An efficient phonovoltaic (pV) material requires a highly energetic optical phonon \( (E_{p,O} \gg k_B T) \) with linewidth dominated by the electron-phonon \((e-p)\) coupling and resonant with its electronic band gap \( (\Delta E_{e,g}) \), as discussed in Paper I [C. Melnick and M. Kaviany, Phys. Rev. B 93, 094302 (2016)]. No current material combines these properties. While graphite (graphene) has the former two, it lacks a band gap. Opening and tuning the band gap in graphite is challenging due to the stability of the Dirac point, e.g., under a uniaxial strain \(<0.25\). We tune its band gap through partial hydrogenation using \textit{ab initio} calculations and find a stable graphane structure with \( \Delta E_{e,g} \approx E_{p,O} \approx 200 \text{ meV}, C_{128}H_{128,24} \). We calculate the \( e-p \) coupling in tuned \( C_{128}H_{128,24} \) and graphene and show that the transition from \( \pi-\pi^* \) (graphene) to \( \sigma-\sigma^* \) (graphane) bands suppresses the electron-phonon coupling, such that optical phonons in \( C_{128}H_{128,24} \) primarily downconvert, and it does not achieve a high figure of merit \( (Z_{pV} < 0.1) \). \textit{Ab initio} phonon-phonon couplings are calculated for graphane and graphene to support this result. Overall, we develop a material with \( E_{p,O} \approx \Delta E_{e,g} \approx k_B T \) and a method for tuning and evaluating pV materials.

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I. INTRODUCTION

Paper I [1] proposes the phonovoltaic (pV) cell shown in Fig. 1, which harvests nonequilibrium (hot) optical phonons across a band gap resonant with or smaller than the optical phonon energy. The degree of local nonequilibrium \( (\eta_C) \), the fraction of hot optical phonons which generate an electron rather than downconvert \( (\gamma^{\star}_{e-p}) \), and the fraction of phonon energy \( (E_{p,O}) \) preserved by the band gap \( (\Delta E_{e,g}) \) limit the pV efficiency. Paper I proposes that the figure of merit \( (Z_{pV}) \) and efficiency \( \eta_{pV} \) are

\[
Z_{pV} = \gamma^{\star}_{e-p} \frac{\Delta E_{e,g}}{E_{p,O}} \lesssim 1, \quad \gamma^{\star}_{e-p} = \frac{\gamma_{e-p}}{\gamma_{e-p} + \gamma_{p-p}},
\]

\[\eta_{pV} \approx \eta_C Z_{pV} \left[1 - 0.75 \exp \left(-\frac{\eta_C \Delta E_{e,g}}{10 k_B T} \right) \right],\]

where \( \eta_C \) is the Carnot limit given by the local electron-phonon \((e-p)\) nonequilibrium, i.e., \( \eta_C = 1 - T_c / T_{p,O} \), and \( \gamma_{e-p} \) and \( \gamma_{p-p} \) are the \( e-p \) and phonon-phonon \((p-p)\) interaction rates, \( T_c \) is the cold, contact temperature, \( T_{p,O} \) is the hot optical phonon temperature, and \( k_B \) is the Boltzmann constant. Thus, the pV requires \( E_{p,O} > \Delta E_{e,g} \approx k_B T \) and that the optical phonon mode has a linewidth dominated by the \( e-p \) coupling (high \( \gamma^{\star}_{e-p} \)).

Few semiconductors exist with an optical phonon mode more energetic than the band gap, and even fewer of these exhibit an optical phonon significantly more energetic than the thermal energy \( (k_B T_c) \) at room temperature. Energetic phonons require strong bonds between light atoms; however, strong bonding localizes electrons and induces a large band gap. Indeed, in an isotropic semiconductor strong bonding typically produces a band gap more than an order of magnitude larger than the optical phonon energy. For example, the strong \( \sigma \) bonds in diamond produce extremely energetic optical phonon modes (164 meV) and a band gap exceeding 5 eV. Conversely, the weak bonds in semimetallic \( Hg_{1-x}Cd_xTe \) (MCT) produce a narrow band gap \( (0 \lesssim \Delta E_{e,g} \lesssim 1 \text{ eV}) \), but no optical phonon more energetic than 45 meV [2]).

Anisotropic structures enable deviation from this trend through an alternate bonding mechanism, e.g., \( \pi \) rather than \( \sigma \) bonding. For example, graphene has a unique combination of electronic and phononic properties due to its \( sp^2 \) hybridized structure, wherein \( \pi \) and \( \pi^* \) bands form near the Fermi surface. Due to the symmetry of graphene, these bands become degenerate at the Dirac point (K) [3,4].

While the active optical phonon modes in graphene have a linewidth dominated by the \( e-p \) coupling \( (\gamma^{\star}_{e-p} > 0.9) \) [5], the lack of a band gap ensures \( Z_{pV} = 0 \). If tuning the band gap of graphite to its optical phonon energy preserved the \( e-p \) and \( p-p \) coupling properties, then the pV figure of merit would exceed 0.8, and its efficiency would greatly exceed that of a thermoelectric generator.

The alternate group IV two-dimensional (2D) crystal silicene has similarly favorable properties [6], and the band gap in silicene has been successfully tuned [7]. However, the energy of the optical phonon modes is limited in crystals comprised of second row and higher elements (due to their higher mass and weaker bonding). Thus, the optical phonon modes in silicene are limited to around 70 meV, and we focus on graphite.

In this study, we tune the band gap of graphite through its partial hydrogenation. The \textit{ab initio} band gap is found for partially hydrogenated graphene cells with variations in the atomic fraction and configuration of hydrogen. A stable structure is found which tunes the band gap to roughly the \( E_{2g} \) graphene phonon mode (200 meV). Additional tuning to the \( E_g \) graphene mode is accomplished through the application of a small, isotropic strain. Then, the phonon and \( e-p \) properties are calculated, and compared with the \( e-p \) and \( p-p \) coupling in graphene and graphene. The transition from \( \pi-\pi^* \) to \( \sigma-\sigma^* \) type bonds is shown to inhibit the \( e-p \) coupling and limit the \( Z_{pV} < 0.1 \). Finally, additional avenues of research are suggested in the search for an efficient pV material.
FIG. 1. (a) The function of a photovoltaic cell comprised of (b) partially hydrogenated graphite. An excited optical phonon population creates electron-hole pairs, and the p-n junction separates them to generate power. The hot optical phonons also downconvert into acoustic phonons and generate entropy. Hydrogenation opens a band gap by transforming the sp\(^2\) hybridized graphene into sp\(^3\) hybridized graphane.

II. TUNING BAND GAP OF GRAPHITE

Opening a band gap in graphene has attracted substantial attention after its initial discovery. Many methods of achieving this goal have been proposed, e.g., functionalization [8], growth on an ordered substrate [9], and the application of a mechanical strain [10–12], electric field [13], or magnetic field [14], or through chemical doping [15]. Of these methods, functionalization is chosen for this study. Relying on a substrate to tune the band gap restricts the pV to a 2D device. Using electric and magnetic fields in a pV prevents its application in, e.g., electronics cooling, where electromagnetic fields disrupt the electronic packages. Chemical doping can substantially shift the Fermi level and preclude its use as a pV material.

That is, despite the initial ab initio results by Ni et al. suggesting that arbitrary uniaxial strains open a band gap in graphene [16], the tight-binding modeling of Pereira et al. showed that strains in excess of 0.2 are required to open a band gap, and they hypothesized that that the ab initio simulations missed the movement of the Dirac point from the K to \(M\) point under strain [10]. This supposition was confirmed by Ni et al., who additionally proposed that under the strain requirements proposed by Pereira et al., the \(\sigma^*\) band becomes the lowest-energy conduction band at \(M\) and may prevent the band gap from opening [17].

Figure 2 presents the ab initio [18] low-energy band structure for variations in the uniaxial strain. It shows that strains exceeding 0.25 are required to open a gap. While the \(\sigma^*\) band does cross over the \(\pi^*\) band to become the first-conduction band, a band gap still opens at \(M\). However, even under strains exceeding 0.3, the band gap remains extremely narrow, in part due to the crossing of \(\sigma^*\) and \(\pi^*\) bands. Such substantial strains are impractical, at best, and fall well beyond the elastic deformation of graphene. Moreover, the application of a tensile strain reduces the energy of the optical phonon modes [19], further limiting strained graphene as a pV material.

In contrast, it has been shown that nonuniaxial strains induce a gauge field which in turn opens a band gap through the quantum Hall effect [11]. However, this approach limits the pV to a 2D device, whereas functionalization is potentially applicable to graphite and enables a 3D cell. Thus, this study focuses on functionalization via the partial hydrogenation of graphene.

III. PARTIALLY HYDROGENATED GRAPHITE

Theoretical investigations of graphene (fully hydrogenated graphene) began before that of graphene [20]. Two stable conformations were discovered, the most stable of which is the chair conformation [Fig. 3(a)], wherein the hydrogen atoms alternate bind above and below carbon atoms in the graphene plane [21]. During hydrogenation, out-of-plane hydrogen atoms transform the local, sp\(^2\) hybridized graphene structure into a sp\(^3\) hybridized graphene structure. The resulting planar sp\(^3\) structure localizes electrons within the \(\sigma\) bond and opens a band gap. Theoretical studies predicted a band gap as high as 3.5–3.7 eV in graphene, depending on the conformation [22]. Although full hydrogen coverage has not been achieved experimentally, experimental and reversible hydrogenation of
graphene has provided evidence for graphane [23] and its phonon properties, which are supported theoretically [24].

As the structural stability of graphane (partially hydrogenated graphene) increases with hydrogen coverage, it is predicted that islands of hydrogenated carbon forms in a particular conformation and that these islands expand as hydrogenation continues [20,22]. An example of such an island is shown in Fig. 3(d). As the hydrogenation progresses, the band gap opens until full coverage is achieved. Thus, the band gap of graphite is tuned by controlling the extent of the hydrogenation.

For the ab initio calculations to follow, the initial configuration chosen for the simulations within the density functional theory (DFT) and the density functional perturbation theory (DFPT) have a substantial impact on the resulting predictions. Thus, constructing a graphane supercell requires care. Indeed, the location of added hydrogen atoms, their conformation, and the number of hydrogen atoms compared to the number of carbon atoms all influence the resulting electronic and phononic properties.

The pV cell requires a non-negligible volume to generate substantial current and power [1]. Thus, a practical pV cell requires the 3D structure of partially hydrogenated graphite rather than the angstrom thickness, 2D structure of graphane. Hydrogenation requires the 3D structure of partially hydrogenated graphite [25]. This study is restricted to the simulation of graphane.

A rectangular graphene supercell of 128 carbon atoms (C128) forms the scaffolding for the simulated graphane cells. Hydrogen atoms are placed onto this scaffold in the chair configuration in fully hydrogenated rings (C128H1×6), as these are expected to be more stable than a partially hydrogenated carbon ring. Structures involving multiple rings (C128H2×6) and larger hydrogenated islands are tested C128H1×N and C128H1×24 is shown in Fig. 4(a). These structures are relaxed within DFT [26], and their formation energies are shown in Fig. 4(a).

While a single hydrogen ring (C128H1×6) is unstable, further and adjacent hydrogenation increases stability (while additional, nonadjacent rings do not). Thus, it remains likely that islands of hydrogen form and then expand outward. These results additionally suggest that the hydrogen prefers to form in even-numbered clumps such that all electrons pair. As expected, stability increases as the internal sp3 structure grows larger than the strained sp2–sp3 interface at the graphene-graphane border.

**IV. ELECTRON PROPERTIES**

The ab initio electronic properties of graphane are presented in Figs. 5 and 6 [27] for a small (a), odd (b), and large (c) number of H atoms. When a small and even number of hydrogen are present, no band gap opens, but the Dirac point moves along the Γ–X line towards X. Only after the hydrogen island occupies a substantial portion of the graphane unit cell does a band gap open (at X), and this band gap exceeds 200 meV, i.e., the most energetic phonon mode in graphite. When an odd number of hydrogen is present, the unpaired electron is trapped at the Fermi surface with a substantial energy gap between this trapping band and the valence and conduction bands.
At low hydrogen concentrations, no band gap opens but the Dirac point moves towards X along the Γ-X high-symmetry line. (b) A trap band forms for an uneven number of hydrogen atoms. (c) At large hydrogen concentrations, however, a narrow band gap opens. Some discontinuities arise from the crossing of the π and σ bands.

These results, in addition to the substantial strains required to open a band gap (see Sec. II), demonstrate the stability of the Dirac point, which has been linked to translation and time invariance (i.e., its symmetry) [28]. This presents a challenge when tuning the band gap to resonate with the optical phonons. However, once sufficient hydrogenation opens a band gap and shifts the topology, arbitrary perturbations tune the band gap.

Indeed, the band gap becomes highly sensitive to the lattice constant (i.e., isotropic strains or thermal expansion) and the addition of additional hydrogen atoms. For example, the band gap in C128H1×24 changes from a 250- to 175-meV band gap under an isotropic strain of −0.02, while the band gap of C128H1×22 is 15 meV larger than C128H1×22. Fine tuning the band gap of graphane through hydrogenation in ab initio simulation is challenging, as large structures (e.g., C128H2×16,2×22) are required to achieve ΔE_{g,φ} ≃ E_{p,0}. Evaluating the phonon and e-p properties in a 600-atom cell is computationally impractical. Thus, this study utilizes small and isotropic strains to fine tune the band gap of C128H1×24 to resonate with the desired optical phonon mode.

V. PHONON PROPERTIES

The phonon Hamiltonian discussed in Paper I [1] is [29,30]

$$\langle \psi \rangle = \langle \psi \rangle_0 + \frac{1}{2!} \sum_{i j x y} \Gamma_{i j x y}^{x y} \epsilon_i \epsilon_j + \frac{1}{3!} \sum_{i j k x y z} \Psi_{i j k x y z}^{x y z} \epsilon_i \epsilon_j \epsilon_k + \ldots,$$

where $\langle \psi_0 \rangle$ is the potential at equilibrium, $\epsilon_i$ is the displacement of atom $i$ from equilibrium in the $x$ direction (Cartesian), and $\Gamma$ and $\Psi$ are the second- (harmonic) and third-order (anharmonic) force constants. The harmonic interactions form the dynamical matrix which determines the phonon frequencies $\omega_{k,\alpha}$ and displacement vector $\epsilon_{k,\alpha}$ for a phonon with polarization $\alpha$ at wave vector $k_p$, where the dynamical matrix is

$$D_{ij}^{xy}(k_p) = \frac{1}{(m_i m_j)^{1/2}} \Gamma_{ij}^{x y} \exp[i k_p \cdot (r_i - r_j)],$$

and $r_i$ and $m_i$ are the position and mass of atom $i$, and its eigenvalues and eigenvectors are $\omega_{k,\alpha}^2$ and $\epsilon_{k,\alpha}$. The amplitude of the resulting atomic displacement for atom $i$ is

$$u_{k,\alpha} = \left( \frac{\hbar}{2m_i \omega_{k,\alpha}} \right)^{1/2} \epsilon_{k,\alpha}.$$

In comparison, the anharmonic forces are responsible for the $p-p$ interactions, e.g., the upconversion of two acoustic phonons into an optical phonon and the downconversion of an optical phonon into two acoustic phonons (hereafter referred to as upconversion and downconversion). These are evaluated for graphene and graphane zone-center phonons in Sec. VI B.

The ab initio dynamical matrices are evaluated within DFPT [31] on a rough mesh of $k_p$ points [32]. These dynamical matrices are Fourier transformed into real space and then interpolated to an arbitrary point in $k_p$ space. The resulting phonon dispersion and density of states for graphene, graphane, and C128H1×24 are shown in Fig. 7.

The active optical phonon modes in graphene are the $E_{2g}$ and $A'_1$ modes, where the $A'_1$ mode is at a Kohn anomaly and couples most strongly to the electronic system. The hydrogenation of graphene substantially changes the phonon properties. Graphane exhibits an extremely energetic optical phonon mode per hydrogen atom, the out-of-plane vibration of hydrogen, as well as the less energetic $E'_{2g}$ (rather than $E_{2g}$) mode. Moreover, the optical modes in graphene tend to have lower group velocities. Graphane exhibits a mix of these modes, some smeared and distorted by the graphene-graphane border.
VI. e-p AND p-p COUPLING

As discussed in Paper I [1], the relaxation of hot optical phonons through the generation of new electrons or acoustic phonons largely determines the performance of a pV cell. The e-p and anharmonic p-p couplings drive these phenomena. This section presents the \textit{ab initio} coupling and kinetics from perturbation theory for both coupling mechanisms.

A. e-p coupling and kinetics

The e-p Hamiltonian is presented in Paper I [1], and a brief description of the resulting e-p matrix element follows. When a phonon with momentum \( k_p \) and polarization \( \alpha \) scatters with an electron of momentum \( k_e \) in band \( i \) (\( |k_e,i\rangle \)), it annihilates that electron and creates an electron in band \( j \) with momentum \( k'_j \) (\( |k'_j,j\rangle \)). The e-p coupling element \( M_{k_i,k'_j,\alpha} \) which follows from this Hamiltonian is

\[
M_{k_i,k'_j,\alpha} = (k'_j,j) \left( \frac{\hbar}{2(m_e \omega_{k'j,\alpha})} \right)^{1/2} \delta_{k'_j,k_{i}\pm k_e},
\]  

where \( \varphi_{KS} \) is the Kohn-Sham potential [31]. The \( \delta \) function conserves momentum during phonon absorption (\( + \)) and emission (\( - \)), such that \( k'_j = k_{i\pm} \pm k_e \).

The interaction rate follows from the Fermi golden rule [29]

\[
\gamma_{e-p}(k_p,\alpha) = \frac{1}{2k_p} \sum_{k_{i,j},\alpha} \frac{2\pi}{\hbar} |M_{k_i,k'_j,\alpha}|^2 \delta(E_e \pm \h\omega_{k'j,\alpha} - E'_{e})
\]

\[
\times \left[ f_e(E_e) - f_e(E_e \pm E'_{e}) \right],
\]

where the \( \delta \) function conserves energy such that \( E'_{e} = E_e \pm \h\omega_{k'j,\alpha}, \) and \( f_e(E_e) \) is the electron occupancy at energy \( E_e \).

Typically, the Fermi-Dirac occupancy applies, i.e.,

\[
f_e = \left[ \exp \left( \frac{E_e - E_F}{k_B T} \right) + 1 \right]^{-1},
\]

where \( E_F \) is the Fermi energy and \( T \) is the temperature. When \( \Delta E_{e,g} \gg k_B T \), the electron occupancy in valence states approaches unity, and the occupancy in conduction states vanishes. This greatly inhibits the e-p interaction rate between two valence (\( f_e \approx f'_e \approx 1 \)) or conduction (\( f_e \approx 0 \)) states. Thus, when \( E_{p,0} > \Delta E_{e,g} \), this ensures that the valence-to-conduction transitions (i.e., generation and recombination events) dominate the valence-to-valence and conduction-to-conduction transitions. Thus, the summation over \( i \) and \( j \) is restricted to generation events, i.e., \( i = v \) and \( j = c \), and the remaining interactions are neglected.

The interaction elements for the \( \Gamma \)-point phonon modes in a C\textsubscript{128}H\textsubscript{1}x24 cell are evaluated within DFPT using QUANTUM ESPRESSO on a \( 6 \times 6 \times 1 \) \( k \)-electron mesh, wherein the matrix elements are calculated as

\[
|M_{k,v,k',\alpha}|^2 = \sum_{i,j,xy} u^{v\alpha}_{i,xy} M_{i,j}^{*} u^{v\alpha}_{j,xy},
\]

where the superscript asterisk denotes the complex conjugate, \( u^{v\alpha}_{i,xy} = (\epsilon\nu)^{v\alpha}_{i,xy} \), and \( M_{i,j} \) is the e-p matrix element associated with a perturbation on atom \( i \) in direction \( x \), i.e.,

\[
M_{ij} = (k_e,v) \frac{\partial \varphi_{KS}}{\partial k_i} |k_e,c\rangle.
\]

These elements, along with the ground-state conduction and valence energy surfaces, are interpolated onto a \( 2000 \times 2000 \times 1 \) \( k \)-electron mesh in the first Brillouin zone (BZ). This fine \( k_e \) mesh enables an accurate calculation of Eq. (6) using a Lagrangian \( \delta \) function with 1 meV of smearing.

Figure 8 shows the e-p coupling between the lowest conduction and highest valence bands in (a) graphene for the \( E_{2g} \) and (b) C\textsubscript{128}H\textsubscript{1}x24 for the \( E_g \) optical phonon modes. After hydrogenation, the interband e-p coupling is substantially reduced near the band edge. Indeed, Fig. 8(c) shows the weak interband coupling in graphene for all optical phonon modes at the band edge (\( X \)), and that the e-p lifetime increases from 0.5 ps in graphene to over 100 ps in graphane. However, the intraband coupling elements shown in Fig. 8(d) are not substantially weakened by the hydrogenation.

The poor overlap between \( \sigma \) and \( \sigma^* \) bands explains these results. Note that near the \( \Gamma \) point in graphene, the \( \sigma \) bands are nearest the Fermi surface, and Fig. 8(a) shows the weak coupling in this region. In comparison, the \( \pi-\pi^* \) band coupling near \( K \) and \( M \) is strong, as these bands overlap significantly.
Hydrogenation opens a band gap by transforming graphene from a $sp^2$ to $sp^3$ hybridized structure. During this transition, the $\sigma$ bands approach the Fermi surface and replace the $\pi$ bands as the lowest-energy bands. At this point, a band gap opens. As the coupling between the $\sigma-\sigma^*$ bands remains weak, so too does the interband coupling in graphene. In contrast, two conduction or valence states overlap completely, such that the intraband coupling remains strong. Thus, the $e-p$ coupling of the $E_{2g}$ mode in graphene is reduced from its value in graphene by approximately the atomic hydrogen fraction, rather than an order of magnitude.

A pV material must achieve stronger $e-p$ coupling in order to succeed as a pV material. As functionalized graphene variants often transition from $sp^2$ to $sp^3$ hybridization, these materials, like graphene, will not achieve a high figure of merit due to the poor overlap between their $\sigma$ and $\sigma^*$ bands. An efficient pV material must achieve stronger $e-p$ coupling, as the anharmonic phonon couplings, in comparison, do not change substantially. This is discussed in the following sections.

B. $p-p$ coupling and kinetics

As discussed in Sec. V, the anharmonic (third-order and higher) interactions are responsible for phonon downconversion. However, fourth-order and higher interactions are typically masked by the third-order coupling [33]. The rate at which a phonon $(k_p, \alpha)$ downconverts into two phonons $(k_p', \alpha')$ and $(k_p'', \alpha'')$ follows from Eq. (2) and the Fermi golden rule, i.e.,

$$\gamma_{p-p}(k_p, \alpha) = \frac{\pi \hbar}{N_k} \left| \frac{1}{16} \sum_{\alpha' \neq \alpha, \alpha''} \left| \psi_{\alpha \alpha'}^{k_p k_p'} \phi_{\alpha''}^{k_p''} \right|^2 \right|^2 \times \delta(\omega_{\alpha p, \alpha' - \omega_{\alpha p, \alpha''} - \omega_{\alpha p, \alpha''}) (f_p' + f_p'' + 1) \right),$$

where $f_p$ is the phonon occupancy, $N_k$ is the number of $k_p$ points used in the integration [34], and the interaction element $\psi_{\alpha \alpha' \alpha''}$ is

$$\psi_{\alpha \alpha' \alpha''} = \sum_{ij} \sum_{k} \sum_{\alpha \beta} \sum_{\alpha} \psi_{ij \alpha' \alpha}^* \psi_{ij \alpha'' \alpha}^* \psi_{ij \alpha \alpha} \psi_{ij \alpha \alpha} \times \exp[i(k_p r_i + k_p' r_j + k_p'' r_k - \delta k_p')].$$

For a $\Gamma$-point phonon, Eq. (10) is simplified, as momentum conservation dictates $k_p = -k_p'$, i.e.,

$$\gamma_{p-p}(\Gamma, \alpha) = \frac{\pi \hbar}{N_k} \left| \frac{1}{16} \sum_{\alpha' \neq \alpha, \alpha''} \left| \psi_{\alpha \alpha'}^{\Gamma k_p} \phi_{\alpha''}^{\Gamma k_p'} \right|^2 \right|^2 \times \delta(\omega_{\alpha p, \alpha' - \omega_{\alpha p, \alpha''} - \omega_{\alpha p, \alpha''}) (f_p' + f_p'' + 1).$$

Restricting the calculation to the lifetime of the zone-center phonon modes greatly reduces the number of third-order coupling elements required to evaluate the downconversion rate and simplifies their calculation. As the third-order force constant calculations are computationally costly, this is extremely important, especially in crystals with more than a few atoms per unit cell.

However, evaluating the third-order force constants in $C_{128H_{124}}$ remains unreasonable even under this restriction. Thus, the $p-p$ lifetimes are evaluated from pure graphene and graphite crystals to estimate the lifetimes of the various targeted optical phonon modes. For both of these structures, the third-order force constants are evaluated within DFPT using the $2n + 1$ formula [35], as implemented within QUANTUM ESPRESSO on an $18 \times 18 \times 1$ $k_p$-point mesh. These dynamical matrix derivatives are Fourier transformed into real space and then interpolated onto a $200 \times 200 \times 1$ mesh of $k_p$ points. Then, the integration in Eq. (12) is calculated using a Lagrangian $\delta$ function with 20-K smearing.

Figure 9(a) shows the the downconversion paths for the graphene $E_{2g}$ and graphene $E_g$ phonon modes, (b) the energy of the phonon modes created, and (c) their lifetimes. Due to the high energy of these modes, the phonon lifetimes begin to change significantly around 500 K. Also shown in Fig. 9(c) is the increased downconversion rates of the zone-center optical phonon modes in graphene, as compared to those in graphene. Indeed, even without the degradation of the electron-phonon coupling, hydrogenation enhances the optical
exhibits significantly enhanced optical phonon downconversion rates. Phonon modes produced, and (c) the scattering rate as a function to \( \sigma \) is of graphene.

The results are shown in Fig. 10. Note that the \( \dot{\gamma}_{e-p} \) coupling must dominate the \( \gamma_{e-p} \) coupling. No current material exists with this combination of properties.

FIG. 10. The fraction of hot optical phonons which generate electrons as they relax (\( \dot{\gamma}_{e,p}^* \)) and \( \nu \), \( \nu \) for graphene and tuned C128H1\( _{24} \) cells. Also shown is the graphene figure of merit if the \( e-p \) and \( p-p \) couplings remained constant as the band gap opens. This figure of merit is substantially larger than that of C128H1\( _{24} \), as the interaction between the \( \sigma \) and \( \sigma^* \) bands is much weaker than that between the \( \pi \) and \( \pi^* \) bands in graphene.

For example, hydrogenation, fluorination, and other functionalization methods which create a \( sp^3 \) structure rely on this transition to open a band gap. If a similar decrease in the interband \( e-p \) coupling occurs across all \( sp^3 \) hybridized structures, as suspected, few options remain [36]. Thus, it is crucial to confirm this hypothesis. However, other unusual and anisotropic materials may combine the material properties crucial to \( \nu \) operation: a highly energetic optical phonon mode resonant with the band gap and with a lifetime dominated by the \( e-p \) coupling.

C. Figure of merit

Using the results from Secs. VI A and VI B, \( \dot{\gamma}_{e,p}^* \) is calculated for the zone-center modes in C128H1\( _{24} \), where \( \dot{\gamma}_{e,p}^* \) is

\[
\dot{\gamma}_{e,p}^*(\Gamma, \alpha) = \frac{\dot{\gamma}_{e,p}(\Gamma, \alpha)}{\dot{\gamma}_{e,p}(\Gamma, \alpha) + \dot{\gamma}_{p,p}(\Gamma, \alpha)}.
\]

The results are shown in Fig. 10. Note that the \( \dot{\gamma}_{p,p}(\Gamma, \alpha) \) used here are from the graphene and graphene calculations.

As shown in Fig. 10, the weak \( e-p \) coupling in partially hydrogenated graphite prevents it from achieving a large \( \dot{\gamma}_{e,p}^* \) and \( Z_{pV} \). This weak coupling is due to the small overlap between its valence (\( \sigma \)) and conduction (\( \sigma^* \)) bands. No similar effect reduces the \( p-p \) interaction strength, such that the \( \dot{\gamma}_{e,p}^* \) achieved by C128H1\( _{24} \) is less than 0.1. As the band gap is tuned to resonate with the optical phonon modes, \( Z_{pV} = \dot{\gamma}_{e,p}^* < 0.1 \).

In graphene, the overlap of \( \pi \) and \( \pi^* \) bands enables a large \( \dot{\gamma}_{e,p}^* > 0.9 \), but the gapless electronic structure ensures \( Z_{pV} = 0 \). If the \( e-p \) and \( p-p \) couplings were to remain unchanged as the band gap opened, however, graphene would achieve \( Z_{pV} > 0.8 \). In such a \( \nu \) cell, the efficiency significantly exceeds that of a TE, even at room temperature and under negligible \( e-p \) nonequilibrium. However, it is challenging to tune the band gap of graphene without transitioning from \( \pi-\pi^* \) to \( \sigma-\sigma^* \) low-energy bands.

FIG. 9. (a) Active downconversion pathways, (b) energy of the phonon modes produced, and (c) the scattering rate as a function of temperature for graphene \( E_{2g} \) and graphene \( E_g \) modes. Graphene exhibits significantly enhanced optical phonon downconversion rates.

PHONOVOIJCIA. II. TUNING BAND GAP TO OPTICAL . . . PHYSICAL REVIEW B 93, 125203 (2016)

VII. CONCLUSIONS

In Paper I [1], a \( \nu \) cell is proposed that harvests a hot population of optical phonons resonant with the band gap and much more energetic than \( k_BT \). For efficient harvest, the \( e-p \) coupling must dominate the \( p-p \) coupling. No current material exists with this combination of properties.

Graphite (graphene), which has energetic optical phonon modes that couple strongly to the electron system, is proposed as a candidate, provided that its band gap is tuned to the optical phonon energy. Uniaxially strained graphene is confirmed to require extreme strain in order to open a band gap via a series of \( ab\) initio calculations, wherein the movement of the Dirac point along the \( K-M \) symmetry line must be carefully tracked. However, the band gap in graphite may be tuned to its optical phonon modes through its partial hydrogenation.

Here, this tuning is accomplished through the systematic \( ab\) initio simulation of partially hydrogenated graphite layers with variations in the atomic fraction and placement of hydrogen. The C128H1\( _{24} \) structure is shown to have a band gap around 250 meV and the arbitrary and isotropic strains enable the fine tuning to the desired optical phonon mode, e.g., the graphene \( E_{2g} \) (200 meV) and the graphene \( E_g \) (174 meV) modes.
However, the $e$-$p$ coupling between the $\sigma$-$\sigma^*$ bands in graphene is significantly weaker than the $\pi$-$\pi^*$ band coupling in graphene. In comparison, the $p$-$p$ interactions in graphene are similar to or stronger than those in graphene, as shown through the 
initio evaluation of graphene and graphene $\Gamma$-point phonon lifetimes. Thus, the $pV$ figure of merit for partially hydrogenated graphite is limited to below 0.1.

The transition from the $\pi$- to $\sigma$-type bands near the Fermi level as a band gap opens in graphene due to, e.g., the application of a uniaxial strain or functionalization is proposed to limit alternate graphene derivatives. However, the reduced-dimensioned graphene structures, e.g., quantum dots and nanoribbons, preserve the $\pi$ and $\pi^*$ bands and may overcome this trend. Additional study remains for other functionalized graphene structures, e.g., fluorographane, and a theoretical explanation of the weak $\sigma$-$\sigma^*$ $e$-$p$ coupling is required, and finding alternate $pV$ materials is crucial.

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[18] The strained graphene unit cells are prepared in QUANTUM ESPRESSO [37–39] by applying a uniaxial strain on graphene in the $x$ direction (zigzag) and then relaxing the cell in the $y$ direction. Then, a self-consistent DFT simulation on a $120 \times 120 \times 1k_y$ mesh provides the charge density. The band structure is then found along the $\Gamma$-$K$-$M$ high-symmetry line on a $1000 \times 1 \times 1k_y$ mesh.
[25] While the cross-plane transport cannot be calculated in graphene simulations, $E_{p\sigma}$, $\Delta E_{\sigma\sigma^*}$, $\sigma_{p\sigma}$, and $\gamma_{p\sigma}$ can be calculated. It is these properties which largely determine the $pV$ cell operation [1].
[26] The density functional theory code VASP [40–43] with plane augmented wave [44, 45] (PAW) pseudopotentials using the generalized gradient approximation (GGA) and Perdew-Burke-Ernzerhof [38, 39] functional (PBE) is used to relax the structures to their minimum energy. A $k_y$ mesh of $6 \times 6 \times 1$ and energy cutoff of 400 meV are used until forces converge within $10^{-4}$ eV/Å and the total energy converges to within $10^{-5}$ eV, where $k_y$ is the electron wave vector.
[27] The density of states for these structures is calculated on a $30 \times 30 \times 1k_y$-point mesh.
[32] Graphene and graphene dynamical matrices are calculated on a $18 \times 18 \times 1k_y$ mesh, while only the $\Gamma$-point dynamical matrix of $C_{128}H_{128}$ is calculated due to computational constraints. However, this $\Gamma$-point calculation contains approximately an $8 \times 8 \times 1$ mesh of atomic force constants.
[36] Other options include low-dimension graphene and the noncovalent functionalization of graphene [46].