Borophene hydride: a stiff 2D material with high thermal conductivity and attractive optical and electronic properties

Bohayra Mortazavi, a Meysam Makaremi, b Masoud Shahrokhi, c Mostafa Raeisi, d Chandra Veer Singh, e, f Timon Rabczuk and Luiz Felipe C. Pereira

1. Introduction

The outstanding properties of graphene1,2 motivated the synthesis of other two-dimensional (2D) materials to discover new physics, with the goal of fabricating advanced materials with enhanced performance. During the last decade, the family of 2D materials has been growing continuously and includes prominent members like hexagonal boron-nitride,3,4 silicene,5,6 germanene,7 stanene,8 transition metal dichalcogenides,9–11 carbon-nitride 2D structures like g-C3N4,12,13 C3N14 and C2N15 and phosphorene.16,17

Two-dimensional (2D) structures of boron atoms, so-called borophene, have recently attracted remarkable attention. In a recent exciting experimental study, a hydrogenated borophene structure was realized. Motivated by this success, we conducted extensive first-principles calculations to explore the mechanical, thermal conduction, electronic and optical responses of borophene hydride. The mechanical response of borophene hydride was found to be anisotropic, with an elastic modulus of 131 N m⁻¹ and a high tensile strength of 19.9 N m⁻¹ along the armchair direction. Notably, it was shown that by applying mechanical loading the metallic electronic character of borophene hydride can be altered to direct bandgap semiconducting, very appealing for application in nanoelectronics. The absorption edge of the imaginary part of the dielectric function was found to occur in the visible range of light for parallel polarization. Finally, it was estimated that this novel 2D structure at room temperature can exhibit high thermal conductivities of 335 W mK⁻¹ and 293 W mK⁻¹ along the zigzag and armchair directions, respectively. Our study confirms that borophene hydride shows an outstanding combination of interesting mechanical, electronic, optical and thermal conduction properties, which are promising for the design of novel nanodevices.

Despite the great experimental success in the fabrication of graphene-like 2D materials, which include a rich, diverse and growing family, only a few 2D materials are monoelemental, such as phosphorene, silicene, germanene and stanene. Notably, these monoelemental 2D materials show out-of-plane buckled structures, different from graphene with its stable flat form. The main reason behind such a structural difference is that these elements present limited ability to vary their bonding nature unlike carbon with its versatile bonding such as sp³, sp² and sp.18 Nevertheless, boron, the neighboring element of carbon placed between metals and non-metals in the periodic table, can also show diverse bonding and as such it is available in a variety of structures ranging from zero-dimensional to three-dimensional.19–20 Interestingly, during the last couple of years, three different 2D boron structures, with both flat21 and buckled22 geometries, have been successfully synthesized through molecular beam epitaxy growth on a silver surface under ultrahigh vacuum conditions. These recent experimental advances in the fabrication of borophene stimulate theoretical studies to explore its applications in various systems such as hydrogen storage, rechargeable metal-ion batteries, superconductors and mechanically robust components.23–44

Research on 2D boron structures is still at a very early stage and there are numerous experimental challenges. Among
these, the greatest obstacle is to develop efficient transferring methods for lifting the borophene sheets from the silver substrate in order to reach isolated nanomembranes. Most recently, experimental realization of 2D hydrogen boride sheets with the empirical formula of B\textsubscript{1}H\textsubscript{1} was successfully achieved by exfoliation and complete ion-exchange between protons and magnesium cations in magnesium diboride.\textsuperscript{45} The fabricated hydrogenated borophene structures were in a multi-layer form and remarkably were not grown on a substrate, which can accordingly serve as a promising sign toward their practical applications. It is worth noting that the existence and stability of such a 2D structure has already been theoretically predicted.\textsuperscript{46} This exciting latest experimental advance\textsuperscript{45} stresses the importance of intensive theoretical studies to establish a comprehensive knowledge about different material properties, very critical for the design of nanodevices. To the best of our knowledge, the mechanical/ failure and thermal conduction properties of the borophene hydride 2D material have not been studied. Moreover, the effects of different loading conditions on the evolution of electronic and optical properties of this recently synthesized hydrogenated borophene via extensive first-principles density functional theory (DFT) calculations.

2. Computational details

The DFT calculations in this study were performed using the Vienna \textit{ab initio} simulation package (VASP).\textsuperscript{47–49} A plane wave basis set with an energy cut-off of 500 eV and the generalized gradient approximation exchange–correlation functional proposed by Perdew–Burke–Ernzerhof\textsuperscript{50} were employed. VMD\textsuperscript{51} and VESTA\textsuperscript{52} packages were also used for the visualization of atomic structures. Periodic boundary conditions were applied along all three Cartesian directions and in order to avoid image–image interactions along the sheets’ normal direction a vacuum layer of 20 Å was employed. To obtain the energy minimized structure, a unit cell with 4 boron and 4 hydrogen atoms was considered. The energy minimized structure was obtained by changing the size of the unit cell and then employing the conjugate gradient method for the geometry optimizations, with termination criteria of 10\textsuperscript{−6} eV and 0.005 eV Å\textsuperscript{−1} for the energy and forces, respectively. We also used 19 × 19 × 1 Monkhorst–Pack\textsuperscript{53} \textit{k}-point mesh size. Similarly to graphene, B\textsubscript{1}H\textsubscript{1} also shows two major orientations, which one can refer to as armchair and zigzag directions in analogy to graphene. We therefore particularly analyzed anisotropy in the mechanical, optical and thermal conductivity responses of single-layer borophene hydride.

Since the dynamical effects such as the temperature are not taken into consideration and periodic boundary conditions were also applied along the planar directions, only a unit-cell modelling\textsuperscript{54} is accurate enough for the evaluation of mechanical properties, and we therefore used a unit-cell consisting of 8 atoms. To evaluate the mechanical properties, we increased the periodic simulation box size along the loading direction in a multistep procedure, every step with a small engineering strain of 0.0005. For the uniaxial loading conditions, upon stretching along the loading direction, the stress along the transverse direction should be negligible. To satisfy this condition, after applying the loading strain, the simulation box size along the transverse direction of loading was changed accordingly in such a way that the transverse stress remained negligible in comparison with that along the loading direction. For the biaxial loading condition, the equal loading strain was applied simultaneously along both planar directions. After applying the changes in the simulation box size, atomic positions were rescaled to avoid any sudden structural changes. We then used the conjugate gradient method for the geometry optimizations, with termination criteria of 10\textsuperscript{−5} eV and 0.005 eV Å\textsuperscript{−1} for the energy and the forces, respectively, in which we used a 15 × 15 × 1 \textit{k}-point mesh size. The final stress values after the termination of the energy minimization process were calculated to obtain stress–strain relations.

The ground state electronic properties were first calculated using the PBE functional. After the calculation of electronic ground states, optical properties such as the imaginary and real parts of the dielectric function were acquired by solving the random phase approximation (RPA)\textsuperscript{55} plus PBE. In the RPA approach, electrons are assumed to respond only to the total electric potential, which is the sum of the external perturbing potential and a screening potential. The external perturbing potential is assumed to oscillate at a single frequency (\textit{ω}), and so through a self-consistent field (SCF)\textsuperscript{56} method. This model yields a dynamic dielectric function denoted \textit{ε}(\textit{ω}). Optical properties are determined by using the dielectric function \textit{ε}(\textit{ω}) = Re ε\textsubscript{\textit{pp}}(\textit{ω}) + iIm ε\textsubscript{\textit{pp}}(\textit{ω}), which is dependent on the electronic structure, and can be obtained from the following relations:\textsuperscript{57,58}

\[
\text{Im } \varepsilon_{\text{pp}}(\omega) = \frac{4\pi^2 e^2}{\Omega} \lim_{q \rightarrow 0} \frac{1}{|q|} \sum_{\epsilon, j} 2w_c \delta(\epsilon_{ij} - \epsilon_{jk} - \omega) \\
\times \left( \langle u_{ik}, e_{pq} | u_{kj} \rangle \langle u_{ik}, e_{pq} | u_{jk} \rangle \right)^* \tag{1}
\]

where \textit{q} is the Bloch vector of the incident wave and \textit{w}_\textit{c} the \textit{k}-point weight. The band indices \textit{i} and \textit{v} are restricted to the conduction and valence band states, respectively. The vectors \textit{e}_\textit{pq} are the unit vectors for the three Cartesian directions and \textit{Ω} is the volume of the unit cell. \textit{U}_{ik} is the cell periodic part of the orbitals at the \textit{k}-point \textit{k}. The real part Re ε\textsubscript{\textit{pp}}(\textit{ω}) can be evaluated from Im ε\textsubscript{\textit{pp}}(\textit{ω}) using the Kramers–Kronig transformation:

\[
\text{Re } \varepsilon_{\text{pp}}(\omega) = 1 + \frac{2}{\pi} \int_0^\infty \frac{\text{Im } \varepsilon_{\text{pp}}(\omega')}{(\omega')^2 - \omega^2 + i\eta} \text{d}\omega' \tag{2}
\]

Local field effects, which correspond to changes in the cell periodic part of the potential, were included in the random phase approximation.

The lattice thermal conductivity of single-layer B\textsubscript{1}H\textsubscript{1} borophene hydride was studied using the ShengBTE\textsuperscript{59} package,
which implements a fully iterative solution to the Boltzmann transport equation. Second order (harmonic) and third-order (anharmonic) interatomic force constants were calculated from first principles calculations based on the DFT. A fully iterative solution, such as the one implemented in ShengBTE, is expected to yield more accurate results than solutions based on single mode relaxation time approximation (SMRTA). Phonon frequencies and the harmonic interatomic force constants were obtained based on the density functional perturbation theory (DFPT) results using the PHONOPY code for a $3 \times 5 \times 1$ super-cell with a $5 \times 5 \times 1$ $k$-point grid. The third-order anharmonic force constants were calculated using the finite displacement approach for the same super-cell and $k$-point mesh size. For the anharmonic force constants, we include the interactions with the tenth nearest-neighbour atoms. The convergence of the thermal conductivity with respect to $q$-points in the full Brillouin zone sampling was confirmed and in order to report the final thermal conduction properties we used a $50 \times 50 \times 1$ $q$-mesh. We also calculated the Born effective charges and dielectric constants using the DFPT method and they were considered in the dynamic matrix as a correction to take into account the long-range electrostatic interactions.

3. Results and discussion

The atomic structure of energy minimized borophene hydride is illustrated in Fig. 1, which shows a graphene-like honeycomb lattice made from boron atoms functionalized by hydrogen atoms. In this structure, the layer of pure boron atoms is sandwiched between two layers of hydrogen atoms, which are decorated on both sides of the B–B bonds. The unit-cell lattice constants were found to be 5.290 Å and 3.018 Å, which match closely with the values 5.299 Å and 2.988 Å reported in the original theoretical work by jiao et al. In Fig. 1, we also plotted the electron localization function (ELF) which takes a value between 0 and 1, where ELF = 1 corresponds to perfect localization. In the lattice of B$_2$H$_4$, two different B–B bonds exist. For the B–B bonds without the hydrogen bridging, the electron localization occurs at the center of B–B bonds, which is characteristic of covalent bonding (Fig. 1, top view). However, for the other B–B bonds bridged by the hydrogen atoms the electrons are localized around the H atoms (Fig. 1, side view). For the energy minimized structure, the bond lengths of B–B bonds without and with hydrogen bridging were found to be 1.720 Å and 1.820 Å, respectively, and the B–H bond length was found to be 1.328 Å.

In Fig. 2a, DFT predictions for the biaxial and uniaxial stress–strain responses of borophene hydride, elongated along the armchair and zigzag directions, are plotted. In all cases, stress–strain responses present an initial linear relationship, which is followed by a nonlinear trend up to the ultimate tensile strength point, a point at which the material illustrates its maximum load bearing. The slope of the first initial linear section of the uniaxial stress–strain response is equal to the elastic modulus. In this work we therefore fitted a line to the uniaxial stress–strain values for strain levels below 0.02 to report the elastic modulus. Based on our modelling results, the elastic moduli of borophene hydride along the armchair and zigzag directions were found to be 131 N m$^{-1}$ and 99 N m$^{-1}$, respectively. For these initial strain levels within the elastic limit, the strain along the traverse direction ($s_t$) with respect to the loading strain ($s_l$) is acceptably constant and can be used to obtain Poisson’s ratio using $-s_t/s_l$. This way, Poisson’s ratio along the armchair and zigzag directions was predicted to be 0.25 and 0.19, respectively. The tensile strength along the armchair and zigzag directions was found to be 19.9 N m$^{-1}$ and 17.8 N m$^{-1}$, respectively. Our results shown in Fig. 2a confirm that both the linear and non-linear parts of the uniaxial stress–strain curves are distinctly different, which suggests anisotropic mechanical response of B$_2$H$_4$ nanosheets.

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Fig. 1 Top and side views of the atomic configuration in single-layer B$_2$H$_4$. The box shown in red shows the B$_2$H$_4$ unit-cell which was used to evaluate the mechanical, electronic and optical properties. We studied the properties along the armchair and zigzag directions as shown. The contours illustrate the electron localization function (ELF) plotted on the two sections.

Fig. 2 (a) Uniaxial and biaxial stress–engineering strain responses of single-layer and free-standing borophene hydride. The change in bond length for the uniaxial loading along the (b) armchair and (c) zigzag directions.
in which, along the armchair and zigzag directions, the structure shows higher rigidity and stretchability, respectively.

As is clear, the higher elastic modulus and tensile strength along the armchair direction can be partly attributed to the stiffening effect of B–H bonds that are exactly oriented along the armchair direction. To understand better the underlying mechanism that results in anisotropic tensile response of borophene hydride, we analyzed the evolution of bond lengths. Fig. 2b and c compares the changes in bond length as a function of strain for the uniaxial loading along the armchair and zigzag directions, respectively. In this system, there are three different bonds, B–B covalent bonds (oriented along the zigzag direction), B–B bonds along the armchair direction (which includes hydrogen bridging) and B–H bonds. For the stretching along the armchair direction, two bonds are exactly along the loading direction and are directly involved in load bearing such that by increasing the engineering strain level these bonds increase substantially, which results in higher stress values and elasticity. In this case, the bond oriented along the transverse direction of loading also increases. On the other side, for uniaxial stretching along the zigzag direction, only the covalent B–B bonds are almost oriented along the loading direction and the other bonds in the system are exactly along the transverse direction of loading. In this case, according to the results illustrated in Fig. 2c, by increasing the strain level the B–B covalent bond keeps increasing and at the same time the other B–B bond first only slightly stretches and at higher strains of ~0.15 starts to contract. In this case, notably the B–H bond length changes remain below ~2%, which further confirms that this bond plays no major role in load bearing along the zigzag direction. Interestingly, the covalent B–B bonds show high stretchability as compared with the other bonds in the B$_2$H$_4$ structure, which allows the structure to endure at higher strain levels when stretching along the zigzag direction.

The results shown in Fig. 2a for the uniaxial strain along the armchair direction reveal that the structure exhibits an unusual yield point at strain levels around 0.19 in which a slight drop in stress values is observable. On further increasing the strain levels, the stress values start to increase again to reach the maximum tensile strength point at strain levels around 0.24. Our bond analysis depicted in Fig. 2b clearly confirms that this yield point in the stress–strain response is basically due to the bond contractions. If one compares the bond lengths at this initial yield point and tensile strength point, very close values are observable. Our structural analysis reveals that at this yield point the structure contracts considerably along the sheet transverse direction, which helps the material to flow more easily along the loading direction such that it relieves the bond stresses by bond contraction. In Fig. 2a, the stress values along the armchair and zigzag directions for the biaxial loading are also plotted, which similarly show higher stress values along the armchair direction.

Next we shifted our attention to investigating the electronic and optical properties of monolayer borophene hydride. In this case, we particularly analyzed the possibility of tuning of the electronic and optical properties by uniaxial or biaxial loading. First, in order to probe the electronic properties, the band structure along the high symmetry directions and the total electronic density of states (EDOS) are calculated using the PBE functional. Fig. 3a illustrates the band structure and EDOS of strain-free borophene hydride. As shown in this figure, this structure yields metallic character and moreover our predicted band structure matches excellently with the original theoretical work.46 Fig. 3b–d, respectively, illustrate the evolution of band structures and the corresponding EDOS of borophene hydride under biaxial and uniaxial loading along the armchair and zigzag directions, with an engineering tensile strain of 0.2. Based on our results shown in Fig. 3 for this monolayer, metal–semiconductor phase transition occurs when biaxial or uniaxial tensile loading along the zigzag is applied. In contrast, the metallic electronic character was completely preserved for the sample under uniaxial tensile loading along the armchair. Interestingly, the formed band-gap is larger for the uniaxial tensile loading along the zigzag direction as compared with the biaxial loading. As is shown, for the biaxial and uniaxial loading along the zigzag direction, borophene hydride monolayer yields direct band-gap semiconductor character in which the valence band maximum (VBM) and the conduction band minimum (CBM) lie on the Γ-point. According to our electronic structure results based on the PBE functional, for the samples under a tensile strain of 0.2 for the biaxial straining and uniaxial loading along the zigzag direction the band-gap is predicted to be 0.12 eV and 0.38 eV, respectively.

Since the PBE functional underestimates the band-gap values, we also computed the EDOS using the HSE06 62 hybrid functional with a 10 × 10 × 1 k-point mesh size. The band-gap values predicted by the PBE and HSE06 methods for strained monolayers are reported in Table 1. Because of the fact that the HSE06 method provides more accurate predictions for the band-gap, 63 in Fig. 4 we specifically analyze the evolution of band-gap on the basis of the HSE06 results. As is clear, as compared with the uniaxial loading along the zigzag direction, the band-gap opening occurs in a more uniform pattern for the biaxial loading. This observation highlights that biaxial straining can be considered as a more promising route for the engineering of electronic band-gap in borophene hydride. It is worth reminding that for the flat and buckled pristine borophene nanomembranes, mechanical loading could not lead to band-gap opening.14 Presenting a small band-gap semiconductor electronic character is a very promising feature for the application of this novel 2D structure in nanoelectronics.

We next analyzed the optical response of this novel 2D material. The imaginary and real parts of the dielectric function of strain-free and strained monolayers for the parallel and perpendicular polarized directions (E∥x, E∥y and E∥z) are illustrated in Fig. 5. In this case for the strained systems, we only considered the structure with a large strain of 0.2 as it shows more considerable effects on the electronic response. For a strain-free structure, the absorption edge of Im ε$_\omega$(ω) is at 1.90, 2.15 and 4.60 eV for E∥x, E∥y and E∥z, respectively, which are in the visible range of light for parallel polarization. These
results indicate that these 2D systems can absorb visible light. Moreover, there are small peaks in the low frequency regime for the parallel polarizations, which is completely missing in the perpendicular one. The value of the static dielectric constant (the real part of the dielectric constant at zero energy) is 1.96, 1.72 and 1.62 for $E_k^x$, $E_k^y$ and $E_k^z$, respectively. Since our previous electronic calculations indicated that the band-gap of this system can be increased by exerting biaxial or uniaxial loading along the zigzag direction, we also calculated the Im $\varepsilon$ and Re $\varepsilon$ for the strained structures. In Fig. 5, the anisotropic response for parallel and perpendicular polarized directions to the planes can be easily observed for all the strained structures. It can be seen that for parallel polarization, by applying uniaxial loading along the zigzag direction the adsorption edge of Im $\varepsilon$ in the low-frequency regime shifts to higher energies, which is known as a blue shift. On the other side, a shift to lower energies (red shift) is observable by applying biaxial strain or uniaxial loading along the armchair. In addition, by applying biaxial or uniaxial loading along the zigzag the value of the static dielectric constant in all polarizations increases.

### Table 1

| Structure               | Band-gap | Re $\varepsilon$ (0) |
|-------------------------|----------|----------------------|
|                         | PBE      | HSE06                |
| Strain = 0.0            | Metallic | Metallic             |
| Strain = 0.1, biaxial   | 0.04     | 0.82                 |
| Strain = 0.2, biaxial   | 0.12     | 1.26                 |
| Strain = 0.1, uniaxial armchair | Metallic | Metallic            |
| Strain = 0.2, uniaxial armchair | Metallic | Metallic            |
| Strain = 0.1, uniaxial zigzag | Metallic | Metallic            |
| Strain = 0.2, uniaxial zigzag | 0.38     | 1.6                  |

Fig. 3 Band structure and total EDOS of unstrained and strained single-layer borophene hydride predicted by the PBE functional. For the strained structures, the applied engineering strain is equal to 0.2. The Fermi energy is aligned to zero.

Fig. 4 Electronic band-gap of single-layer borophene hydride as a function of the applied engineering strain as predicted by the HSE06 functional.
while for uniaxial loading along the armchair it increases along $E_x$ and $E_z$ and decreases along $E_y$. Table 1 also summarizes the value of the static dielectric constant for all the explored cases.

We finally analyze the lattice (phononic) thermal transport of borophene hydride. The phonon dispersion curve of borophene hydride is presented in Fig. 6, which shows excellent agreement with the earlier investigation. Around the center of the Brillouin zone ($\Gamma$ point) we notice three acoustic modes, two of them with linear dispersion and a third one with parabolic dispersion, characteristic of 2D materials. We also note a gap in the phonon bands, in the interval from approximately 30 to 42 THz, characteristic of the mass difference in the chemical species present, B and H. The vibrational density of states is also presented in the figure, projected on B atoms and H atoms, respectively. In the lower frequency range, i.e. below the phonon gap, the phonon density of states (PDOS) is dominated by B atoms, whereas in the higher frequency range (above the gap) it is dominated by H atoms. This behaviour is due to the mass difference, since hydrogen atoms have lower mass and therefore tend to vibrate with higher frequencies.

Our results also indicate no sign of imaginary (negative) frequencies in the phonon dispersion, which accordingly confirms the dynamical stability of this attractive 2D material.

Next, we consider the phonon group velocities for all vibrational modes as depicted in Fig. 7. In this case, we used a $q$-point path identical to the one employed in the calculation of phonon dispersion shown in Fig. 6. Such a presentation can be very illustrative to observe the group velocity associated with the acoustic modes. This way, inside the region of the phonon gap, the group velocity is undefined. Based on our results shown in Fig. 7, the group velocities of the acoustic modes achieve values in excess of $10^4$ m s$^{-1}$, which indicates a probable high thermal conductivity and long effective phonon mean free paths. We calculated the lattice thermal conductivity along two in-plane directions, armchair and zigzag, in analogy with graphene, in the temperature range from 200 K to 800 K and the results are illustrated in Fig. 8. In agreement with a previous study, the van der Waals diameter of the boron...
atom (4.16 Å) was taken as the thickness of single-layer borophene. We note an anisotropy in the conductivity, which is particularly pronounced in the lower temperature regime. Over the temperature range considered the conductivity along the zigzag direction is larger than the one along the armchair direction. Nonetheless, the thermal conductivity along both directions is inversely proportional to temperature, $T$, as expected when phonon–phonon scattering dominates over phonon-defect and phonon-boundary scattering. Interestingly, the observed anisotropy in the thermal conductivity is the inverse of that for elastic modulus. The origin of anisotropy in both load transfer and phonon transport can be partially attributed to the B–H bonds, which yield contrasting effects: a stiffening effect for load transfer and a scattering effect for phonon transport. Nevertheless, based on our first-principles results single-layer borophene hydride at room temperature can exhibit a high thermal conductivity of 335 W mK$^{-1}$ and 293 W mK$^{-1}$ along the zigzag and armchair directions, respectively. These values are by around 50% higher than the earlier prediction for $Pmmn$ borophene structure which is reasonable since the $Pmmn$ borophene shows a buckled atomic structure.

We analyze the dependence of the cumulative lattice thermal conductivity with respect to the maximum phonon mean free path in the samples, which corresponds to the system length along the heat transport direction. From the behaviour of the cumulative conductivity with the system length we can estimate an effective phonon mean free path along each direction in a range of temperatures, using the following expression:67,68

$$\kappa(L) = \kappa_{\infty}/[1 + (\Lambda_{\text{eff}}/L)^{\alpha}]$$  \hspace{1cm} (3)

Here, $\Lambda_{\text{eff}}$ is the effective phonon mean free path, $L$ is the length and $\kappa_{\infty}$ is the length independent thermal conductivity.

In the above equation we introduced an extra parameter, $\alpha$, temperature-dependent exponent "$\alpha$", which modifies the influence of the system length on the calculated conductivity for each temperature. In Fig. 9 we show the cumulative lattice thermal conductivity as a function of maximum phonon mean free path for both the studied directions at 300 K, 500 K and 700 K. The data points are calculated by using ShengBTE, while the continuous lines are fits based on the equation above. With the inclusion of the extra exponent "$\alpha$", we see an agreement of almost four decades between the calculation data and the fit. Note that previous studies have always considered $\alpha = 1$, but in our case it decreases with temperature from 1.33 to 1.25 for the armchair direction and from 1.50 to 1.40 along the zigzag direction. Nonetheless, it is interesting to note that $\alpha$ is only weakly dependent on the temperature.

Finally, we investigate the dependence of $\Lambda_{\text{eff}}$ on temperature. In contrast with the lattice thermal conductivity, there is no clear prediction of the dependence of the effective phonon mean free path on temperature. Adjusting different functional forms to the data points from 200 K to 800 K we found that $\Lambda_{\text{eff}} \sim T^{-2}$, as shown in Fig. 9, inset. To the best of our knowledge this is the first time that this kind of behaviour has been observed. Furthermore, the behaviour does not depend on the heat current direction, just as in the case of the temperature dependence of lattice thermal conductivity.

4. Conclusion

Most recently, a hydrogenated borophene structure with the empirical formula $\text{B}_7\text{H}_4$ was realized. In this work we elaborately explored the mechanical response, thermal conductivity, electronic and optical responses of borophene hydride using first-principles density functional theory calculations. The
mechanical properties of borophene hydride were found to be anisotropic in which the elastic moduli along the armchair and zigzag directions were predicted to be 131 N m$^{-1}$ and 99 N m$^{-1}$, respectively. It was also shown that borophene hydride can yield remarkable tensile strengths of 19.9 N m$^{-1}$ and 17.8 N m$^{-1}$ along the armchair and zigzag directions, respectively. As an interesting finding, it was shown that by applying biaxial tensile strains or uniaxial tensile loading along the zigzag direction the metallic electronic character of borophene hydride can be changed to direct band-gap semiconducting character, which is a highly desirable factor for application in nanoelectronics. Our results therefore confirm that tensile strains can be considered as a promising route for the engineering of the electronics properties of borophene hydride. For the optical properties of this novel 2D material, we found that the absorption edge of the imaginary part of the dielectric function occurs in the visible range of light for parallel polarization. Based on our first-principles results, it was predicted that single-layer borophene hydride at room temperature can exhibit high thermal conductivities of 335 W mK$^{-1}$ and 293 W mK$^{-1}$ along the zigzag and armchair directions, respectively. The high thermal conductivity of borophene hydride is very appealing for application in thermal management systems and enhancing the thermal conductivity of polymeric materials. Based on our results the anisotropy in the thermal conductivity is the inverse of that observed for the elastic response. It was concluded that the B–H bonds can partially act as the origin of anisotropy in both mechanical and thermal conduction properties; they simultaneously enhance the load bearing of the structure and increase the phonon scattering rate. This investigation provides comprehensive information concerning the mechanical, electronic, optical and thermal conduction properties of borophene hydride such that it can be useful for future theoretical and experimental studies.

Conflicts of interest

The authors declare no conflict of interest.

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