

<https://doi.org/10.1038/s41524-024-01399-z>

Enhanced high harmonic efficiency through phonon-assisted photodoping effect



Jin Zhang^{1,2,6}✉, Ofer Neufeld^{2,6}✉, Nicolas Tancogne-Dejean²✉, I-Te Lu², Hannes Hübener², Umberto De Giovannini^{2,3} & Angel Rubio^{2,4,5}✉

High-harmonic generation (HHG) has emerged as a central technique in attosecond science and strong-field physics, providing a tool for investigating ultrafast dynamics. However, the microscopic mechanism of HHG in solids is still under debate, and it is unclear how it is modified in the ubiquitous presence of phonons. Here we theoretically investigate the role of collectively coherent vibrations in HHG in a wide range of solids (e.g., hBN, graphite, 2H-MoS₂, and diamond). We predict that phonon-assisted high harmonic yields can be significantly enhanced, compared to the phonon-free case – up to a factor of ~20 for a transverse optical phonon in bulk hBN. We also show that the emitted harmonics strongly depend on the character of the pumped vibrational modes. Through state-of-the-art *ab initio* calculations, we elucidate the physical origin of the HHG yield enhancement – phonon-assisted photoinduced carrier doping, which plays a paramount role in both intraband and interband electron dynamics. Our research illuminates a clear pathway toward comprehending phonon-mediated nonlinear optical processes within materials, offering a powerful tool to deliberately engineer and govern solid-state high harmonics.

High harmonic generation (HHG), initially observed in rare gas atoms, has recently attracted substantial interest in the attosecond and strong-field processes in solids^{1–5}. HHG provides a unique tool to probe optical susceptibility, band structures, and electron dynamics of solids and liquids^{6–12}. In terms of applications, solid-state HHG provides an effective coherent light source for extreme ultraviolet radiation, making it of significant importance to understand and control the nonlinear optical phenomena to engineer the emitted harmonics characteristics^{13–19}. The underlying mechanism responsible for HHG in solids mostly involves two emission pathways: i) light emission from the accelerated electrons within a band under an external driving field, also known as intraband process and ii) light emission from electron-hole recombination, that is, interband process^{14–17}. These two pathways depend on both material and external driving-field properties, e.g. the band structure, carrier density, laser polarization, and ellipticity, which have been used for effectively tuning the spectral and temporal properties of the HHG emission, as well as probing electron dynamics in real time^{8,18–24}.

Periodic solids host collective lattice vibrations, or phonons, that are widely present either thermally, incoherently or coherently, which can modify the electronic, optical, and magnetic properties of materials^{25–37}. Since solid HHG is sensitive to dynamical variations in the electronic density and nuclei potential, one could manipulate, by exciting phonons, the emitted harmonics characteristics¹⁵. Such use of an internal degree of freedom in the lattice has so far not been systematically explored as a control method for HHG from solids. Recently, Ginsberg et al. illustrated the impact of the phonons on the second and third harmonic generation, and showed that the emitted yields are sensitive to lattice vibrations in hexagonal-Boron-Nitride (hBN)²⁵. It was also shown that lower-order harmonics can be sensitive to coherent phonon dynamics in vanadium dioxide²⁶ and in quartz [<https://www.nature.com/articles/s41566-024-01457-4>], and that the carrier envelope-phase of the driving field can play an additional role in phonon HHG spectroscopy²⁷. In this regard, coherently excited phonons provide a promising strategy to engineer or probe ultrafast nonlinear dynamics in solids. However, the scope of how to utilize collective lattice dynamics to

¹Laboratory of Theoretical and Computational Nanoscience, National Center for Nanoscience and Technology, Chinese Academy of Sciences, 100190 Beijing, China. ²Max Planck Institute for the Structure and Dynamics of Matter and Center for Free-Electron Laser Science, Luruper Chaussee 149, 22761 Hamburg, Germany. ³Università degli Studi di Palermo, Dipartimento di Fisica e Chimica—Emilio Segrè, Palermo, I-90123, Italy. ⁴Center for Computational Quantum Physics (CCQ), The Flatiron Institute, 162 Fifth avenue, New York, NY, 10010, USA. ⁵Nano-Bio Spectroscopy Group, Universidad del País Vasco UPV/EHU- 20018 San Sebastián, San Sebastián, Spain. ⁶These authors contributed equally: Jin Zhang, Ofer Neufeld. ✉e-mail: jinzhang@nanoctr.cn; ofer.neufeld@mpsd.mpg.de; nicolas.tancogne-dejean@mpsd.mpg.de; angel.rubio@mpsd.mpg.de

control HHG in different materials is largely explored; more importantly, the underlying microscopic mechanism for such phonon-assisted HHG has not been investigated in detail.

Here, we show how coherent optical phonons affect HHG from several prototype layered materials (graphite, 2H-MoS₂, and hBN) and covalently-bonded solids (diamond) using time-dependent density-functional theory (TDDFT) coupled to classical nuclei motion^{38–45}. The phonon-assisted HHG spectra from solids strongly depend on both the specific solid and the characteristics of the excited phonon modes. In bulk hBN, for example, the simulated high-harmonic yields can be significantly enhanced by a factor of ~20 in the presence of transverse optical (TO) phonon with E_{1u} mode at 7.3 μm . We reveal that the lattice vibrations effectively help promote the excited carrier dynamics and boost the photoinduced doping, which is the source of the harmonic yield enhancement. Our work implies that phonon pumping provides a new degree of freedom to optimize HHG from condensed matter.

Results and discussion

Phonon-assisted HHG in bulk hBN

In the family of layered materials, both the strong in-plane bonding and the weak van der Waals interactions between layers provide a variety of degrees of freedom in lattice vibrations and electronic structures. In this regard, we first focus on the effect of the optical phonons on HHG in several layered solids (e.g., hBN, graphite, and 2H-MoS₂). The phonons are prepared by setting an initial displacement on the ions along the eigenmode direction (see Methods). We then propagate the electron-phonon coupled dynamics within the framework of TDDFT for the electrons coupled to the Ehrenfest dynamics for the classical ions³⁸. We validate this approach, by comparing the results to the case where the collective lattice motion was initiated following interactions with a phonon-mode-resonant THz laser pulse.

To demonstrate the role of phonons in the HHG emission from solids, we first consider an in-plane TO phonon with E_{1u} mode at Γ point in hBN. Such phonon mode can induce broken inversion symmetry when strongly excited by a laser pulse with a wavelength of 7.3 μm (~170 meV)²⁵. The atomic displacements of the mode in the adjacent hBN layers are in opposite directions within the hBN plane, leading to a net change in the polarizations, as shown in the inset of Fig. 1a. Phonon modes in hBN are prepared by setting an initial velocity on the ions along the eigenmode displacement direction by 1% of the lattice parameter and we then propagate classical Ehrenfest dynamics together with TDDFT. To check the method, we also

applied pump laser pulses with the same frequencies and polarizations of phonon modes of hBN and tracked the atomic displacements. Our calculations confirm the two methods are equivalent in the simulations of high-harmonic generation, thus mimicking the experimental condition whereby coherent phonons are excited by another laser pulse.

Figure 1a shows that excited phonons can significantly modify the HHG spectrum of hBN. For the phonon-free system, clean harmonic peaks are observed up to the 29th order; that is, for the pump laser with a wavelength of 800 nm (~1.55 eV), the estimated cutoff energy for the HHG plateau is 45 eV ($\approx 29 \times 1.55$ eV). In the presence of a coherent TO mode (E_{1u} at the Γ point, Supplementary Fig. S1), the intensity of the harmonics up to the 7th order is enhanced compared to the phonon-free case, while harmonic orders beyond 7 can no longer be distinguished. As expected, even-order harmonics (the 2nd, 4th and 6th orders) appear due to the phonon-induced symmetry breaking in the adjacent layers^{13,45}. We only focus on the phonon mode at Γ point because of energy and momentum conservation. Furthermore, a $2 \times 2 \times 1$ hBN supercell is employed to confirm that the spectrum emitted from the supercell is equivalent to that emitted from the unit cell, indicating that the unit cell is enough to illustrate the phonon-mediated harmonics, as shown in Supplementary Fig. S2.

As a striking point, the spectra obtained in the presence of excited phonons are noisy and do not display a clear odd-harmonic structure, for example, see Fig. 1a–c. This results from the phonon HHG sidebands, which are generated due to the presence of two fundamental frequencies: i) the driving laser field frequency (generally generating high harmonics), and ii) the frequency of the phonon motion itself, generating sidebands. In hBN, the yields of some even-order harmonics are comparable to that of odd-order harmonics. This is mainly attributed to the breaking of inversion symmetry caused by collective lattice vibrations (phonons), which is supported by recent experiments²⁵.

As an example of this, a peak at an energy of 0.17 eV appears in Fig. 1a, which corresponds to the energy of the vibrations of the E_{1u} mode in hBN (where electrons adiabatically follow the lattice). For a more general perspective, we also explore the HHG spectrum with a driven longitudinal optical mode with the same frequency (see Supplementary Fig. S3). The corresponding HHG spectra show a similar trend to that for the excited TO case, where there is however a comparatively smaller increase in the intensity of the even-order harmonics. This also demonstrates that the spectra contain mode-selective information that might be useful for identifying the exact optically active phonon.

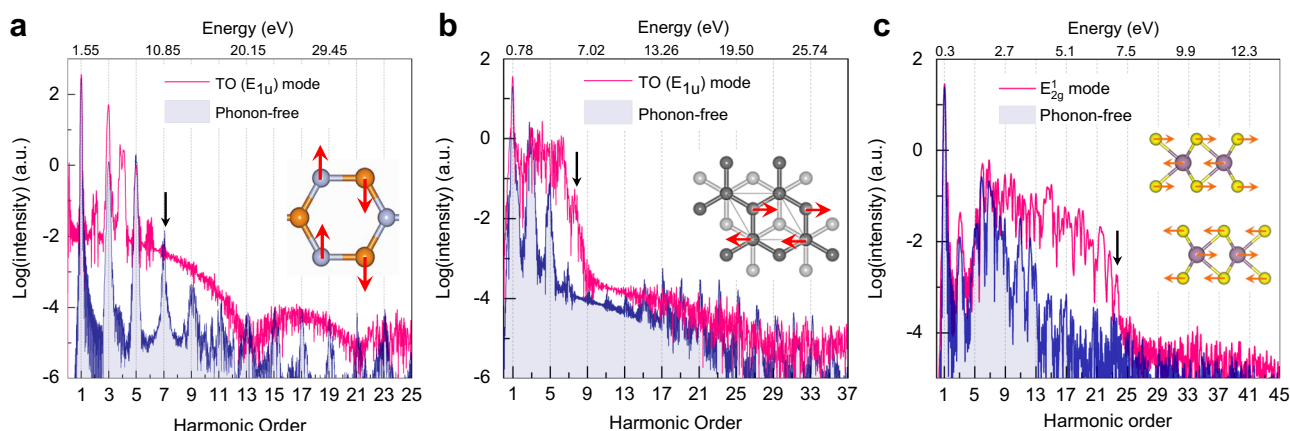


Fig. 1 | HHG spectra from layered materials assisted with phonons. **a** HHG spectrum for hBN with the excitation of the TO (E_{1u} mode at Γ point, red curve) and with the frozen ions (blue curve). We employ a laser pulse with a wavelength of $\lambda = 800$ nm (i.e., $\hbar\omega = 1.55$ eV) and $I_0 = 1 \times 10^{13}$ W cm⁻². The adjacent hBN layers stack laterally with the AA' configuration of a hexagonal structure. **b** HHG spectrum for graphite with the excitation of the TO (E_{1u} mode at Γ point, red curve) and with the frozen ions (blue curve). The laser wavelength is $\lambda = 1.6$ μm ($\hbar\omega = 0.78$ eV) and the intensity is $I_0 = 1 \times 10^{11}$ W cm⁻². The adjacent carbon layers stack laterally with

the AB configuration of a hexagonal structure. **c** HHG spectrum for 2H-MoS₂ with the excitation of the TO (E_{2g}^1 mode at Γ point, red curve) and with the frozen ions (blue curve). The laser wavelength is $\lambda = 4.133$ μm ($\hbar\omega = 0.3$ eV) and the intensity is $I_0 = 1 \times 10^{11}$ W cm⁻². The insets show the atomic displacements of the corresponding modes. The black arrows mark the corresponding cutoff energies of the phonon-mediated spectra. The harmonic intensity is probed along the direction parallel with the laser polarization of the corresponding applied laser pulse.

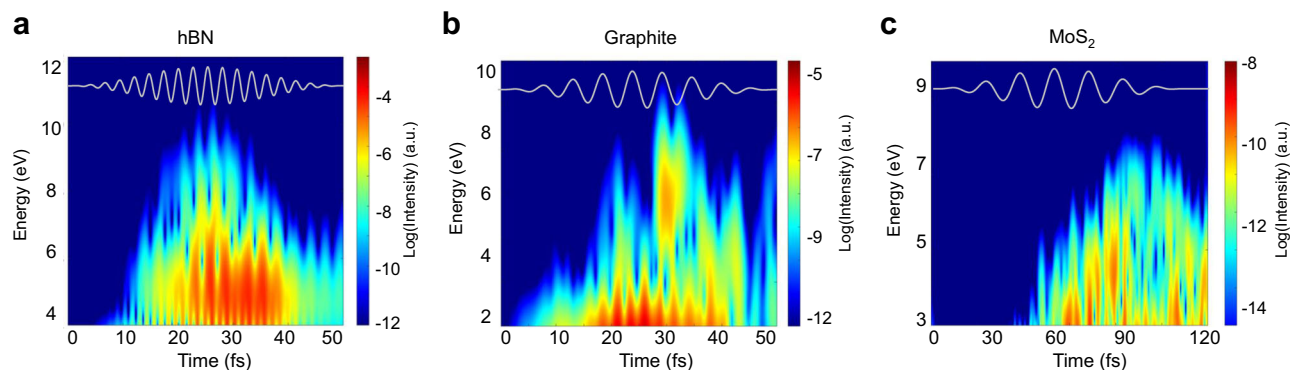


Fig. 2 | Laser-induced total currents and time-frequency analysis of harmonic emissions. **a** Time-frequency analysis of harmonic emissions for hBN with the TO (E_{1u}) mode. At different times (x-axis), the intensity illustrates the emitted harmonics with corresponding energy (y-axis). **b** The same quantity as (a) for graphite

with TO (E_{1u}) mode. **c** The same quantity as (a) for MoS₂ with the E_{2g}^1 mode. The vector potentials of the applied laser pulses are shown in (a–c) (white curves), respectively.

We next perform a more detailed analysis of the harmonic yield changes due to excited optical phonons in hBN. For simplicity, we consider the total yield that is computed by integrating the HHG yield between 4 and 12 eV (roughly from the 3rd to the 8th harmonics). As a result, a 17-fold enhancement is observed for the TO (E_{1u}) mode while a 16-fold increase is obtained for the LO (E_{1u}) mode. In short, by pre-exciting optical phonons, the total harmonic yields can be enhanced by more than an order of magnitude. It should be noted that a laser pulse with a carrier wavelength of 7.3 μm can excite the IR-active TO (E_{1u}) mode directly²⁵. See Supplementary Fig. S4 for the details on the effect of the phonon amplitude and Supplementary Fig. S5 for the case with thermal phonons at 300 K.

Phonon-assisted HHG in graphite and 2H-MoS₂

To generalize these findings, we further study other layered materials with distinct electronic structures, e.g., graphite with zero bandgap and semiconducting 2H-MoS₂. Furthermore, we extend our simulations to a non-layered system, diamond (Supplementary Fig. S6). From these materials, we find a clear physical picture of the harmonic emission mediated by optical phonons as will be discussed below.

For graphite, the infrared-active TO (E_{1u}) mode with a frequency of $\sim 1600\text{ cm}^{-1}$ (corresponding to 0.2 eV)⁴⁶ hosts atomic displacements with antiphase motion in the two adjacent graphene planes. Figure 1b presents the HHG spectra with and without an excited TO mode (E_{1u} at the Γ point). For a pump laser pulse with a wavelength of 1.6 μm , the phonon-free spectrum exhibits clear peaks up to the 5th order, which is equivalent to a cutoff energy of 3.87 eV. With the E_{1u} phonon excited, the yield for the harmonics up to the 7th order is enhanced by a factor of 3.7. Similar to hBN, no clear harmonic peaks are visible due to the excited sidebands.

For 2H-MoS₂, we focus on the Raman-active TO (E_{2g}^1) mode at the Γ point (with an energy of 0.3 eV) with in-plane atomic displacements. In Fig. 1c, for the phonon-free case, harmonic peaks up to the 11th order (corresponding to the cutoff energy of ~ 3.3 eV) are clearly observed. With a pre-excited E_{2g}^1 mode, the HHG cutoff energy extends to 6.9 eV (the 23rd order). Moreover, the harmonic efficiency below the cutoff energy increases, and the associated harmonic yield doubles compared to the phonon-free case (computed by integrating the spectrum between 1 and 7 eV). To illustrate the role of different modes, we also study the effects of a pre-excited out-of-plane A_{1g} mode on the HHG spectra (Supplementary Fig. S7). We find that the phonon-induced modification in the plateau from the A_{1g} mode is smaller compared to the E_{2g}^1 mode. This comparison indicates that different modes affect the strong-field induced electron dynamics in different ways, and that this effect can be probed by HHG.

To explore the temporal characteristics of the electron dynamics, we perform a time-frequency analysis below; such analysis can illustrate how the HHG response builds up in time and allows us estimating the emission and return times of the electron dynamics, the kinetic energies, and their role

in the harmonic spectrum^{47,48}. This analysis can further provide a comprehensive understanding of the ultrafast processes involved in high-harmonic generation, which is useful for deciphering the mechanism through which phonons enhance the HHG yield.

Figure 2 exhibits the time-frequency analysis of harmonic emission for the bulk materials with corresponding excited phonons. For hBN, we can make out short and long HHG electron trajectories, as have been well established^{21,22}. However, we observe that the specific characteristics become slightly attenuated along the laser pulse, corresponding to the instantaneous changes in the lattice structure induced by the collective lattice motion. In other words, the HHG emission is asymmetric between adjacent laser cycles, which also contributes to the noisier spectra observed with no apparent odd-harmonic peaks because temporal translation symmetry is broken²⁷ and phononic sidebands are generated²⁸. Largely similar results are observed for graphite and 2H-MoS₂ (Fig. 2b, c), though here the phonon-induced modification to the electronic dynamics is even more significant, and we can make out particular laser cycles that dominantly contribute to the response in certain spectral regions. For instance, in graphite the HHG emission from ~ 3 to 6 eV is dominantly contributed by one laser cycle after the envelope peak power [around 30 fs, see Fig. 2b]. The excited lattice vibrations increase the portion of electrons excited to the conduction bands by strong pulses, therefore, introducing phonon-enhanced carrier concentrations. As we will show below, one main effect of the pre-excited phonon dynamics on the electronic response is to allow enhanced excitations to the conduction bands, therefore, introducing phonon-enhanced carrier concentrations that increase the HHG yield.

Intensity dependence of phonon-assisted HHG spectra

Next, we focus on how the enhancement of the HHG spectra due to optical phonons depends on the laser pulse characteristics^{14–16}. We use hBN as an example and compute its harmonic yields in the presence and absence of optical phonons under three pump intensities (I_0) (see Fig. 3). For the phonon-free case, we observe clean harmonic peaks with significantly larger efficiencies for a larger intensity of $I_0 = 5 \times 10^{12}\text{ W cm}^{-2}$. As for the TO (E_{1u}) phonon case, the harmonic efficiencies increase significantly with the laser intensity (see below for the comparison of the harmonic yields). In addition, the cutoff energy of the harmonic plateau increases from the 7th to the 11th order (equivalent to an energy of 17.1 eV). In this regard, the phonon seems to lead to a reduction in the energy cutoff.

From the HHG spectra of 2H-MoS₂ [Fig. 1c], we find a small change in the harmonic yield for harmonics with energies below the bandgap (~ 0.9 eV in our simulations). In contrast, we observe a significant change in the harmonic intensity for the 9th–21st orders with energies above the bandgap. Such results indicate that the interband mechanism for HHG is more sensitive to the collective lattice movements. This is further supported by the sidebands in the HHG spectrum, demonstrating that the

Fig. 3 | Strength dependence of the phonon-assisted harmonic spectra. **a** Laser strength dependence of the harmonic yields for bulk hBN with the frozen ions. For the probe pulse, we employ an in-plane driving electric field with a wavelength of $\lambda = 800$ nm with different intensities (I_0) and the polarization is parallel to the vibrational mode of TO (E_{1u}) mode. **b** The same quantity as **(a)** for the conditions with the excited TO (E_{1u}) mode.

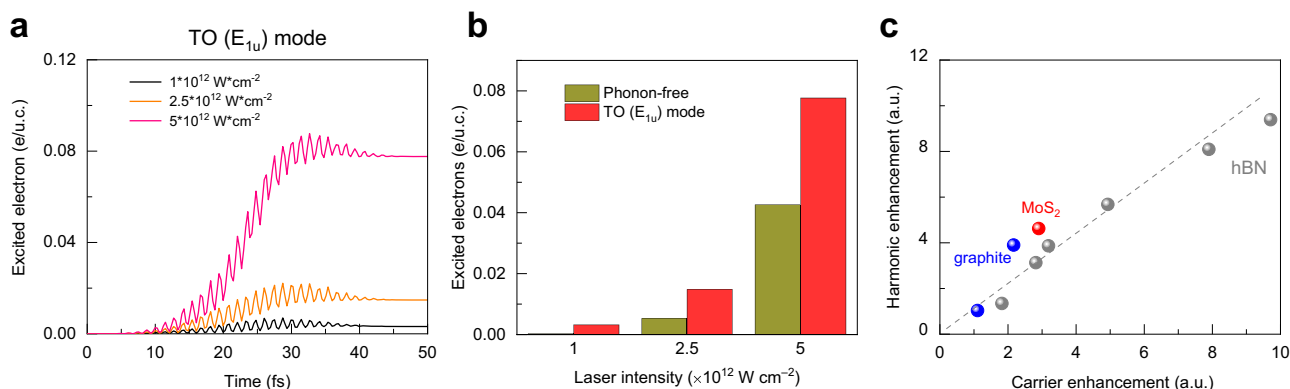
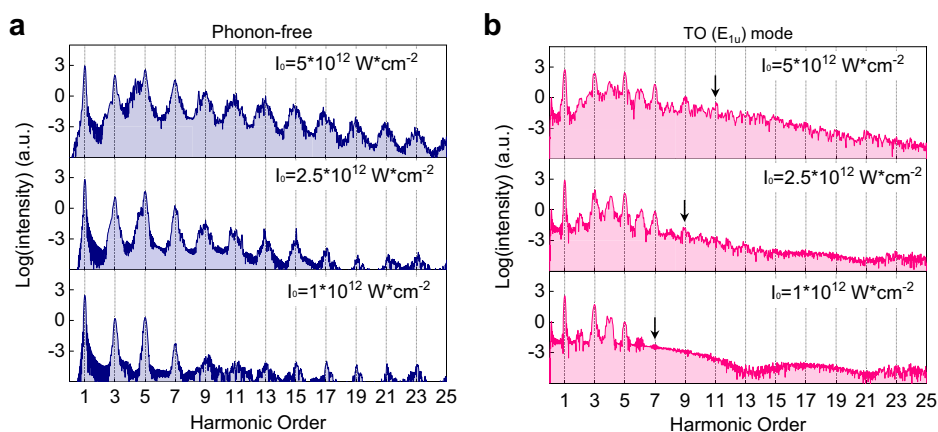


Fig. 4 | Photon-induced doping effects in enhanced harmonic efficiency.

a Number of excited electrons as a function of time in hBN with the excited TO (E_{1u}) mode under different laser intensities. **b** Number of excited electrons for the phonon-free case pulses and the excited phonon case in bulk hBN for different laser intensities at the end of the laser. The excited electrons indicate the number of photoexcited electrons after the laser pulse ($t = 50$ fs) as shown in **(a)**. **c** The

dependence of harmonic enhancement ($\frac{Y_{\text{Phonon}}}{Y_{\text{Frozen-ions}}}$) on photoinduced carrier enhancement ($\frac{N_{\text{Phonon}}}{N_{\text{Frozen-ions}}}$). The harmonic yields are integrated over the harmonic plateaus for the corresponding materials. Different points are from various laser powers, and the initial phonon displacements are kept the same. Blue and red dots indicate the dependence of harmonic yield enhancement on the carrier enhancement for graphite and MoS_2 , respectively. The gray line is a guide for the eyes.

interband transition plays a dominant role when the phonons are evoked.

We note that the HHG spectra obtained in the presence of the TO excited optical phonons are more structured and do not display a clear odd-only harmonic structure. This in fact results mostly from the presence of phonon-induced sidebands, which are generated due to the presence of electron and phonon frequencies [See SI for more details]. The sideband effect can also well explain the dip at the even harmonic position in our simulations. The energy width of two split peaks of even harmonics is nearly twice of the energy of the TO mode, indicating that the first sidebands are more intense than the even-harmonic response. While phonons may enhance the overall signal, the smoothing effect on higher harmonic peaks can have important implications for the spectral characteristics, efficiency, and interpretation of EUV generation processes. Understanding and mitigating these effects are crucial for advancing the field and optimizing the performance of EUV sources for various applications.

Phonon-enhanced photodoping effect

To explain the phonon-mediated harmonic yields, we inspect the photon-induced carrier doping in the systems, i.e., the time evolution of the number of excited electrons. Figure 4a shows the number of excited electrons in hBN as a function of time for three different laser pulse intensities. Compared to the phonon-free case, more charge carriers are excited from valence bands to conduction bands [Fig. 4b and Supplementary Fig. S8]. For instance, for the

intensity of $I_0 = 1 \times 10^{12} \text{ W cm}^{-2}$, the number of the excited electrons due to the TO (E_{1u}) mode is 8.5 times larger than that for the phonon-free case. The enhancement of photodoping implies a strong nonlinear interaction between the coherent phonons and the electronic states. The amplification effect suggests that coherent phonons are not just a perturbative factor but play a significant role in modulating the carrier density through electron-phonon dynamical couplings.

To find a quantitative relationship between the photodoping carrier density and the phonon-enhanced harmonic yield, we first use hBN as an example and compute its harmonic yield evaluated using the plateau region between 4 and 12 eV. Figure 4c shows a nearly linear relation between the harmonic enhancement and the photoexcited carrier enhancement for hBN. Here, the harmonic yield and the photoexcited carrier enhancement are defined as $\frac{Y_{\text{Phonon}}}{Y_{\text{Frozen-ions}}}$ and $\frac{N_{\text{Phonon}}}{N_{\text{Frozen-ions}}}$, respectively, where Y_{Phonon} ($Y_{\text{Frozen-ions}}$) and N_{Phonon} ($N_{\text{Frozen-ions}}$) are the harmonic yield and the number of excited carriers with (without) the excited phonons, respectively. Our results illustrate that phonons can modulate light-induced carrier excitations, which in turn significantly increase the harmonic yields. We also obtain similar trends for the excitation dynamics in graphite and 2H- MoS_2 (Supplementary Fig. S9). With the excited phonon, the photoinduced carrier concentration increases by 2.1 (2.5) times for graphite (MoS_2), compared to the phonon-free case. Putting all the above materials together, we obtain a universal curve between the phonon-enhanced harmonic efficiency and the laser-enhanced carrier density [Fig. 4c]. In addition, we excited both TO

(7300 nm) and LO (6200 nm) modes to confirm the scaling. Our analysis on the harmonic yield and carrier enhancement follows the same trend with the single mode excited cases [Supplementary Fig. S10]. Our results indicate that the photodoping carrier density in solids is strongly correlated to the phonon excitation, and is thus one of the crucial ingredients in phonon-assisted HHG.

The mechanism can be interpreted as follows: photocarriers are produced by generating excited electron-hole pairs when a laser field induces interband Zener tunnelling with significant probability. The enhancement of photodoping implies a strong nonlinear interaction between the coherent phonons and the electronic state. The coherent phonons can interact with electronic states, influencing carrier dynamics. The amplification effect suggests that coherent phonons are not just a perturbative factor but play a significant role in modulating the carrier density. This is indeed distinct from the linear perturbation, which typically would result in much smaller changes. The mechanisms could be explained by strong nonlinear electron-phonon coupling and dynamic band modulation [Supplementary Fig. S11]. However, the exact contribution of interband Zener tunneling is difficult to extract because all interactions are entangled in the total photoinduced current and harmonics.

As observed, the photoinduced carrier concentration is effectively enhanced from collective lattice vibrations and corresponding phonon-modulated electronic structures. On the one hand, phonon-induced carrier density enhancement would boost the harmonic yields from the intraband mechanism, naturally attributed to Bloch oscillation. On the other hand, the enhanced carriers provide more channels and increase the possibility of electron-hole recombination. Therefore, the contribution from interband transitions is improved as well. Previously, the phonon-assisted HHG process was reported to be sensitive to the pump-probe delay and carrier-envelope phase of driving fields²⁷. The enhancement or suppression in the harmonic efficiency observed here is in-line with those results and is associated with instantaneous changes in the lattice structure (bond compression or stretching) and modulation of the electronic structures, which should also be delay-dependent.

Furthermore, we compare our results in context with some relevant experimental studies. Wang et al.²³ showed that the suppression of the interband HHG yield in ZnO is attributed to the pre-photodoping effect, which depletes the carriers in the valence band and leads to fewer electrons contributing to emitting harmonics. In another study, Nishidome et al.²⁴ demonstrated that the harmonic yields of the single-walled carbon nanotube can be tuned by injecting carriers into the nanotube via electrolyte gating. We find that the phonon-assisted photodoping carriers are coherent with the laser pulses, and more excited electrons are involved to emit harmonics with phonons. Our work, together with the above studies²³, points out that more excited carriers lead to stronger harmonic yields, which is attributed to the phonon-assisted tunneling to the conduction bands. In this work, we do not investigate the effects on HHG yields due to the pump-probe delay or carrier-envelope-phase, which could play an additional role in controlling phonon-assisted HHG yields (depending on the laser parameter regime)²⁷. Our results emphasize the importance of ionic movements in nonlinear optics, which can be extended to a situation where multiple phonons are involved in HHG.

In summary, we have investigated the effects of coherent optical phonons on HHG from typical solids using TDDFT coupled to classical ionic dynamics. The excitation of optical phonons enhances high-order harmonic yields, but less clear high-order harmonic peaks are observed due to the generation of phononic sidebands. We identified that the underlying mechanism for the enhancement of the harmonic yield is that phonons can effectively assist promoting photoexcited carriers, which results in increasing harmonic emission. Overall, our results provide a new platform to study strong-field physics in solids, and reveal the importance of lattice degrees of freedom for controlling harmonic emission.

Methods

TDDFT calculations

The time evolution of the wave functions and the evaluation of the time-dependent electronic current were computed by propagating the Kohn–Sham equations in real space and real time, as implemented in the Octopus code^{40–42} with the adiabatic LDA functional⁴⁴. All calculations were performed using fully relativistic Hartwigsen, Goedecker, and Hutter (HGH) pseudopotentials⁴⁵. In our simulations, the laser pulses are treated using the velocity gauge and the induced vector fields are imposed to be time-dependent but homogeneous in space, and we use a sin-square pulse envelope.

The full harmonic spectrum is computed directly from the total electronic current density $\mathbf{j}(\mathbf{r}, t)$ as shown in Fig. 1.

$$HHG(\omega) = \left| FT \left(\frac{\partial}{\partial t} \int d^3r \mathbf{j}(\mathbf{r}, t) \right) \right|^2, \quad (1)$$

where FT denotes the Fourier transform. We note that both electronic and ionic currents contribute to the HHG spectra in experimental measurements. In our simulations, the ionic current from lattice vibrations contributes negligibly, compared with the electronic current and is therefore neglected (it is dominantly expected to contribute only at the phonon resonance).

The time-frequency analysis is obtained by performing a windowed Fourier transform of the current,

$$G(\omega, t_0) = \left| \int dt \partial_t \mathbf{j}(t) e^{-\frac{(t-t_0)^2}{\sigma^2}} e^{-i\omega t} \right|^2, \quad (2)$$

The number of excited electrons (N_{ex}) is calculated by projecting the time-evolved wavefunctions ($|\psi_{n,\mathbf{k}}(t)\rangle$) on the basis of the ground-state wavefunctions ($|\psi_{n',\mathbf{k}}^{GS}\rangle$).

$$N_{ex}(t) = N_e - \frac{1}{N_{\mathbf{k}}} \sum_{n,n'}^{occ} \sum_{\mathbf{k}}^{BZ} \left| \langle \psi_{n,\mathbf{k}}(t) | \psi_{n',\mathbf{k}}^{GS} \rangle \right|^2, \quad (3)$$

where N_{ex} is the total number of electrons and $N_{\mathbf{k}}$ is number of $\mathbf{k}$ -points used to sample the Brillouin zone (BZ). The sum over band indices n and n' go over all occupied states. It should be noted that ground-state wavefunctions are dependent on the time-propagated atomic coordinates. In this regard, self-consistent calculations are performed using the time-dependent atomic structures every 0.3 fs to obtain the ground-state wavefunctions for projections.

Simulations of HHG from bulk hBN

The real-space cell was sampled with a grid spacing of 0.4 Bohr and the Brillouin zone was sampled with a $42 \times 42 \times 21$ $\mathbf{k}$ -point grid to sample the BZ, which yielded converged results for hBN. The boron nitride bond length is taken here as the experimental value of 1.445 Å. Within the LDA functional, the bandgaps of hBN are 4.6 eV. We consider a laser pulse of 25-fs duration at full-width half-maximum (FWHM) with a sin-square envelope, and the carrier wavelength λ is 800 nm, corresponding to 1.55 eV, polarized along polarized along x axis.

Simulations of HHG from graphite and bulk MoS₂

For graphite, the experimental lattice constant is used (1.42 Å) in our simulations with a $42 \times 42 \times 21$ $\mathbf{k}$ -point grid to sample the BZ. The real-space cell was sampled with a grid spacing of 0.4 Bohr. The laser pulse of 25-fs duration at FWHM is considered with a sin-square envelope, and the carrier wavelength λ is 1600 nm, corresponding to 0.78 eV. We used a peak intensity of $I_0 = 10^{11}$ Wcm⁻².

For bulk MoS₂, we used a real-space spacing of 0.35 Bohr and a mesh of $24 \times 24 \times 12$ to sample the BZ of MoS₂. The lattice constant of MoS₂ is 3.16 Å. A carrier wavelength of $\lambda = 4133$ nm was used, corresponding to a carrier photon energy of 0.3 eV. The peak intensity in vacuum is $I_0 = 10^{11}$

Wcm^{-2} . In our simulations, a longer laser pulse (FWHM = 50 fs) is employed, in order to fully cover the lattice vibrations of E_{2g}^1 phonon mode at Γ point. The phonon modes of MoS_2 is prepared by starting by the time-evolution from a distorted atomic configuration along the phonon modes of 4% of the lattice constant because smaller displacements result in no significant modulations.

Data availability

The data that support the findings of this study are available from the corresponding author upon request. Source data are provided with this paper.

Received: 29 January 2024; Accepted: 18 August 2024;

Published online: 02 September 2024

References

- Ghimire, S. & Reis, D. A. High-harmonic generation from solids. *Nat. Phys.* **15**, 10–16 (2019).
- Ferray, M. et al. Multiple-harmonic conversion of 1064 nm radiation in rare gases. *J. Phys. B Mol. Opt. Phys.* **21**, L31 (1988).
- Luu, T. T. et al. Extreme ultraviolet high-harmonic spectroscopy of solids. *Nature* **521**, 498 (2015).
- Schmid, C. P. et al. Tunable non-integer high-harmonic generation in a topological insulator. *Nature* **593**, 385 (2021).
- von Hoegen, A., Mankowsky, R., Fechner, M., Först, M. & Cavalleri, A. Probing the interatomic potential of solids with strong-field nonlinear phononics. *Nature* **555**, 79–82 (2018).
- Uzan, A. J. et al. Attosecond spectral singularities in solid-state high-harmonic generation. *Nat. Photon.* **14**, 183–187 (2020).
- Vampa, G. et al. All-optical reconstruction of crystal band structure. *Phys. Rev. Lett.* **115**, 193603 (2015).
- Yoshikawa, N., Tamaya, T. & Tanaka, K. High-harmonic generation in graphene enhanced by elliptically polarized light excitation. *Science* **356**, 736 (2017).
- Liu, H. et al. High-harmonic generation from an atomically thin semiconductor. *Nat. Phys.* **13**, 262 (2017).
- You, Y. S., Reis, D. A. & Ghimire, S. Anisotropic high-harmonic generation in bulk crystals. *Nat. Phys.* **13**, 345–349 (2017).
- Ferré, A. et al. Multi-channel electronic and vibrational dynamics in polyatomic resonant high-order harmonic generation. *Nat. Commun.* **6**, 5952 (2015).
- Mondal, A. et al. Probing the low-energy electron-scattering dynamics in liquids with high-harmonic spectroscopy. *Nat. Phys.* **19**, 1813–1820 (2023).
- Neufeld, O., Podolsky, D. & Cohen, O. Floquet group theory and its application to selection rules in harmonic generation. *Nat. Commun.* **10**, 405 (2019).
- Li, L., Lan, P., Zhu, X. & Lu, P. Huygens-fresnel picture for high harmonic generation in solids. *Phys. Rev. Lett.* **127**, 223201 (2021).
- Tancogne-Dejean, N., Mücke, O. D., Kärtner, F. X. & Rubio, A. Impact of the electronic band structure in high-harmonic generation spectra of solids. *Phys. Rev. Lett.* **118**, 087403 (2017).
- Tancogne-Dejean, N., Mücke, O. D., Kärtner, F. X. & Rubio, A. Ellipticity dependence of high-harmonic generation in solids originating from coupled intraband and interband dynamics. *Nat. Commun.* **8**, 745 (2017).
- Klemke, N. et al. Polarization-state-resolved high-harmonic spectroscopy of solids. *Nat. Commun.* **10**, 1319 (2019).
- Ndabashimiye, G. et al. Solid-state harmonics beyond the atomic limit. *Nature* **534**, 520–523 (2016).
- Pertot, Y. et al. Time-resolved x-ray absorption spectroscopy with a water window high-harmonic source. *Science* **355**, 264–267 (2017).
- Schubert, O. et al. Sub-cycle control of terahertz high-harmonic generation by dynamical Bloch oscillations. *Nat. Photon.* **8**, 119–123 (2014).
- Le Breton, G., Rubio, A. & Tancogne-Dejean, N. High-harmonic generation from few-layer hexagonal boron nitride: Evolution from monolayer to bulk response. *Phys. Rev. B* **98**, 165308 (2018).
- Tancogne-Dejean, N. & Rubio, A. Atomic-like high-harmonic generation from two-dimensional materials. *Sci. Adv.* **4**, eaao5207 (2018).
- Wang, Z. et al. The roles of photo-carrier doping and driving wavelength in high harmonic generation from a semiconductor. *Nat. Commun.* **8**, 1686 (2017).
- Nishidome, H. et al. Control of high-harmonic generation by tuning the electronic structure and carrier injection. *Nano Lett.* **20**, 6215–6221 (2020).
- Ginsberg, J. S. et al. Phonon-Enhanced Nonlinearities in Hexagonal Boron Nitride. *Nat. Comm.* <https://doi.org/10.1038/s41467-023-43501-x> (2023).
- Bionta, M. R. et al. Tracking ultrafast solid-state dynamics using high harmonic spectroscopy. *Phys. Rev. Res.* **3**, 023250 (2021).
- Neufeld, O., Zhang, J., De Giovannini, U., Hübener, H. & Rubio, A. Probing phonon dynamics with multidimensional high harmonic carrier-envelope-phase spectroscopy. *Proc. Natl Acad. Sci. USA* **119**, e2204219119 (2022).
- Rana, N., Mrudul, M. S., Kartashov, D., Ivanov, M. & Dixit, G. High-harmonic spectroscopy of coherent lattice dynamics in graphene. *Phys. Rev. B* **106**, 064303 (2022).
- Yao, K. et al. Enhanced tunable second harmonic generation from twistable interfaces and vertical superlattices in boron nitride homostructures. *Sci. Adv.* **7**, eaabe8691 (2021).
- Neufeld, O., Tancogne-Dejean, N., Hübener, H., De Giovannini, U. & Rubio, A. *Phys. Rev. X* **13**, 031011 (2023).
- Biswas, A. et al. Non-Linear Optics at Twist Interfaces in h-BN/SiC Heterostructures. *Adv. Mater.* **35**, 2304624 (2023).
- Nova, T. F., Disa, A. S., Fechner, M. & Cavalleri, A. *Science* **364**, 1075–1079 (2019).
- Först, M. et al. Nonlinear phononics as an ultrafast route to lattice control. *Nat. Phys.* **7**, 854–856 (2011).
- Zhang, J. et al. High-harmonic spectroscopy probes lattice dynamics. *Nat. Photonics*, <https://doi.org/10.1038/s41566-024-01457-4> (2024).
- Wall, S. et al. Ultrafast changes in lattice symmetry probed by coherent phonons. *Nat. Commun.* **3**, 721 (2012).
- Shin, D. et al. Phonon-driven spin-Floquet magneto-valleytronics in MoS_2 . *Nat. Commun.* **9**, 638 (2018).
- De Giovannini, U., Hübener, H., Sato, S. & Rubio, A. Direct measurement of electron-phonon coupling with time-resolved ARPES. *Phys. Rev. Lett.* **125**, 136401 (2020).
- Andrade, X. et al. Modified Ehrenfest formalism for efficient large-scale ab initio molecular dynamics. *J. Chem. Theory Comput.* **5**, 728–742 (2009).
- He, L. et al. Monitoring ultrafast vibrational dynamics of isotopic molecules with frequency modulation of high-order harmonics. *Nat. Commun.* **9**, 1108 (2018).
- Castro, A. et al. Octopus: a tool for the application of time-dependent density functional theory. *Phys. Status Solidi* **243**, 2465 (2006).
- Andrade, X. et al. Real-space grids and the Octopus code as tools for the development of new simulation approaches for electronic systems. *Phys. Chem. Chem. Phys.* **17**, 31371–31396 (2015).
- Tancogne-Dejean, N. et al. Octopus, a computational framework for exploring light-driven phenomena and quantum dynamics in extended and finite systems. *J. Chem. Phys.* **152**, 124119 (2020).
- Onida, G., Reining, L. & Rubio, A. Electronic excitations: density-functional versus many-body Green's-function approaches. *Rev. Mod. Phys.* **74**, 601–659 (2002).
- Perdew, J. P., Burke, K. & Ernzerhof, M. Generalized gradient approximation made simple. *Phys. Rev. Lett.* **77**, 3865–3868 (1996).

45. Hartwigsen, C., Goedecker, S. & Hutter, J. Relativistic separable dual-space Gaussian pseudopotentials from H to Rn. *Phys. Rev. B* **58**, 3641–3662 (1998).
46. Giura, P. et al. Temperature evolution of infrared- and Raman-active phonons in graphite. *Phys. Rev. B* **86**, 121404 (2012).
47. Yue, L. & Gaarde, M. Introduction to theory of high-harmonic generation in solids: tutorial. *J. Opt. Soc. Am. B* **39**, 535–555 (2022).
48. Morisson Faria, C., Dörr, M., Becker, W. & Sandner, W. Time-frequency analysis of two-color high-harmonic generation. *Phys. Rev. A* **60**, 1377 (1999).

Acknowledgements

We thank Jared Scott Ginsberg, Lede Xian and Dongbin Shin for helpful discussions. This work was supported by the Cluster of Excellence Advanced Imaging of Matter (AIM), Grupos Consolidados (IT1249-19) and SFB925. The Flatiron Institute is a division of the Simons Foundation. J.Z. acknowledges funding from the European Union's Horizon 2020 research and innovation program under the Marie Skłodowska-Curie grant agreement No. 886291 (PeSD-NeSL). O.N. and I.L. gratefully acknowledge the support of the Humboldt foundation. O.N. gratefully acknowledges the support of a Schmidt Science Fellowship.

Author contributions

Jin Zhang, Ofer Neufeld and Angel Rubio conceived and designed the study. Jin Zhang performed the first-principles calculations, data analysis, investigation, and wrote the initial manuscript, all with consultation from Ofer Neufeld, Nicolas Tancogne-Dejean, I-Te Lu, Hannes Hübener and Umberto De Giovannini. All authors participated in discussing and editing the manuscript.

Funding

Open Access funding enabled and organized by Projekt DEAL.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at <https://doi.org/10.1038/s41524-024-01399-z>.

Correspondence and requests for materials should be addressed to Jin Zhang, Ofer Neufeld, Nicolas Tancogne-Dejean or Angel Rubio.

Reprints and permissions information is available at <http://www.nature.com/reprints>

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit <http://creativecommons.org/licenses/by/4.0/>.

© The Author(s) 2024, corrected publication 2024