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Electronic and magnetic properties of single-layer MPX₃ metal phosphorous trichalcogenides

Bheema Lingam Chittari,^{1,2} Youngju Park,² Dongkyu Lee,² Moonsup Han.² Allan H. MacDonald.³ Euvheon Hwang.^{1,*} and Jeil Jung^{2,†}

¹SKKU Advanced Institute of Nanotechnology, Sungkyunkwan University, Suwon, 16419, Korea ²Department of Physics, University of Seoul, Seoul 02504, Korea

³Department of Physics, The University of Texas at Austin, Austin, Texas 78712, USA

We survey the electronic structure and magnetic properties of two dimensional (2D) MPX₃ (M= V, Cr, Mn, Fe, Co, Ni, Cu, Zn, and X = S, Se, Te) transition metal chalcogenophosphates to shed light on their potential role as single-layer van der Waals materials that possess magnetic order. Our *ab initio* calculations predict that most of these single-layer materials are antiferromagnetic semiconductors. The band gaps of the antiferromagnetic states decrease as the atomic number of the chalcogen atom increases (from S to Se to Te), leading in some cases to half-metallic ferromagnetic states or to non-magnetic metallic states. We find that the competition between antiferromagnetic and ferromagnetic states can be substantially influenced by gating and by strain engineering. The sensitive interdependence we find between magnetic, structural, and electronic properties establishes the potential of this 2D materials class for applications in spintronics.

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I. INTRODUCTION

Following seminal studies that reported on the exfoliation of a stable single-layer graphene and on transport measurements with nearly ideal Dirac fermion fingerprints,^{1,2} research on ultrathin two dimensional (2D) materials has emerged during the last decade as one of the most active research topics in condensed matter physics. Two-dimensional materials are interesting in part because their properties can be modified in situ by adjusting gate voltages. Recently, the focus of 2D materials research has expanded beyond graphene to include other layered van der Waals materials with a variety of distinct physical properties.³ For example, one popular class of 2D materials is the transition metal dichalcogenides (TMDC)⁴ which includes metals, semiconductors with exceptionally strong light-matter coupling,⁵ and broken symmetry electronic states including ones with charge density wave and superconducting order.⁶⁻⁹ Magnetic order has, however, not vet been established in any two-dimensional material. Roomtemperature magnetism in a single-layer material is an extremely attractive materials target because of the expectation that it might provide unprecedented electrical control of magnetism and enable new classes of information processing devices that incorporate non-volatile memory elements more intimately.

One strategy to search theoretically for magnetism in isolated two-dimensional van der Waals materials is to explore the magnetic properties of single-layers exfoliated from a bulk material that exhibits robust magnetic order. Following this approach, recent theoretical studies have proposed a number of potential magnetic single-layer van der Waals materials, including group-V based dichalcogenides,¹¹ FeBr₃, the chromium based ternary tritellurides $CrSiTe_3$ and $CrGeTe_3$, $^{12-19}$ CrX_3 trihalides, 20,21 and $MnPX_3$ ternary chalcogenides. 22,23 The vanadium based dichalcogenides, VX_2 (X=S, Se) were proposed first and are predicted to have strain tunable ferromagnetic phases.¹¹ The trihalides CrX₃ (X = F, Cl, Br, D^{21} are new classes of semiconducting ferromagnet with Curie temperature predictions of $T_{\rm C} < 100$ K.

The bulk ternary tritellurides $CrATe_3$ (A = Si, Ge)¹³ were predicted within LDA to be ferromagnetic with small band gaps of 0.04 and 0.06 eV respectively. Their few layers limits have recently been studied in temperature dependent transport experiments.¹⁴ The anisotropy along the c-axis and the dynamic correlations in the ab-plane seen by elastic and inelastic neutron scattering are characteristic of 2D magnetism.¹⁵ In the single layer limit CrSiTe₃ is a semiconductor with a GGA gap of 0.4 eV,¹⁶ substantially larger than its bulk value, and has a negative thermal expansion.¹⁷ Reports differ on the most stable magnetic phase between ferromagnetic^{13,16,17} or antiferromagnetic.¹⁸ For larger atomic number compounds like CrGeTe₃ and CrSnTe₃, DFT predicts ferromagnetic semiconducting phases with Curie temperatures between 80-170K.14,18,19

We focus here on single-layer materials formed from compounds in the transition metal phosphorous trichalcogenide (MPX₃) family, which are known to exhibit magnetism for M=Mn and for a number of other metal atom species. Metal phosphorous trichalcogenides are cousins of CrSiTe₃, but are so far less studied for 2D magnetism. According to one recent study monolayer MnPSe3 and MnPS3 exhibit Neel antiferromagnetism²² and valley-dependent optical properties.²³ The structures and magnetic properties of some bulk compounds from this family have already been extensively studied.²⁴⁻⁴⁹ Because of the van der Waals character of these materials, one focus of bulk materials research is to characterize ion intercalation properties.^{50–54} For 2D magnetic materials to be most useful in device applications it is desirable to seek pathways to increase the critical temperature at which they order magnetically. We compare theoretical predictions for a variety of late 3d transition metals (M= V, Cr, Mn, Fe, Co, Ni, Cu, Zn) and consider all there chalcogen atoms (X = S, Se,Te) in an effort to explore the magnetic phases that can be expected as fully as possible. We study how the electronic bands are modified when the magnetic state undergoes a transition from antiferromagnetic to ferromagnetic, or from magnetic to non-magnetic. Our results confirm the expected strong interdependence between magnetism and structural properties, for example lattice constant and crystal symmetry, and explain a surprisingly strong dependence of exchange interaction strengths on electron density and strain. Because these materials may have relatively strong correlations, *ab initio* density-functional theory is not able to make quantitatively reliable predictions for all properties. Nevertheless our survey provides considerable insight into materials-property trends, and into the potential for engineering the magnetic properties of these materials using field effects and strain engineering.

Our paper is structured in the following manner. We start in Sec. II by briefly summarizing some specific details of our first principles electronic structure calculations. In Sec. III we discuss our results for ground-state properties including structure, magnetic properties, and electronic band structures and densities-of-states. Sec. IV is devoted to an analysis of the carrier-density dependence of the magnetic ground state, and to a study of the influence of strain on the magnetic phase diagram. Finally we close the paper in Sec.V with a summary and discussion of our results.

II. AB INITIO CALCULATION DETAILS

The study of the ground-state electronic structure and magnetic properties in this work has been carried out using plane-wave density functional theory as implemented in Quantum Espresso.⁵⁵ We have used the Rappe-Rabe-Kaxiras-Joannopoulos ultrasoft (RRKJUS) pseudoptentials for the semi-local Perdew-Burke-Ernzerhof (PBE) generalized gradient approximation (GGA)56 together with the VdW-D2 correction⁵⁷. We choose the GGA+D2 as a reference calculation because of the overall improvement of the GGA over the LDA⁵⁸ for covalent bond description, and add the longer ranged D2 correction to improve the description of binding between the layers. (The GGA typically performs poorly for van der Waals bonds.⁵⁹) The magnetic solutions have also been compared with calculations employing the DFT+U scheme, using the same value of U = 4 eV and alternatively using onsite repulsion U values that saturate the magnitude of the band gap. Comparisons were made with hybrid HSE+D2 functionals⁶⁰ to assess non-local exchange effects that could further influence the magnetic ground states. All structures were optimized without constraints until the forces on each atom reached 10^{-5} Ry/au. The self-consistency criteria for total energies has been set at 10^{-10} Ry and momentum space integrals were performed using a regularly spaced k-point sampling density of $16 \times 16 \times 1$ for the triangular lattice case and $16 \times 8 \times 1$ for the rectangular lattice case, with a plane wave energy cutoff between 60 to 90 Ry. For the HSE+D2 calculation we used a coarser effective k-point sampling of $4 \times 4 \times 1$. The out-of-plane vertical size of the periodic supercell was chosen to be 25 Å, which typically leaves an adequate vacuum spacing greater than 10 Å between two-dimensional layers.

FIG. 1: (Color online) Schematic structure of the transition metal phosphorous trichalcogenide MPX₃ compounds with $(P_2X_6)^{4-1}$ bipyramids that enclose metal atoms. a. Atomic structure of a MPX₃ monolayer. The rectangular supercell referred to in the main text is indicated by black dashed lines and the smaller triangular lattice unit cell is represented by red solid lines. Each triangular unit cell contains a transition metal atoms(M) honeycomb sublattice. The phosphorus (P) dimers are perpendicular to the plane at the center of each honeycomb lattice hexagon, and three sulphur atoms are bound to each of the P atoms. The top and bottom chalcogen trimers have a relative in-plane twist of 60° degrees. **b.** The lowest energy 3D-bulk structure consist of individual layers stacked in an ABC sequence. We present the structure from three viewpoints, a side view, a top in which the phosphorous atoms within a single layer lie on top of each other, and a view in which the transition metal atoms of adjacent layers lie on top of each other.

III. STRUCTURAL AND MAGNETIC PROPERTIES

The atomic structure of MPX₃ transition metal chalcogenophosphate layers is anchored by $(P_2X_6)^{4-}$ bipyramids arranged in a triangular lattice that provide enclosures for transition metal atoms. (See Fig. 1 for a schematic illustration of the single layer unit cell and the bulk atomic structure.) The bulk crystals consists of ABC-stacked single layer assemblies that are held together by van der Waals forces. Although the atomic structures of single layer transition metal phosphorous trichalcogenide crystals are similar to those of bulk crystals, small changes appear in response to the absence of the interlayer coupling, with

distortions in the ground-state crystal geometries correlated mainly with the magnetic phase. The analysis of magnetic properties is simplified by the fact that the magnetic moments develop almost entirely at the metal atom sites.

We calculate the magnetic ground-state and meta-stable magnetic configurations by identifying the energy extrema obtained by iterating self-consistent field equations to convergence starting from magnetic initial conditions that can be classified as either Néel antiferromagnetic (AFM), or ferromagnetic (FM), or nonmagnetic (NM) states. The calculation indicates that most 2D MPX₃ crystals are semiconductors with localized magnetic moments that are ordered antiferromagnetically. We find that the use of chalcogen atoms with larger atomic numbers tends to yield smaller energy gaps between valence and conduction bands. In the following we present an analysis of the structural and magnetic properties of representative 3d transition metal MPX₃ trichalcogenides, and discuss their underlying electronic band structures.



FIG. 2: (Color online) Summary of key structure metrics (in-plane lattice constant, P-P bond length, M-M bond length and P-X bond length) in single-layer MPX₃ compounds. Bond lengths increase with the size of the chalcogen atom (S- square, Se - circle, Te - star) and a strong systematic dependence on whether the magnetic state is NM (non-magnetic - orange), FM (ferromagnetic - blue), or antiferromagnetic (AFM-green), but have a more complex non-monotonic dependence on metal atomic number,

A. Structural properties

The MPX₃ compounds that we study consist of V, Cr, Mn, Fe, Co, Ni, Cu, and Zn transition metal atoms with 3d valence electrons, combined with three different chalcogen atoms S, Se, and Te. We thus expand our study beyond the most common crystals in this class, the MnPS₃, FePS₃, CoPS₃, NiPS₃, ZnPS₃ thiophosphates whose bulk structure had been explored in past experiments.^{24,26–41,51–54} We have optimized MPX₃ lattice structures using the unit cell shown in Fig. 1 for single layers and assuming ABC stacking for bulk. Given this framework, structures are characterized by the value of the inplane lattice constant a, the (P-P) distance between the phosphorous atoms, the (M-M) distance between metal atoms, and the (P-X) distance between phosphorous atoms and chalcogen atoms. The variation in the magnitude of these bond lengths as a function of the metal and chalcogen atom species, and magnetic configuration is illustrated in Fig. 2, and summarized numerically in Table II in the Supplemental Material. Results are presented for all metastable magnetic states. The structures have a simple dependence on chalcogen atom size, with larger in-plane lattice constants and monolayer thicknesses (P-P distances) for larger chalcogens. However, the dependence of structure on metal atom atomic number is not straightforward. Experiments in the bulk MPS₃ observed a close correlation between the radius of the metal cation and the P-P distance of the thiophosphate bipyramid.²⁴ The nonmonotonic variation of the metal cation radius with respect to atomic number is reflected in the P-P distance trend. However, our calculations show that bond lengths correlate more strongly with magnetic state than with metal atomic number.

P-P distances range between 2.15-2.3 Å while M-M bond lengths vary by about 5%. The M-M bond lengths within an hexagon can be unequal, distorting the hexagonal lattice arrangement of the metal atoms as illustrated in Fig. S1 in the Supplemental Material⁶¹. In general ferromagnetic states have the largest lattice constants, and antiferromagnets have intermediate lattice constants. The fact that the structural properties of these crystals are correlated with their magnetic configurations suggests the possibility of controlling magnetic properties by straining the lattice, as we will discuss more in detail in a later section.

The comparison between lattice parameters calculated with GGA+D2 and the experimental bulk structure is summarized in Table I. The comparison of the GGA and LDA lattice parameters in Table I of the Supplemental Material indicates important discrepancies between the GGA and LDA, and is consistent with the tendency of the LDA to overbind covalent bonds. The stronger LDA bonds leads to a global reduction of lattice constant and to a stronger distortion of the metal honeycombs. Within the LDA we find optimized lattice parameters that are (2-5%) shorter than in bulk experiments. The GGA agreement is good (0.5-1.6%) for MnPS₃, FePS₃, CoPS₃, NiPS₃ and ZnPS₃.



FIG. 3: (Color online) Top view of different magnetic ordering arrangements (a) ferromagnetic, (b) Néel antiferromagnet, (c) zigzag antiferromagnet and (d) stripy antiferromagnetic. In MPX₃ compounds, the magnetic moments reside primarily on the metal atoms which have a honeycomb structure in each layer. The black and white spheres represent up and down spins, respectively. The red circles identify nearest neighbors (NN) of the central metal site, the navy blue circles (dashed-dotted) identify the second nearest neighbors, and the light blue circles identify the third nearest neighbors. The magnetic energy landscape can be approximated by assigning Heisenberg coupling constants J_1 , J_2 and J_3 to metal atom pairs with these three separations.

B. Magnetic properties

Experimental studies of the magnetic properties of bulk Mn, Fe, Co, Ni thiophosphates have found antiferromagnetic ground states with Néel temperatures ranging between $82K \sim 155K$.^{24,25} Here we find that magnetism persists in single layer MPX₃ compounds. The magnetic moments develop mainly at the localized metal atom sites, except for a noticeable spin polarization that develops on the phosphorous and sulphur atom sites in the ferromagnetic configuration.

The late 3d transition elements Cr, Mn, Fe, Co and Ni stand out in the periodic table as elements that tend to order magnetically. The bonding arrangements of particular compounds can however enhance or suppress magnetism. In 2D MPX₃ crystals, transition metal atoms are contained within phosphorous trichalcogenide bipyramidal cages, and have weak direct hybridization with other transition metal atoms. The exchange interactions between the metal atoms are therefore mainly mediated by indirect exchange through the intermediate chalcogen and P atoms. Magnetic interactions can be extracted from *ab initio* electronic structure calculations by comparing the ground state energies of different magnetic configurations. We compare the energies of antiferromagnetic, ferromagnetic, and nonmagnetic states in V, Cr, Mn, Fe, Ni based compounds in Table II, where we find that the AFM phase is normally favored over the FM phase.

Because our calculations show that the magnetic moments are concentrated at the metal atoms sites we can characterize the magnetic properties of 2D MPX₃ compounds by mapping the energy landscape to an effective classical spin Hamiltonian on a honeycomb lattice:

$$H = \sum_{\langle ij \rangle} J_{ij} \vec{S}_i \cdot \vec{S}_j = \frac{1}{2} \sum_{i \neq j} J_{ij} \vec{S}_i \cdot \vec{S}_j \tag{1}$$

where \vec{S}_i is the total spin magnetic moment of the atomic site *i*, J_{ii} is the exchange coupling parameters between two local spins, and the prefactor 1/2 accounts for the double-counting. The estimated magnetic anisotropy energies that we obtained from non-collinear magnetization calculations are on the order of $\sim 160 \ \mu eV$ per formula unit for the FM compounds CrPS₃, CrPSe₃ and on the order of $\sim 1000 \ \mu eV$ per formula unit for the AFM compounds MnPS₃, MnPSe₃ with magnetization favored perpendicular to the plane. For this reason, in making the T_c estimates that we describe below, we take the Ising limit of this spin-Hamiltonian. By evaluating the three independent energy differences between the four mag-netic configurations^{18,24,62} illustrated in Fig. 3, ferromagnetic (FM), Néel (AFM), zigzag AFM (zAFM), and stripy AFM (sAFM), and assuming that the magnetic interactions are short range, we can extract the nearest neighbor (J_1) , second neighbor (J_2) , and third neighbor (J_3) coupling constants:¹⁸

$$E_{\rm FM} - E_{\rm AFM} = 3(J_1 + J_3)\vec{S}_A \cdot \vec{S}_B$$
 (2)

$$E_{\text{zAFM}} - E_{\text{sAFM}} = (J_1 - 3J_3) \dot{S}_A \cdot \dot{S}_B \tag{3}$$

$$E_{\rm FM} + E_{\rm AFM} - E_{\rm zAFM} - E_{\rm sAFM} = 8J_2 \dot{S}_A \cdot \dot{S}_A \quad (4)$$

where \vec{S}_X is the average spin magnetic moment on the honeycomb sublattice X. The total energies of the zAFM and sAFM magnetic configurations, which must be calculated using a rectangular supercell, usually have higher energies than the average of the AFM solutions and FM solutions. The larger rectangular unit cell required for describing more complex zAFM and sAFM spin configurations imposes certain symmetries that restricts the relaxation of the lattices when compared to the triangular unit cell. For the case of FePS₃, however, the zAFM has the lowest energy. The average magnetic moment S at each lattice site obtained within the GGA+D2 and GGA+D2+U are listed in Table III.

Single-layer magnetic ordering temperatures T_c were estimated by running Monte Carlo simulations of the threecoupling-constant effective models using the Metropolis algorithm in lattice sizes up to $N=32\times64$ with periodic boundary conditions,^{63–65} and verified against calculations performed using Wang-Landau Monte Carlo sampling algorithm in smaller $N=8\times16$ lattices.⁶⁶ We calculated the heat capacity $C = k\beta^2 (\langle E^2 \rangle - \langle E \rangle^2)$ as a function of temperature and identified its diverging point as the Neel and Curie temperatures. See the Supplemental Material⁶¹ for the plot of the representative results for the temperature dependent heat capacity.

The calculated average values of the magnetic moments in the AFM configuration vary widely, assuming the values 2.01, 3.14, 4.16, 3.2, 0.8 and 1.14 μ_B for V, Cr, Mn, Fe, Co and Ni based sulphides. (See Table III). The magnitudes of the magnetic moments at the metal atoms generally have relatively small differences between different magnetic configurations, between 3%~10%, while larger variations are found in CoPS₃ and NiPSe₃. Within GGA+D2 approximation we find that the magnetic moments in 2D MPX₃ develop almost entirely at the metal atom sites. In comparison the phosphorous and chalcogens in the bulk structure do acquire a small spin polarization that can be attributed to interlayer coupling. The use of onsite U introduces a small spin polarization enhancement at nonmetal atom sites.

C. Band structure and density of states

Understanding the electronic properties of 2D MPX₃ layers is an essential stepping stone on the path toward possible integration in nanodevices. For spintronic applications of magnetic 2D materials, which seek to couple charge and spin degrees of freedom, it is desirable to understand how the electronic structure depends on the type of magnetic order. The spin resolved band structures of the lowest energy spin configuration, calculated within the GGA+D2 approximation, are presented in Fig. 4, and the associated densities-of-states are presented in Fig. 5, and the projected partial density of states (PDOS) results for various MPX₃ compounds presented in Fig. 6. Single layer MPX₃ with M = V, Cr, Mn, Fe, Co, Ni, Cu, Zn transition metal atoms and X = S, Se, Te chalcogen atoms are considered. Band structures for higher energy spin configurations are included in the Supplemental Material⁶¹.

We find that AFMs are almost always gapped semiconductors, that the FM states are metallic, and that the NM phases can be either semiconducting or metallic. It follows that the MPX₃ class of materials includes almost all of the behaviors being studied in current spintronics research, including importantly both antiferromagnets and ferromagnets and both metals and insulators.

The AFM band structures for V and Mn based compounds in Fig. 4 show semiconductor behavior and reveal conduction band edges that are near the K-points, whereas the valence band edges are between Γ and K. This valence bands near the Fermi energy are energetically separated from deeper lying valence bands. We notice from the density of states plot in Fig. 5 that there is strong sensitivity of the electronic structure to the choice of electron-electron interaction model, both for the HSE+D2 approximation containing non-local exchange and for on-site repulsion enhancement introduced through a Hubbard U parameter. The analysis of the orbital projected partial density of states in Fig. 6 for the AFM compounds reveals that the conduction band edges have an important contribution from the s and p orbitals of the P atom while for the valence band edges the chalcogen atom orbitals have an importance presence. Other atoms do contribute near the band edges, e. g. for the V based compounds the d-orbitals are an important fraction of the valence band edge while for the Mn based compounds they lie at deeper energies. We can thus expect that surface functionalization and variations in the carrier density at the surface chalcogens will have a more immediate impact in the valence bands than the conduction bands. The AFM compounds containing Fe and Ni result in non-magnetic solutions when heavier chalcogen atoms are used within our GGA+D2 calculation. The case of FePS₃ has as lowest energy configuration the zigzag AFM phase.

For MPX₃ compounds the FM configurations are metallic as a general rule. Within our GGA+D2 approximation only the Cr based chalcogenophosphates have FM ground states, while they favor the AFM phase when the onsite U is added, see the Supplemental Material for GGA+D2+U results. We find that the FM configurations are often meta-stable local minima solutions in MPX₃ compounds whose lowest energy solutions are AFM. The FM configurations are found in metallic phase at charge neutrality and give rise to half-metallic solutions for MnPS₃ and NiPS₃. The half-metallicity could be achieved also in MnPSe₃ when carrier doped away from neutrality. For the MnPS₃ and MnPSe₃ compounds we can observe a distinct population of the carriers for the hole and electron sides where doping populate orbitals located at the chalcogen and P atoms respectively.

The electronic properties of NM configurations are also presented in Figs. 4-5. The NM layers could be potentially interesting if magnetic phases could be induced by forming vertical heterojunctions with magnetic materials. Unexpectedly, one typically magnetic element (Co) turns out to be nonmagnetic in MPX₃ compounds according to our GGA+D2 based calculations. As shown in Fig. 4 the band structure of these non-magnetic materials can be either metallic (Co, Ni, Cu based compounds), semiconducting (Fe), or be a wide gap insulator (Zn).

We now turn our attention to the magnitude of the band gaps of the semiconducting and insulating 2D MPX₃ chalcogenophosphates calculated within the GGA approximation, and using other approximations that partially account for Coulomb correlation through non-local exchange or a Hubbard U, are shown in Table IV. These gaps are consistently larger when calculated using the hybrid HSE+D2 approximation which includes non-local exchange in the energy functional. A smaller gap enhancement is obtained using a +U interaction correction. Band gaps are gradually reduced when S is replaced by heavier chalcogen atoms.

The corresponding density of states (DOS), plotted in Fig. 5, shows the strong influence on the ground-state electronic structure when Coulomb correlations are included. Note in particular the impact of the non-local Coulomb exchange included in the HSE approximation. This suggests that the physics of MPX₃ compounds can be dominated by correlation effects and modeling will be most successful when we rely on effective models that feed from experimental input or high level *ab initio* calculations.

The orbital content of the valence and conduction band edges that are most relevant for studies of carrier-density dependent magnetic properties can be extracted from the orbital



FIG. 4: (Color online) GGA+D2 band structures for single layer MPX₃ compounds in their lowest energy magnetic configuration for M = V, Cr, Mn, Fe, Co, Ni, Cu, and Zn transition metal atoms and X = S, Se, Te chalcogen atoms. The plotted band structures were calculated using the triangular structural unit cell, except for the case of FePS₃ which is predicted to have a larger periodicity zAFM (see text) magnetic structure that has a triangular unit cell with a doubled lattice constant. The bands are violet for AFM configurations, violet and orange for the up and down split spin bands in FM configurations, and green in NM phases. We note that the AFM phases have semiconducting band gaps and that the FM phases are metallic, while the non-magnetic phases can be either metallic or semiconducting. An overall reduction of the band gaps is observed when we use heavier chalcogen atoms, in keeping with the reduction of the covalent bond energy in larger shell orbitals.

projected partial density of states (PDOS) results for various MPX₃ compounds presented in Fig. 6. Depending on the specific composition and on the magnetic configuration of the material under consideration, the valence and conduction band edge orbitals can be dominated by metal, phosphorous or chalcogen atoms. A detailed study on the influence of the Coulomb interaction model on the localization properties of the wave functions and their influence on the exchange coupling in MPX₃ compounds will be presented elsewhere.

IV. TUNABILITY OF MAGNETIC PROPERTIES

Two dimensional magnetic materials are of interest primarily because of the prospect that their properties might be more effectively altered by tuning parameters that are available *in*



FIG. 5: (Color online) Total density of states (DOS) for the magnetic ground-states in Fig. 4, calculated using three different exchangecorrelation energy functional calculations GGA+D2, HSE+D2 and GGA+D2+U. We have placed the valence band edge at E = 0. Overall band gap enhancements are found for the semiconducting phases when additional electron-electron interactions are approximated as a nonlocal exchange corrections in HSE+D2 and as onsite repulsion corrections in GGA+U. Both HSE and GGA+U enhance the band gaps. Other features in the DOS plots also depend on the DFT exchange-correlation functional approximation.



FIG. 6: (Color online) Orbitally projected partial density of states (PDOS) calculated for self-consistently converged ground-state magnetic configurations. The zero of energy is chosen at the valence band edge. From the PDOS of the various MPX_3 compounds we observe that the orbital content of the valence and conduction band edges vary widely. For the compounds showing carrier-density dependent magnetism the states closest to the Fermi energy generally have an important contribution from the p-orbitals of the chalcogen and P atoms.

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FIG. 7: (Color online) Carrier density dependent total energy differences per MPX₃ formula unit between the AFM and FM states of V, Cr, Mn, Fe, Ni based single layer trichalcogenides. The prominent cusps in these curves appear because the antiferromagnetic state has a band gap whereas the ferromagnetic state does not have a band gap. The AFM ground-states favored near charge neutrality can be switched to FM states at accessible carrier densities in V, Mn, Ni, and Fe based compounds. For CrPX₃ with X = Se or Te, it is possible to switch from a FM to an AFM state at finite n-doping. Carrier densities of up to a few ~ 10^{14} electrons per cm² should be in principle accessible through ionic liquid or gel gating. A carrier density of 0.1 electrons per MPX₃ formula unit corresponds to ~ 6×10^{13} electrons per cm² when the lattice constant is ~ $6\mathring{A}$.

situ. Two potentially important control knobs that can be exploited experimentally in two-dimensional-material based nano-devices are carrier-density and strain. The dependence of magnetic properties on carrier density is particularly interesting because it provides a convenient route for electrical manipulation of magnetic properties. In this section we explore the possibility of tailoring the electronic and magnetic properties of MPX₃ ultrathin layers by adjusting carrier density or by subjecting the MPX₃ layers to external strains.

A. Field-effect modification of magnetic properties

The goal of modifying the magnetic properties of a material simply by applying a gate voltage is a holy grail in magnetoelectronics because it could allow magnetically stored information to be written at negligible energy cost. Electric field control of magnetic order has been demonstrated in a number of materials. In ferromagnetic semiconductors the mechanism is carrier density variation in thin films which leads to a modification of the magnetic exchange interaction and magnetic anisotropy. In thin films of ferromagnetic metals, gate voltages can vary the Fermi level position at the interface, which governs the magnetic anisotropy of the metal.^{67–69}

We leave to future work a complete microscopic analysis of the response of a MPX₃ layer, acting as one electrode of a capacitor, to a gate voltage. Here we capture the most important response by simply examining the dependence of magnetic properties on carrier density. In doing so we neglect the possible role of charge polarization within a MPX₃ layer. Fig. 7 summarizes the theoretically predicted trends in competition between AFM and FM states in 2D MPX₃ compounds with M = V, Cr, Mn, Fe, Ni and X = S, Se, Te. We find that when the magnetic ground state is AFM (M=V, Mn, Ni, Fe), transitions to FM states are driven by sufficiently large electron or hole carrier densities. The origin of this trend is easy to understand. Because the FM state is gapless the energy changes associated with adding one electron and removing one electron are identical. In the gapped AFM states they differ by the energy gap E_{gap} . It follows that the energy difference per area unit between antiferromagnetic and ferromagnetic states $\delta E \equiv E_{AFM} - E_F$ is given at low carrier densities by

$$\delta E(n) = \delta E_0 + (E_{gap}/2 + \delta \mu) n$$

$$\delta E(p) = \delta E_0 + (E_{gap}/2 - \delta \mu) p$$
(5)

where *n* is the carrier density of *n*-type samples, *p* is the carrier density of *p*-type samples, δE_0 is the energy difference per area unit between antiferromagnetic and ferromagnetic states in neutral MPX₃ sheets, and $\delta\mu$ is the difference between the mid-gap energy of the antiferromagnetic semiconductors and the chemical potential of the ferromagnetic metals. Introducing *n*-carriers is most effective in driving a transition from antiferromagnetic to ferromagnetic states when $\delta\mu$ is positive. whereas introducing *p*-carriers is most effective when $\delta\mu$ is negative. By comparing with Fig. 7 we conclude that $\delta\mu$ is small in most cases, but large and negative in MnPS₃ and MnPSe₃. Because the energy difference per formula unit between ferro and antiferromagnetic states is much smaller than the energy gap, a transition between antiferromagnetic and ferromagnetic states can be driven by carrier density changes per formula unit that are much smaller than one - especially so when $\delta\mu$ plays a favorable role. In particular we see in Fig. 7 that a transition between ferromagnetic and antiferromagnetic states are predicted in MnPS3 and MnPSe3 at hole carrier densities that are $\sim 10^{14}$ cm⁻², which corresponds to about ~ 0.16 electrons per formula unit. Density changes of this magnitude can be achieved by ionic liquid or gel gating. Since this size of carrier density is sufficient to completely change the character of the magnetic order, we can expect substantial changes in magnetic energy landscapes at much smaller carrier densities. Transitions between ferromagnetic and antiferromagnetic states are predicted in most MPX₃ compounds. Our calculations show that the FM solution is the preferred stable magnetic configuration in almost all cases when the system is subject to large electron or hole densities in the range above a few times $\pm 10^{14}$ cm⁻². The DOS and PDOS shown in Figs. 5-6 in the main text and in Figs. 5-10 in the Supplemental Material may suggest that a Stoner ferromagnetic instability is at play when the peaks near the band edges are sufficiently prominent. Our calculations therefore motivate efforts

to find materials which can be used to establish good electrical contacts to MPX₃ compounds, and in particular to the valence bands of MnPS₃ and MnPSe₃. For the Cr based compounds, whose behavior is different, the ground states at zero carrier density is FM and we find that a transition to an AFM state can occur for *n*-doped systems.

B. Strain-tunable magnetic properties

The membrane-like flexibility of ultrathin 2D materials makes them suitable platforms for tailoring the electronic structure properties by means of strains. Representative examples on effects of strains in 2D materials properties that have been discussed in recent literature include the observation of Landau-level like density of states near high curvature graphene bubbles⁷⁰, or the commensuration moire strains that opens up a band gap at the primary Dirac point in graphene on hexagonal boron nitride for sufficiently long moire patterns^{71–73}. The sensitive variation of the different M-M, P-P, M-X bond lengths as a function of magnetic configuration we have presented earlier in Sec. III suggests that strains can be used as switches to trigger magnetic phase transitions or alter the stability of the magnetic phases by modifying the total energy difference with respect to the nonmagnetic phases. Here we have carried out calculations of the total energies for different magnetic phases to explore the influence of expansive or compressive in-plane biaxial and uniaxial strains and the effects of external pressure along the outof plane axis in 2D MPX₃ materials. The compressive and expansive biaxial strains have been modeled through uniform scaling of the rectangular unit cell in Fig. 1 and likewise the uniaxial strains are modeled by scaling the unit cell either in the zigzag or armchair directions. We notice that generally the effects of uniform biaxial versus uniaxial strains introduce modifications in the magnetic phase energy difference total energies that are qualitatively similar for the compressive or expansive strains, and that the effects are stronger for biaxial strains when compared to uniaxial strains which do not show a noticeable difference between the zigzag or armchair directions. Pressure strains along the out of plane vertical c-axis have been applied by artificially modifying the P-P distance and we find for MnPS₃ a clear transition from AFM to FM with the total energy difference $E_{AFM} - E_{FM}$ changing from -93 meV to 268 meV per formula unit in the presence of a -4% bond length shortening strain for the P-P distance. Relatively large strains are required to alter the magnetic properties in compounds like MnPS₃ but other systems such as CrPS₃, FePS₃, FePSe₃ NiPS₃ and VPTe₃ the magnetic phase transitions can be achieved for smaller strains as they have relatively smaller total energy differences between the magnetic phases. We find a particularly sensitive transition near charge neutrality for FePS₃, FePSe₃ and VPTe₃ where strains on the order of $\sim 1\%$ is enough to switch between different magnetic configurations.



FIG. 8: (Color online) Influence of in-plane strain on the magnetic configurations of selected MPX₃ compounds. Magnetic phase transitions are introduced by in-plane biaxial compressive and expansive strains for several magnetic MPX₃ compounds at zero carrier density. Particularly sensitive strain dependence is seen for FePS₃, FePSe₃ and VPTe₃ where small strains on the order of 1% of the lattice constant can switch the magnetic phases.

V. SUMMARY AND DISCUSSION

In this paper we have carried out an *ab initio* study of the MPX₃ transition metal phosphorous trichalcogenide class of two-dimensional materials. We have considered different combinations of 3d metals (M = V, Cr, Mn, Fe, Co, Ni, Cu, Zn) and chalcogens (X = S, Se, Te) in an effort to identify materials that are promising for spintronics based on twodimensional materials. Our calculations suggest that magnetic phases are common in the single-layer limit of these van der Waals materials, and that the configuration of the magnetic state depends systematically on the transition metal/ chalcogen element combination. We find that semiconducting Néel antiferromagnetic states are most common; and predict AFM phases for V, Mn, Fe, and Ni based compounds. A metallic ferromagnetic states is found in Cr based compounds, and non-magnetic states are found with both semiconducting and metallic electronic structure, while introduction of U tends to stabilize the AFM magnetic phase. As expected on the basis of the weaker covalent bonds of larger atomic orbitals we find that for semiconducting antiferromagnetic materials, replacing a smaller chalcogen atom with larger chalcogen atom reduces the energy band gap and as a consequence also the difference between the ground state energies of FM and AFM states. Interestingly we do not find magnetic states for CoPX₃, even though Co is typically a magnetic atom.

The electronic structures predicted by density functional theory for these materials are sensitive to the choice of exchange-correlation energy functional. For example, there are substantial discrepancies in predicted densities-of-states between the standard semi-local GGA, approximations with a local U correction, and hybrid functionals with non-local exchange. This sensitivity of the optimized ground-state results to the choice of the approximation scheme makes it desirable to benchmark the results against experiment in order to establish which approximation is most reliably predictive. The fact that the predictions of density-functional theory are qualitatively sensitive to the exchange-correlation approximation employed, indicates that the MPX₃ compounds are correlated.

An interesting interdependence between magnetic order and atomic structure was found, with typical variations on the order of a few percents in the lattice constant and interatomic bond lengths leading to structural distortions depending on the magnetic configuration of the system. The bond length variations were traced through M-M, P-P, P-X bond distances associated with the distortions in the honeycomb array of the transition metal atoms and the distance between the atomic centers in the $(P_2X_6)^{4-}$ bipyramids. We leave for future work the analysis on the role of spin-orbit coupling effects and optical properties arising from lattice symmetry breaking coupled to the onset of magnetism. Sizeable differences between the GGA and the LDA solutions is suggestive of the delicate balance between the different chemical bonds for configuring the optimized structure of these compounds which in turn are affected by the onset of magnetism. In the antiferromagnetic state, the strength of the exchange interactions is expected to vary inversely with the band gap; approximations that underestimate the band gap will overestimate exchange interaction strengths. We expect that the LDA likely does underestimate band gaps, as it commonly does, and therefore the exchange interactions it implies may well be overestimates. In 2D MPX₃ a variety of different stable and meta-stable magnetic configurations were found. Semiconducting states with a band gap are typically antiferromagnetic phases in either Néel, zigzag, or stripy configuration and the metallic states are typically ferromagnetic phases. The critical temperatures for magnetic phases in the single layer limit are generally expected to be smaller than in the bulk due to the reduction in the number of close neighbor exchange interactions, although changes in the degree of itineracy and shifts in relative band positions can also play a role. We analyzed the magnetic phases of the 2D MPX₃ compounds by building an effective model Hamiltonian with exchange coupling parameters extracted by mapping the total energies from our *ab* initio calculations onto an effective classical spin model. The Curie and Néel temperatures T_C, T_N are obtained by using a statistical analysis based on the Metropolis algorithm⁶⁴. The calculated Néel temperatures of the antiferromagnetic comControl of magnetic phases by varying the electric field in a field effect transistor device is a particularly appealing strategy for 2D magnetic materials. Our calculations indicate that a transition between antiferromagnetic and ferromagnetic phases can be achieved by inducing carrier densities in 2D MPX₃ compounds that are in the range that can be induced by a field effect. We find that a magnetic phase transition from a FM state to an AFM state can be induced in metallic CrPSe₃ and CrPTe₃ compounds, which are close to the phase boundary, by gating to large n-type carrier densities. Similarly AFM to FM transitions can be achieved in VPTe₃, MnPTe₃, and FePS₃. For materials exhibiting magnetic phases we find that using the heavy chalcogen Te can reduce the carrier densities required for magnetic transitions.

The interdependence between atomic and electronic structure suggest that strains can be employed to tune magnetic phases. We have found that the ground state magnetic configuration can undergo phase transitions driven by in-plane compression or expansion of the lattice constants or by as c-axis pressure. The magnitude of the required strains vary greatly, with values below 1% for compounds such as FePS₃, FePSe₃ and VPTe₃, and larger strains on the order of or far greater than 4% are required to trigger a FM-AFM magnetic phase transition in CrPS₃ and NiPS₃. Phase transitions for even larger strains on the order of \sim 9% are observed in VPSe₃, MnPS₃, NiPS₃, NiPSe₃ whose non-strained configuration has a robust gapped antiferromagnetic phase.

Based on our calculations, we conclude that single layer MPX₃ transition metal thiophosphates are interesting candidate materials for 2D spintronics. Their properties, including their magnetic transition temperatures, can be adjusted by the application of external strains or by modifying the carrier densities in field effect transistor devices. The sensitivity of these systems to variations in system parameters, such as composition stoichiometry, details of the interface, and the exchange coupling of the magnetic properties with external fields, offers ample room for future research that seek new functionalities.

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TABLE I: Comparison between experimental and theoretical bulk lattice parameters, and theoretical two dimensional lattice parameters for transition metal phosphorous trichalcogenide MPX₃ compounds. These results were obtained using GGA+D2 and are expressed in Å; a is the in-plane lattice constant and c' is the layer thickness, *i..e* it is the distance between the planes containing the three chalcogen atoms in a single MPX₃ layer. The values in parenthesis below the bulk c'(Å) values are the interlayer distances, *i.e.* the vertical distances between the top P atom of one layer and the bottom P atom of an adjacent layer.

MPX ₃	Х	NM		FM		AFM	
		a(Å)	c'(Å)	a(Å)	c'(Å)	a(Å)	c'(Å)
	S	5.737	3.142	5.880	3.166	5.846	3.170
	S(bulk)	5.742	3.280	5.918	3.273	5.845	3.294
	((2.518)		(2.585)		(2.655)
	S(Expt)		(,		(,		()
VPX ₂	Se	6.123	3.266	6.230	3.204	6.214	3.318
	Te	6 885	3 328	6 935	3 4 1 8	6 980	3 4 3 1
	5	5 851	3.071	5 914	3.032	5 885	3 336
	S(bulk)	5 862	3 1 1 6	5 913	3 185	5 934	3 301
	S(bulk)	5.002	(2.624)	5.715	(2.506)	5.754	(2.465)
	S(Evnt)		(2.024)		(2.300)		(2.403)
CrDV.	S(Expt)	6 025	2 267	6 2 2 0	2 100	6 2 2 5	2 225
CIFA3	<u>э</u> е Та	6.055	2.307	6.520	2 2 2 2 0	6.525	2.222
	10	0.330	3.072	0.000	3.330	0.804	3.300
	3	5.780	2.952	0.023	3.230	5.997	3.240
	S(bulk)	5.787	2.875	6.018	3.125	5.993	3.268
			(2.456)		(2.567)	< 0 - <	(2.562)
	S(Expt)					6.076	
$MnPX_3$	Se	6.045	3.182	6.340	3.416	6.334	3.402
	Te	6.530	3.444	6.805	3.550	6.874	3.535
	S	5.730	2.808	5.891	3.150	5.958	3.180
	S(bulk)	5.736	2.969	5.874	3.313	5.921	3.217
			(2.386)		(2.840)		(4.368)
	S(Expt)					5.934	
FePX ₃	Se	6.130	2.911	6.134	3.347	6.239	3.214
	Te	6.676	3.038				
	S	5.763	2.790				
	S(bulk)	5.784	2.950				
			(4.250)				
	S(Expt)	5.901					
CoPX ₃	Se	6.186	2.850				
5	Te	6.772	3.057				
	S	5.766	2.974	5.792	3.064	5.783	3.042
	S(bulk)	5.763	2.995	5.789	3.062	5.779	3052
	~()		(2.431)		(2, 449)		(2.480)
	S(Expt)		(())	5.813	(
NiPX-	Se	6 132	3 086	6 1 3 4	3 100	6 140	3 1 4 2
in ny	Te	6 700	3 208	0.151	5.100	0.110	5.112
	<u> </u>	5 837	3.430				
	S(bulk)	5 8/3	2.45				
	S(DUIK)	5.645	(2.002)				
	S(Evet)		(3.903)				
CuDV	S(Expt)	6 151	2 5 2 2				
CuPA3	3C T-	6 711	3.332 2.617				
	re	0./11	3.017				
	S	5.966	3.230				
	S(bulk)	5.973	2.564				
			(3.025)				
	S(Expt)	5.971					
ZnPX ₃	Se	6.323	3.379				
	Te	6.871	3.597				

TABLE II: Total energy relative to that of the lowest energy magnetic configuration calculated using the GGA+D2 approximation for the ferromagnetic (FM), antiferromagnetic (AFM) and non-magnetic (NM) states of single layer transition metal phosphorous trichalcogenides MPX₃. The absence of an entry for a magnetic configuration means that the corresponding state is not metastable. Energies are expressed in terms of meV/unit cell. Our first principles calculations suggest stable magnetic phases for V, Cr, Mn, Fe, Ni metal compounds. The rectangular supercells and structural triangular unit cells used in these calculations are illustrated in Fig. 1. The smaller triangular unit cell is used to compare the NM, FM and Neel AFM phases and the larger rectangular unit cell is needed when we also compare with the total energies of zAFM, sAFM phases. The work functions are in eV units for the smaller triangular unit cell.

MPX ₃	Х		Re	ctangu	ılar		Tr	ar	Work function (eV)			
		NM	FM	AFM	zAFM	sAFM	NM	FM	AFM	NM	FM	AFM
VPX ₃	S	1142.8	643.2	0.0	246.9	132.7	1083.8	695.8	0.0	3.56	3.26	3.42
	Se	906.9	275.8	0.0	164.6	87.78	1348.1	458.8	0.0	2.82	3.57	3.58
	Te	0.0	399.3	323.9	371.4	327.9	1296.5	183.1	0.0	3.08	3.88	3.85
	S	1454.5	0.0	91.65	30.62	63.23	1182.5	0.0	73.68	3.40	3.07	3.62
CrPX ₃	Se	1514.6	0.0	146.1	57.06	522.7	1293.7	0.0	315.4	3.98	3.52	3.65
	Te	1373.4	0.0	242.9	81.31	12.06	1482.6	0.0	264.2	3.74	3.55	3.41
	S	1576.2	186.3	0.0	60.68	74.59	1964.1	205.4	0.0	3.74	4.41	3.98
MnPX ₃	Se	1356.6	156.7	0.0	48.96	68.98	1417.3	166.4	0.0	3.98	4.07	3.91
	Te	0.0					840.0	12.44	0.0	3.68	3.66	3.77
	S	141.5	286.8	115.8	0.0	289.0	44.87	0.0	308.3	3.41	4.13	4.23
FePX ₃	Se	0.0					0.0	165.0	324.5	3.93	3.73	3.86
	Te	0.0					0.0			3.32		
	S	0.0					0.0			3.84		
CoPX ₃	Se	0.0					0.0			4.33		
	Te	0.0					0.0			3.50		
	S	507.5	276.7	64.16	0.0	343.5	425.1	213.4	0.0	4.63	4.91	5.08
NiPX ₃	Se	324.8	196.2	0.0	383.2	278.2	168.0	158.8	0.0	4.79	4.54	4.44
	Te	0.0					0.0			2.58		
	S	0.0					0.0			4.96		
CuPX ₃	Se	0.0					0.0			5.15		
	Te	0.0					0.0			2.96		
	S	0.0					0.0			3.97		
ZnPX ₃	Se	0.0					0.0			4.47		
	Te	0.0					0.0			1.11		

TABLE III: Magnetic moments in Bohr magneton μ_B per metal atom for bulk and single layer magnetic MPX₃ structures, the three nearest neighbor exchange coupling strengths J_i in meV implied by the Heisenberg model mapping, and Monte Carlo estimates of the single-layer critical temperatures based on the Ising limit of the classical spin model. The critical temperature is identified with the maximum in the derivative of magnetization with respect to temperature. The Ising limit estimate is motivated by the strong perpendicular anisotropy of these materials. The *ab initio* calculations were performed within GGA+D2, as well as GGA+D2+U using a constant site repulsion at the metal atoms of U=4eV. The magnetization is calculated as the difference between up and down local spin resolved charge densities as implemented in Quantum Espresso.

MPX ₃	Х	Bulk			GGA+D2						GGA+D2 + U								
		FM	AFM	FM	AFM	zAFM	sAFM	J_1	J_2	J_3	T _c	FM	AFM	zAFM	sAFM	J_1	J_2	J_3	T _c
	S	2.03	2.00	1.92	2.01	1.68	1.76	24.5	4.26	3.24	1100	2.96	2.93	2.95	2.93	3.93	0.147	0.143	570
VPX ₃	Se			2.02	2.13	1.77	2.14	10.2	0.340	0.437	760	2.56	2.99	3.01	2.98	2.81	0.132	0.170	400
	Te			2.22	2.16	2.06	2.19	9.63	-3.59	-7.07	570	3.10	3.08	3.13	3.08	5.10	-1.68	-3.51	560
	S	2.83	3.07	2.85	3.14	2.37	2.34	-1.73	-0.015	0.029	280	3.94	4.03	3.93	4.11	1.34	-0.064	-0.490	130
CrPX ₃	Se			2.77	2.83	2.45	2.38	-9.82	-3.49	6.62	1080	4.14	4.03	4.04	4.11	0.608	-0.254	-3.05	250
	Te			2.92	2.88	2.82	2.48	-4.89	0.006	-0.087	750	4.25	4.17	4.17	4.21	0.849	0.333	-2.10	900
MnPX ₃	S	4.24	4.15	4.26	4.16	3.82	3.83	1.21	0.180	0.536	500	4.56	4.54	4.54	4.55	0.369	0.0284	0.152	200
	Se			4.25	4.20	4.24	3.82	0.958	0.136	0.506	450	4.56	4.54	4.55	4.56	0.231	0.021	0.141	150
FePX ₃	S	3.32	3.14	3.33	3.20	3.24	2.48	-1.59	-0.179	3.00	690	3.48	3.46	3.48	3.47	0.575	0.140	0.69	260
NiPX ₃	S	1.25	1.13	1.26	1.14	1.12	1.20	-11.3	-0.115	36.0	1110	1.21	1.17	1.22	1.19	-4.11	1.95	17.4	560
	Se			0.54	0.94	0.12	0.99	127	-82.3	-67.6	2580	1.14	1.08	1.22	1.14	-3.11	1.04	18.2	510

TABLE IV: Band gaps of MPX₃ compounds predicted to have antiferromagnetic semiconductor ground states. The band gaps are listed in eV energy units and have values that depend substantially on the DFT exchange-correlation approximation employed. Values are listed for GGA+D2, HSE+D2 hybrid functional and GGA+D2+U approximations. The magnitude of U was chosen to maximize the band gap. As expected the HSE+D2 functional yields the largest band gaps.

	Х	GGA+D2	HSE+D2	GGA+D2+U
	S	1.23	2.50	2.32 (U = 3.25)
VPX ₃	Se	0.73	2.10	1.67 (U = 3.25)
	Te	0.41	1.57	1.00 (U = 3.25)
	S	1.36	3.05	2.34 (U = 5.0)
MnPX ₃	Se	0.98	2.24	1.70 (U = 5.0)
	Te	0.38	1.06	0.69 (U = 5.0)
	S	0.57	2.07	2.26 (U = 5.3)
FePX ₃	Se	0.20	1.48	1.19 (U = 5.3)
	Te	0.13	1.35	0.59 (U = 5.3)
	S	0.84	2.83	1.74 (U = 6.45)
NiPX ₃	Se	0.59	2.22	1.31 (U = 6.45)
	Te	-	0.27	-
	S	0.01	1.76	0.08 (U = 1.0)
CuPX ₃	Se	-	0.56	-
	S	2.19	3.11	2.23 (U = 5.4)
$ZnPX_3$	Se	1.38	2.19	1.41 (U = 5.4)
	Te	0.73	1.26	0.74 (U = 5.4)

- * Electronic address: heh82320gmail.com
- [†] Electronic address: jeil.jung@gmail.com
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