degeneracy into account, gives a slight upshift of the K edge, but only for the case of $Z = 5$. Dramatically different from all four model predictions, our QMD results (blue diamonds) show significant upshifts for $\rho_{\text{CH}} > 25 \text{ g/cm}^3$. Note that our QMD calculations do not assume what ionization level is reached for each density studied. The ATOMIC model also predicts the K -edge upshifting (caused by the Pauli blocking term discussed earlier) but by a greater amount than the QMD results. For example, at $\rho_{\text{CH}} = 100 \text{ g/cm}^3$ and $T = 125\,000 \text{ K}$ the carbon K edge from our *ab initio* calculation gives $E_K = 422.9 \text{ eV}$ (QMD), in contrast to model predictions of $E_K = 235.9 \text{ eV}$ (Stewart-Pyatt), 76.4 eV (Ecker-Kröll), 136.1 eV (modified ion sphere), 226.6 eV (Crowley), and 613.5 eV (ATOMIC), respectively.

To fully understand the physics behind the K -edge upshifting resulting from our *ab initio* calculations, we have developed a “single mixture in a box” (SMIAB) model for strongly coupled fully degenerate plasmas. The SMIAB model consists of three components: (1) the continuum-lowering effect on the $1s$ -electron binding energy E_{1s} of targeted carbon ions; (2) the average degree of ionization Z of the mixture determined by an average-atom model [54]; and (3) the rise of the electronic Fermi surface because of compression, which can be calculated by $E_F = \hbar^2/2m(3\pi^2 Z \rho_{\text{CH}}/A_{\text{CH}})^{2/3}$ in the case concerned, where $\hbar = h/2\pi$ is the reduced Planck constant, $A_{\text{CH}} = (A_C + A_H)/2$ is the averaged atomic weight of the single mixture (the “basic unit” of polystyrene), and ρ_{CH} is the mass density. To determine the $1s$ -electron binding energy E_{1s} of the carbon ion in a dense polystyrene (ρ_{CH}),

we placed the single mixture (one carbon atom and one hydrogen atom) inside a cubic box having a volume of $V = L^3 = A_{\text{CH}}/\rho_{\text{CH}}$. Using periodic boundary conditions, this simple SMIAB picture mimics the microscopic environment of each individual carbon atom experienced in such dense plasmas. We then calculated the band energy of E_{1s} by self-consistently solving the following Kohn-Sham equation for all N_b electronic orbitals (in atomic units):

$$\left\{-\frac{1}{2}\nabla^2 + V_N(r) + V_H[\rho_e](r) + V_{xc}[\rho_e](r)\right\}\psi_i(r) = E_i\psi_i(r), \quad (7)$$

where V_N represents the electron interactions with both carbon and hydrogen nuclei, and the electron density and the Hartree term are defined as

$$\rho_e(r) = \sum_{i=1}^{N_b} |\psi_i(r)|^2; \quad V_H[\rho_e](r) = \int \frac{\rho_e(r')}{|r-r'|} dr'. \quad (8)$$

Since the exchange-correlation term V_{xc} and the Hartree term depend on the electron density $\rho_e(r)$, which is again a function of $\psi_i(r)$, the above Kohn-Sham equation can be solved self-consistently. The PBE exchange-correlation functional is also adopted in the above E_{1s} calculations.

To determine the average ionization Z , we employed the Fermi-Dirac average-atom model with local-density approximation for the exchange-correlation functional. We have applied the equal-pressure mixing rule [55] to gauge the ionization degree from each individual species. For example, for the chosen temperature of $T = 125\,000 \text{ K}$ and a CH density ρ_{CH} , we vary the densities of pure carbon (ρ_C) and pure hydrogen (ρ_H) in the average-atom model to satisfy the following two conditions simultaneously: (1) the equal-pressure condition, e.g., $P_C = P_H$; and (2) the additive-volume condition, i.e., $\rho_{\text{CH}} = A_{\text{CH}}/(A_H/\rho_H + A_C/\rho_C)$. With these matching conditions, the average-atom model gives the ionization degree (Z_C and Z_H) for each species. Therefore, the average ionization of equimolar CH can be expressed as $Z = (Z_C + Z_H)/2$. The resulting Z is then used to calculate the elevated Fermi surface. Finally, the K -edge locations are computed for each CH density by $E_K = E_F + E_C - E_{1s} + \Delta E$ (note the continuum is also set to $E_C = 0$). As an example, the SMIAB results are listed in Table I for CH at different densities but having the same temperature ($T = 125\,000 \text{ K}$), which is also plotted

TABLE I. The predicted carbon K edges (with $\Delta E = 11.7\text{-eV}$ shift) by the single mixture in a box (SMIAB) model for polystyrene (CH) at different densities with a fixed temperature of $T = 125\,000 \text{ K}$. Z and E_F are the average charge of CH and the electron Fermi energy, respectively.

Density (g/cm^3)	E_{1s} (eV)	Z	E_F (eV)	E_K (eV)
1	-268.2	1.20	8.7	288.6
4	-245.5	1.47	24.3	281.5
10	-220.1	1.79	51.0	282.8
25	-179.5	2.12	105.8	297.0
50	-142.2	2.37	179.7	333.6
100	-93.5	2.59	302.7	407.9
150	-58.5	2.69	407.4	477.6

in Fig. 8 by the blue solid line. As with the QMD results, we added $\Delta E = 11.7$ eV to the SMIAB-predicted carbon K edge, accounting for the inaccuracy of the PBE functional for energy-gap calculations.

From Fig. 8 one can see that this simple SMIAB model does a much better job than all other continuum-lowering models at explaining the QMD results (within an error of ± 15 eV), indicating that the SMIAB with periodic boundary conditions can predict the continuum lowering of atomic ions affected by surrounding ions/electrons in a dense material. Moreover, the rising of the Fermi surface, estimated from the average-atom model, reproduces well the essence of electron degeneracy in such systems while the existing models failed to predict the K -edge locations in such strongly coupled and fully degenerate dense materials. A similar model [56] also explains the K -edge shifting in extremely dense pure carbon plasmas. We anticipate that the simple SMIAB model will find important applications in studying HED properties of extremely dense materials. For example, one can use this simple model to infer density of highly compressed materials from experimental K -edge measurements. It is noted that the traditional continuum-lowering models, including Ecker-Kröll, Stewart-Pyatt, and modified ion sphere, *did not include the Pauli blocking (degeneracy) effect*; adding the Pauli blocking effect from our average-atom model to these continuum-lowering models still results in different K -edge locations from our QMD calculations and SMIAB model predictions.

C. Opacity comparisons between QMD and other models

For ICF and HED applications, we have extended our QMD calculations of opacity to cover a wide range of CH in the warm-dense-matter regime. Specifically, we have scanned the CH density from $\rho_{\text{CH}} = 0.1$ g/cm³ to $\rho_{\text{CH}} = 100$ g/cm³ with the temperature range of $T = 2000$ to 1 000 000 K. To make a global opacity table, we first compare the QMD-calculated opacity with model predictions. Two opacity models are compared here: (1) the AOT [31] and (2) the ATOMIC model, [32] which contains improvements to the free-free opacity contribution compared to earlier work [49]. ATOMIC is designed to produce opacities over very large ranges of temperatures and densities and is most commonly used for astrophysical modeling at low and moderate densities as well as in radiation-hydrodynamics codes for simulations of ICF and HED experiments. Still, it is of interest to explore how these ATOMIC opacities compare at the high (compressed) densities studied here.

ATOMIC is a multipurpose opacity and kinetics code, [50] which has been recently used to generate new Los Alamos OPLIB tables for hydrogen through zinc [49] that employ atomic data from the Los Alamos suite of atomic physics codes [51] and an equation of state (known as ChemEOS) based on a chemical picture [52]. Recently, improvements to the free-free component of the opacity in ATOMIC have been implemented [32] based on enforcement of the conductivity sum rule, [57] which resulted in excellent agreement between ATOMIC calculations, DFT-molecular dynamics (MD) simulations, [47] and pseudo-atom-potential calculations [58] for the opacity of deuterium. For the CH opacity studied here,

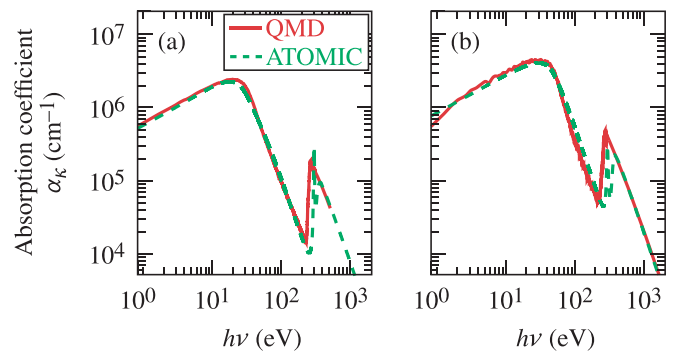


FIG. 9. Comparisons of the absorption coefficient α_K of polystyrene as a function of photon energy $h\nu$ between our QMD calculations and LANL's newly improved ATOMIC model are made for densities of (a) $\rho = 4$ g/cm³ and (b) $\rho = 10$ g/cm³, both at $T = 125\,000$ K.

a multielement procedure [59] was used within ATOMIC that iterates to ensure a self-consistent free-electron density for a specified temperature and mass density of the constituent elements.

We first compare the QMD results (red lines) with the ATOMIC predictions (green dashed lines) in Figs. 9 and 10 for CH densities varying from $\rho_{\text{CH}} = 4$ g/cm³ to $\rho_{\text{CH}} = 100$ g/cm³ at $T = 125\,000$ K, in which the absorption coefficient α_K is plotted as a function of x-ray photon energy $h\nu$. At relatively low densities ($\rho_{\text{CH}} = 4$ g/cm³ and $\rho_{\text{CH}} = 10$ g/cm³), Fig. 9 shows that the improved ATOMIC model agrees overall with QMD in the entire photon energy range. Some detailed differences near the K edge are noticed; for example, the ATOMIC model shows the existence of the $1s \rightarrow 2p$ peak (before the K edge) while QMD gives smooth K edges without distinct bound-bound transition peaks. We argue that the density effect, represented *naturally* in the *ab initio* QMD calculations, has made the $2p$ states of carbon emerge into the continuum. Again, the observed K -edge energy deficit (ΔE) in QMD is caused by the use of the PBE exchange-correlation functional (discussed above). When the CH density is further increased to $\rho_{\text{CH}} = 50$ g/cm³ [Fig. 10(a)] and $\rho_{\text{CH}} = 100$ g/cm³ [Fig. 10(b)], the carbon K -edge shifts are significantly different between QMD and ATOMIC predictions. For example, Fig. 10(a) indicates that for $\rho_{\text{CH}} = 50$ g/cm³, the QMD-predicted carbon K edge is located at $E_K = 349.8$ eV,

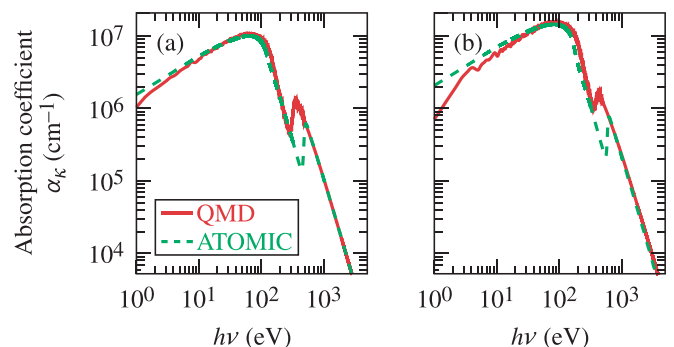


FIG. 10. Similar to Fig. 9 but for CH densities of (a) $\rho = 50$ g/cm³ and (b) $\rho = 100$ g/cm³ both at $T = 125\,000$ K.

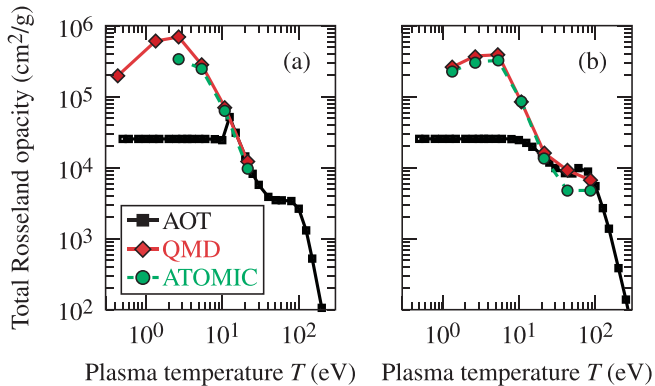


FIG. 11. Comparisons of Rosselland mean opacity (K_R) as a function of CH temperature, among our QMD calculations, the LANL’s ATOMIC model, and the cold-opacity-patched astrophysics opacity table (AOT), are made for CH densities of (a) $\rho = 0.5 \text{ g/cm}^3$ and (b) $\rho = 4.0 \text{ g/cm}^3$.

while, in contrast, the ATOMIC model gives $E_K = 510.8 \text{ eV}$. The difference in the carbon K -edge shift between QMD and ATOMIC is further increased for the highest density ($\rho_{\text{CH}} = 100 \text{ g/cm}^3$). As previously discussed, the ATOMIC model includes the competing effects of IPD (from an occupation probability consideration [49]) that lower the K edge, and the Fermi surface rising [50] that increases the K edge. Both of these effects are density dependent, and it is not yet clear which effect is responsible for the difference found with the QMD results.

Next, we compare the total/gray Rosselland mean opacity K_R of CH in Figs. 11 and 12, calculated by Eq. (6) for different CH densities. In these figures, the total Rosselland mean opacity is plotted as a function of CH temperature. Figures 11(a) and 11(b) show the cases for $\rho_{\text{CH}} = 0.5 \text{ g/cm}^3$ and $\rho_{\text{CH}} = 4.0 \text{ g/cm}^3$, respectively. Besides the QMD and ATOMIC results, we have also added the AOT predictions. Historically, the AOT model had been used to simulate ICF and HED experiments [60–63]. The data in the AOT table were limited to certain temperatures for dense materials, below which the cold opacity was usually “patched in” to enable radiation-hydro simulations. Figure 11(a) indicates that for relatively low CH density ($\rho_{\text{CH}} = 0.5 \text{ g/cm}^3$), all three (QMD,

ATOMIC, and AOT) predictions agree with each other as the temperature increases; for $T < 10 \text{ eV}$, the cold-opacity-patched AOT underestimates K_R by more than an order of magnitude in comparison with QMD and ATOMIC. The latter two results are almost identical to each other for a temperature as low as $\sim 5.39 \text{ eV}$, although a factor of ~ 2 difference is observed at $T = 2.69 \text{ eV}$. For low temperatures and moderate densities, convergence of the ChemEOS model is often more difficult to obtain. Also, the role of negative-ion formation and its effect both on the EOS and opacity require further scrutiny. At $\rho_{\text{CH}} = 4.0 \text{ g/cm}^3$, Fig. 11(b) shows a similar situation: the improved ATOMIC model gives overall good agreement with QMD predictions, although some differences are seen for $T = 40\text{--}100 \text{ eV}$.

For higher densities of $\rho_{\text{CH}} = 10 \text{ g/cm}^3$ and $\rho_{\text{CH}} = 50 \text{ g/cm}^3$, Fig. 12 shows a trend similar to that in Fig. 11. Namely, the cold opacity patched to AOT generally underestimates the Rosselland opacity of warm, dense CH by a factor of ~ 10 at low temperatures. The improved ATOMIC model gives a good account of the enhanced opacity in QMD calculations for $\rho_{\text{CH}} \leq 10 \text{ g/cm}^3$. Again, some differences between QMD and ATOMIC are still seen for the relatively high- T regime ($T \sim 40\text{--}100 \text{ eV}$). This may be attributed to the detailed differences near the carbon K edge (see Figs. 9 and 10) since the Rosselland mean opacity weighs more around $h\nu \approx 4kT$ [i.e., the peak of $\partial B(\omega, T)/\partial T$]. Consequently, Fig. 12(b) further shows a larger K_R difference between QMD and ATOMIC for the higher CH density of $\rho_{\text{CH}} = 50 \text{ g/cm}^3$. This is consistent with the K -edge shifting difference seen in Fig. 10(a). Again, the K_R difference is within a factor of ~ 2 , between QMD and ATOMIC, but overall higher than the cold opacity patched to AOT.

Finally, we compare the grouped Rosselland mean opacities between QMD and AOT in Figs. 13 and 14, for CH densities of $\rho_{\text{CH}} = 4 \text{ g/cm}^3$ and $\rho_{\text{CH}} = 50 \text{ g/cm}^3$, respectively. As we discussed above, AOT has been extensively used for radiation-hydrodynamics simulations of ICF/HED experiments.

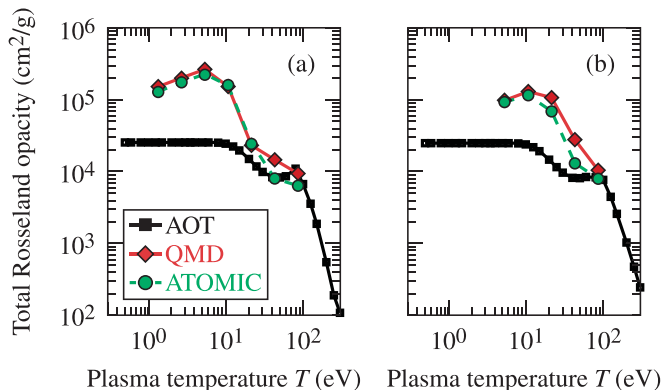


FIG. 12. Similar to Fig. 11 but for CH densities of (a) $\rho = 10 \text{ g/cm}^3$ and (b) $\rho = 50 \text{ g/cm}^3$.

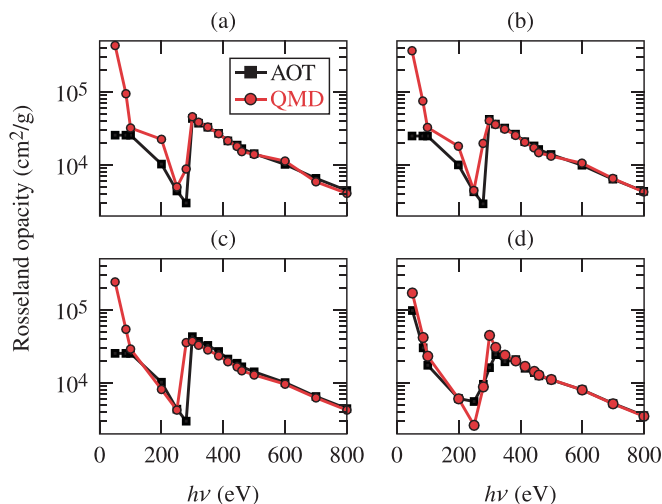


FIG. 13. The grouped Rosselland opacities, from both our QMD calculations and AOT, are plotted as a function of x-ray photon energy ($h\nu$) for a fixed CH density of $\rho = 4 \text{ g/cm}^3$, but at different temperatures of (a) $T = 62\,500 \text{ K}$, (b) $T = 125\,000 \text{ K}$, (c) $T = 500\,000 \text{ K}$, and (d) $T = 1\,000\,000 \text{ K}$.

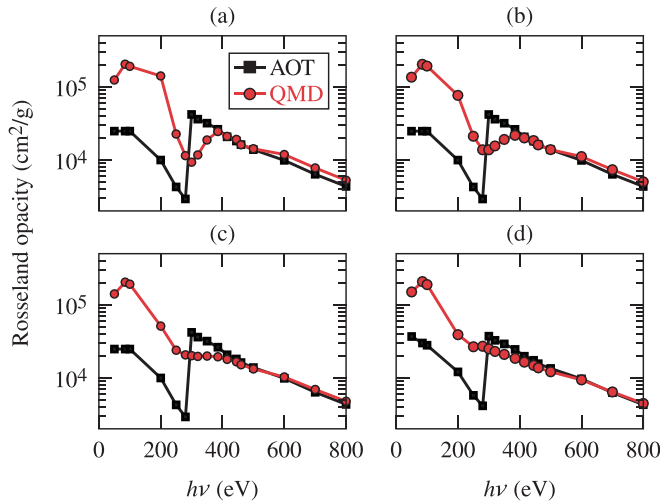


FIG. 14. The grouped Rosseland opacities, from both our QMD calculations and AOT, are plotted as a function of x-ray photon energy ($h\nu$) for a much higher CH density of $\rho = 50 \text{ g/cm}^3$ and at different temperatures of (a) $T = 125\,000 \text{ K}$, (b) $T = 250\,000 \text{ K}$, (c) $T = 500\,000 \text{ K}$, and (d) $T = 1\,000\,000 \text{ K}$.

Forty-eight photon-energy groups are normally used for CH simulations. Therefore, we compare our QMD results with AOT in these figures with the same 48 energy groups. In Fig. 13, the four panels display the grouped Rosseland mean opacities as a function of photon energy in each group for temperatures varying from (a) $T = 62\,500 \text{ K}$ to (d) $T = 1\,000\,000 \text{ K}$. One sees from Fig. 13 that for $T < 500\,000 \text{ K}$, the first and second energy groups ($h\nu \leq 100 \text{ eV}$) are largely different between QMD and AOT; the opacities for high-energy groups above the carbon K edge are almost identical between the two models, although some small differences are seen near the K -edge groups. For a higher density of $\rho_{\text{CH}} = 50 \text{ g/cm}^3$, Fig. 14 shows significant differences between QMD and AOT: (1) The QMD opacities of low-energy groups below K edge ($h\nu \leq 285 \text{ eV}$) are $10\times$ higher than the cold opacity patched to AOT; and (2) the K -edge shifting predicted by QMD is totally absent in AOT. As the temperature increases, our QMD results indicate that the carbon K edge gradually becomes less pronounced and eventually disappears at $T = 1\,000\,000 \text{ K}$ in such a dense polystyrene, shown by Fig. 14(d). This is expected since the $1s$ -core electrons become free (ionized) as T and ρ increase. On the contrary, the AOT model persistently gives the sharp and unshifted K edge of carbon. We expect that these differences will affect radiation transport of low-energy ($h\nu \leq 400 \text{ eV}$) photons in ICF/HED simulations.

IV. CONCLUSION

We have used the DFT-based QMD method to perform *ab initio* studies on the x-ray absorption properties of highly compressed CH. These first-principles QMD studies are made possible by using the all-electron pure Coulombic potentials, which are in contrast to most frozen-core pseudopotential DFT calculations. We found that as CH is compressed to larger and larger densities, the carbon K edge is upshifted significantly, which was missing in traditional IPD models.

The ATOMIC model, which includes IPD effects and the rising of the Fermi surface, also predicts the K -edge upshifting, but to a larger degree than the QMD calculations. In addition, we have developed a physics-based SMIAB model to understand the observed carbon K -edge shifting from our QMD calculations. The simple SMIAB picture reveals that for strongly coupled and fully degenerate dense CH, the carbon K edge is essentially determined by both the lowering of the ionization potential of carbon ion and the rising of the electronic Fermi surface resulting from the compression in dense CH. SMIAB gives good agreement with the *ab initio* results from detailed and large-scale QMD calculations. Such a simple SMIAB model might provide guidance for inferring density/temperature conditions from K -edge-shifting measurements in extremely dense materials.

These *ab initio* calculations have been expanded to a wide range of CH densities and temperatures. The resulting FPOT was compared with two extensively used opacity models (AOT and ATOMIC). We found that the cold-opacity-patched AOT generally underestimated the CH opacity by an order of magnitude for warm, dense polystyrene while the improved ATOMIC model gave a good account for the enhanced opacity seen in QMD for relatively dense CH ($\rho_{\text{CH}} \leq 10 \text{ g/cm}^3$) in the low-temperature regime ($T < 20 \text{ eV}$). As the CH density further increases to $\rho_{\text{CH}} \geq 50 \text{ g/cm}^3$, however, the ATOMIC model gives a factor of ~ 2 lower opacity when compared to our QMD results because of an overestimate of the shift of the K edge in ATOMIC. Finally, the comparisons between our QMD results with the extensively used AOT clearly indicate that the Rosseland mean opacities in low-photon energy groups (below the K edge) can differ by a factor of ~ 10 . We anticipate that these detailed differences will affect the radiation transport of soft x-ray photons in simulations of ICF and HED experiments. We hope such detailed studies will facilitate experimental measurements, in particular, of the observable K -edge shifting in extremely dense materials.

Note added. After our manuscript was submitted, an interesting research paper [64] on K_α , K_β , and K -edge shifts of Fe in dense Be plasmas was published, in which the DFT-based model (MUZE: similar to our SMIAB discussed here) properly explained the observed line/edge shifting in experiments.

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