An *ab initio* Study on the Mechanical Stability, Spin-Dependent Electronic Properties, Molecular Orbital Predictions, and Optical Features of Antiperovskite A₃InN (A = Co, Ni)

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ABSTRACT: Structural, mechanical, spin-dependent electronic, magnetic, and optical properties of antiperovskite nitrides A₃InN (A = Co, Ni) along with molecular orbital diagram are investigated here by using an *ab initio* density functional theory (DFT). The mechanical stability, deformation, damage tolerance and ductile nature of A₃InN are confirmed from elastic calculations. Different mechanical anisotropy factors are also discussed in detail. The spin dependent electronic properties such as the band structure and density of states (DOS) of A₃InN are studied and, the dispersion curves and DOS at Fermi level are different for up and down spins only in case of Co₃InN. These calculations also suggest that Co₃InN and Ni₃InN behave as ferromagnetic and nonmagnetic, respectively. The induced total magnetic moment of Co₃InN is found 2.735 μB/cell in our calculation. Mulliken bond population analysis shows that the atomic bonds of A₃InN are contributed by both ionic and covalent bonds. Molecular orbital diagrams of A₃InN antiperovskites are proposed by analyzing orbital projected band structures. The formation of a molecular orbital energy diagram for Co₃InN is similar to Ni₃InN with respect to hybridization and orbital sequencing. However, the various ground-state optical functions such as real and imaginary parts of the dielectric constant, optical conductivity, reflectivity, refractive index, absorption coefficient, and loss function of A₃InN are studied with implications. The reflectivity spectra reveal that A₃InN reflects >45% of incident electromagnetic radiations in both the visible and ultraviolet region, which is an ideal feature of coating material for avoiding solar heating.

1. INTRODUCTION

Antiperovskite transition-metal nitrides (ATMNs) have become familiar to scientists and researchers in the last several decades.¹⁻³ ATMNs demonstrate a broad range of attractive and tunable physical properties such as the Invar-like effect, itinerant antiferromagnetism, giant magneto-resistance, superconductivity, damage tolerance, spin-glass-like activities, strong spin–lattice coupling characteristics, and topological electronic behavior.¹⁻⁴⁻⁸ Perovskite compounds are denoted by the formula ABX₃, where A and B are cations and X is an anion. In antiperovskite compounds, the general formula is reversed, so that the X sites are occupied by a cation, while A and B sites are occupied by different types of anions. Typical ATMNs adopt the crystal structure AₓBX with space group Pnma (no. 221), where A is a transition metal; B is a divalent or trivalent element; and X is nitrogen.⁹⁻¹¹ B. V. Beznosikov predicted more than 80 nitride compounds with antiperovskite structure.⁹ The generation of large negative thermal expansion in Ge-doped antiperovskite manganese nitrides MnₓXN (X = Cu, Zn, Ga) is reported by Takagi et al.¹⁰ Ferro- and paramagnetic orders are observed in ATMNs when the transition elements are substituted by group 13 metals.¹¹,¹² It is reported that the nonstoichiometry affects the magnetic properties in Ni-rich antiperovskite carbides depending on Ni/C atomic ratios.¹³⁻¹⁵ However, the nonmagnetic ground state is found in many cases for Ni-based ternary carbides due to the reduced Stoner factor and C–Ni bonding nature.¹⁶⁻¹⁸ Although there are predictions about some highly stable two-dimensional (2D) hypercoordinate materials such as Cu–Si and Cu–Ge alloy films, NiₓGe and NiₓSi monolayers, aluminum boride (AlB₆) nanosheets with interest-
ing physical properties, and diverse applications,\textsuperscript{19–26} here we consider only the bulk antiperovskite $A_3$InN in our calculation.

The nickel (Ni)-based antiperovskite nitrides gained considerable interest due to the discovery of superconductivity in Ni$_5$CuN and Ni$_5$ZnN with $T_c$ around 3 K.\textsuperscript{27,28} Ni$_4$CdN exhibits very soft and weak ferromagnetism, and Ni$_4$InN exhibits a spin–glass-like behavior.\textsuperscript{20,30} Different groups synthesized Co$_3$InN and Ni$_3$InN by different methods.\textsuperscript{1–4} Due to the diversity of the physical properties of the antiperovskite materials and limited knowledge on the cobalt- and Ni-based nitrides, first-principle calculations for Co$_3$InN and Ni$_3$InN were carried out in order to study Young’s moduli, Poisson’s ratio, and spin-polarized band structure and projected density of states (pDOS) only.\textsuperscript{3} To extend these previous works, we thus focus here on exposing different physical properties of Co- and Ni-based antiperovskite $A_3$InN ($A = \text{Co, Ni}$) nitrides using an \textit{ab initio} density functional theory (DFT) method. The spin-polarized structural, elastic, and mechanical stability and the effect of spin in electronic behavior, magnetic nature, bonding character, and optical properties of antiperovskites $A_3$InN are elucidated in detail. In addition, to expose the strength and extent of anisotropy of $A_3$InN, we focused on the Vickers hardness and different anisotropy indices, respectively. The anisotropic mechanical behavior and elastic responses of $A_3$InN are presented by two- and three-dimensional graphical presentations. To take full advantage of the electronic properties of $A_3$InN in addition to spin-polarized bands and pDOS, we represent here the results of magnetic moments (total and individual atoms), charge density distribution mapping, and spin-polarized Fermi surface topologies. The Fermi surface can provide information for predicting the physical properties of a metal.

The molecular orbital (MO) theory is regarded as an effective tool to determine the molecular structure and bonding nature of a compound. It takes the idea of overlapping between different atomic orbitals. The distribution of different MO energy levels is represented by a MO diagram. It also gives insight about bonding, bond order, and change of molecular behavior with ionization. A lot of reports are published on the MO diagram for ternary $X_2YZ$ Heusler compounds,\textsuperscript{31,32} where $X$ and $Y$ are transition metals and $Z$ is a main group element. Compared to $X_2YZ$ Heusler compounds, the study of $d$-orbital hybridization and bonding order for $A_3BX$-based antiperovskite has not been explored yet to the best of our knowledge.

Moreover, we predicted the orbital hybridization among different atoms based on orbital-projected band diagrams and unfolding orbital degeneracy of $A_3$InN which may help to better understand the magnetic behavior of these compounds.

Finally, the interaction of $A_3$InN with electro-magnetic radiation is expressed by means of some nonlinear optical constants such as reflectance, absorption coefficient, refractive index, complex dielectric constants, optical conductivity, and loss function. As far as we know, there are no experimental or theoretical data available on the Vickers hardness, anisotropic elastic indices, Fermi surface topology, and nonlinear optical properties of $A_3$InN, which are focused here. We strongly believe this study will be useful for using $A_3$InN in the arena of engineering applications.

2. METHODS OF CALCULATIONS

The ground-state physical properties of the $A_3$InN ($A = \text{Co and Ni}$) are simulated using \textit{ab initio} density functional theory (DFT) via the CASTEP code.\textsuperscript{35} The Vanderbilt-type ultrasoft pseudopotential method is employed to treat the outermost electrons as valence electrons for each atom in $A_3$InN. The Perdew–Burke–Ernzerhof (PBE) parametrization within the generalized gradient approximation (GGA) is applied to treat the exchange and correlation interactions.\textsuperscript{34} The plane-wave cutoff energy of 500 eV is used to expand the Eigen functions of the valence and nearly valence electrons. The Monkhorst–Pack grid of $18 \times 18 \times 18$ $k$-points is used to investigate the Brillouin zone. The Broyden–Fletcher–Goldfarb–Shanno (BFGS) algorithm is employed to optimize the geometry optimization through minimizing the total energy and the internal forces.\textsuperscript{35} To optimize the geometry conditions, the tolerances for total energy, maximum force, maximum stress, and maximum atomic displacement are set to less than $5 \times 10^{-6}$ eV per atom, $0.01$ eV/Å, $0.02$ GPa, and $5 \times 10^{-3}$ Å, respectively. The single-crystal elastic tensors ($C_{ij}$) are calculated by the “stress–strain” method embodied into the CASTEP code in which a set of finite identical deformations is applied. The resultant stress is calculated after optimization of the internal degrees of freedom, and a maximum amplitude of 0.003 Å is chosen for each strain having four steps. The atomic population analysis is performed by means of the conventional Mulliken formalism. The electron charge density difference and Fermi surface are calculated by setting the $k$-point separation to less than $0.01$ Å$^{-1}$ with $26 \times 26 \times 26$ grids. The optical properties of $A_3$InN are calculated using the same code. It is important to note that spin polarization is considered in all calculations. Orbital-projected band diagrams of these compounds $A_3$InN ($A = \text{Co and Ni}$) are also evaluated using the Quantum ESPRESSO\textsuperscript{36} code with the above motioned parameters.

3. RESULTS AND DISCUSSION

3.1. Structural Properties. Bulk $A_3$InN ($A = \text{Co and Ni}$) is crystalized as a cubic system in space group $Pm3m$ (no. 221).\textsuperscript{3} The crystal structure of antiperovskite $A_3$InN nitrides in the above-mentioned space group is shown in Figure 1. The structural unit cell of $A_3$InN, which adopts an octahedral coordinate, consists of six $A$ atoms, one In atom, and one N atom. The Wyckoff positions are at $(0, 1/2, 1/2)$, $(0, 0, 0)$, and $(1/2, 1/2, 1/2)$ for $A$, In, and N atoms, respectively. The equilibrium crystal structures of $A_3$InN are optimized at the minimum total energy. The calculated lattice parameter and unit cell volume of $A_3$InN are tabulated in Table 1. It is observed that the lattice constants of $A_3$InN which are very close ($<1.07\%$)
compared to the experimental values\(^1,4\) indicate the highest level of accuracy of our study.

Table 1. Comparison between the Theoretically Calculated and Experimental Values of Unit Cell Parameters of A\(_3\)InN (A = Co, Ni)\(^4\)

| phase    | a (Å) | V (Å\(^3\)) | method               | reference          |
|----------|-------|-------------|----------------------|---------------------|
| Co\(_3\)InN | 3.877 | 58.287      | DFT (GGA)            | this work           |
| Ni\(_3\)InN | 3.901 | 59.361      | DFT (GGA)            | this work           |
| Co\(_3\)InN | 3.855 | 57.289      | DFT (GGA)            | 3                   |
| Ni\(_3\)InN | 3.852 | 57.286      | DFT (GGA)            | 3                   |
| Ni\(_3\)InN | 3.882 | 58.501      | DFT (GGA)            | 3                   |
| Ni\(_3\)InN | 3.784 | 54.181      | DFT (LDA)            | FM                  |
| Co\(_3\)InN | 3.8518| 57.146      | electrochemical      | 1                   |
| Ni\(_3\)InN | 3.8599| 57.509      | electrochemical      | 1                   |
| Co\(_3\)InN | 3.8541| 57.249      | solid–gas reactions  | 4                   |
| Ni\(_3\)InN | 3.8445| 56.822      | solid–gas reactions  | 4                   |

\(^4\)GGA = generalized gradient approximation. LDA = local density approximation. FM = ferromagnetic. NM = Nonmagnetic.

3.2. Mechanical Properties. The mechanical properties are important because they describe different characteristics such as elasticity, plasticity, strength, hardness, ductility, and brittleness of a material. Among these properties, elastic constants correlate the dynamical behavior of a material with its mechanical properties. The single-crystal elastic tensors (\(C_{ij}\)) of newly synthesized A\(_3\)InN (A = Co, Ni) are calculated to provide a deep insight into the mechanical stability and stiffness of these materials. In general, materials with cubic symmetry have three elastic constants, namely, \(C_{11}\), \(C_{12}\), and \(C_{44}\). The constant \(C_{11}\) is related to tensile, whereas \(C_{12}\) and \(C_{44}\) are associated to shear. The values of \(C_{11}\), \(C_{12}\), and \(C_{44}\) are listed in Table 2. The elastic constants of A\(_3\)InN satisfy the Born stability criteria \(C_{11} > 0, C_{11} - C_{12} > 0, C_{44} > 0\), and \((C_{11} + 2C_{12}) > 0\) signifies the mechanical stability for these cubic crystals. It has also been seen that the constants follow \(C_{11} > C_{12} > C_{44}\) which indicates the anisotropic nature, whereas the low value of \(C_{44}\) is allied to the shear deformation and damage-tolerant nature of A\(_3\)InN. The different elastic moduli (\(B\), \(G\), \(Y\), and \(v\)) are also calculated from individual elastic constants. The bulk modulus \((B)\) of A\(_3\)InN is given by

\[
B = \frac{C_{11} + 2C_{12}}{3}
\]

(1)

The shear modulus \((G)\) of a crystalline solid varies with direction and is defined by the Voigt–Reuss–Hill (VRH) equation as

\[
G_V = \frac{1}{5}(C_{11} - C_{12} + 3C_{44})
\]

(2)

\[
G_R = \frac{5(C_{11} - C_{12})C_{44}}{4C_{44} + 3(C_{11} - C_{12})}
\]

(3)

where \(G_V\) and \(G_R\) are the Voigt and Reuss bonds, respectively. The arithmetic average of \(G_V\) and \(G_R\) gives the shear modulus of A\(_3\)InN.

\[
G = \frac{G_V + G_R}{2}
\]

(4)

The isotropic polycrystalline aggregate values for Young’s modulus \((E)\) and Poisson’s ratio \((\nu)\) are obtained using the following equations as

\[
Y = \frac{9GB}{(G + 3B)}
\]

(5)

\[
\nu = \frac{3B - 2G}{2(3B + G)}
\]

(6)

The values of \(B\), \(G\), \(Y\), and \(\nu\) are tabulated in Table 2. In general, the hardness of a compound is measured by \(B\) and \(G\). The values of \(B\) (\(G\)) are found to be 197 (93) GPa and 182 (63) GPa for Co\(_3\)InN and Ni\(_3\)InN, respectively, indicating higher hardness of Co\(_3\)InN. Moreover, the high value of the bulk modulus with low shear modulus reveals the damage-tolerant, quasi-ductile, easily machinable, and stiff nature of A\(_3\)InN. On the other hand, Co\(_3\)InN is found to be stiffer than Ni\(_3\)InN due to the higher value of \(Y\). Another three important factors, Poisson’s ratio \((\nu)\), Pugh’s ratio \((G/B)\), and Cauchy pressure \((C_{12} - C_{44})\), have also been calculated and shown in Table 2. From Table 2, it is seen that \(\nu\) is greater than Frantsevich’s criterion\(^39\) value of 0.26 and \(G/B < 0.57\). These parameters suggest that A\(_3\)InN is ductile. The elastic nature of A\(_3\)InN is also verified by the Cauchy pressure.\(^39\) The negative value of Cauchy pressure indicates the brittle nature of the compound, while a positive value endorses the quasi-ductile nature. In addition, the values of \(B\), \(G\), \(Y\), and \(G/B\), and \((C_{12} - C_{44})\) are compared with other antiperovskites and found to be consistent, which is shown in Table 2.

The Vickers hardness \((H_V)\) is highly related to the elastic constants of a material. We have calculated the \(H_V\) of A\(_3\)InN using the different approximations proposed by Teter et al.,\(^40\) \((H_V)_{Teter} = 0.151\) G, Tian et al.,\(^41\) \((H_V)_{Tian} = 0.92(G/B)^{0.3585}\), and Chen et al.,\(^42\) \((H_V)_{Chen} = 2\left((G/B)^2\right)^{0.3585}\). The calculated values of \(H_V\) are presented in Table 3. It is noticeably seen that Co\(_3\)InN has higher \(H_V\) than Ni\(_3\)InN which may be due to a higher elastic modulus and bond strength.

Mechanical anisotropy is closely related to different essential physical processes of a material. To visualize the level of anisotropic mechanical behavior and elastic responses of A\(_3\)InN, the Young’s modulus \((Y)\), linear compressibility \((\beta)\), shear modulus \((G)\), and Poisson’s ratio \((\nu)\) are plotted in contour plots.

Table 2. Comparison of Elastic Properties between A\(_3\)InN (A = Co, Ni) and Other Antiperovskites

| compound | \(C_{11}\) | \(C_{12}\) | \(C_{44}\) | \(B\) | \(G\) | \(Y\) | \(\nu\) | \(G/B\) | \(C_{12} - C_{44}\) | ref |
|----------|------------|------------|------------|-----|-----|-----|-----|-----|-------------|-----|
| Co\(_3\)InN | 333 | 129 | 87 | 197 | 93 | 241 | 0.29 | 0.472 | 42 | this work |
| Co\(_3\)InN | 317 | 126 | 95 | 190 | 95 | 224 | 0.28 | 0.50 | 31 | 3\(^a\) |
| Ni\(_3\)InN | 283 | 131 | 55 | 182 | 63 | 170 | 0.34 | 0.346 | 76 | this work |
| Ni\(_3\)InN | 274 | 131 | 60 | 178 | 64 | 172 | 0.34 | 0.359 | 71 | 3\(^a\) |
| Ti\(_3\)TiN | 196 | 131 | 52 | 153 | 43 | 118 | 0.40 | 0.276 | 79 | 8\(^a\) |
| Ni\(_3\)SnN | 266 | 132 | 41 | 177 | 50 | 137 | 0.37 | 0.282 | 91 | 8\(^a\) |
| Co\(_3\)AlC | 451 | 119 | 86 | 230 | 112 | 290 | 0.29 | 0.487 | 38 | 3\(^a\) |

\(^a\)Theoretical.
of three- (3D) and two-dimensional (2D) presentations (Figures 2 and 3) using the ELATE code. From Figures 2 and 3, it is demonstrated that all the constants except linear compressibility are deviated from the complete spherical (3D) and circular (2D) shapes, indicating the anisotropic nature of these compounds. However, the deviation from the spherical and circular shapes is the same in all directions (xy, xz, and yz planes) but a bit higher for Co3InN compared to Ni3InN. The maximum and minimum values of Y, β, G, and ν are enlisted in Table 4.

The elastic anisotropy of cubic A3InN can be expressed using the Zener anisotropy factor46 A2 as

\[ A_2 = \frac{2G(1 + \nu)}{Y} = \frac{2C_{44}}{C_{11} - C_{12}} \]  

(7)

The Zener anisotropy factor is related to share anisotropy, and A2 = 1 for a perfect elastically isotropic material.

An alternative measure of the elastic anisotropy (A2) for the cubic compound is proposed by Chung and Buessem47 as

\[ A_G = \frac{3(A_2 - 1)^2}{3(A_2 - 1)^2 + 25A_L} \]  

(8)

The anisotropy factor A2 is expressed in percentage, and A2 = 0 indicates the isotropic nature of a material.

The universal elastic anisotropy index (A2)48 involves all the individual components of the elasticity tensor of a material which may be defined for a cubic crystal as

\[ A_U = 5\frac{G_{V}}{G_{R}} - 5 \]  

(9)

The logarithmic universal Euclidean anisotropy (A2)49 for a cubic compound is related to the Voigt and Reuss bounds on the bulk modulus simplifies as

\[ A_L = \sqrt{3} \ln \left( \frac{G_{V}}{G_{R}} \right) \]  

(10)

The orientation of Poisson’s ratio is specified in two directions; the lateral contraction/extension in one direction and the corresponding extension/contraction in a normal direction. In general, the extreme values of Poisson’s ratio may occur due to strain along (110) and the corresponding orthogonal strain along (001) and (110) for a cubic compound.50 The extrema of Poisson’s ratio in these particular directions is expressed as

\[ \nu(110, 001) = \frac{2A_2C_{12}}{3B + A_2C_{11}} \]  

(11)

The values of A2, A2, and A2, \( \nu(110, 001) \) and \( \nu(110, 1\bar{1}0) \), are listed in Table 5. From the table it is clear that the values of different anisotropy parameters except A2 are higher for Co3InN than Ni3InN. The zero value of A2 and A2 signifies the higher isotropic nature of Co3InN than Ni3InN. However, these anisotropy parameters confirm that Co3InN is more anisotropic in compression, whereas Ni3InN is more anisotropic in shear. The extrema of Poisson’s ratio is found, \( \nu(110, 1\bar{1}0) > \nu(110, 001) \), which reveals that the orthogonal strain may occur along this particular direction for A3InN.

3.3. Band Structure and Density of States (DOS). The electronic band structure and density of states (DOS) play an imperative role to visualize several optoelectrical and magnetic properties of a crystal at the microscopic level. The spin-polarized electronic band structures of A3InN (A = Co, Ni) are calculated for spin-up (↑) and spin-down (↓) using high-symmetry k-points along the path X–R–M–G–R in the first Brillouin zone. The results are displayed in Figure 4(a)–(c).

From Figure 4, it is seen that the dispersion curves for up and down spins are different for Co3InN. Two bands for up and two bands for down spin cross the E_F which is shown in Figures 4(a) and 4(b). On the contrary, the dispersion curves for both spins are the same for Ni3InN. Two bands cross the E_F, as seen in Figure 4(c). The dispersion curve for down spin is not shown due to the spin-symmetric nature of Ni3InN. Both Co3InN and Ni3InN exhibit metallic nature, as the conduction and valence bands notably overlap at the Fermi level.

To explore the origin of metallic and magnetic nature, the spin-polarized total density of states (DOS) and orbital-resolved partial DOS (pDOS) are calculated for A3InN. The spin-dependent DOS and pDOS of Co3InN are shown in Figure 5. It is clearly seen that the DOS at the Fermi level is nonzero for both spin channels [Figure 5(a)], revealing the metallic nature of Co3InN with zero energy gap. The observed DOS values are found at 1.176 states/eV and ~2.893 states/eV at E_F for the spin-up and spin-down channel, respectively. Thus, the DOS of Co3InN is contributed by both spins but significantly ~2.5 times higher for the spin-down channel. The contribution of individual atoms in Co3InN that occupied the electronic states per unit energy is determined by the pDOS shown in Figure 5(b)–(d).

It is seen that the valence band of Co3InN is roughly separated into three sub-bands, ranging from (~9.1 to ~5) eV, (~5 to ~0.5) eV, and (~0.5 to E_F) eV, showing some sharp peaks. These peaks arise due to the hybridization of different electronic states. The lower energy band (~9.1 to ~5) eV is contributed by Co-3d, Co-3p, In-5s, In-5p, and N-2p orbitals, whereas the rest of the band is donated by Co-3d, In-5p, and N-2p orbitals. The electrons in the Co-3d orbital contribute strongly in the conduction mechanism of the Co3InN phase for both spin channels. However, the contribution of Co-3p, Co-4s, In-5s, In-5p, and N-2p orbitals is minor. In addition, the DOS and pDOS peaks are seen to be blue-shifted for the spin-down channel, which indicates the presence of strong spin–orbit coupling (SOC) in Co3InN.

The spin-dependent DOS of Ni3InN is shown in Figure 6(a)–(d). The nonzero values of DOS [Figure 6(a)] for both spin channels indicate the metallic nature of Ni3InN. It is notable that the DOS and pDOS patterns of Ni3InN are identical for up and down channels in contrast to the case of Co3InN.
The calculated value of the DOS at the Fermi level is 1.04 states/eV for both spins in Ni$_3$InN. The DOS of Ni$_3$InN is mainly contributed by Ni-3d orbitals, whereas the Ni-3p, In-5s, In-5p, and N-2p orbitals contribute faintly as seen from Figure 6(b). The valence band of Ni$_3$InN is roughly separated into three sub-bands as Co$_3$InN. The lower (~9.1 to -4.80) eV, middle (~4.80 to -0.792) eV, and top sub-bands that cross $E_F$ result from the hybridization of Ni-3d, Ni-3p, In-5s, and N-2p orbitals, Ni-3d, Ni-3p, In-5p, and N-2p orbitals, and Ni-3d, Ni-3p, In-5s, In-5p, and N-2p orbitals, respectively. From the pDOS [Figures 5(c) and 6(c)], it is clearly seen that the contribution of an In atom is feeble at $E_F$, which indicates the bare minimum involvement of the In atom in metal-like conductions of both Co$_3$InN and Ni$_3$InN.}

### 3.4. Magnetic Properties and Molecular Orbital Analysis

The magnetic moments for total and individual...
atoms are calculated from the integrated DOS at the Fermi level. The magnetic moment in Co$_3$InN may arise due to shifts of the pDOS for spin-up and -down channels. The induced total magnetic moment of Co$_3$InN is 2.735 $\mu_B$/cell, which is contributed by Co, N, and In atoms. The local magnetic moment is calculated as 0.9163 $\mu_B$, 0.0919 $\mu_B$, and $-0.1651$ $\mu_B$ for Co, N, and In atoms, respectively. On the contrary, the total magnetic moment for Ni$_3$InN is calculated as $0.526 \times 10^{-8}$ $\mu_B$/cell, which is negligible. It is noteworthy that we have also considered the magnetic unit cell with both ferromagnetic and antiferromagnetic configurations.

Figure 3. Three and two-dimensional contour diagrams of (a) Young’s modulus ($Y$), (b) linear compressibility ($\beta$), (c) shear modulus ($G$), and (d) Poisson’s ($\nu$) ratio for Ni$_3$InN antiperovskite.
antiferromagnetic ordering of Co(Ni) atoms to find out the nature of A3InN. Here it should be mentioned that a single unit cell of A3InN (cf. Figure 1) is enough for both ferromagnetic (where all A atoms are in the same spin orientation) and antiferromagnetic settings (A atoms in alternate layers are with opposite spin orientations) and considered as a magnetic unit cell. Our calculations suggest that Co3InN (Ni 3InN) is ferromagnetic (nonmagnetic) in nature.

In order to further understand the magnetic nature and the contribution of s, p, and d orbitals fully in the band structure of A3InN, we pictured the orbital hybridization among different atoms based on orbital-projected band diagrams. Comparing Figures 5(b) and 6(b), it is clear that for empty states in the pDOS Co-3d states have a significant contribution only for the down-spin channel but a nominal contribution for the up spin of Co-3d and for both spin channels of Ni-3d. These findings of pDOS characteristics are similar to an earlier study. For the next part, we consider only the down-spin channel.

Orbital-projected band diagrams along Γ−X−R−M−Γ−R directions for Co3InN and Ni3InN are evaluated using the Table 4. Minimum and Maximum Limit of Y, β, G, and ν for A3InN (A = Co, Ni)

| compound   | Young’s modulus (GPa) | linear compressibility (TPa−1) | shear modulus (GPa) | Poisson’s ratio |
|------------|------------------------|--------------------------------|---------------------|-----------------|
| Y max      | Y min                  | β max                          | G max              | ν max           | ν min           | ref     |
| Co3InN     | 228.79                 | 260.99                         | 1.6893             | 1.6893          | 87.543          | 101.99  | 0.25286          | 0.34833 | this work |
| Ni3InN     | 151.05                 | 200.93                         | 1.8309             | 1.8309          | 55.462          | 76.338  | 0.25331          | 0.45183 | this work |
| Ti2TIN     | 91.04                  | 140.09                         | 1.8309             | 1.8309          | 41              | 67      | 0.23              | 0.53    | 8       |
| Ni5SnN     | 114.17                 | 178.44                         | 1.8309             | 1.8309          | 86              | 166     | 0.13              | 0.49    | 8       |
| Co2AlC     | 229.37                 | 401.31                         | 1.8309             | 1.8309          | 83              | 83      | 0.25              | 0.34    | 43      |
| Ti5AlC     | 151                    | 151                            | 1.8309             | 1.8309          | 58              | 58      | 0.3               | 0.3     | 45      |

The results of other antiperovskite compounds are listed for comparison.

Table 5. Anisotropy Parameters and Extrema of Poisson’s Ratio for A3InN (A = Co, Ni)

| compound   | A_Z | A_G | A_U | A_L | ν(110, 001) | ν(110, 110) | ref     |
|------------|-----|-----|-----|-----|-------------|-------------|---------|
| Co3InN     | 0.85| 0.003| 0.00 | 0.00 | 0.250       | 0.352       | this work |
| Ni3InN     | 0.72| 0.129| 0.161| 0.071| 0.247       | 0.456       | this work |
| Ti2TIN     | 0.015| 0.015| 0.015| 0.015| 0.247       | 0.456       | 51      |
| Fe2AlC     | 0.093| 0.093| 0.093| 0.093| 0.247       | 0.456       | 51      |
| Sm2AlC     | 0.375| 6.00| 6.00| 6.00| 0.247       | 0.456       | 51      |

The results of other antiperovskite compounds are listed for comparison.

Figure 4. Electronic band structures with (a) spin-up and (b) spin-down channels for Co3InN and (c) the band structure for only the spin-up channel for Ni3InN ternary antiperovskites. The pattern of the band structure of Ni3InN for the down-spin channel is identical with that for the spin-up channel and is not shown here.
Quantum ESPRESSO\textsuperscript{36} code and presented in fat band representation in Figures 7 and 8, respectively. In these figures, the vertical width of the belts (dispersion lines) represents spectral weights of (a) Co/Ni-e\textsubscript{g} (d\textit{z}\textsuperscript{2}, d\textit{x}\textsuperscript{2}−d\textit{y}\textsuperscript{2}), (b) Co/Ni-t\textsubscript{2g} (d\textit{xz}, d\textit{yz}, d\textit{xy}), (c) In-t\textsubscript{1u} (p\textit{x}, p\textit{y}, p\textit{z}), (d) N-t\textsubscript{1u} (p\textit{x}, p\textit{y}, p\textit{z}), and (e) In\textit{a}\textsubscript{1g} (s) orbitals, respectively. For better clarity, orbital-projected bands are not superimposed on normal band dispersion curves. Figures 7 and 8 represent normal band dispersion curves where the orbital degeneracy at different energy levels is identified by red color orbital symbols on the vertical line along the high-
symmetry point $\Gamma$. The Fermi energy is set to zero and shown by a dashed black line. The inset of Figure 7(a) is for two separate calculations considering Co-$d_{x^2-y^2}$ and Co-$d_{z^2}$ orbitals, respectively, and the vertical width of the belt confirms that the 2-fold degeneracy of $e_g$ is broken, and it splits in two orbitals $d_{x^2-y^2}$ ($B_{1g}$) and $d_{z^2}$ ($A_{1g}$), respectively, where the energy of $d_{x^2-y^2}$ is greater than that of $d_{z^2}$. The findings are similar for Ni$_3$InN where 2-fold degeneracy of $e_g$ is also broken with $d_{x^2-y^2}>d_{z^2}$ and not shown in the inset of Figure 8(a).

From Figures 7 and 8, it is seen that the band structure of Co$_3$InN (Ni$_3$InN) is projected into Co (Ni)-$e_g$ ($d_{x^2-y^2}$) and $t_{2g}$ ($d_{xy}$, $d_{yz}$, $d_{xz}$), In ($N$)-$t_{1u}$ ($p_x$, $p_y$, $p_z$), and $a_{1g}$ ($s$) orbitals. The eigen states at the $\Gamma$ point in the Brillouin zone are considered to explain the orbital hybridization process. Figure 9(b) and 9(c) represents the crystal field splitting and the possible hybridization between $d$-orbitals of Co–Co/Ni–Ni and $s$- and $p$-orbitals of In/N atoms. These hybridizations arise due to interactions of the bonding and the antibonding states of the orbitals.

Figure 7. Orbital projected band structure of Co$_3$InN for spin-down channels in a fat band representation along $\Gamma-X-R-M-\Gamma-R$ directions. Vertical width of the belts (dispersion curves) represents spectral weights of (a) Co-$e_g$, (b) Co-$t_{2g}$, (c) In-$t_{1u}$, (d) N-$t_{1u}$, and (e) In-$a_{1g}$ orbitals, respectively. For better clarity, orbital projected bands are not superimposed on normal band dispersion curves. (f) Represents normal band dispersion curves where the orbital degeneracy at different energy levels is identified by red color symbols on the vertical line along the high-symmetry point $\Gamma$. The Fermi energy is set to zero and shown by a dashed black line. The inset of (a) is for two separate calculations considering Co-$d_{x^2-y^2}$ and Co-$d_{z^2}$ orbitals, respectively, and the vertical width of the belt confirms that the 2-fold degeneracy of $e_g$ is broken, and it splits in two orbitals $d_{x^2-y^2}$ ($B_{1g}$) and $d_{z^2}$ ($A_{1g}$), respectively, where the energy of $d_{x^2-y^2}$ is greater than that of $d_{z^2}$.

Antiperovskite A$_3$InN ($A =$ Co, Ni) compounds with Hermann–Mauguin notation $m\overline{3}m$ belong to octahedral ($O_h$) symmetry. Within Co$_3$InN (Ni$_3$InN), N and In atoms are coordinated by 6 and 12 Co (Ni) atoms, respectively, forming a Co$_6$N (Ni$_6$N) octahedron and Co$_{12}$In (Ni$_{12}$In) cuboctahedron, respectively. In A$_3$InN ($A =$ Co, Ni), In and N atoms occupy Wycko positions $a$ ($0,0,0$) and $b$ ($1/2,1/2,1/2$) with the site symmetry of $O_h$, and 3 $A$ atoms occupy Wycko positions $c$ ($0,1/2,1/2$) with the site symmetry $D_{4h}$, respectively.

Here we used $e_g$ ($d_{x^2-y^2}$, $t_{2g}$ ($d_{xy}$, $d_{yz}$, $d_{xz}$), $t_{1u}$ ($p_x$, $p_y$, $p_z$), and $a_{1g}$ ($s$) symbols of $O_h$ symmetry for orbitals shown in the first bracket. We also used $A_{1g}$ ($d_{z^2}$), $B_{1g}$ ($d_{x^2-y^2}$), $B_{2g}$ ($d_{xy}$), and $E_g$ ($d_{xy}$, $d_{xz}$) symbols of $D_{4h}$ symmetry for orbitals shown in the first bracket in the molecular orbital diagram for considering $A-A$ hybridization between 3$d$ orbitals of two $A$ atoms with local site symmetry $D_{4h}$ [cf. Figure 9(a)].

Based on the orbital-projected band diagrams (Figures 7 and 8), we predicted a possible hybridization scheme which is explained below. Figure 9(b) and 9(c) is molecular orbital
diagrams between spin-down orbitals sitting at different sites in the case of the Co$_3$InN and Ni$_3$InN antiperovskites, respectively. Here we used a similar approach as used in earlier studies to construct the MO diagrams of other perovskites with different crystal structures (such as X$_2$YZ).\textsuperscript{32,53,54} Here the hybridizations between the two different A (A = Co, Ni) atoms with D$_{4h}$ site symmetry are considered first [cf. Figure 9(a)]; afterward A−A and an N (site symmetry O$_h$) atoms are hybridized with another A and an In (site symmetry O$_h$) atoms, respectively. The site-symmetry approach establishes symmetry relations between the localized atomic electron states and crystalline states; then the localized states transform following the overall crystal symmetry. 

Here, the hybridization and combination of double-degenerated $E_g$ ($d_{xz}, d_{yz}$) and single-degenerated $B_{2g}$ ($d_{xy}$) of A−A and A with symmetry D$_{4h}$ transformed according to O$_h$ symmetry to produce double-degenerated bonding $e_g$ orbitals and antibonding $e_g^*$ orbitals. Due to the unavailability of more sites with D$_{4h}$ symmetry to hybridize the antibonding states of A−A: $E_g^*$ ($d_{xz}, d_{yz}$), $A_{1g}^*$ ($d_{z^2}$), $B_{2g}^*$ ($d_{xy}$), and $B_{1g}^*$ ($d_{x^2−y^2}$) remain as nonbonding in the resulting MO orbital diagram [cf. Figures 9(b) and 9(c)]. However, double-degenerated $E_g$ ($d_{xz}, d_{yz}$) and single-degenerated $B_{2g}$ ($d_{xy}$) of A−A with symmetry D$_{4h}$ combined and transformed according to O$_h$ symmetry to produce the nonbonding triple-degenerated $t_{2g}$ state. Interestingly, two remaining nonbonding orbitals $A_{1g}$ ($d_{z^2}$) and $B_{1g}$ ($d_{x^2−y^2}$) do not combine to create a double-degenerated nonbonding $e_g$ state.

As a whole for Co$_3$InN, we postulate in addition to bonding (a$_{1g}$, t$_{1u}$, t$_{2g}$, and e$_g$) and antibonding ($e_g^*$, $t_{2g}^*$, and t$_{1u}^*$) states the hybridizations of Co−Co [Figure 9(a)] and N with another Co, and an In atom also produces one nonbonding 3-fold t$_{1u}$ and one nonbonding $e_g$ state with broken 2-fold degeneracy [cf. Figure 9].

Figure 8. Orbital-projected band structure of Ni$_3$InN for the spin-down channel in fat band representation along Γ−X−R−M−Γ−R directions. Vertical width of the belts (dispersion curves) represents spectral weights of (a) Ni−$e_g$ (b) Ni−t$_{2g}$ (c) In−t$_{1u}$ (d) N−t$_{1u}$ and (e) In−a$_{1g}$ orbitals, respectively. For better clarity, orbital-projected bands are not superimposed on normal band dispersion curves. (f) Represents normal band dispersion curves where the orbital degeneracy at different energy levels is identified by red color symbols on the vertical line along high symmetry point Γ. The Fermi energy is set to zero and shown by a dashed black line. The 2-fold degeneracy of $e_g$ is broken, and it splitting in two orbitals $d_{x^2−y^2}$ ($B_{1g}$) and $d_{z^2}$ ($A_{1g}$), respectively, as marked in (f), confirmed by two separate calculations (not shown here) considering Ni−$d_{x^2−y^2}$ and Ni−$d_{z^2}$ orbitals, respectively (for details see text).
In the molecular orbital diagram, it is also clear that the $e_g$ state splits in two orbitals $d_{z^2-r^2}$ ($B_{1g}$) and $d_{x^2}$ ($A_{1g}$), respectively, where $d_{z^2-r^2} > d_{x^2}$, as confirmed by separate calculations considering Co-d$_{z^2-r^2}$ and Co-d$_{x^2}$ orbitals separately, compared with the inset of Figure 7(a). Such splitting might be due to local Jahn–Teller distortion, which we believe is a local z-out distortion of $O_h$ complexes which results $d_{z^2-r^2} > d_{x^2}$. In the molecular orbital diagram of Co$_3$InN, one antibonding $t_{2g}$ state is above $E_F$, followed by nonbonding $d_{z^2}$ and $d_{x^2-y^2}$ states toward higher energy.

The $p$ orbitals of N and In atoms hybridize and form bonding and antibonding states of $t_{1u}$ orbitals, whereas $s$ orbitals hybridize and form bonding and antibonding $a_{1g}$ orbitals. The antibonding state $a_{1g}$ is above the considered energy window and does not appear in this diagram. The triple-degenerated antibonding state $a_{1g}$ is similar to Co$_3$InN with respect to hybridization and orbital sequencing. However, the orbital positions with respect to $E_F$ and separations [compare Figure 9(b) with Figure 9(c)] between them are different. An earlier DFT study postulated that Ni-based ternary nitrides with cubic antiperovskite structure are nonmagnetic due to hybridization between Ni-3d and N-2p states. Our results presented above do not support such a hybridization scheme. Interestingly, in the molecular orbital diagram for Ni$_3$InN [Figure 9(c)], $E_F$ lies between two nonbonding $d_{z^2}$ and $d_{x^2-y^2}$ states. These two nonbonding states in the immediate vicinity of $E_F$ might be the reason for almost zero magnetism in Ni$_3$InN.

### 3.5. Mulliken Populations Analysis

The nature of chemical bonding via the charge transfer mechanism in a compound can be predicted using the Mulliken atomic population analysis. The Mulliken atomic and bond overlap populations of A$_3$InN are tabulated in Table 6.

From Table 6, it is seen that N atoms contain negative Mulliken charge, while it is positive for A (Co, Ni) and In atoms. Hence, charges transferred from A/In to N atoms, which indicates the presence of ionic bonding in A$_3$InN. The ionic nature of A$_3$InN can be restricted by the electronegativities of the atomic species. The distribution of carrier density in different bonds and a quantitative measure of bonding and antibonding strengths in A$_3$InN are described by the Mulliken bond overlap population (BOP) analysis. In general, the positive and negative BOP stands for the bonding and antibonding nature of atoms, respectively. The high and low values of BOP point out the increase of covalent and ionic bond nature, respectively, between two atoms. In this study, the N–A bonds possess a higher degree of covalency with bonding nature. The

### Table 6. Mulliken Atomic and Bond Overlap Populations of A$_3$InN (A = Co, Ni)

| phase | atoms | Mulliken atomic population | Mulliken bond overlap population |
|-------|-------|-----------------------------|---------------------------------|
|       |       | $s$ | $p$ | $d$ | total | charge ($e$) | bond | $n^*$ | $P^*$ | $d^*$ (Å) |
| Co$_3$InN | N | 1.68 | 4.02 | 0.0 | 5.71 | -0.71 | N–Co | 03 | 2.01 | 1.94 |
| | Co | 1.29 | 2.01 | 23.43 | 26.73 | 0.27 | Co–In | 03 | 0.21 | 2.74 |
| | In | 0.56 | 2.02 | 9.97 | 12.55 | 0.45 | Co–Co | 03 | -1.17 | 2.74 |
| Ni$_3$InN | N | 1.65 | 4.00 | 0.0 | 5.65 | -0.65 | N–Ni | 03 | 1.92 | 1.95 |
| | Ni | 1.20 | 2.16 | 26.43 | 29.79 | 0.21 | Ni–In | 03 | -0.03 | 2.76 |
| | In | 0.65 | 1.93 | 9.97 | 12.56 | 0.44 | Ni–Ni | 03 | -0.57 | 2.76 |

$n^*$, $P^*$, and $d^*$ denote the bond number, bond overlap population, and bond length, respectively.
Co–In bond is also covalent, whereas A–A bonds show ionic nature.

3.6. Charge Density Distribution Mapping and the Fermi Surface. The bonding nature of $A_3InN$ ($A = Co, Ni$) is investigated by the charge density difference of unlike atomic sites for further insights. The charge density mapping of $A_3InN$ is shown in Figure 10 on the (001) crystallographic plane. The high and low charge density of electrons is indicated by red and blue colors of the adjacent scale bar, respectively. In general, the greater accumulation of charges (positive value) is favorable for the formation of covalent bonds between two atoms, while the negative value or the lower accretion of charges indicates the formation of ionic binding. The contour maps of electron charge density show higher accumulation of charges in (A, In) than (A, A) atoms, which is consistent with the Mulliken bond population analysis. Moreover, the spherical charge distributions around A atoms indicate the ionic nature of chemical bonds in $A_3InN$.

The conception of Fermi surface is regarded as the heart for understanding the state of a compound with metallic nature. The dynamical properties of a material largely depend on the position and shape of the Fermi surface with respect to the Brillouin zone (BZ). The Fermi surface topology of the ternary nitride perovskite $A_3InN$ is evaluated in the equilibrium structure with spin-polarized conditions at zero pressure. It is important to note that the effect of spin channels is absent on the Fermi surface of Ni$_3$InN. This spin-independent nature of the Fermi surface for Ni$_3$InN is also supported by band dispersion study. The Fermi surface topologies for spin-up, spin-down, total channel of Co$_3$InN, and total channel of Ni$_3$InN are shown in Figure 11(a)–(d), respectively, and explained separately below.
Co$_3$InN with spin-up channel: (1) Half of an oblate-spheroid-shaped open surface between high-symmetry points X, and R [marked by H1 in Figure 11(a)] within the BZ is a hole pocket. A total of 24 such hole pockets are visible at the 6 zone faces. The conventional equation of the oblate spheroid is written as
\[\frac{x^2}{a^2} + \frac{y^2}{b^2} + \frac{z^2}{c^2} = 1\]
where \(x, y,\) and \(z\)-axes intercepts are \(a, a,\) and \(c,\) respectively, and \(a > c\). One hole pocket at the right face of the BZ cube is marked to show the \(x, y, z\) directions and their \(a, a, c\) intercepts. From Figure 11(a), it is evident that the \(z\)-direction is considered along the high-symmetry direction \(X - R\). The ratio \(a/c\) is calculated as \(\sim 3.25:2\) \((3.25c = 2a)\). The center of the hole pocket is situated exactly at the middle of the two high-symmetry points \(X\) and \(R\). The open surface of a hole pocket that cuts the BZ face creates approximately an elliptical shape with the major axis along the \(y\)-direction. The elongated 4 of such a major axis on a single BZ face forms a perfect square which is \(45^\circ\) rotated \([\text{shown by dotted line, Figure 11(a)}]\) and inscribed within the square of the BZ face. Here it should be mentioned that these hole pockets are not exactly ideal oblate-spheroid-shaped. (2) In addition, at the zone edges, the open surfaces centered at \(M\) seen within the BZ are electron pockets \([\text{marked by E1 in Figure 11(a)}]\). They enclose approximately rectangular areas at the adjacent sides of zone faces. It seems if two such surfaces are attached, it will create a dome shape with an approximately square base. A total of 12 such electron pockets are visible in Figure 11(a).

Ni$_3$InN with spin-down channel: (3) Closed surface around the \(G\) (gamma) point which very nearly resembles an octahedron with curved edges as observed in Figure 11(b), marked by H2 as a hole pocket. (4) In addition, one fourth of an oblate-spheroid-shaped \((a > c)\) open surface centered at \(M\) observed at the BZ edge is an electron pocket, marked by E2 in Figure 11(b). The open surfaces of an electron pocket that cuts the adjacent sides of zone faces create half \((\text{cut along the minor axis})\) of an elliptical shape \((\text{with equal size, at both sides})\) with the major axis along a high-symmetry MX direction. One electron pocket at the left-upper edge of BZ is marked to show the \(x, y, z\) directions and their \(a, a, c\) intercepts of an oblate spheroid. Here, the ratio \(a/c\) is calculated as \(\sim 3:2\) \((3c = 2a)\). A total of 12 such electron pockets are visible in Figure 11(b).

Co$_3$InN with total-spin channel: As both the electron pockets \([\text{explained in (2) and (4)}]\) for both up- and down-spin channels are centered at same point \(M\), the Fermi surface nesting is expected as evident in Figure 11(c), where the open surface of an electron pocket \((\text{originated from up spin})\) in turn is nested within another one-fourth portion of a larger oblate-spheroid-shaped open surface \((\text{of another electron pocket \((\text{originated from down spin})\))}\).

Ni$_3$InN with total-spin channel: Here a similar explanation holds as Co$_3$InN with the spin-up channel explained earlier and shown in Figure 11(d). However, here a major difference is that the volume of hole \((\text{electron})\) pockets is smaller \((\text{larger})\) compared to Co$_3$InN with a spin-up channel. The observations are consistent with the band diagram analysis.
In conclusion, the Fermi surface of A3InN is composed of multiple electron- and hole-type sheets. The nonspherical shape of Fermi sheets indicates the metallic conductivity of both Co3InN and Ni3InN. The multi Fermi sheets of A3InN are formed due to low dispersion of A-3d, In-5p, and N-2p states, as revealed by pDOS studies. The observed Fermi surface nesting may have an effect on the magnetic order and phonon softening in Co3InN.39

3.7. Optical Properties. The optical properties of A3InN are calculated as a function of incident photons along the [100] direction. A phenomenological damping constant of 0.05 eV, the free-electron plasma frequency of 10 eV, and Gaussian smearing of 0.5 eV are used for the inclusion of intraband transitions for nonlinear optical constant calculations. The absorption coefficient (\( \alpha \)) determines how far the light of a certain wavelength can penetrate into a material before being absorbed. The variation of the absorption coefficient \( \alpha \) with photon energy is shown in Figure 12(a). From this figure, it is clearly seen that the optical absorption starts from zero photon energy, which confirms the absence of optical band gap in A3InN, complementing the metallic nature obtained in the electronic structure calculations. The absorption coefficient increases sharply in the visible spectrum region (after \( \sim 2.5 \) eV) and displays a high value for a wide energy range (\( \sim 18 \) eV). Hence, A3InN could be used as a promising absorber of electromagnetic radiation for both visible and ultraviolet (UV) regions.

The reflectivity spectra (R) of A3InN are shown in Figure 12(b). It is seen that A3InN reflects more that 98% of light at zero photon energy. After that, the reflectivity decreases gradually with increasing photon energy. However, the average reflectivity of A3InN is >45% in the visible and UV spectral region, which indicates the suitability of A3InN as a coating material to avoid solar heating.

The optical conductivity (\( \sigma \)) starts at zero photon energy [Figure 12(c)], which reconfirms the metallic nature of A3InN. The response of \( \sigma \) with photon energy is slightly blue-shifted for Co3InN compared to Ni3InN. There is a sharp peak at \( \sim (4.2-4.5) \) eV which might have arisen due to the interband transitions of charge carriers from occupied to unoccupied orbitals in A3InN.

The loss function of a material is interconnected with absorption and reflection. It describes the energy loss of charge carrier traveling in a material. The energy loss function (\( \delta \)) is calculated for A3InN and shown in Figure 12(d). There is no loss peak observed in the energy range of \( (0-15) \) eV due to large absorption of electromagnetic radiation. The highest loss peaks appeared for Co3InN and Ni3InN at \( \sim 22.80 \) eV and \( \sim 22.94 \) eV, respectively, which represent the plasma resonance owing to collective charge excitation. The corresponding frequency at which the highest peak of the loss spectrum occurs is called the plasma frequency \( (\omega_p) \) of the material. Below \( \omega_p \), A3InN is expected to reflect the electric field screening by electrons; otherwise, it is transparent to the incident photon.

The appearance of \( \omega_p \) in the UV range makes A3InN reflective in the visible spectral range. The refractive index is a complex parameter, composed of the real and imaginary parts of the complex index of refraction. The real part of the refractive index \( (n) \) demonstrates the phase velocity of the electromagnetic wave inside a material. The frequency dependence of the refractive index is shown in Figure 12(e). It is seen that \( n \) is higher at low photon energy (<5 eV), which decreases with the increasing photon energy. Both Co3InN and Ni3InN display a sharp peak at \( \sim 3.78 \) eV and \( \sim 4.06 \) eV, respectively, due to the intraband transition of electrons.

The imaginary part of the complex index of refraction is known as the extinction coefficient \( (k) \), which measures the amount of attenuation of the incident light when traveling through the material. The extinction coefficient of A3InN is plotted in Figure 12(f). It is seen that A3InN exhibits a large static value of \( k \), which decreases gradually with higher photon energy. The high static value of \( k \) is indicative of metallic conduction in A3InN.

The macroscopic electronic response of a material is explained by means of the complex dielectric constant as \( \varepsilon(\omega) = \varepsilon_1(\omega) + i\varepsilon_2(\omega) \), where \( \varepsilon_1(\omega) \) and \( \varepsilon_2(\omega) \) are the real and imaginary part of the dielectric function. The response of the real and imaginary part of the dielectric constant with photon energy is plotted in Figure 12(g) and 12(h), respectively. From Figure 12(g), it is seen that A3InN displays negative values of \( \varepsilon_1 \) at very low energies, after which \( \varepsilon_1 \) vanishes. The response of \( \varepsilon_1 \) at low energy is shown in the inset of Figure 12(g). The negative values of \( \varepsilon_1 \) at low photon energies signify the Drude-like behavior of A3InN. On the other hand, A3InN displays positive values of \( \varepsilon_2 \) and drops down to zero sharply at very low energies [Figure 12(h)]. The variation of \( \varepsilon_2 \) at low energies is shown in the inset of Figure 12(h). Both \( \varepsilon_1 \) and \( \varepsilon_2 \) disappear at very low photon energy, which signifies the high metallic nature of A3InN.

4. CONCLUSIONS

In summary, we have investigated the structural, mechanical, spin-dependent electronic and molecular orbital predictions, and magnetic and optical properties of antiperovskites nitriles A3InN (A = Co, Ni) using the first-principles DFT calculations combined with the spin polarization effect. The calculated values of different elastic moduli show the damage-tolerant, quasi-ductile, easily machinable, and stiff nature of A3InN. Poisson’s ratio, Pugh’s ratio, and Cauchy pressure calculations suggest that A3InN is ductile in nature. The values of elastic anisotropy indices reveal that Co3InN is more anisotropic in compression, while Ni3InN is more anisotropic in shear. The effect of spin channels is observed in the band structure, DOS, and Fermi surface of Co3InN only. The Fermi surface of A3InN is composed of multiple nonspherical electron and hole type sheets, and the observed Fermi surface nesting in Co3InN might contribute to charge density wave formation and also to enhance the effect of electron–phonon coupling. The induced total magnetic moment of Co3InN is predicted as 2.735 \( \mu_B/\text{cell} \), whereas Ni3InN shows nominal magnetic moments. Molecular orbital diagrams of A3InN antiperovskites are drawn by analyzing orbital-projected band structures. It is exciting to note that the hybridizations of different energy states are identical for both Co3InN and Ni3InN. Therefore, it is expected that it will follow the similar trend for A3BX compounds.

The bonding nature between atoms of A3InN is thoroughly explained with the aid of Mulliken atomic populations and charge density calculations. The above-mentioned properties reveal that these compounds display the combined bonding nature like ionic and covalent. The variations of the various optical constants such as real and imaginary parts of the dielectric constant, refractive indices, reflectivity, absorption coefficient, and loss function with the energy of incident radiation show metallic behavior, complementing the outcomes of electronic band structure and DOS calculations. The reflectivity spectra show that A3InN displays superior reflectivity.
in the visible and UV spectral region, which indicates that A3InN compounds have potential significance to be used as coating materials to evade the solar heat. We hope this work will provide helpful data for the family of antiperovskite materials for further investigation through experiments and theories.

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**Notes**

The authors declare no competing financial interest.

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