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Cite as: J. Appl. Phys. 129, 085102 (2021); https://doi.org/10.1063/5.0043623
Submitted: 10 January 2021 . Accepted: 10 February 2021 . Published Online: 23 February 2021

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Topological effects of phonons in GaN and AlGaN: A potential perspective for tuning phonon transport

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ABSTRACT
Tuning thermal transport in semiconductor nanostructures is of great significance for thermal management in information and power electronics. With excellent transport properties, such as ballistic transport, immunity to point defects and disorders, and forbidden backscattering, topological phonon surface states show remarkable potential in addressing this issue. Herein, topological phonon analyses are performed on hexagonal wurtzite gallium nitride (GaN) to check the topological characteristics of phonons. And other nitrides of the same family, i.e., AlN and AlGaN crystals, are also calculated from a topological phonon phase transition perspective. With the aid of first-principle calculations and topological phonon theory, Weyl phonon states, which host surfaces states without backscattering, are investigated for all these materials. The results show that there is no nontrivial topological phonon state in GaN. However, by introducing Al atoms, i.e., in wurtzite type AlN and AlGaN, more than one Weyl phonon point is found, confirmed by obvious topological characteristics, including non-zero integer topological charges, source/sink in Berry curvature distributions, surface local density of states, and surface arcs. As AlN and AlGaN are typical materials in AlGaN/GaN heterostructure based electronics, the existence of topological phonon states in them will benefit thermal management by facilitating the design of one-way interfacial phonon transport without backscattering.

I. INTRODUCTION
Gallium nitride (GaN) with a wurtzite structure, a representative of wide bandgap semiconductors, is currently one of the most important semiconductor materials for information and power electronics. With the increase in working power and integration density, near-junction thermal management is becoming more and more important, including accurate thermal simulations, optimization of thermal transport (i.e., phonon transport in dielectric materials), and efficient heat dissipation. As a typical GaN-based electronic, the GaN high electron mobility transistor (HEMT) has received much attention in the past decade. Theoretical and experimental analyses show that thermal resistance in HEMTs is mainly concentrated in the near-junction region, i.e., the channel layer. Based on hybrid simulations combining phonon Monte Carlo methods and Fourier's law calculations, a deeper understanding of near-junction thermal transport is realized. The approximation of one-dimensional thermal transport used in many types of research is inappropriate for HEMT thermal analyses. The actual thermal transport process is three-dimensional, where heat is generated by a dot-like heat source and spreads to plane-like areas crossing the channel layer and several interfaces. Hence, thermal transport design methods that can couple with the detailed transport process will be more valuable, while reducing the thickness of the channel layer or increasing the thermal conductivity of the substrate may be less effective. Particularly, enhancing interfacial phonon thermal transport parallel to the interface is desirable as it can reduce thermal spreading resistance significantly.

Many methods have been proposed to manipulate phonons and facilitate phonon transport. The typical regimes include
phonon interface/boundary scattering under the phonon ballistic-diffusive regime,\textsuperscript{7–10} phonon coherence from phonon wave properties,\textsuperscript{11} surface phonon polarization from phonon-photon coupling,\textsuperscript{12} phonon localization,\textsuperscript{13} and phonon response to strain, external fields, and phase changes.\textsuperscript{14–19} In general, these methods are built based on the current understanding of phonons, which are regarded as both particles and waves, with a Bose–Einstein distribution, yet without an electric charge, spin, or circular polarization. Thus, a deeper understanding of phonon physics will promulgate further developments in materials and transport science. The topological effects of phonons,\textsuperscript{20–22} which host non-dissipative surface phonon states or without backscattering, promise to provide potential perspectives for tuning phonon transport.

Topological phonon theory has been developed\textsuperscript{20–22} inspired by analogy with topological band theory in electrons\textsuperscript{23–25} and investigations on the phonon Hall effect.\textsuperscript{22,26} Similarly, topological matter physics elucidates a new understanding of electron properties with wave function’s topological structures. Phonon physics also benefit significantly from phonon band topology. Besides phonon dispersion relations and scatterings in classical phonon theory, the phonon wave function’s topological structures can further divide phonons into different phases; with that, excellent phonon surface states are present in topological nontrivial phonon systems. Following the topological photon (Bosons) research of Haldane and co-workers,\textsuperscript{27,28} investigations into the topological effects of phonons in lattice systems with time-reversal symmetry breaking began a decade ago. Zhang et al.\textsuperscript{22,26} first introduced the topological theory to explain the phonon Hall effect. Liu et al.\textsuperscript{20,21} developed the systematic topological phonon theory by adopting the physics in electron topological band theory and proposed possible lattice models with nontrivial topological phonon states (TPSs) as well as effective Hamiltonian models. Similar to the Su–Schrieffer–Heeger (SSH) model for electronic systems, simple phonon topological models were studied in low-dimensional crystal, quasicrystal, and amorphous systems.\textsuperscript{20–22} Though there are many differences between phonons and electrons, topological theory can be well used in phonons and other boson systems with the aid of lattice symmetry. In recent years, accompanied by advances in semi-metal states in electron systems, the Weyl phonon state, a new topological phonon state, received much attention, and several kinds of Weyl phonon states have been reported in real materials.\textsuperscript{34–42}

From the perspective of tuning phonon transport, especially phonon transport in specific semiconductor structures, it is critical to realize and apply topological nontrivial phonon states in certain materials. In this work, the topological effects of phonons in hexagonal wurtzite GaN and its related materials used in GaN-based HEMTs are analyzed with first-principle calculations and the topological phonon theory to check the existence of TPSs, especially Weyl phonon states. It is found that wurtzite AlN and AlGaN contain nontrivial TPSs, specifically Weyl phonons, though the phonon states in GaN are all topologically trivial. Since AlN and AlGaN are both important...
wide bandgap semiconductors consisting of an AlGaN/GaN heterostructure, the finding of Weyl phonons that host surface phonon states without backscattering promises to be helpful to understand phonon properties and phonon transport tuning in GaN-based electronics.

II. TOPOLOGICAL PHONON THEORY AND CALCULATION METHODS

The first topological state in a time-reversal invariant system is the quantum spin Hall insulator.23–25,43 The time-invariant conjugated double degenerate states are preserved natively by the Kramer theorem since the electron is a spin-1/2 particle. Fu44 extended this kind of topological state by introducing crystal lattice symmetry. By combining point group symmetry in the lattice and time-reversal symmetry, double degenerate states can also be realized in a spin-less system represented by pseudospin. Since there is no spin for phonons, the most important task to realize the topological nontrivial insulator phonon state is to create pseudospin, which refers to a topological crystalline insulator. Requirements for topological insulator phonon states are demanding since the crystal symmetry used for bulk states should also be present in the corresponding surfaces, which is one of the reasons why real materials containing topological insulator phonon states are still not reported in the literature. Topological semi-metals such as Dirac, Weyl, triple degenerate nodal point,
and nodal line and ring fermions are new topological states in semi-metal systems.\textsuperscript{45,46} The corresponding Weyl points in three-dimensional phonon systems only require double degenerate band crossing in the bulk system without a requirement for surface lattice symmetry, raising more possibilities of TPSs in real lattices.\textsuperscript{20,34,35,37-42,47} For phonons in phase with topological insulator states and Weyl states, the topological nontrivial surface states are bound states immune to disorder and point defects (two-dimensional topological insulator states) with forbidden backscattering (three-dimensional topological insulator states and Weyl states), hosting robust one-way propagation characteristics protected by bulk topological characteristics of phonons.

In studies of topological states, the definition of topological quantities is crucial and depends on a detailed understanding of the underlying physics. In the crystal lattice, the lattice vibration of associated atoms is described by a set of dynamic equations based on Newton’s law of motion. By adopting the harmonic approximation and lattice wave solutions, an eigenvalue problem can be obtained, i.e., the phonon dynamic equation. Information of harmonic phonons is fully contained in the equation.

**FIG. 4.** The LDOS for the (0001) surface along the high symmetry paths in surface Brillouin zone calculated by the iterative Green’s function method in (a) and (b) and directly solving the eigenstate equation of phonon of a slab structure in (c). The blue and red lines indicate the surface states on the top and bottom surfaces of a slab structure, consistent with those illustrated in (a) and (b).
FIG. 5. Phonon dispersion of wurtzite AlN. (a) The phonon dispersion of AlN with and without NAC. (b) The phonon dispersion obtained by standard phonon calculations and reproduced by Wannier tight-binding model Hamiltonian. (c) The phonon dispersion along the path including two different Weyl points marked as WP$_1$ and WP$_2$ is not affected by NAC. (d) and (e) are enlarged views of two Weyl points in (c).
\[ D(k)u(k) = \omega^2(k)u(k), \]  
(1)

where \( D \) is dynamic matrices, \( u \) is an eigenstate, and \( \omega \) is the phonon frequency. They are all functions of the phonon wave vector \( k \). While more attention is paid to phonon eigenvalues, i.e., phonon dispersion relations, as mentioned above, much information is hidden in eigenvectors, like the topological structure of eigenvectors. By properly defining a phonon wave function, topological quantities used in electron systems can also be defined in phonon systems, where the Berry connection is defined as

\[ A_{nk} = -i\langle u_{nk} | \nabla_k | u_{nk} \rangle, \]  
(2)

Then, Berry curvature is defined as

\[ \Omega_{nk} = \nabla_k \times A_{nk}, \]  
where \( n \) is the band index, and the Berry phase is defined as integral of Berry connection along a path or loop

\[ \gamma_n = \int L A_{nk}/dL. \]  
(3)

FIG. 6. Weyl points in the Brillouin zone at plane \( k_z = 0 \) shown by the black hexagon. The dots with different shapes represent different Weyl points, where the rhombic dots represent Weyl point 1 and its equivalent points as shown in Fig. 5(c), and the circle dots represent Weyl point 2 and its equivalent points. The dots in red mean Weyl points with positive chirality equal to 1, while those in blue mean Weyl points with negative chirality equal to \(-1\). Weyl points linked by lines represent a couple of Weyl points.

FIG. 7. [(a) and (b)] The Wannier charge center evolution for a couple of Weyl points \((0.2989, 0.1249, 0.0000)\) and \((0.4239, -0.1246, 0.0000)\), with positive chirality 1 and negative chirality \(-1\), respectively. [(c) and (d)] The Wannier charge center evolution for a couple of Weyl points \((0.3549, -0.1169, 0.0000)\) and \((0.3549, -0.2380, 0.0000)\), with positive chirality 1 and negative chirality \(-1\), respectively.
Topological invariant is the basic physical quantity to identify the topological properties of the system directly. For the Weyl points focused on in this work, the topological invariant is the topological charge, i.e., the Chern number. The three-dimensional condition is defined in a closed surface containing the Weyl point as

\[
C = \frac{1}{2\pi} \int_{\Omega} \cdot d\mathbf{k}.
\]  

(4)

Currently, there are mainly two kinds of Weyl points, including the single Weyl point with Chern number equal to +1 and the double Weyl point with Chern number equal to +2. Most recently, Weyl points beyond these two kinds have also been reported. Weyl point characteristics can be illustrated by the evolution of the Wannier charge center, the distribution of the Berry curvature, and surface states, besides topological charge calculations. At the Weyl point, a source or sink can be observed in the Berry curvature distribution in a plane containing this Weyl point. Surface states are direct evidence of nontrivial TPSs, in which the surface arc is the characteristic of Weyl points. In detail, the iterative Green’s function method implemented in open-source software WannierTools is adopted to calculate the surface local density of states (LDOS)

FIG. 8. The Berry curvature distribution \((\Omega_x, \Omega_y)\) in the \(k_x-k_y\) plane with \(k_z=0\) for (a) fourth and (c) eighth phonon band. (b) and (d) are enlarged views of the zones encircled by blue rectangles in (a) and (c), respectively. The source and the sink can be found from the distributions. The red and blue dots represent the source and the sink of Berry curvature distribution, illustrating positive and negative chirality. The black hexagon represents the \(k_z=0\) plane of the Brillouin zone of wurtzite AlN.
with a phonon tight-binding (TB) Hamiltonian from second-order interatomic force constants (IFCs) and the Wilson-loop method is used to calculate Wannier charge center evolutions. The basic quantities, including second-order IFCs and phonon eigenvectors, are calculated by adopting the frozen phonon method from first principles accompanied by the open-source software Phonopy. All first-principle calculations are performed within the framework of density functional theory as implemented in the Vienna ab-initio Simulation Package (VASP). Projective augmented wave pseudopotential and the generalized gradient approximation in the Perdew–Burke–Ernzerhof form for the exchange-correlation functional are adopted. The kinetic energy cutoff for a plane wave basis of 800 eV is employed, and the Brillouin zone is sampled using converged $12 \times 12 \times 9$ Gamma-centered $k$-mesh grids. The conventional unit cells are relaxed until the residual stress and the maximum forces acting on each atom are smaller than $10^{-2}$ kbar and $10^{-7}$ eV/Å. The supercells for wurtzite GaN and AlN are

![Diagram of Weyl surface arcs](image)

**FIG. 9.** Weyl surface arcs. The projection of the iso-frequency plane of surface states on the (0001) surface Brillouin zone at a frequency equal to (a) 12.23 THz and (b) 19.57 THz. The red and blue dots represent Weyl points with positive chirality 1 and negative chirality −1, respectively, which are linked by phonon surface arcs. The black hexagon indicates the surface Brillouin zone of the (0001) surface in wurtzite AlN. (b) and (d) are enlarged views of the zones encircled by blue rectangles in (a) and (c), respectively.
both built on conventional unit cells with size $4 \times 4 \times 3$ and $4 \times 4 \times 4$, respectively. For AlGaN, the size depends on the number of atoms in the unit cell, where a $3 \times 3 \times 3$ supercell is used for an eight-atom unit cell and a $4 \times 4 \times 4$ supercell is used for a four-atom unit cell. Born effective charges and dielectric constants are calculated using density functional perturbation theory to consider the effects of the long-range Coulomb interaction on second-order IFCs, which is often called non-analytical correction (NAC) and mainly affects phonon states near the Brillouin zone center, i.e., the Gamma point.

III. RESULTS AND DISCUSSION

A. Topological effects of phonons in GaN

Wurtzite structure is the most common lattice structure for GaN as it is thermodynamically stable. As illustrated in Fig. 1(a), the wurtzite structure is a hexagonal structure with lattice constants $a$ and $c$, and internal parameter $u$, belonging to space group $P6_3mc$ (No. 186). The wurtzite structure host 12 lattice symmetry operations, including threefold rotation symmetry, twofold, and sixfold non-symmorphic screw rotation symmetry, mirror symmetry, and screw mirror symmetry. The Brillouin zone is also shown in Fig. 1(b), where high symmetry points and paths are labeled.

As mentioned above, it is hard to satisfy the requirements for the topological insulator states of phonons, as shown in the wurtzite structure in the following discussion. Also, surface phonons of nodal line (ring) states do not contain excellent transport properties such as forbidden backscattering. Thus, Weyl point states are focused on in this work. Detecting topological nontrivial phonon states starts from analyses of phonon dispersions, especially phonon degeneracy at specific points, lines, and planes. A detailed discussion on the identification of potential topological phonon states can be seen from the supplementary material. The phonon dispersions of GaN are shown in Fig. 2. It is noted ahead of the detailed discussion that NAC is still not properly included in the phonon TB Hamiltonian for materials with strong polarization, which is under development. Thus, phonon dispersion relations without NAC are used in this work. And phonon bands near the Gamma point are then not considered in the topological analyses. As seen from Fig. 2, the circled phonon band crossing points (BCPs) are not affected by NAC for slight differences in the magnitude of phonon frequencies. For all calculations in this work, we have confirmed that the BCPs discussed are not affected by NAC, except the last example of Al$_3$Ga$_4$. Also, including NAC will not bring in new BCPs in general since it will introduce LO-TO splitting at the Gamma point. Turning attention to phonon dispersion of GaN, due to the large mass ratio between Ga and N atoms, a large gap exists in phonon dispersion relations, which may reduce many BCPs.

For topological insulator states, two pairs of phonon degenerate phonon modes are needed at the same wave vector in both the bulk and surface Brillouin zones. However, only sixfold screw rotation symmetry exists, instead of sixfold rotation symmetry, which cannot support topological insulator states. To check Weyl phonon points in GaN, all phonon BCPs at high symmetry paths and planes (including planes $k_y=0$, $k_x=0.5$, $k_z=0$, and $k_x=0$ in Cartesian coordinates) in the Brillouin zone are searched by calculating topological charges and Berry curvature distribution. Since no topological characteristics of phonons are found after these
FIG. 11. Phonon dispersion of wurtzite AlGaN$_2$. (a) The phonon dispersion of AlGaN$_2$ with and without NAC. (b) The phonon dispersion obtained by standard phonon calculations and reproduced by Wannier tight-binding model Hamiltonian. (c) The phonon dispersion along the path including two different Weyl points marked as WP$_1$ and WP$_2$ is not affected by NAC. (d) and (e) are enlarged views of two Weyl points in (c), where the slight gap in the case without NAC results from the numerical errors and that the Weyl points are difficult to locate very accurately since they are not at high symmetry paths.
In this section, we further check GaN related structures, including wurtzite GaN with tensile strain along and perpendicular to the polar axis, III-A nitrides (AlN and BN), and AlGaN crystal. Discussion on AlGaN alloys can be seen in the supplementary material. It is found that single Weyl phonons exist in AlN and four types of AlGaN when Al atoms are introduced into GaN, while no Weyl phonon is found in other systems. Particularly, discussions on wurtzite AlN and wurtzite-like AlGaN (space group P3m1, No. 156) are more valuable since these two materials are discussed in Refs. 35 and 54, conclusions and inferences show that the wurtzite GaN is a trivial topological phonon system. Herein, taking the 14th BCP as an example, trivial surface states at the (0001) surface are presented and provide an intuitive understanding of the calculations, which can also be regarded as comparisons with the following nontrivial results. In electronic components, wurtzite nitrides are generally grown along the polar axis, i.e., the (0001) direction. Therefore, the (0001) surface is selected in this work for surface state calculations. As seen from Fig. 3, in the projection of the iso-frequency plane on the (0001) surface, instead of a surface arc of Weyl points, there are only uninterrupted annular shapes. The LDOS of the surface along high symmetry paths in the surface Brillouin zone is shown in Fig. 4. The surface states are calculated by both the iterative Green’s function method and directly solving the eigenstates equation of phonons of a slab structure. Both results are consistent. It is noted here that there are two kinds of (0001) surfaces, including a Ga surface and an N surface, which are equivalent to bottom and top surfaces, respectively. In the following discussions, the surfaces with significant topological characteristics are focused on while the surface type is not distinguished particularly.

B. Topological phonon phase transition by introducing Al

In Refs. 35 and 54, conclusions and inferences show that the Weyl complex and type II Weyl phonons can exist in the wurtzite structure (Space group P63mc, No. 186) at the high symmetry points Gamma (A) and K (H)54 and the high symmetry plane instead of the high symmetry path.54 Hence, topological phonon phase transitions may be realized in GaN of wurtzite structure as we have verified that there is no Weyl point in this system. The classical model in topological phonon theory, i.e., SSH-like models for phonons72-74 and the effective Hamiltonian model for topological insulator phonon states20 indicates that second-order IFCs are critical parameters in determining topological properties of the phonon system and the requirement for lattice symmetry is necessary but not sufficient.

In this section, we further check GaN related structures, including wurtzite GaN with tensile strain along and perpendicular to the polar axis, III-A nitrides (AlN and BN), and AlGaN crystal. Discussion on AlGaN alloys can be seen in the supplementary material. It is found that single Weyl phonons exist in AlN and four types of AlGaN when Al atoms are introduced into GaN, while no Weyl phonon is found in other systems. Particularly, discussions on wurtzite AlN and wurtzite-like AlGaN (space group P3m1, No. 156) are more valuable since these two materials are used in real electronics. The wurtzite-like AlGaN can be formed by replacing one Ga atom with one Al atom in wurtzite GaN. Therefore, in the following, we focus on wurtzite AlN and wurtzite-like AlGaN. For other AlGaN systems, we only introduce Al3GaN4 (space group P-43m, No. 215) as an example.

The phonon dispersions of AlN are shown in Fig. 5, where phonon dispersions of AlN along high symmetry paths are shown in Figs. 5(a) and 5(b). In Figs. 5(c)-5(e), nontrivial band crossing points with and without considering NAC are both presented, and the figures show that the crossing points are not affected by NAC. Slight frequency changes may occur, resulting in a small displacement of the Weyl point with NAC, explaining the slight gap in the NAC case. The systematic topological phonon analyses here are the same as those in Sec. III A. The positions of Weyl phonons in the Brillouin zone are labeled in Fig. 6. Nontrivial topological charges are calculated using the Wilson-loop method and confirmed by Wannier charge center evolution (Fig. 7). Besides, the Berry curvature distribution where the source and the sink are present at Weyl points with positive and negative chirality is shown in Fig. 8, in which one source and one sink are marked with red and blue dots, respectively. Surface arcs in the projection of the iso-frequency plane on the (0001) surfaces (Fig. 9) and topological surface states illustrated by surface LDOS at specific paths linking two Weyl phonons with opposite chirality (Fig. 10) provide solid evidence for nontrivial TPBs. Though there are rich BCPs along the high symmetry path, no Weyl point is found in them. Two different Weyl points (equivalent points are not counted here) in AlN exist in the high symmetry path, the kz=0 plane with kx=0. For the first kind of Weyl phonon, the points with positive chirality 1 and negative chirality −1 are (0.2989, 0.1249, 0.0000) and (0.4239, −0.1246, 0.0000) in fractional coordinates, and their respective equivalent points, which are all shown in Fig. 6. The fourth and fifth bands cross at this point with a frequency of 12.23 THz. For the second kind of Weyl point, the points with chirality 1 and −1 are (0.3549, −0.1169, 0.0000) and (0.3549, −0.2380, 0.0000), and their equivalent points in the Brillouin zone based on lattice symmetry. These Weyl points are formed by the crossing of the eighth
and ninth bands at a frequency of 19.57 THz. In Figs. 9(b) and 9(d), the projection of the iso-frequency plane at the (0001) surface is enlarged to clearly show the surface arcs linking a couple of Weyl phonons. The surface states for the second kind of Weyl point are not as clean as those in the first kind, as there are also trivial surface states near this point with a frequency around 19.57 THz, i.e., the lower single line below 19.6 THz in Fig. 10(b).

$\text{Al}_x\text{Ga}_{1-x}\text{N}$ is the most common alloy of GaN, an important material in the AlGaN/GaN heterostructure. Experimentally, the proportions of Al and Ga atoms can be controlled. However, it may not be easy to calculate phonon properties from first principles for structures with a large unit cell or real alloy structures. In the public material database, there are seven types of AlGaN crystal structures, including structures with space group P3m1 (No. 156), P-4m2 (No. 115), and three other structures with very low symmetry. AlGaN with space group P3m1 is a structure corresponding to wurtzite GaN and more similar to $\text{Al}_x\text{Ga}_{1-x}\text{N}$ used in the AlGaN/GaN heterostructure. Here, the topological phonon properties of this AlGaN crystal are discussed.

Phonon dispersions of AlGaN are shown in Fig. 11, where phonon dispersions along high symmetry paths are shown in Figs. 11(a) and 11(b). There are 12 bands since there are four atoms in each unit cell. The calculations show that there are also two different single Weyl points in AlGaN, which are located at a high symmetry plane, the $k_x-k_y$ plane with $k_z=0$. The phonon dispersions along the path across the two kinds of Weyl points are also shown in Fig. 11(c). The first kind of Weyl point is formed by the seventh and eighth bands at (0.4225, 0.0776, 0.0000), and P-4m2 (No. 115), and three other structures with very low symmetry. AlGaN with space group P3m1 is a structure corresponding to wurtzite GaN and more similar to $\text{Al}_x\text{Ga}_{1-x}\text{N}$ used in the AlGaN/GaN heterostructure. Here, the topological phonon properties of this AlGaN crystal are discussed.

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Phonon dispersions of AlGaN are shown in Fig. 11, where phonon dispersions along high symmetry paths are shown in Figs. 11(a) and 11(b). There are 12 bands since there are four atoms in each unit cell. The calculations show that there are also two different single Weyl points in AlGaN, which are located at a high symmetry plane, the $k_x-k_y$ plane with $k_z=0$. The phonon dispersions along the path across the two kinds of Weyl points are also shown in Fig. 11(c). The first kind of Weyl point is formed by the seventh and eighth bands at (0.4225, 0.0776, 0.0000), and P-4m2 (No. 115), and three other structures with very low symmetry. AlGaN with space group P3m1 is a structure corresponding to wurtzite GaN and more similar to $\text{Al}_x\text{Ga}_{1-x}\text{N}$ used in the AlGaN/GaN heterostructure. Here, the topological phonon properties of this AlGaN crystal are discussed.

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0.0000) in fractional coordinates in the Brillouin zone at a frequency of 17.41 THz. And, the second one is formed by the crossing of the 9th and 10th bands at (0.3097, −0.1668, 0.0000) at a frequency of 18.79 THz. The positions of Weyl points in the Brillouin zone are illustrated in Fig. 12. Nontrivial Wannier charge center evolutions and Berry curvature distribution are illustrated in Figs. 13 and 14. Unlike Weyl points in AlN, a Weyl point of the first kind is linked by a surface arc with the other one outside the first Brillouin zone, as shown in Figs. 12 and 15(a) and 15(b). Actually, the equivalence of Weyl points in AlGaN is conserved by lattice symmetry and time-reversal invariance symmetry. In Fig. 15, distinct surface arcs between Weyl points are illustrated for both kinds of Weyl phonons. Also, both the surface states are distinguishable as that there is no other surface state near frequency 17.41 THz and 18.79 THz (Fig. 16).

Besides in AlGaN with space group P3m1, Weyl points are also detected in other AlGaN structures. However, these structures may not be practical for applications in electronics. Therefore, one Weyl point in Al$_3$GaN$_4$ with space group P-43m (No. 215) is introduced here as a representative. Al$_3$GaN$_4$ is a cubic structure, where eight atoms including three Al, one Ga, and four N consist of one primitive unit cell. This structure is similar to the zincblende structure, as three Al atoms replace three Ga atoms in zincblende GaN.
FIG. 15. Weyl surface arcs. The projection of the iso-frequency plane of surface states on the (0001) surface at the frequency (a) 17.41 THz and (b) 18.79 THz. The red and blue dots represent Weyl points with positive chirality 1 and negative chirality $-1$, respectively, which are connected by a phonon surface arc. The black hexagon indicates the surface Brillouin zone of the (0001) surface in wurtzite AlGaN$_2$. (b) and (d) are enlarged views of the zones encircled by blue rectangles in (a) and (c), respectively.

FIG. 16. The LDOS for the (0001) surface along the selected path linked a couple of Weyl points. (a) The first kind of Weyl points and (b) the second kind of Weyl points in AlGaN$_2$. The single line between a couple of Weyl points indicates the topological surface states, where $P_1$ and $P_2$ are on the extension line connecting two Weyl points.
conventional unit cell. The Brillouin zone \( \text{Al}_3\text{Ga}_4 \) is also cubic. The phonon dispersions along high symmetry paths are shown in Fig. 17.

The calculations show that several single Weyl phonons exist in the \( k_z = 0 \) and \( k_z = 0.5 \) planes of the Brillouin zone. The Weyl point discussed here is formed by the crossing of the sixth and seventh bands at wave vector \((0.0941, -0.5000, 0.0000)\) at a frequency of 9.82 THz, located at the high symmetry path X-M, as illustrated in Fig. 18. The positions of the Weyl point and its equivalent point are shown in Fig. 18 in the \( k_z = 0 \) plane. The characteristics of the Weyl phonons are first verified by nontrivial Wannier charge center evolutions and Berry curvature distribution, as illustrated in Figs. 19 and 20. It is noted here this kind of Weyl point also exists in the \( k_x = 0 \) and \( k_y = 0 \) planes based on the lattice symmetry. In Figs. 21 and 22, surface arcs and topological surface states can be observed where the surface arcs are relatively long since Weyl points with opposite chirality are far apart. However, surface states in this structure on the (0001) surface are ample, resulting in the overlap of trivial and nontrivial topological surface states. Particularly, it is noted that this kind of Weyl

**FIG. 17.** Phonon dispersion of wurtzite \( \text{Al}_3\text{Ga}_4 \). (a) The phonon dispersion of \( \text{Al}_3\text{Ga}_4 \) with and without NAC. (b) The phonon dispersion obtained by standard phonon calculations and reproduced by Wannier tight-binding model Hamiltonian. (c) The phonon dispersion along the path including the Weyl point marked as WP1 is affected by NAC. (d) The enlarged view of the Weyl point in (c), which indicates that the Weyl point disappears with NAC.

**FIG. 18.** Weyl points in the Brillouin zone at plane \( k_z = 0 \) are shown by the black rectangle. The dots represent the Weyl point shown in Fig. 17(c) and its equivalent points. The dots in red are Weyl points with positive chirality 1, while those in blue are Weyl points with negative chirality −1. Each Weyl point is coupled with the nearest one with opposite chirality.
point in Al$_3$GaN$_4$ may not exist in reality, since the Weyl points disappear after considering NAC, which is clearly shown in Fig. 17(d). After including NAC in phonon dispersion calculations, the band crossing point in the corresponding position disappears. Though generally NAC mainly affects the phonon dispersion near the Brillouin zone, it also affects the phonon dispersion at the Brillouin zone boundary for Al$_3$GaN$_4$. As seen from Fig. 17(c), this band crossing point in the case without NAC may move to high symmetry point X after considering NAC, and the Weyl phonon state may still be preserved, which can be confirmed when calculation methods are available for cases with NAC.
IV. CONCLUSIONS AND PERSPECTIVES

In summary, we performed systematical analyses on the topological effects of phonons in wurtzite GaN with the aid of first-principle calculations and topological phonon theory by concentrating on Weyl point states and discussed the topological phonon phase transition in generalized GaN systems by introducing Al atoms, including AlN and AlGaN. The results show that wurtzite GaN is a trivial topological phonon system where no topological insulator phonon state and Weyl phonon state is found. However, by introducing Al atoms, Weyl phonon states are found in AlN and AlGaN of different structures, and they can be regarded as a topological phonon phase transition in AlGaN systems. By calculating topological charges, Wannier charge center evolutions, Berry curvature distributions, the projection of iso-frequency plane on the surface, and surface LDOS, two kinds of single Weyl phonon states in AlN, two kinds of single Weyl phonon states in AlGaN2 with space group P3m1, and one single Weyl phonon state in cubic Al2GaN4 (which may not exist in reality due to NAC) are confirmed.

Topological effects of phonons affect phonon transport by inducing topological surface phonon states with novel transport characteristics that are robust and protected by the topological properties of the bulk system. Since Weyl phonon surface states are present in the gaps of trivial surface states, which can be regarded as isolated and bound states, and only half of the nontrivial states exist in opposite surfaces (arc states), scatterings of topological surface phonon states promise to be reduced significantly, especially backscattering. As typical materials in AlGaN/GaN heterostructures, wurtzite AlN and AlGaN2 with space group P3m1 promise to host topological surface states without backscattering as they are confirmed to be topological phonon nontrivial systems with open Weyl surface arcs. Future research could directly examine these surface/interface states and the scattering characteristics between topological surface phonon states and trivial surface phonon states by molecular dynamic simulations. While AlN’s thermal conductivity is relatively high, the contribution from topological surface phonon states may be insignificant. However, as mentioned in Sec. I, interfacial thermal transport parallel to the AlGaN/GaN interface is critical for the spreading of heat generated from the near-junction region. Hence, through effective design, the excellent transport properties of Weyl phonon states promise to improve the thermal management in information and power electronics.

SUPPLEMENTARY MATERIAL

See the supplementary material for discussions on the identification of potential topological phonons and AlGaN alloys.

ACKNOWLEDGMENTS

We appreciated the help from Dr. Yizhou Liu (Weizmann Institute of Science, Israel) and Professor Rui Wang (Chongqing University, China) in understanding the topological physics of phonons. This work was supported by the National Natural Science Foundation of China (NNSFC) (Grant Nos. 51825601 and U20A20301).

DATA AVAILABILITY

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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