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Theoretical study on the physical properties of synthesized SrMO_3 ($M = \text{Hf}$ and Pt) oxide perovskites using DFT

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We investigated the physical behavior of SrMO_3 ($M = \text{Hf}$ and Pt) compounds, which are strontium-based oxide perovskites. We utilized the WIEN2k software to simulate and investigate their physical properties. The structural stability of SrHfO_3 and SrPtO_3 was verified using the Birch–Murnaghan equation of states for optimization. We also checked the elastic stability through the computation of elastic constants using the IRelast software. Our results indicate the stability of these compounds and showed their anisotropic, ductility, scratch-resistive, and plastic strain-resistant characteristics. Using the TB-mBJ potential approach, we determined that SrHfO_3 is an insulator, whereas SrPtO_3 is a metal in nature. The density of states computations was used to find the band structure as well as the contribution of different electronic states. Optical property research was conducted using the band gap energies of these substances. Our findings suggest that these crystals have low energy absorption and reflectivity of up to 65%, making them suitable for use in high-frequency UV devices. Specifically, SrHfO_3 is more transparent before the energy point 2.80 eV, while the compound SrPtO_3 after 6.50 eV to 12.0 eV and SrHfO_3 from 12.0 and 14.0 eV. This study represents the first DFT-based investigation of these discussed crystals according to the best of our knowledge.

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1. Introduction

Life requires energy, yet the public rarely has access to it in large numbers. Because perovskite is the crystal utilized to produce and store energy, researchers are still interested in it. Therefore, they are working to create new perovskites with improved characteristics. Despite being essential for human survival, energy is a limited resource that is available to everyone. Because perovskite is the crystal utilized to produce and store energy, researchers are still interested in it. Thus, they are attempting to develop new perovskite compounds with enhanced properties. Each perovskite structure consists of twelve entities of cation “A” and six atoms of cation “B” attached

to an anion. Insulators, conductors, and semiconductors are the three distinct properties of oxide-perovskites. In addition to their excellent electrical properties, these compounds are also capable of possessing a stable crystalline structure. Applications of oxide perovskites that support sustainability and renewable energy are being investigated. They are employed in solid oxide fuel cells, for instance, which provide effective energy conversion with little emissions. Perovskite-based catalysts can also reduce environmental problems, including air pollution. Perovskite materials are being investigated for cutting-edge data storage and processing technologies such as spintronics and magneto-optics because of their distinctive magnetic and electrical properties. The exploration of oxide perovskites presents the possibility of finding unique materials with novel properties, which may result in innovations in a number of disciplines. It has also been pointed out that these materials have applications in lithography, opto-electrics, batteries, and lenses.^{1–5} In particular, oxide perovskites have been the subject of a plethora of research because of the continued interest in learning more about their unique features. Most studies have found them to be mechanically stable and elastically anisotropic.^{6–9} ABO_3 crystals have many uses,^{10–12} including in extremely efficient solar constituents, and are superior for automobile energy storage, electronics, and optics. Stable oxide perovskites are created when oxygen is coupled with inorganic,

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organic, or transition metals. The best options are wide band-gap oxide perovskites. Because of their high potential and broad energy band gap, such molecules can be used in the fabrication of vacuum ultraviolet (VUV) and ultraviolet (UV₃)-operating glass.^{13,14} Ref. 15–24 show that several recent studies on oxide perovskite have been described. Because of its large direct band-gap and ensembles of the imaginary component of the insulating features in the ultra-violet region, BaCsO₃ was found to be appropriate for optoelectronic devices by Harmel *et al.*,¹⁸ who utilized DFT to investigate the characteristics of barium-based oxide perovskite. The work of Daniel *et al.*¹⁹ after looking at a few properties of LiBaO₃, it became clear that it may be used to store energy. Compounds with band gaps larger than 3.10 eV will function well in the UV region of the electromagnetic spectrum. SrHfO₃ and SrPtO₃ are promising candidates for electronic applications because of their nonmetallic properties and their position among the best semiconductors. In this study, we used DFT and the FP-LAPW method to analyze some of the important features of SrMO₃ (M = Hf and Pt) of oxide perovskites in order to use it as a tool for future research. Structure, elasticity, electricity, and optics are all part of these features.

2. Computational methodology

The FP-LAPW method was reused in conjunction with the WIEN2k program to calculate the aforementioned characteristics, and the results have been reported.²⁵ Because the TB-mBJ method may be reused to determine optical and electrical characteristics and the density of states (DOS), it was chosen for this study.²⁶ The exch-correlation potential and GGA are useful techniques for calculating structural and elastic parameters.²⁷ Since this research looks into multiple FP-LAPW base functions, the smallest possible radius for the “muffin tin spheres (RMT)” was chosen as 8.0. Here, we used K_{max} in our model, where the maximum k points in the plane wave expansion represent the magnitude of the convergence required for practical applications. The RMT values were 2.38 au for M = (Hf and Pt) crystals, 1.73 au for O, and 1.67 au for Sr. Increasing the spherical harmonics inside the “muffin-tin-spheres” to $l_{\text{max}} = 11.0$ and decreasing the Fourier expanded charge density to $G_{\text{max}} = 12.0$ au. For the self-consistent field calculations to be considered convergent, the total energy must fall in the energy series of 0.001 Ry. The energy vs. volume curve was analyzed utilizing the Birch–Murnaghan equation of states.^{28,29} The elastic constant utilized in calculating the structural features was calculated with the help of the IRelast program.²⁹ The visual properties of the selected crystals were also determined using the dielectric function.^{30,31}

3. Results and discussion

The outcomes produced by our Tb-Mbj potential techniques were thoroughly scientifically analyzed, as described in this section. Using this method, we will discuss the structural and optical features.

3.1 Structural properties

We investigated the characteristics of SrMO₃ crystals, where M can be either Hf or Pt. These crystals belong to the space group ($Pm\bar{3}m\# 221$) and have a single molecule, cubic unit cell structure, conforming to the ABO₃ formula, revealing the structure of oxide perovskite. The arrangement of atoms within the unit cell follows a specific pattern, with Sr atoms located at (0, 0, 0) positions, M (M = Hf and Pt), and oxygen atoms at (0, 1/2, 1/2), (1/2, 0, 1/2), and (1/2, 1/2, 0), and (0.50, 0.50, 0.50), respectively, as shown in Fig. 1. To predict the arrangement behavior of the chosen crystals, we used Birch–Murnaghan’s equation of state to compute the volume optimization. This allowed us to determine the total energy surrounding V_0 , which is the volume of the unit cell at equilibrium conditions. By reducing the unit cell’s overall energy, we obtained the lowest stable unit cell. We then analytically calculated the data from Birch–Murnaghan’s equation of state to predict the lowest state properties of the unit cell, as shown in Fig. 2. These properties include the steadiness lattice numbers a_0 , the bulk modulus B , and its pressure derivative B' . The true the minimal ground state of the specific chemical is given by E_0 , for which the volume receives its lowest value, V_0 . The structures of crystals with higher negative energy are more stable. Table 1 presents the optimized structural parameters, such as a_0 , E_0 , V_0 , B_0 , and B' , which are necessary for structural comparison. The outcomes confirm that the bulk modulus decreases as the lattice numbers increase, which is consistent with the general trend of this technique. The accuracy and precision of our computed results are demonstrated by these findings. By examining the optimization graphs, we found that SrPtO₃ has more negative energy than SrHfO₃, which supports the idea that SrPtO₃ is more structurally stable, as seen in Table 1. In summary, we have analyzed the structural properties of SrMO₃ crystals and provided detailed information about their unit cell structure, volume optimization, ground state properties, and stability. We hope that these findings will be useful for future research in this field.



Fig. 1 The conventional cubic crystal structure of ternary molecules SrMO₃ (M = Hf, Pt).

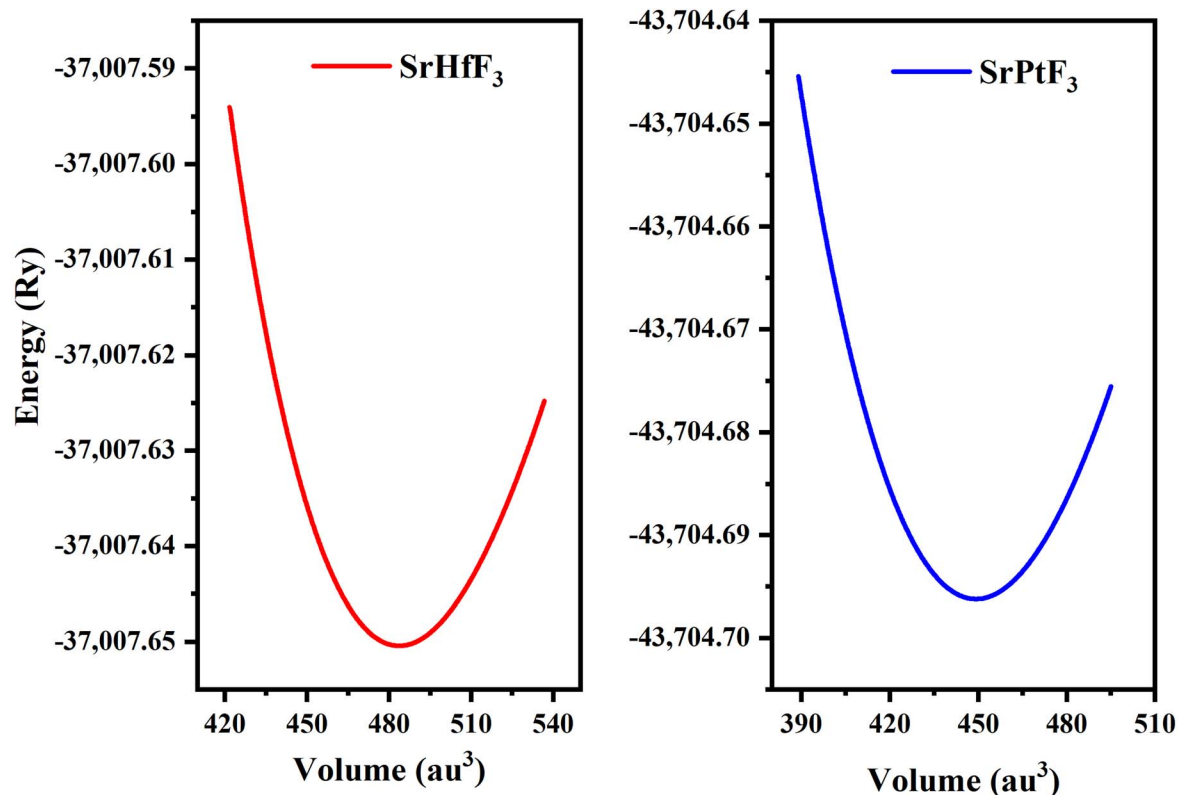


Fig. 2 SrMO_3 ($M = \text{Hf, Pt}$) crystal optimization curve fitted by Birch–Murnaghan's equation of states.

Table 1 Optimized crystal unit cell computed data for SrMO_3 ($M = \text{Hf}$ and Pt) molecules

Compounds	a_0 (Å)	B (GPa)	B'	V_0 (a.u. ³)	E_0 (Ry)
SrHfO_3	4.15	159.49	5.01	483.68	−37007.65
SrPtO_3	4.05	150.20	3.39	449.22	−43704.70

3.2 Electronic properties

We used the Wien2k tool to simulate different parameters of the unit cell and obtained the band structure diagrams, density of states (DOS), and charge arrangement which enabled us to examine the electronic positions of SrMO_3 ($M = \text{Hf}$ and Pt) complexes. It is important to highlight that LDA and GGA calculations significantly underestimate the crucial band separation of semiconductors and dielectrics.^{32,33} It is because its fundamental structure is inadequate for reproducing the exchange–correlation energy or its charge derivative with any degree of certainty. As a means of addressing the band gap underestimation, the modified Becke–Johnson potential (TB-MBJ) was selected because of its widespread application in recent research.^{17,34,35} The energy-band structures for SrMO_3 ($M = \text{Hf, Pt}$) are shown in Fig. 3, and they were generated along the higher symmetry axis of the Brillion zone. E_F for the Fermi barrier, the separation line between the two bands. The valence assembly is illustrated below the Fermi line, whereas the conduction gang is above this level. It is evident from Fig. 3 that SrPtO_3 is metal because of the presence of a transition state

between the valence and conduction bands, but SrHfO_3 is an insulator because of its 5.0 eV indirect energy band gap ($M-\Gamma$). Fig. 4 provides the total and partial densities of the states calculated, from which we obtained a clear image of the energy state contribution of various states in the lower and upper bands. The Fermi line is drawn vertically between the conduction and valence bands and it is assumed that this line represents a barrier with zero energy. The peaks reveal the availability of the electronic energy levels and offer important information about the electronic characteristics and behaviors of the compound. The material's capacity to absorb and emit light at a particular energy is directly correlated with the greatest peaks in the DOS. The permitted energy transitions that contribute to the material's characteristics are shown by these peaks. Each molecule's valence band is located on the left side, while its conduction band is positioned on the right side, relative to the Fermi line. Below are the contributions of the various states at the aforementioned tiers. In the valence band of the SrHfO_3 crystal, the Hf-d state contributes the most energy, with a peak at −3.9 eV and subsequent peaks at −3.10 eV, −1.90 eV, 1.0 eV, and 0.2 eV. Sr-tot also contributes within the same eV range, from −3.90 to 0.30. Its values are as follows: 0.10 at −3.80 eV, 0.30 at −3.20 eV, 0.10 at 1.8 eV, 0.2 at −1.20e V, and 0.20 at −0.50 eV. At −3.80, −3.10, and 1.80 eV, the other values in the graph for the same state of Sr-d are 0.10, 0.20, and 0.05, respectively. The O-p state contributes marginally to the same state. The Sr-d state contributes most to the conduction band of the identical compound between 5.30 and 6.20 eV, with the



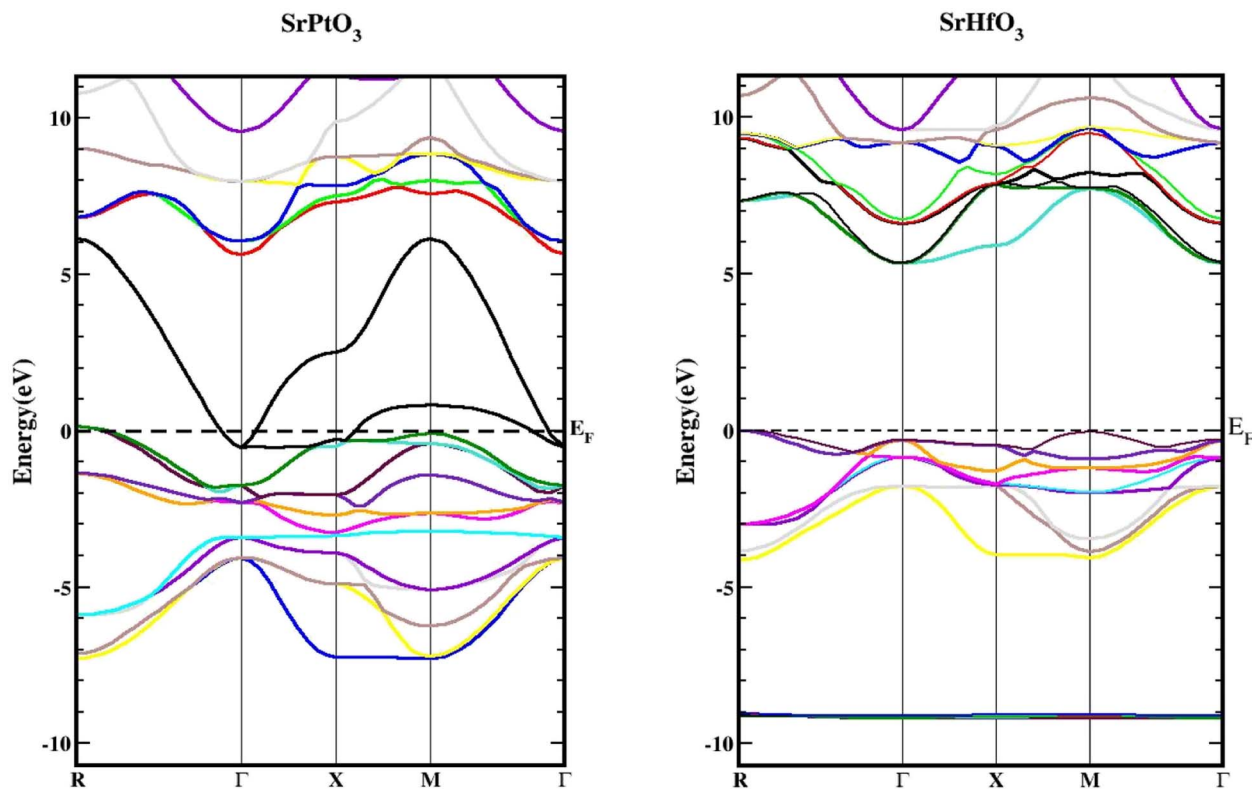


Fig. 3 Band structures of SrMO_3 ($M = \text{Hf, Pt}$) using TB-mBJ approximation.

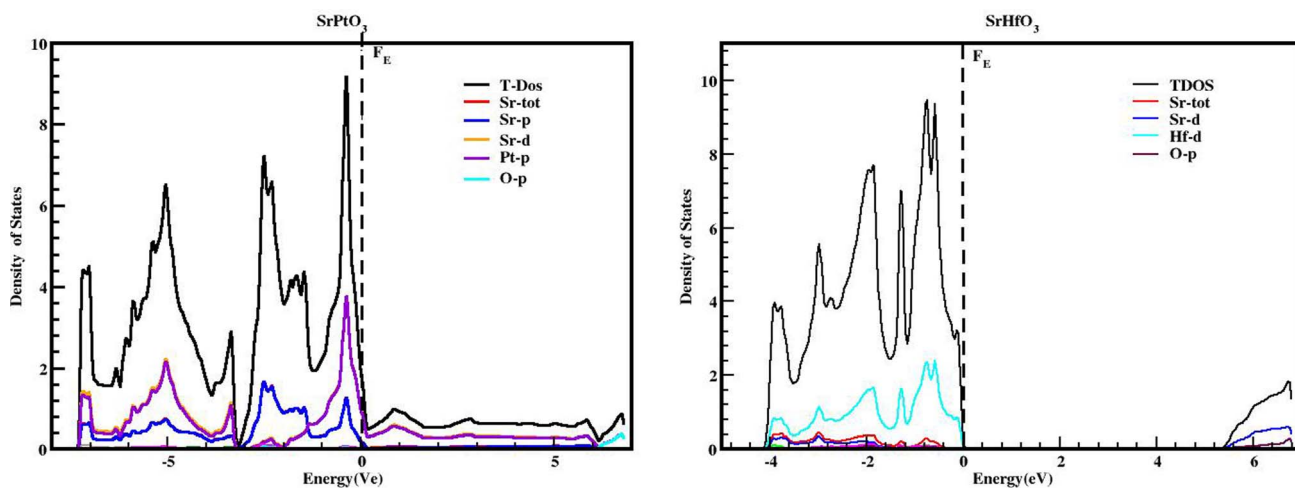


Fig. 4 TDOS and PDOS of SrMO_3 ($M = \text{Hf, Pt}$) crystals obtained using the TB-mBJ approach.

largest peak of 0.50 at 6.10 eV. In the same band, the O-p state also contributes very little as well. The valence band of the second compound (SrPtO_3) contains a number of states with varying degrees of participation, such as the Pt-p state, which exhibits peaks at -6.20 , -5.0 , -4.0 , and 0.2 eV. Similarly, the Sr-p state contributes, with an energy range of -6.0 eV to 0.0 eV. Peaks were observed at -6.0 , -5.0 , -3.0 , -2.0 , and 0.20 eV with energies of 0.20 , 0.30 , 1.40 , 0.20 , and 1.30 , respectively. The Pt-p state contributes energy between 0 and 6.80 eV to the

conduction band, with an approximation of 0.1 remaining in this region.

3.3 Elastic properties

The ability of the material to return to its original state after being stretched is known as elasticity. The stability and hardness of the crystal can be inferred from the results of these computations. Without applying any external force, the compounds' elastic characteristics were determined by

Table 2 Important constants of SrMO₃ (M = Hf, Pt) molecules

Compounds	SrHfO ₃	SrPtO ₃
C_{11}	329.41	189.71
C_{12}	84.63	134.87
C_{44}	65.91	44.75
G	84.67	36.77
A	0.538	1.632
V	0.266	0.384
B/G	1.88	4.09

determining the stress tensor components for small deformation and then adding energy in accordance with the lattice distortion that maintained volume.³⁶ Elastic constants were calculated using the Wien2k and IRelast programs, respectively, with full attention paid to the particulars of the cubic system. We may learn important details about every cubic crystal system from the values of the elastic constants C_{11} , C_{12} , and C_{44} . Table 2 lists some elastic constants that are not related to one another. For a cubic crystal to be mechanically stable, the elastic constants must satisfy the following conditions: to ensure that $C_{11} > C_{12}$, $C_{11} > 0$, $C_{44} > 0$, $C_{11} + 2C_{12} > 0$, and $B > 0$, the following criteria must be satisfied.³⁴ The mechanical stability of our compounds was inferred from the fact that the calculated elastic constants met all of the aforementioned requirements. Based on our findings, SrHfO₃ is typically stiffer than SrPtO₃, with a C_{11} value of 329.41 GPa compared with 189.71 GPa for SrPtO₃. An additional factor, the anisotropic constant (abbreviated as “A”), describes a crystal’s propensity to form microscopic cracks. With this information in hand, engineers can examine the tiny response of a crystal to stress. Using the elastic constants we determined previously, we can calculate the values of A for the compounds we selected using the formula below³⁷

$$A = 2 \times C_{44}/C_{11} - C_{12} \quad (1)$$

If $A = 1$, then the material is isotropic; otherwise, it is anisotropic. In Table 2, we show the “A” values we determined for both of our compounds. The value of “A” for both of these substances is not equal to 1, making them anisotropic; the variance shows how much anisotropy exists. The calculated value of A for the molecule SrHfO₃ is 0.538, while the value for the molecule SrPtO₃ is 1.632, demonstrating that the molecule SrHfO₃ is less isotropic than the molecule SrPtO₃. The elastic constants used to determine other crucial parameters, such as Young’s modulus (represented by “E”), shear modulus (represented by “G”), and Poisson’s ratio (represented by “ν”), are summarized in Table 2. The equations used to determine these values are as follows:^{38–40}

$$E = \frac{9 \times B \times G}{G + 3 \times B} \quad (2)$$

$$\nu = \frac{3 \times B - 2 \times G}{2(G + 2 \times B)} \quad (3)$$

$$G_v = \frac{C_{11} - C_{12} + 3 \times C_{44}}{5} \quad (4)$$

$$G_R = \frac{5 \times C_{44}(C_{11} - C_{12})}{4 \times C_{44} + 3C_{11} - C_{12}} \quad (5)$$

Ductility and brittleness are two of the most important characteristics to look for when analyzing a crystal’s structure. The ductility of a crystal can be described by the parameter known as Cauchy’s pressure, which is given as $(C_{11} - C_{44})$.^{41–44} If the difference between these constants is a positive numerical value, then the material will be ductile; otherwise, it will be brittle. Due to their positive Cauchy’s pressures, 144.95 GPa for SrPtO₃ and 263.51 GPa for SrHfO₃, both of our crystals exhibit ductile properties. On the basis of these calculated values, we conclude that the SrMO₃ (M = Hf, Pt) crystals we selected are ductile, anisotropic, robust, and fracture-resistant from a mechanical standpoint. The modern technological world finds many uses for qualities such as these.

3.4 Optical properties

Utilizing input photons with energies ranging from zero to fourteen electron volts (eV), the anticipated equilibrium lattice constant, and the dielectric function, we were able to compute the optical characteristics of the selected crystals.

3.5 Dielectric function

The dielectric function is denoted by “ $\epsilon(\omega)$ ” and given by equation $\epsilon(\omega) = \epsilon_1(\omega) + i\epsilon_2(\omega)$. The first half represents the real part, while the second is the imaginary part. Fig. 5 depicts the photon energy dispersion by the crystals and their electronic polarizability. According to Fig. 5, the value of the real part of the dielectric constant, $\epsilon_1(\omega)$, for the compound SrHfO₃ was found to be 2.5 at 0 eV, and the maximum value for the same compound was found to be 5.7 at 7.8 eV. According to the Penn model,⁴⁴ if the static dielectric function has a greater value, the band gap energy will be smaller. Also, the real part of the compound SrPtO₃ has a static dielectric function is 9, and its value fluctuates from negative to positive values. In the same Fig. 5, we can see the computed value of the imaginary part of the dielectric function $\epsilon_2(\omega)$ for both of our selected crystals for energies up to 14.0 eV. The first critical peak in the SrHfO₃ and SrPtO₃ dielectric permittivity spectra was found to occur at around 8.0 eV with the average value of 5.5. At the x -symmetry node, the absorption edge causes a sharp optical transition from the lower to the higher band. As soon as the energy barrier is broken, the curve starts to rise and flatten out. For energy sources that are less polluting than 2.0 eV, both the real and imaginary components of dielectric permittivity operate in the same manner.

3.5.1 Refractive index. The refractive index, represented by $n(\omega)$, conductivity, exemplified by $\sigma(\omega)$, absorption coefficient, represented by $I(\omega)$, and reflectivity, represented by $R(\omega)$ may all be determined from the values of $\epsilon_1(\omega)$ and $\epsilon_2(\omega)$, respectively, for any crystal. The calculated values for the refractive index of



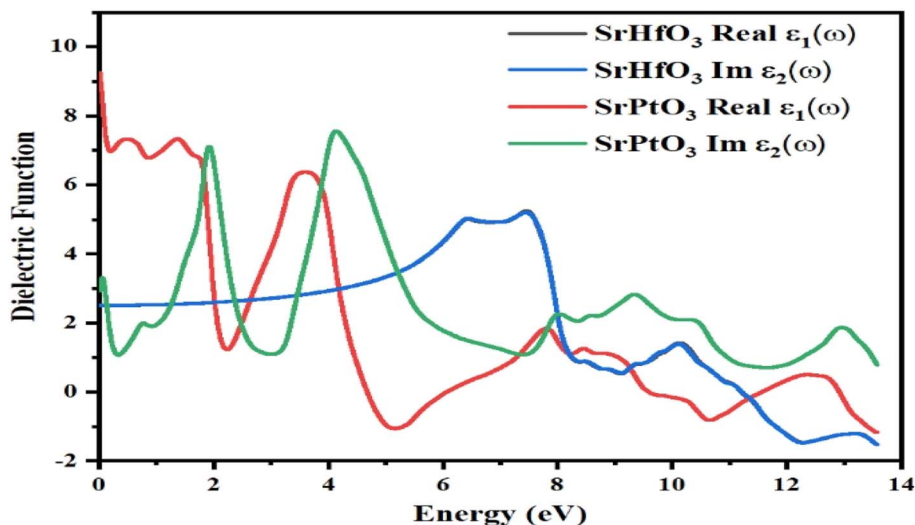


Fig. 5 Dielectric function $\epsilon(\omega)$ of SrMO_3 ($M = \text{Hf, Pt}$).

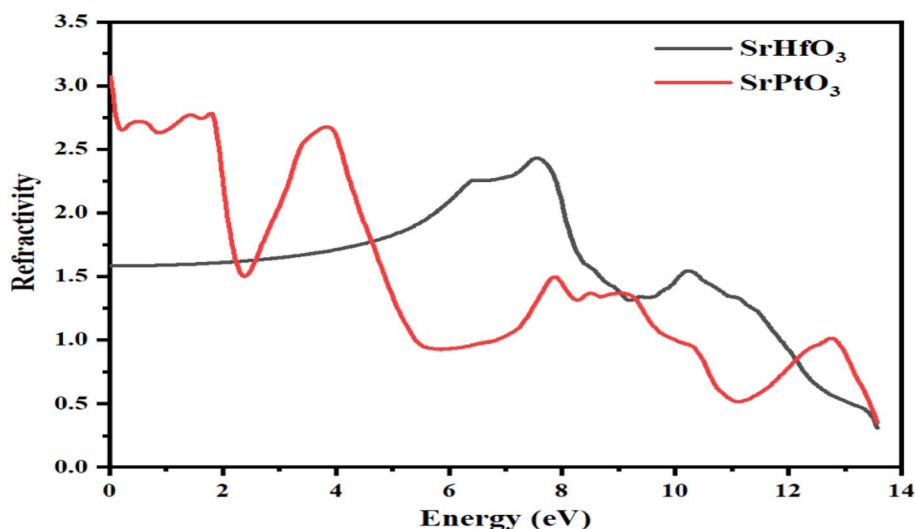


Fig. 6 Calculated refractive index of c SrMO_3 ($M = \text{Hf and Pt}$).

SrMO_3 ($M = \text{Hf and Pt}$) crystals are displayed in Fig. 6. Fig. 6 clearly depicts the refractive index at 0.0 eV, represented by $n(0)$, which is 3.1 for the SrPtO_3 crystal and 1.3 for the SrHfO_3 crystal. The dielectric constants of the two substances are very different throughout the board, as depicted in the graph. There are several prominent peaks in the refractive index of SrPtO_3 at 2, 4, 8, 9, and 13 eV, including 2.5, 2.4, 1.2, and 1. The maximum values for the refractive index of SrPtO_3 are 2.4 and 1.2 at 7.8 and 10.4 eV, respectively. An examination of the calculated data revealed that SrPtO_3 had a refractive index greater than that of SrHfO_3 from zero to 4.8 eV and after that, the refractive index of SrHfO_3 had a greater value until 12 eV. In the energy range from 12 eV to 14 eV, once again, SrPtO_3 takes the turn of greater value of the refractive index. It was also experimentally observed that the substances with a high concentration of electrons tend to be more resistant to change. As a result, the refractive index of any compound will increase if its electron density is increased by

any means. The refractive index is a crucial parameter for determining the degree of refraction of light because it is particularly useful in photoelectric applications. In a region where the refractive index is greater than one, is because those photons are slowed as they enter a substance because of the interactions with the crystal's electrons.

3.6 Absorption coefficient

The absorption coefficient is represented by $I(\omega)$, which indicates the absorption of falling photons on a particular compound. Fig. 7 shows the absorption coefficients of both SrHfO_3 and SrPtO_3 crystals. As evident from the figure, the value of absorption coefficient of the compounds SrHfO_3 and SrPtO_3 at zero electron volt is zero. It is also clear from Fig. 7 that it is zero for the crystal SrHfO_3 until 5 eV and remains fluctuating for the crystal SrPtO_3 in the same energy range with a non-zero

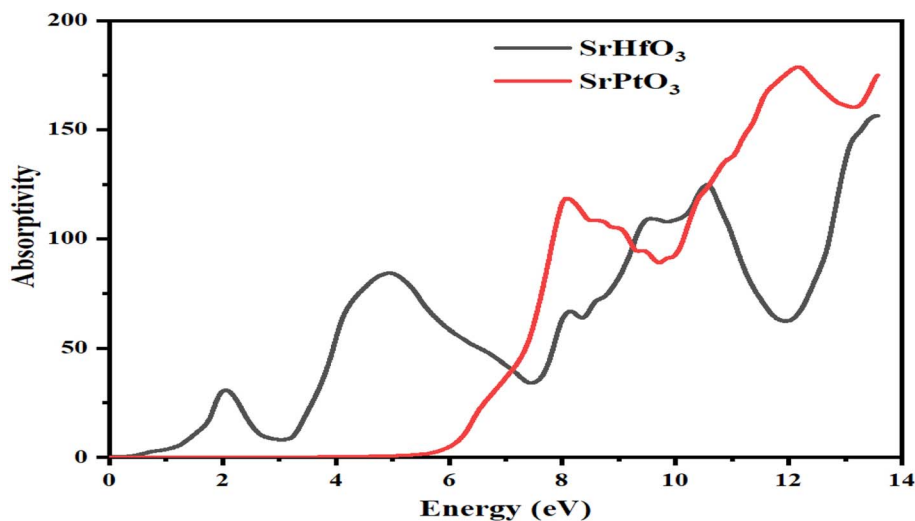


Fig. 7 Calculated absorption coefficient of SrMO_3 crystals ($M = \text{Hf}$ and Pt).

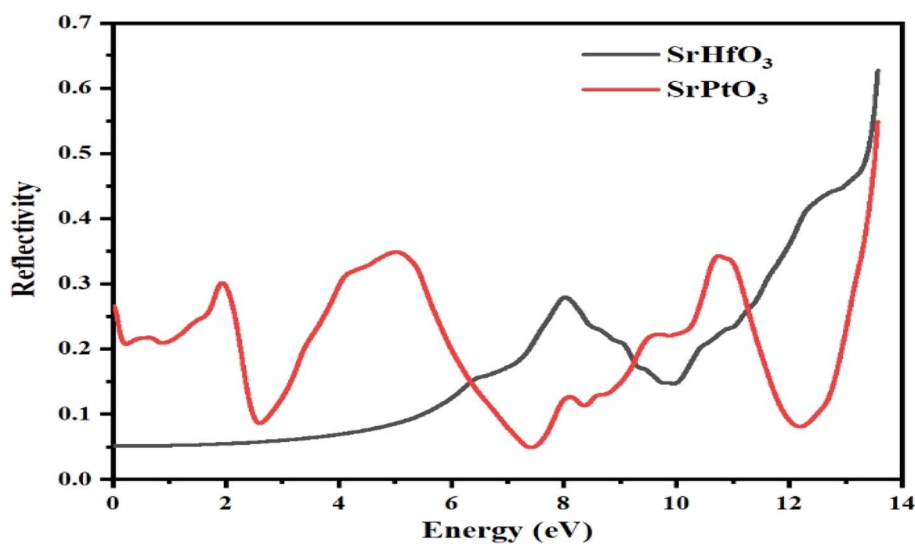


Fig. 8 Reflectivity $R(\omega)$ of incident light from SrMO_3 ($M = \text{Hf}$ and Pt) crystals.

value. After 6.5 eV, if we compare the absorption coefficients of both of our crystals, we can say that the compound SrHfO_3 is a better absorber than SrPtO_3 .

3.7 Reflectivity

Fig. 8 depicts the reflectivity $R(\omega)$ of our selected compounds within the selected energy range. The reflectivity $R(\omega)$ of a particular substance is the degree to which that substance reflects light incident on it. It is calculated from the dielectric function. The threshold values for SrHfO_3 and SrPtO_3 are 0.05 and 0.28, respectively; neither is zero. Both of our selected compounds showed non-zero reflectivity throughout the selected energy range. However, its value for SrHfO_3 was increasing with the increase in energy and reached its maximum value of 0.8 at 14 eV, while the reflectivity of SrPtO_3 fluctuated. For the latter, its minimum value was 0.05 at 7.5 eV

and its maximum value was 0.53 at 14 eV. One can say that an energy of less than 6.5 eV for SrPtO_3 indicates it is a good reflector compared to the SrHfO_3 compound. Fig. 8 summarizes the entire mess, and for effective lens materials, it is recommended to use transparent materials, which in this case is SrHfO_3 .

3.8 Optical conductivity

The optical conductivity is shown in Fig. 9 are also known as the frequency–amplitude relation, describes the connection between the magnitude of an induced electric field and the rate at which it occurs in a given material. This property is used to characterize the way in which a material internally conducts photons. Using the dielectric function, the photon conductivity is displayed in Fig. 9. The graph shows that the photon conductivity of both of our selected crystals is zero at 0 eV, and



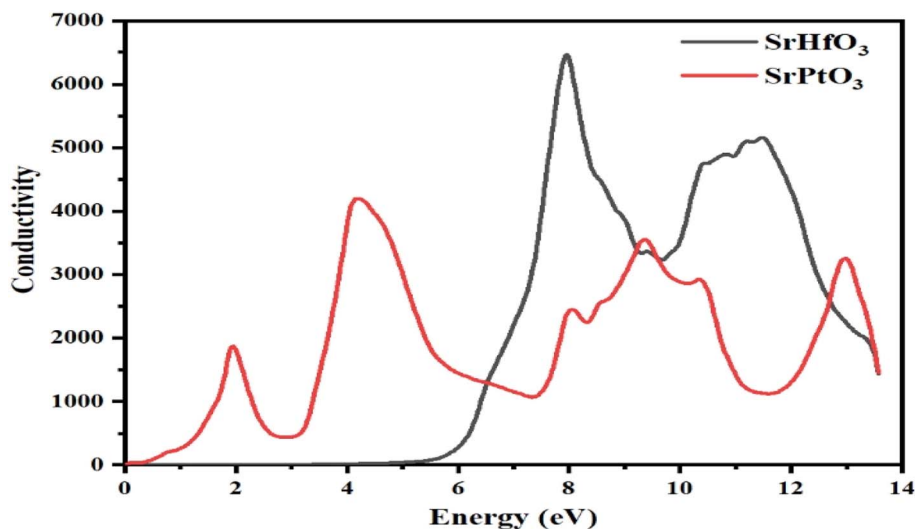


Fig. 9 The optical conduction $\sigma(\omega)$ through SrMO_3 ($M = \text{Hf}$ and Pt) compounds.

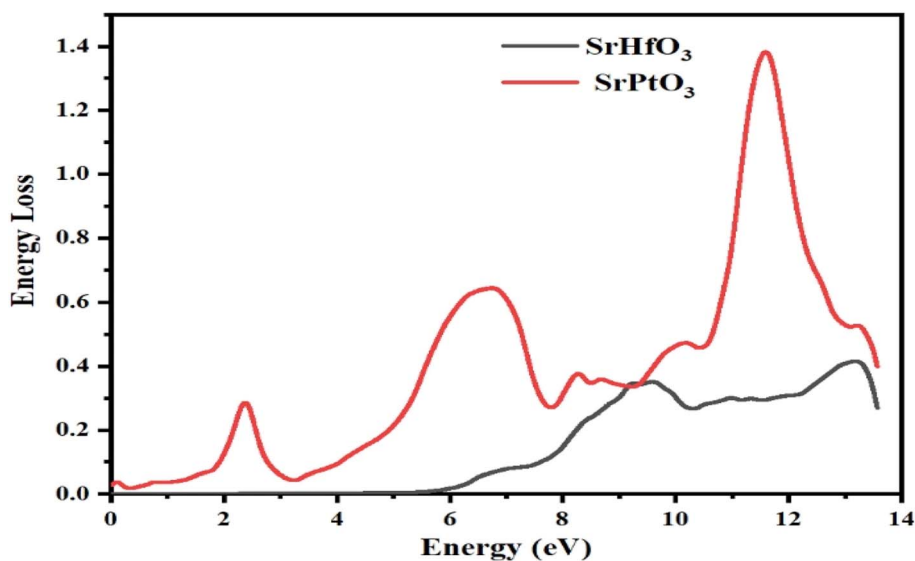


Fig. 10 Representation of the function for the loss of the optical energy $L(\omega)$ of SrMO_3 ($M = \text{Hf}$, Pt).

for SrHfO_3 , it stayed at 0 until 5 eV, and then its value increased to the maximum value of 6800 at 8.3 eV. After 0 eV, the photon conductivity of SrPtO_3 increases and reaches a maximum value of 4200 at 4.2 eV. Before 6.5 eV, SrPtO_3 shows good photon conductivity, and then the second crystal, however, both of our selected compounds have better photon conductivity at higher energy.

3.9 Energy loss function (ELF)

In this section, we discuss the energy an electron loses as it travels through a certain material, which is described by the electron loss function (ELF). The same function can be used to define intra-band, inter-band, and plasmon dependency. Fig. 10 explains the energy loss by an electron as it encounters the selected crystals. According to the aforementioned graph, one

of our target compounds SrHfO_3 has zero ELF values between 0 and roughly 6 eV, while SrPtO_3 ELF remained non-zero up to 14 eV. SrHfO_3 's value increases from 6 eV to a peak of 0.6 at 9 and 13.7 eV. SrPtO_3 ELF value is all the way greater than that of the compound SrHfO_3 with the maxima of 1.4 at 12.3 eV. This means that in comparison at all energies, the molecule SrPtO_3 absorbs more electron energy.

4. Conclusion

We calculated numerous properties, including elastic, structural, optical, and electronic properties of SrMO_3 ($M = \text{Hf}$ and Pt). Because of their extreme harmony and coherence among the results calculated for these compounds, our results are more precise and accurate. Our oxide crystals SrMO_3 ($M = \text{Hf}$

and Pt) are cubic and structurally stable according to the results of the structural parameters. The fundamental elastic constant, anisotropy factor, Poisson ratio, ductility, Cauchy's pressure, shear modulus, Pugh ratio, and Young modulus were among the elastic parameters predicted by the IRelast software. We conclude that both are ductile, anisotropic, elastically stable, and scratch-resistant based on these fundamental elastic features. These findings assure us that these crystals can be utilized in several modern electrical gadgets. Using the TB-mbj approximation, we investigated the fundamental electronic characteristics of the chosen crystals. Additionally, it is determined that SrPtO₃ and SrHfO₃ are conductors and insulators, respectively. The TDOS and PDO details are as follows, SrHfO₃ valence band primarily consists of O-tot, whose energy extends from −4.4 to 0 eV, with the largest peak occurring at 2.8, which corresponds to −1.0 eV. The same states also exhibit peaks at energies of 1.8, 2, and 2.6 at −1.5, −1.2, and −0.6 eV, respectively. The Sr-s state makes up the second-highest contribution in the same band, with a value of 2.0 at 5.8 eV. The Sr-p state makes up a large portion of the conduction band in the same crystal, contributing at its highest levels between 3.2 and 5 eV, with the biggest peak occurring at 3.3 eV. Sr-p and Sr-d states cross the Fermi level and are dominant and enhance the conductivity of the compounds. The valence band of the second compound (SrPtO₃) contains a number of states with varying degrees of participation, such as the Pt-p state, which exhibits peaks at −6.20, −5, −4, and 0.2 eV. Similarly, the Sr-p state contributes with an energy range of −6 eV to 0 eV. Peaks were observed at −6, −5, −3, −2, and 0.20 eV with energies of 0.20, 0.30, 1.40, 1.20, and 1.30, respectively. The Pt-p state contributes energy between 0 and 6.80 eV to the conduction band, with an approximation of 0.1 remaining in this region. The optical properties were studied using the band gap energies of these substances. Our findings suggest that these crystals have low energy absorption and reflectivity of up to 65%, making them suitable for use in high-frequency UV devices.

Conflicts of interest

The authors declare no conflict of interest.

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