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MPX_3 van der Waals magnets under pressure (M = Mn, Ni, V, Fe, Co, Cd; X = S, Se)

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2 ABSTRACT

van der Waals antiferromagnets with chemical formula MPX_3 (M = V, Mn, Fe, Co, Ni, Cd; X =3 S, Se) are superb platforms for exploring the fundamental properties of complex chalcogenides, 4 revealing their structure-property relations, and unraveling the physics of confinement. Pressure 5 is extremely effective as an external stimulus, able to tune properties and drive new states of 6 matter. In this review, we summarize experimental and theoretical progress to date with special 7 emphasis on the structural, magnetic, and optical properties of the MPX_3 family of materials. 8 Under compression, these compounds host inter-layer sliding and insulator-to-metal transitions 9 accompanied by dramatic volume reduction and spin state collapse, piezochromism, possible 10 polar metal and orbital Mott phases, as well as superconductivity. Some responses are already 11 providing the basis for spintronic, magneto-optic, and thermoelectric devices. We propose that 12 strain may drive similar functionality in these materials. 13

14 Keywords: van der Waals solids, complex chalcogenides, external stimuli, compression

1 INTRODUCTION

Transition metal phosphorous trichalcogenides, MPX_3 , where *M* represents a transition metal ion, and *X* is either S or Se, are a family of semiconducting van der Waals magnets that were originally discovered in

17 1894 (Friedel, 1894), but were not studied intensively until the 1970s (Klingen et al., 1973). Originally,

18 interest in these materials was sparked by their low dimensional magnetic properties (Le Flem et al., 1982)

19 and ability to form intercalation compounds (JOHNSON and Jacobson, 1982), but more recently these

20 materials have attracted attention as single-layer magnets (Burch et al., 2018) and as highly tunable systems

21 that undergo bandwidth-controlled Mott transitions under applied pressure (Kim et al., 2019).

MPX₃ materials typically crystallize in two space groups, C2/m (No. 12) for the sulfur analogs and $R\overline{3}$ 22 23 (No. 148) for the selenium-containing compounds. In both cases, the *M* ions form a honeycomb net, and at the center of each hexagon there is a P-P dimer that extends perpendicular to the net. S or Se anions 24 terminate each MPX_3 layer on both sides, and there is a van der Waals gap between the layers, leading to 25 small cleavage energies and easy exfoliability. Single layers of the MPS₃ and MPSe₃ materials can also be 26 viewed as a honeycomb net of M^{2+} ions coordinated by distorted octahedra of S/Se anions, and P-P dimers 27 centering the hexagons and forming a $[P_2S_6]^{-4}$ anionic sublattice. To emphasize the importance of the 28 P-P dimers, the chemical formula of MPX_3 materials is sometimes written as $M_2P_2X_6$, but this is incorrect 29 nomenclature as the chemical formula is not supposed to reflect structural features (Hiroi, 2008). These 30 31 structural motifs lead to three different types of chemical bonding: ionic, covalent, and van der Waals which makes understanding the electronic structure of this family of materials quite challenging (Dedkov 32 et al., 2023). 33

In general, MPS_3 materials with M = V, Mn, Fe, Co, and Ni are reported to be antiferromagnetic Mott insulators with variable semiconducting energy gaps and magnetic structures. CdPS₃, in contrast, has a filled *d* shell and is semiconducting and nonmagnetic. The Mottness and magnetism of MPS_3 compounds originates from their partially-filled *d*-orbital shells; this, in addition to the presence of a van der Waals gap, suggests that the MPS_3 series is an excellent platform for the study of two-dimensional correlated electron phenomena and magnetism.

40 Tuning the physical properties of these compounds is a highly contemporary research subject, allowing for the exploration of new states of matter and functionalities. Toward this end, researchers are using 41 42 chemical substitution, doping and intercalation, as well as various external perturbations such as electric field, light, strong magnetic fields, and strain (Wang et al., 2018a; Pei et al., 2022). For instance, chemical 43 44 substitution on the M and X sites provides a viable means to control d-orbital filling and bandwidth, 45 respectively, which are both important parameters to control Mottness, magnetism, and spin-orbital-lattice 46 coupling. Even better, many of MPS_3 compounds have both X = S and = Se limits as stable systems, which allows continuous tuning of lattice parameter and bandwidth via synthesizing MPS_xSe_{3-x} samples 47 48 (Basnet et al., 2022b). Even differences in X-site chemistry sometimes gives rise to interesting contrast; for 49 example, the presence and absence of superconductivity in pressurized FePSe₃ and FePS₃, respectively. At the same time, doping and intercalation control carrier concentration and impart new properties (Basnet 50 51 et al., 2022a; Tezze et al., 2022).

Among these external stimuli, pressure is a highly effective tool for tuning the physical properties of MPX_3 materials. The interlayer van der Waals (vdW) interaction is weak; thus, the interlayer spacing can be effectively modulated by pressure. As the layers are forced closer together, layers slide with respect to each other, changing their relative orientation. Furthermore, pressure modifies the *c/a* ratio, reflecting the more compressible interlayer distance than the *a-b* plane. Bond lengths and angles within the chalcogenide layers are also modified under compression.

With the collapse of the van der Waals gap upon pressure, interlayer orbital overlap increases, and the materials become more three-dimensional with enhanced bandwidth. Hence, pressure becomes a tuning knob to control both the crossover from two-dimensional to three-dimensional regimes in addition to a bandwidth-controlled Mott transition. With varying *d*-orbital occupancies and the resulting different orbital degrees of freedoms (i.e. t_{2g} vs. e_g), pressure can induce drastically different responses of lattice structure, as well as electronic and magnetic properties depending on the chemistry of the metal center and chalcogen anion. 65 Secondly, in addition to bandwidth and dimensionality tuning, pressure provides another path to tweak 66 materials properties via 'layer sliding' in the MPX_3 compounds (and in other layered vdW systems). 67 Because of the monoclinic structure of many MPS₃ compounds, even hydrostatic pressure can create 68 shear strain that induces sliding of adjacent layers and variation of stacking order. Differences in layer 69 stacking can create contrasting responses in electronic and magnetic properties, as recently suggested in 1T-TaS₂ with the "star of David" charge density wave formation. (Butler et al., 2020) The presence of 70 this layer-sliding in the MPS₃ series under pressure, as we will see below, creates complicated yet rich 71 72 structural, electronic, and magnetic behaviors. Therefore, pressure becomes a valuable tool in unlocking 73 tremendously rich physical characteristics hidden within the MPX₃ series.

Over the past few years the evolution of the structural, magnetic, and electronic properties of MPX_3 compounds under compression have been extensively studied. Given the rapid progress in this field, the time is ripe for a short review.

In the sections below, we first discuss Mn, Fe, and Ni MPX_3 materials, which have been the topics of the most number of studies. We then discuss $V_{0.9}PS_3$ and $CoPS_3$, which have been much less studied. Lastly, we discuss the non-magnetic compounds, CdPS₃ and CdPSe₃, and the structure-property relations that can be unraveled in these systems. The final section of the review contains future perspectives on this fascinating family of materials.

2 PROPERTIES OF MPX₃ MATERIALS UNDER PRESSURE

82 2.1 MnPS₃ and MnPSe₃

83 MnPS₃ is the prototype in this family of materials and is therefore extensively investigated. This system 84 crystallizes in a monoclinic C2/m structure whereas MnPSe₃ displays a trigonal $R\overline{3}$ arrangement. The 85 P–P dimer is an important structural building block as is the P₂S₆⁴⁻ cluster. These elements are crucial 86 in defining the thickness of the chalcogen layer. Table 1 summarizes the layer thickness and van der 87 Waals gap for MnPS₃ and MnPSe₃ and compares them to other members of this series. The C2/m vs. $R\overline{3}$ 88 structural difference between sulfur and selenium analogs plays out in other members of the *M*PX₃ family 89 of materials as well [Fig. 1 + Tables 2 and 3].

Early neutron diffraction studies of MnPS₃ uncovered collinear N/'eel-type AFM order at $T_N = 78$ K,(Le 90 Flem et al., 1982; Kurosawa et al., 1983b; Joy and Vasudevan, 1992; Wildes et al., 1994) although there were 91 92 discrepancies as to the direction of the magnetic propagation vector. A recent study revisited this problem using single crystal neutron diffraction and polarimetry.(Ressouche et al., 2010) This work revealed 93 94 a propagation vector $\mathbf{k} = [000]$ as well as magnetoelectric coupling and ferrotoroidicity. (Ressouche 95 et al., 2010) By comparison, $T_N = 74$ K for MnPSe₃. (Wiedenmann et al., 1981; Le Flem et al., 1982) A similar reduction of the transition temperature is observed for $FePX_3$ when the X is modified from 96 97 sulfur to selenium. Neutron scattering of MnPSe₃ reveals an identical propagation vector ($\mathbf{k} = [000]$). (Wiedenmann et al., 1981; Bhutani et al., 2020) Photoemission and x-ray absorption spectroscopy confirms 98 a high spin Mn²⁺ configuration in MnPSe₃.(Fujii et al., 2022) Each Mn center is antiferromagnetically 99 coupled to its three nearest neighbors.(Kurosawa et al., 1983a) One surprising finding is the strong 100 101 magnetic interaction between third nearest neighbors. The spin wave excitations have also been studied by Raman and neutron scattering in these materials. Major findings include: hybridization of the two-magnon 102 103 excitation with phonons (which impacts the spin wave decoherence pathway) and absence of nonreciprocity 104 in zone boundary magnons (which provides experimental limits that can be tested against theoretical predictions).(Sun et al., 2019; Vaclavkova et al., 2020; Mai et al., 2021; Wildes et al., 2021) 105

106 Several different but complementary efforts were required to unravel the sequence of events that takes 107 place under pressure. In the first experimental work, Wang et. al. uncovered a simultaneous spin-state crossover and metal-insulator transition in both MnPS₃ and MnPSe₃ [Fig. 2] as well as zigzag chain 108 formation of the Mn atoms in the metallic phase. (Wang et al., 2016) The critical pressures (P_c 's) for 109 these processes were reported to be 30 and 25 GPa for MnPS₃ and MnPSe₃, respectively. First-principles 110 electronic structure calculations using DFT + U and DFT + DMFT techniques reproduced the insulator-111 to-metal transition with simultaneous spin-state transition but with a Mn-Mn dimerization at 64 GPa, 112 (Kim et al., 2019) although it was hinted that some structural components might be missing. This early 113 experimental and theoretical work did not consider layer sliding, which turns out to modify the picture in 114 115 important ways.

In a significant conceptual advance, Coak et. al. discovered an intermediate pressure transition in the 116 series of MPS_3 ($M = V_{0.9}$, Mn, Fe, Ni) materials.(Coak et al., 2020) This transition involves coherent 117 118 layer sliding to create an alignment of P-P dimers along the layer-normal direction. This insight enabled a number of subsequent advancements. For instance, infrared, Raman scattering, and optical measurements as 119 120 a function of pressure combined with first-principles calculations reveal that MnPS₃ undergoes a C2/m (LP) 121 $\rightarrow P\bar{3}1m$ (HP-I, insulating, around 10 GPa) $\rightarrow C2/m$ (HP-II, metallic, around 30 GPa) with a layer-sliding 122 between the LP and HP-I phases, similar to observed in FePS₃ [Fig. 1 and Table 2]. The HP-I to HP-II transition accompanies spin-state transition, collapse of MnS₆ volume, and a sudden collapse of interlayer 123 124 distance with an layer-normal P-chain formation, consistent to observations reported in FePS₃ [Figs. 1, 2, and 3]. The trigonal-to-monoclinic symmetry lowering at the HP-I \rightarrow HP-II transition is thought to 125 originate from the low-spin d^5 orbital nature of the Mn²⁺ ions, contrary to the case of FePS₃ with the 126 same layer sliding but with C2/m (LP) $\rightarrow C2/m$ (HP-I, insulating after the layer sliding) $\rightarrow P\bar{3}1m$ (HP-II, 127 128 metallic) transitions.

We note in passing that theory tends to over-estimate the band gap and pressure of the insulator-to-metal transition in these materials.(Grasso et al., 1991; Harms et al., 2020; Yan et al., 2023) Gap estimation is a well-known challenge for first-principles density functional theory-based approaches, and weak interactions with shallow potentials are notoriously difficult to calculate with accuracy. Nevertheless, these techniques nicely reveal the semiconducting behavior of the aligned phosphorous dimer phase and the metallic character of the aligned phosphorous chain phase [Fig. 4].

From the optical properties point of view, MnPS₃ is a semiconductor with a 2.7 eV band gap. It hosts 135 spin-charge coupling across the magnetic ordering transition and a set of crystal field excitations that have 136 been widely investigated. (Boerio-Goates et al., 1981; Grasso et al., 1991; Harms et al., 2020; Yan et al., 137 2023; Park et al., 2024) There is strong p-d hybridization which activates the d-to-d on-site excitations of 138 the Mn²⁺ centers. There is also a broad near infrared emission band assigned as ${}^{4}T_{1} \rightarrow {}^{6}A_{1}$ with a half life 139 of approximately 50 µsec at 77 K.(Boerio-Goates et al., 1981) The ratio of the crystal field splitting to the 140 Racah parameter (10Dq/B) is unusually large and may explain the strongly red-shifted light emission in 141 this system. That the band gap of $MnPS_3$ decreases systematically under pressure suggests that this system 142 hosts piezochromism.(Harms et al., 2020) It turns out that MnPS₃ is green at ambient conditions, and as 143 shown in Fig. 5, it is yellowish-green at 3 GPa, bright red at 8 GPa, and black by 20 GPa. The latter occurs 144 because the band gap shifts out of the visible wavelength range, moving systematically toward closure at 145 a rate of approximately -50 meV/GPa. This effect is guenched by the appearance of the insulator-metal 146 transition. This high level of linear responsivity under pressure is unusual, and piezochromism of this 147 type has applications in pressure sensing, actuation, as well as artificial skin. At this time, it is not known 148 whether strain can drive similar effects in the bulk material, although in few- and single-sheet MnPS₃ there 149

150 is a C2/m to $P\bar{3}1m$ transition that may involve surface strain effects (Neal et al., 2019, 2020) and biaxial 151 strain is predicted to impact the magnetic ground state in MnPSe₃ monolayers.(Pei et al., 2018)

Thermal properties of these materials are also of interest from a fundamental point of view and for heat management purposes in a device. Kargar *et. al.* find in- and out-of-plane thermal conductivities of $6.3 \pm$ 154 1.7 and 1.1 ± 0.2 W/m·K, respectively.(Kargar et al., 2019) These values are higher than what is found in 155 the Fe analog.

156 2.2 FePS₃ and FePSe₃

157 FePS₃ and FePSe₃ have been reported as layered antiferromagnets with $T_{\rm N}$ s around 120 K. (Le Flem et al., 1982; Joy and Vasudevan, 1992). FePS₃ has gained significant interest recently for its Ising-like 158 antiferromagnetic ordering in the single-layer limit. (Lee et al., 2016) The d^6 configuration of the Fe²⁺ 159 charge status shows a high-spin configuration S = 2 with the partially-filled t_{2g} complex, implying 160 161 anisotropic nature of magnetism as revealed by experimental and theoretical observations. (Nauman et al., 162 2021; Zhang et al., 2021) In the presence of external pressure, like Mn-based siblings, spin-state transition 163 and the resulting complex electronic and structural responses are expected in FePS₃ and isostructural 164 FePSe₃. Accordingly, the responses of magnetic, electronic, and structural properties of FePS₃ and FePSe₃ 165 have been actively investigated.

Recent x-ray, neutron scattering, and optical spectroscopy studies on the effect of hydrostatic pressure on 166 167 FePS₃ reveal a sequence of structural phase transition, (Haines et al., 2018; Coak et al., 2021; Harms et al., 2022b) analogous to MnPS₃ and $V_{1-x}PS_3$ (Harms et al., 2020; Coak et al., 2019); a gradual layer-sliding 168 crossover from the monoclinic ambient (C2/m space group) to the pseudo-hexagonal intermediate phase 169 (HP-I, C2/m) around 2~5 GPa, and the following $C2/m \rightarrow P\bar{3}1m$ (HP-II) transition accompanied by the 170 insulator-to-metal transition at 14 GPa (see Fig. 1). (Coak et al., 2021) The ambient and HP-I phases 171 of FePS₃ have been reported to be insulating and antiferromagnetic with a Néel-type order. Figures 6a 172 and **b** show the magnetic configurations of the ambient and HP-I phases from neutron data, respectively. 173 (Coak et al., 2021). Figure 6c describes the evolution of the T_N as a function of pressure. (Wang et al., 174 2018b; Haines et al., 2018; Coak et al., 2021) The gradual enhancement of the T_N can be attributed to 175 the increased kinetic energy scale due to the closer Fe-Fe distances and the resulting enhanced magnetic 176 exchange interactions. 177

The second transition (HP-I \rightarrow HP-II) accompanies the breaking of the P-P dimer structure and the formation of interlayer P-chain perpendicular to the Fe layers, in addition to a sudden collapse of the interlayer distance and volume, (Haines et al., 2018) similar to the case of MnPS₃. (Harms et al., 2020) Note that these observations are consistent with separate first-principles electronic structure calculation results, (Zheng et al., 2019; Deng et al., 2023) where the origin of the coexistence of the ambient and HP-I phases around 2~5 GPa is suggested to be the result of the shear-induced energy barrier between the ambient and HP-I phases (see Fig. 7a). (Deng et al., 2023)

From both experimental and theoretical studies the Mott-insulating antiferromagnetic phase is reported 185 to remain robust in the ambient and HP-I phases, reflecting the quasi-two-dimensional nature. Haines 186 et al. (2018); Zheng et al. (2019) With the formation of the interlayer P-chain in the HP-II phase this 187 two-dimensional character breaks down, and so does the insulating behavior. The sudden collapse of the 188 interlayer distance and cell volume at the HP-I \rightarrow II transition implies a spin-state transition (see Fig. 3). 189 Indeed, an early study on pressurized phases of FePS₃ and FePSe₃ reported collapses of magnetic moments 190 and transitions to low-spin phases around 13 and 8 GPa, respectively, Wang et al. (2018b) and additionally, 191 the DFT study of Zheng *et al.* also reports the spin-state transition from the high-spin S = 2 to the low-spin 192

S = 0 configuration of the Fe ions at the HP-I \rightarrow II transition. (Zheng et al., 2019) Interestingly, on the 193 other hand, a high-pressure neutron diffraction data by Coak et al. suggests the presence of short-range 194 magnetic order in the HP-II phase, persisting even up to room temperature (see Fig. 6c). Coak et al. (2021) 195 From a detailed electronic structure calculation study, Deng *et al.* argues that this puzzling behavior may 196 be explained by the potential coexistence of two metastable metallic states; the high-spin one with the 197 intra-layer P₂ dimer structure remains intact comparing the short-range order(2D-like), while the P₂ dimer 198 breaks to form the out-of-plane P chain structure (3D-like) in the other low-spin state. (Deng et al., 2023) 199 A separate dynamical mean-field theory study on FePS₃ also suggests the coexistence of the two structural 200 phases, each one hosting the normal Fermi liquid-like character (with the interlayer P-chain) and the 201 202 incoherent 'Hund's metal' character (with the intralayer P-dimer) in the high-pressure regime, where the Hund's metal phase keeps the local moments intact and induces bad-metallic behavior due to magnetic 203 fluctuations. (Kim et al., 2022) 204

It is interesting to note that, in Kim *et al.*, the Hund's metal phase with the orbital-selective correlation effects occurs only with a non-hydrostatic pressure, especially when the pressure perpendicular to the Fe layers are stronger than the in-plane counterparts. This originates from the direction-selective orbital effect under pressure, where the t_{2g} orbitals of Fe ions becomes more susceptible to the out-of-plane stress compared to e_g . (Kim et al., 2022) This implies a subtle role of anisotropy and inhomogeneity in pressure, which may often occur in high-pressure experimental situations depending on the choices of pressure-transmitting medium. (Matsuoka et al., 2021).

212 Lastly we comment on FePSe₃. This system shares a crystal structure and magnetic ground state with FePS₃, (Le Flem et al., 1982; Wiedenmann et al., 1981) and expected from theoretical calculations to 213 214 show many common properties, such as layer-sliding transition to HP-I, volume collapse at the HP-I \rightarrow II 215 transition accompanying the simultaneous spin-state and insulator-metal transitions. (Zheng et al., 2019) 216 Experimentally the second transition is observed to occur at a lower pressure of around 8 GPa than that 217 of FePS₃, which can be attributed to the larger hybridization effects from Se ions. (Wang et al., 2018b) 218 Unlike FePS₃, superconductivity of unknown origin has been reported in the HP-II metallic phase of FePSe₃ below 2.5K (see Fig. 7c and d). (Wang et al., 2018b) Structural refinement in the same study 219 220 suggests the formation of the zigzag chains within the Fe honeycomb layer in the HP-II phase of FePSe₃, 221 hinting the presence of orbital physics from high-spin configuration at Fe ions and potential unconventional 222 superconductivity.

223 2.3 NiPS₃ and NiPSe₃

224 NiPS₃ and the Se analog provide another exciting platform for exploring new states of matter under external stimuli and developing different types of structure-property relationships. The magnetic ordering 225 temperatures are $T_{\rm N}$ = 155 and 206 K, respectively (Le Flem et al., 1982; Joy and Vasudevan, 1992; 226 Chandrasekharan and Vasudevan, 1994), so we see that in this case, heavier anions act to raise the 227 Neél temperature. In the low temperature phase of NiPS₃, the magnetic propagation vector $\mathbf{k} = [010]$. 228 Analysis of the integrated intensity of the (010) magnetic Bragg peak as a function of temperature reveals a 229 critical exponent for the magnetization $2\beta = 0.30$ away from the transition temperature, consistent with 230 a two-dimensional universality class, and 0.30 near $T_{\rm N}$, suggesting the importance of three-dimensional 231 interactions in this regime (Wildes et al., 2015). There are a number of low energy excitations in this system 232 from which competing exchange interactions can be extracted (Wildes et al., 2023; Mehlawat et al., 2022), 233 and active pumping of orbital resonances can also manipulate magnetic order in this quasi-two-dimensional 234 van der Waals antiferromagnet (Afanasiev et al., 2021). Solid solutions in the form of Ni_{1-x}Mn_xPS₃ (0 \leq 235

 $x \le 1$) can also tune the spin flop transition from approximately 7 to 2 T due to modulation of the magnetic 236 237 anisotropy. (Basnet et al., 2021) Intercalation is also very effective in this regard. It can even drive the ground state ferrimagnetic. (Basnet et al., 2022a; Tezze et al., 2022) The magnetic properties of NiPS₃ are 238 239 clearly very sensitive to chemical pressure, suggesting that there is more to learn under physical pressure. 240 The latter is obviously a much cleaner technique as well. Light is also an important external stimuli in these materials. NiPS₃, for instance, sports laser-induced demagnetization with responsivity (lifetime) on the 241 order of 10 ps.Kuntu et al. (2023) Photo-induced quenching of magnetic order is likely to be very sensitive 242 243 to pressure.

244 Early high pressure x-ray diffraction work combined with first-principles calculations suggested strong 245 similarities between NiPS₃ and the M = Mn and V analogs, with LP-to-HP-I and the HP-I-to-HP-II transitions near 15 and 27 GPa with an insulator-to-metal transition near 20 GPa (Ma et al., 2021). More 246 247 recent efforts combining transport and x-ray diffraction reveal a series of different crystal structures under 248 pressure, time-dependent and somewhat sluggish character to several of the transitions, and an insulator-249 to-metal transition near 30 GPa. (Matsuoka et al., 2021) Systematic studies of hydrostatic pressure vs. 250 strain demonstrates that the critical pressure of the insulator-to metal transition (as identified by transport measurements) is reduced from 30 to 12.5 GPa under uniaxial strain (Cui et al., 2021) - an observation that 251 252 again emphasizes the need for hydrostatic pressure conditions. This confusing and somewhat contradictory 253 picture was resolved by bringing together synchrotron-based infrared absorption, Raman scattering, x-ray 254 diffraction, and first-principles calculations under pressure to show that in fact NiPS₃ is quite different from 255 the other MPS_3 materials (Harms et al., 2022a) with a series of five different critical pressures. The set of symmetry progressions in this material is complex (involving $C2/m \rightarrow P3 \rightarrow P3m1 \rightarrow P3m1 \rightarrow P3 \rightarrow P1$ 256 space groups) from monoclinic to trigonal to noncentrosymmetric trigonal to triclinic phases (Figs. 8 and 257 258 9). The insulator-to-metal transition takes place near 23 GPa as evidenced by the development of a Drude 259 response + screening of the phonons in the infrared spectrum. This definition of the insulator-to-metal 260 transition obviously places the critical pressure below that obtained from a transport measurement (23 261 vs. 30 GPa), but the development of Drude signature in the electrodynamic response is an exquisitely 262 sensitive indicator of metallicity. In any case, it's important to realize that several of the aforementioned 263 space groups lack an inversion center, suggesting that NiPS₃ may be a polar metal under compression. 264 (Harms et al., 2022a) Whether the material is switchable (ferroelectric) or not switchable (pyroelectric) 265 under these conditions (in the P3m1 phase above 23 GPa) is unexplored at present. In principle, any electric 266 polarization should increase as the space group progresses from $P3m1 \rightarrow P3 \rightarrow P1$. (Harms et al., 2022a; 267 Aoyama et al., 2014) Real physical examples of polar metals are rare, so it's useful to consider how NiPS₃ hosts such a state when it has not been sighted (as of yet) in related MPX₃ systems. The current working 268 269 hypothesis is that because NiPS₃ does not favor a $P\bar{3}m1$ -type structure or layer-normal phosphorous chain formation, it may engage in negative charge transfer with hole formation at the S sites. (Harms et al., 270 271 2022a). The optical properties of NiPS₃ have also been of sustained interest (Kim et al., 2018; Kang et al., 272 2020; Lane and Zhu, 2020) - both in single crystal and thin film form - although to our knowledge, there have been no studies under pressure or strain. Interestingly, electric field also drives an insulator-to-metal 273 274 transition in the monolayer (Lane and Zhu, 2020), suggesting that pressure and electric field phase space 275 may share more similarities that previously supposed.

NiPSe₃ is different, primarily because it does not have a complicated set of symmetry progressions under compression. Instead, there is a relatively simple structural sequence as shown in Fig. 10 and Table 3. (Harms et al., 2022a; Sun et al., 2023) More recent efforts involving combined synchrotron x-ray diffraction and first-principles theory confirm these two structural transitions: LP to HP1 (where the honeycomb layers slide relative to each other) and HP1 to HP2 (where the structure has more three-dimensional character) (Sun et al., 2023). There is also a concomitant magnetic crossover at $P_{C,1}$ where the direction of the antiferromagnetic moment switches from out-of-plane to in-plane . Complementary resistance measurements indicate a bandwidth-controlled Mott insulator-metal transition near 8.0 GPa. The latter coincides with the emergence of superconductivity with $T_c \approx 4.8$ K.(Sun et al., 2023) This leads to a remarkable and complex pressure-temperature phase diagram hosting coexisting zigzag antiferromagnetic order and superconductivity [Fig. 11].

Finally, we point out that transition metal dichalcogenides such as MoS₂ and WS₂ and related 287 nanostructures are employed for solid state lubrication due to their ultra-low friction coefficients -288 especially under high shear conditions.(Musfeldt et al., 2020) In an important initiative, Deng et. al 289 are starting to explore the mechanical properties of terenary chalcogenides like NiPS3 using lateral force 290 microscopy. (Deng et al., Friction, under review, (2024) These measurements reveal a friction coefficient 291 of approximately 4.5×10^{-3} under 5×10^{-3} Pa with load up to 768 nN. These findings are discussed in 292 terms of the extremely weak interlayer interactions. Table 1 summarizes other candidates with low layer 293 thickness/van der Waals gap ratios including CdPS₃ and FePSe₃ that should be tested. 294

295 2.4 V_{1-x} PS₃ (x < 1)

The vanadium phosphate synthesis has been achieved with the vanadium deficiency in $V_{1-x}PS_3$ which 296 can be explained as due to valence mixing on the vanadium site between V^{2+} and V^{3+} states (Ouvrard 297 et al., 1985b; Ichimura and Sano, 1991). The LP-phase has an electrical conductivity higher than 10^{-5} 298 Scm^{-1} , while MnPS₃, NIPS₃ and ZnPS₃ crystals lower than 10^{-9} Scm⁻¹ at room temperature. The high 299 conductivity of the former crystals is attributed to their mixed valency on the metal and/or the S and 300 P sites, which has been revealed by x-ray photoelectron spectroscopy. (Ichimura and Sano, 1991) The 301 302 $LP \rightarrow HP$ -I transition starts at 2.6 GPa followed by the gradual transition to HP-I completing near 8.0 GPa. (Coak et al., 2019) The layers of V_{0.9}PS₃ shift relative to each other in a sliding motion of $\sim a/3$ 303 along the a-axis such that the S atoms become arranged in a hexagonal closed packing layout between 304 the layers (Table 2, and Fig.1). (Coak et al., 2019) In HP-I, P atoms are slightly distorted along the a-axis 305 (the x coordinate value is 0.0074 at 17.7 GPa) of the unit cell, and this distortion results in the same 306 C2/m symmetry. (Coak et al., 2019) The volume per formula unit ($V_{f.u.}$) decreases continuously without a 307 sudden or discontinuous changes across the gradual LP \rightarrow HP-I transformation (Fig. 3). (Coak et al., 2019) 308 The electrical resistance decreases significantly and continuously with increasing pressure, and then the 309 310 insulator-to-metal transition occurs near 12 GPa above the pressure where the LP \rightarrow HP-I transformation completes. (Coak et al., 2019) Thus, the isostructural insulator-to-metal transition is concluded, suggesting 311 a second-order transition. (Coak et al., 2019) From the magnetoresistance data, Coak et al. claim the ITM 312 involves the antiferromagnetic to paramagnetic transition. (Coak et al., 2019) The R vs. T curve in the 313 314 metallic phase shows an upturn at low temperatures. (Coak et al., 2019) Coak et al. suggest that Kondo effect due to the vanadium deficiency and disordered valence mixing on the vanadium sites. (Coak et al., 315 2019) The synthesis of VPSe₃ has not been reported to the knowledge of the authors. 316

317 2.5 CoPS₃ and CoPSe₃

Like other structural siblings, $CoPS_3$ has gained significant interests recently for the study of twodimensional magnetism. Since Co^{2+} ion is in the d^7 configuration with a hole in the t_{2g} shell, significant effects of spin-orbit coupling on magnetism is expected. Indeed, recent theoretical studies suggested an interesting way to realize the so-called Kitaev's exchange interactions in cobalt-based compounds, (Kitaev, 2006; Liu and Khaliullin, 2018; Sano et al., 2018) which were originally considered in heavier transition metal systems. (Jackeli and Khaliullin, 2009) Several Co-based systems including CoPS₃ have been studied for the search of the Kitaev-induced frustrated magnetism. (Kim et al., 2020, 2021; Lin et al., 2021; Zhang et al., 2023; Kim et al., 2023) Considering the presence of metastable spin states and significant spin-orbit coupling in Co ions, (Yamaguchi et al., 1996; Zobel et al., 2002) it is expected that pressure may induce intriguing evolution of electronic and magnetic properties of CoPS₃.

 $CoPS_3$ crystallizes in C2/m structure at ambient pressure as other MPS_3 family at ambient pressure. 328 CoPS₃ exhibits AFM at ambient pressure with a $T_{\rm N} = 122$ K and a Weiss temperature of $\theta = -166$ 329 K. (Ouvrard et al., 1982; Wildes et al., 2017) Single-crystal neutron diffraction shows that the magnetic 330 propagation vector is $\mathbf{k} = [010]$ with the moments mostly along the *a*-axis and with a small component 331 along the c-axis. (Ouvrard et al., 1982; Wildes et al., 2017) This magnetic structure is in sharp contrast to 332 MnPS₃ and FePS₃, as observed for NiPS₃. Its effective moment is 4.9 $\mu_{\rm B}$, slightly larger than the expected 333 value for a pure spin moment of a Co^{2+} cation (Ouvrard et al., 1982), implying orbital contribution to the 334 magnetization as mentioned above. In addition, an inelastic neutron scattering study suggests a XXZ-type 335 336 antiferromagnetic order in CoPS₃, signalling the presence of the spin-orbit coupling-induced magnetic anisotropy. (Kim et al., 2020) 337

High-pressure properties of CoPS₃ have not been experimentally reported until recently, probably 338 due to the significant difficulty in the synthesis and single crystal growth of this compound. A first-339 340 principles calculation from an article dedicated to the pressure effects in CoPS₃ predicts a pressure-driven isostructural Mott transition accompanied by a spin crossover. (Gu et al., 2021) XRD and Raman scattering 341 measurements reveal that CoPS₃ exhibits C2/m (LP) $\rightarrow P\overline{3}$ (HP-I) structural transformation (Figure 1 and 342 Table 2) at 7 GPa and room temperature accompanied by the small drop (2.9%) of the volume per formula 343 unit (Figure 3). (Matsuoka et al., 2023) This volume collapse and comparing FePS₃ and MnPS₃, it is 344 concluded that CoPS₃ exhibits the spin crossover ($S = 3/2 \rightarrow 1/2$) accompanied by the metallization. 345 On the other hand, the 2.9% of volume collapse is much smaller than that of FePS₃ (10.6%) and MnPS₃ 346 (19.7%) (Figure 3 and 3). (Matsuoka et al., 2023; Wang et al., 2016; Haines et al., 2018; Wang et al., 2018b) 347 The ionic radii of high- (HS, 0.89 Å) and low-spin (LS, 0.79 Å) Co^{2+} ions make the HS \rightarrow LS radius 348 reduction 11.2%, which is not much smaller than that of Mn^{2+} (HS: 0.97 Å, LS: 0.81 Å, 16.5%) and Fe²⁺ 349 (HS: 0.92 Å, LS: 0.75 Å, 18.5%) (Shannon, 1976). Thus, the slight volume reduction of CoSP₃ cannot 350 be explained solely by the difference between HS and LS radii. It is also noticeable that the volume per 351 formula unit and c-axis in the $P\overline{3}$ phase show a steeper compression between 7 GPa and 12 GPa followed 352 by moderate compression above 12 GPa, indicating a sign of structural stabilization. (Matsuoka et al., 2023) 353 These series of changes in compression behavior suggest an electronic transition in the $P\overline{3}$ phase. Upon 354 355 decompression at room temperature, the $P\overline{3}$ phase remains down to 2 GPa revealing a large hysteresis in 356 the volume per formula unit and c-axis. (Matsuoka et al., 2023)

357 Concomitantly with the structural transition, the electrical resistance decreases significantly and $CoPS_3$ becomes metallic. (Matsuoka et al., 2023) The positive Hall resistivity reveals this metallic $CoPS_3$ is a hole-358 359 dominant conductor with multiple conduction bands, and its non-saturating behavior against the external magnetic field, indicating the absence of the anomalous Hall effect, suggests that the metallic CoPS₃ is not 360 ferromagnetic (Figure 12). (Matsuoka et al., 2023) The linear magnetoresistance $(\Delta \rho_{xy}(B) / \rho_{xx}(B = 0))$, 361 where $\Delta \rho_{xx}(B) = \rho_{xx}(B) - \rho_{xx}(B = 0)$, along with the small volume collapse at the metallization, 362 suggest the incomplete high-spin \rightarrow low-spin transition in the metallic phase (Figure 12). (Matsuoka et al., 363 364 2023) It is expected that the metallic $CoPS_3$ possibly possesses an inhomogeneous magnetic moment distribution and short-range magnetic ordering. (Matsuoka et al., 2023) Furthermore, the magnetoresistance 365 366 changes from linear to more quadratic magnetic field dependent as pressure increases. (Matsuoka et al., 367 2023) Considering the suggested electronic transition from the volume per formula unit vs. pressure slope across 12 GPa (Figure 3), it can be thought that the spin crossover proceeds with pressure up to 12 GPa and 368

369 stops or progress moderately above 12 GPa in the $P\overline{3}$ phase (Matsuoka et al., 2023). 370

371 2.6 CdPS₃ and CdPSe₃

CdPS₃ crystallizes in the C2/m structure with the monoclinic angle $\beta = 107.27^{\circ}$, possessing an insulating 372 electrical character like other MPX₃ family members (Ouvrard et al., 1985a; Niu et al., 2022). The Raman 373 scattering measurements combined with ab-initio calculation reveal that CdPS₃ exhibits two structural 374 transformations the LP-phase $(C2/m) \rightarrow$ HP-I phase (R3) that initiates near 0.11 GPa and completes near 375 1.1 GPa GPa (Niu et al., 2022). The HP-I ($R\overline{3}$) \rightarrow HP-II ($R\overline{3}$) unedergoes from 7.35 GPa to 10.2 GPa (Niu 376 et al., 2022). The $C2/m \rightarrow R\overline{3}$ transition is obtained by sliding a single layer of the LP-phase along the 377 B axis for a distance of about $\frac{b}{6}$ and simultaneously sliding it for approximately $\frac{a}{3}$ along the a axis. This 378 results in the C2/m unit's β angle changing from 107.27° to approximately 90° (Niu et al., 2022). The HP-I 379 $(R\overline{3})$ to HP-II $(R\overline{3})$ transition is instead caused by the distortion of CdS6 polyhedrons (Niu et al., 2022). 380 The ab-initio calculation suggests a small volume reduction (-2.05%) with the LP-I \rightarrow HP-I transformation 381 (Niu et al., 2022). On the other hand, the volume vs. P curves of HP-I and HP-II almost overlap across 382 a wide pressure range (2 - 22 GPa), reflecting the transformation nature (Niu et al., 2022). The *ab-initio* 383 calculation also suggests the further transformation from HP-II to HP-III (P31m) at approximately 25 GPa 384 (Niu et al., 2022). Compared with other MPM₃, it is noticeable that the LP-phase of CdPS₃ is relatively 385 more sensitive to external pressure than other MPX_3 reported so far (Table 2). This would suggest that even 386 a slight strain can be an effective stimulus to drive the appearance of a new structural phase of CdPS₃. 387

3 SUMMARY AND FUTURE PROSPECTS

Complex chalcogenides display properties and states of matter that are incredibly sensitive to external 388 stimuli. This particular review is inspired by opportunities to employ pressure to unveil elusive states of 389 matter and uncover charge-structure-function relationships. Compression is particularly well-suited to 390 tuning the properties of chalcogenides because, in addition to changing bond lengths and angles, it provides 391 deterministic control of the c/a structural ratio and the van der Waals gap. By controlling the environment 392 around the metal center, compression also controls spin. As emphasized throughout this review, strain is 393 likely to offer complementary advantages. At the same time, expanding the chemical phase space of the 394 MPX₃ system to include solid solutions like (Zn,Cd)PS₃ or (Ni,Co)PS₃, heavy 4- and 5d- metal centers 395 to enhance spin-orbit coupling, the CrPS₄ member of this family, and dual sublattice analogs such as 396 bimetallic CuInP₂S₆ and AgInP₂S₆ provides exciting new platforms for exploring ferroelectricity, new 397 types of photoinduced behavior, ionic motion, and even charge pumping under compression. 398

CONFLICT OF INTEREST STATEMENT

The authors declare that the research was conducted without any commercial or financial relationships that could be construed as a potential conflict of interest.

AUTHOR CONTRIBUTIONS

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Figure 1. Crystal structures of MPS_3 (M = V, Mn, Fe, Ni, Co, Cd) famility of materials in the low- and highpressure phases visualized using VESTA3(Momma and Izumi, 2011) based upon reported experimental and theoretical results. (Coak et al., 2019; Ouvrard et al., 1985a; Harms et al., 2020; Haines et al., 2018; Matsuoka et al., 2023; Harms et al., 2022a) The bottom right shows layer sliding in C2/m symmetry with decreasing monoclinic angle β under pressure. When $\beta = 90^{\circ}$, the honeycomb pattern of metal ions overlaps between adjacent layers, and the system transforms to a trigonal high pressure phase.

FIGURE CAPTIONS



Figure 2. Magnetic moment and spin state as a function of pressure for MnPS₃, MnPSe₃, FePS₃, and FePSe₃ relative to properties at ambient conditions. Data taken from (Wang et al., 2016) and (Wang et al., 2018b).



Figure 3. Volume per formula unit as a function of pressure. (a) MPS_3 (M = V, Mn, Fe, Co, Ni). (b) MnPS₃ and FePS₃ with MnPSe₃ and FePSe₃ for comparison. The circles denote the low (or ambient) pressure phase, which is semiconducting, including the HP-I phases for MnPS₃ and FePS₃. The squares indicate high pressure metallic phases. The solid lines guide the eyes.



Figure 4. Spectral functions from DFT+DMFT calculations showing the *k*-resolved and orbital-projected spectral functions calculated with the aligned phosporous dimer (APD) and aligned phosphorous chain (APC), respectively.(Harms et al., 2020)



Figure 5. Photographs of piezochromic MnPS₃ inside the diamond anvil cell at several characteristic pressures and also after release at room temperature. The scale bar (left panel) is 200 μ m.(Harms et al., 2020).



Figure 6. Crystal and magnetic structures of the ambient and the first high-pressure (HP-I) phases of FePS₃. (a) The magnetic cell of FePS₃ with the $\mathbf{k} = [0, 1, 1/2]$ propagation vector - overall antiferromagnetic order, consisting of a zig-zag chain antiferromagnetically coupled to their neighbors along a-axis and between the planes (Lançon et al., 2016). (b) The magnetic unit cell with $\mathbf{k} = [0, 1, 0]$ propagation vector, where the planes are ferromagnetically aligned (Coak et al., 2021). (c) Pressure-temperature phase diagram of FePS₃ from magnetization measurements and neutron scattering experiments. Blue circles, red crosses (and squares) are from magnetization measurements and integration of neutron spectra, respectively. Dashed lines are fits to the data points, and vertical lines mark structural transitions. Inset shows normalized magnetization versus temperature with increasing pressure. The plot is from (Coak et al., 2021).



Figure 7. High-pressure electronic states of FePS₃ and superconducting FePSe₃ (**a**) Enthalpy difference from DFT calculations as a function of the monoclinic β angle in the *C*2/*m* structure for the ambient (LP) and HP-I AFM phases of FePS₃ at 1.0 GPa. Panel from Deng et al. (2023). (**b**) Heat map of magnetic peaks at 80 K in FePS₃, showin the presence of magnetic signals in the HP-II (P > 15 GPa) phase. Panel from Coak et al. (2021). (**c**) The temperature dependence of the electrical resistance of FePSe₃ at 29.6 GPa under magnetic fields of 0, 0.1, 0.25, 0.5, 1, 2, and 5 T. Inset shows the field dependence of T_c for FePSe₃ at 29.6 GPa. (**d**) Temperature–pressure phase diagram of FePSe₃. Solid circles: the pressure dependence of the onset superconducting transition temperatures (5* T_c), blue triangles: the pressure dependence of Hall coefficient, T_N denoting the Néel temperature. Panel (**c**) and (**d**) from (Wang et al., 2018b).



Figure 8. Vibrational properties of NiPS₃ under compression. (**a-d**) Close-up view of the infrared absorption and frequency vs. pressure plots. (**e-h**) Close-up view of the Raman scattering response and frequency vs. pressure plots. Spectra for the ambient pressure (LP) phase are shown in gray; those for the high pressure phases (HP-I, HP-II, and HP-III) are shown in teal, green, and black, respectively. Closed and open circles represent two independent runs. Note that infrared spectra in the HP-III phase are not included because the phonons are screened by the metallic response. The critical pressures (P_C 's) separate the high pressure phases and are indicated with dashed vertical gray lines. Panels (**a**)-(**h**) from (Harms et al., 2022a)



Figure 9. High pressure x-ray diffraction results of NiPS₃ up to 39 GPa. (**a**) Integrated x-ray diffraction profiles at pressures. The appearance of x-ray diffraction peaks at the given angle are indicated, alongside dashed lines to guide the eye. Critical pressures, dividing individual pressure phases, are indicated by solid blue horizontal lines. The space groups of individual high pressure phases are labeled. The asterisk (*) and plus (+) marks correspond to the diffraction response from NaCl in B1-type structure (below 30 GPa) and B2-type (above 30 GPa). (**b**) and (**c**) The x-ray diffraction images on the detector for HP-I (13 GPa) and HP-III (24 GPa). The peaks from NiPS₃ are enclosed with squares alongside the *hkl* indices for each phase. The gray labels in the panel (**c**) indicate the diffraction from HP-II. The Bragg diffraction from the diamond anvils is masked with red circles.Panels (**a-c**) from (Harms et al., 2022a)



Figure 10. Crystal structures of NiPS₃ and NiPSe₃ in LP ((**a**) and (**d**)), HP-I ((**b**) and (**e**)), and HP-II ((**c**) and (**f**)) phases visualized using VESTA3 (Momma and Izumi, 2011) based on the experimental reports.(Harms et al., 2022a; Sun et al., 2023) As discussed in the text, NiPS₃ displays an unusual sequence under pressure: C2/m with $\beta \sim 108^{\circ} \rightarrow P\overline{3} \rightarrow P\overline{3}m1 \rightarrow P31m$ and so on. This structural sequence is not seen in other members of the MPS₃ family. On the other hand, NiPSe₃ is similar to FePS₃ (C2/m with $\beta \sim 108^{\circ} \rightarrow P\overline{3}1m$) and MnPS₃ (C2/m with $\beta \sim 108^{\circ} \rightarrow P\overline{3}1m$). The primary difference between the LP-phases of NiPS₃ and NiPSe₃ is the angle of the P-P dimer. The P-P dimer in NiPS₃ aligns almost perpendicular to the *ab* plane. On the other hand, the dimer in NiPSe₃ aligns mostly in an out-of-plane direction but with an angle.



Figure 11. Phase diagram of NiPSe₃ under pressure. The insets illustrate the magnetic structures, i.e. zigzag-out for the LP phase and zigzag-in for the HP-I phase. $T_{\rm N}$ for AFM order and $T_{\rm C}$ for superconductivity are extracted from resistance measurements and DFT calculations. Created based on data from (Sun et al., 2023).



Figure 12. Transverse magnetotransport data from metallic CoPS₃. (a) The ρ_{xy} at different temperatures. The data at 1.8 K and 20 K at 10 GPa overlap (top panel). The data at 1.8 K and 10 K at 13 GPa overlap (middle panel). (b) The magnetoresistance. The dotted (black) and dashed (blue) lines in the top panel indicate the linear fit, y = Ax + C and quadratic fits $y = Bx^2$ to the data at 10 GPa and 1.8 K. (Matsuoka et al., 2023) Reprinted (Figure 5) with permission from [Matsuoka, T., Rao, R., Susner, M. A., Conner, B. S., Zhang, D., and Mandrus, D., Pressure-induced insulator-to-metal transition in the van der Waals compound CoPS₃, 107, 165125 (2023)).] Copyright (2023) by the American Physical Society.

	$V_{0.78}PS_3$	MnPS ₃	FePS ₃	CoPS ₃	NiPS ₃	CdPS ₃	MnPSe ₃	FePSe ₃
layer thickness (Å)	6.363	6.487	6.423	6.361	6.343	6.546	6.673	6.604
vdw gap (Å)	3.227	3.247	3.250	3.243	3.258	3.218	3.275	3.174
(layer thickness)/(vdW gap)	0.507	0.500	0.506	0.510	0.514	0.492	0.491	0.481

Table 1 Inter-layer distance and van der Waals (vdw) gap of *MPS*₃ and *MPSe*₃. Data are from (Ouvrard et al., 1985b,a; Wiedenmann et al., 1981)

	P (GPa)	Phase	Symmetry	Lattice constants	Ref.
$V_{0.9}PS_3$	1.1	AP	C2/m	a = 5.8436(15) Å	(Coak et al., 2019)
				b = 10.0876(8) Å	
				c = 6.5237(18) Å	
				$\alpha = \gamma = 90^{\circ}, \beta = 107.098(5)^{\circ}$	
	1.1	HP-I	C2/m	a = 5.5469(3) Å	(Coak et al., 2019)
				b = 9.5892(6) Å	
				c = 5.4788(9) Å	
				$\alpha = \gamma = 90^{\circ}, \beta = 90.136(8)^{\circ}$	
$MnPS_3$	0	AP	C2/m	a = 6.077(1) Å,	(Ouvrard et al., 1985a)
				b = 10.524(3)Å	
				c = 6.796(1) Å	
			_	$\alpha = \gamma = 90^{\circ}, \beta = 107.35(2)^{\circ}$	
		HP-I	P31m	a = b = 5.725 A	(Harms et al., 2020)
				c = 5.758 A	
	22			$\alpha = \beta = 90^{\circ}, \gamma = 120^{\circ}$	
	33	HP-II	C2/m	a = 4.858(3) A	(Wang et al., 2016)
				b = 8.171(5) A	
				c = 0.203(3) A	
EaDC	0	۸D	Color	$\frac{\alpha = \beta = 90^{\circ}, \gamma = 107.98^{\circ}}{5.0428(0)^{\circ}}$	$(\mathbf{H}_{a}; \mathbf{r}_{a}, \mathbf{r}_{a}, \mathbf{r}_{a}, 1, 2010)$
rep53	0	AP	C2/m	a = 5.9428(9) A b = 10.200(2) Å	(Hames et al., 2018)
				b = 10.299(2) A c = 6.716(2) Å	
				$\alpha = \alpha = 00^{\circ} \beta = 107.34(2)^{\circ}$	
	10	HP-I	C^{2}/m	$a = 7 = 50$, $\beta = 101.54(2)$ a = 5.7620(12) Å	(Haines et al. 2018)
	10	111 1	02/11	b = 9.988(2) Å	(11411105 01 41., 2010)
				c = 5.803(5) Å	
				$\alpha = \beta = 90^{\circ}, \gamma = 89.33(2)^{\circ}$	
	18	HP-II	$P\overline{3}m1$	a = b = 5.699(4) Å	(Haines et al., 2018)
				c = 4.818(3) Å	
				$\alpha = \beta = 90^{\circ}, \gamma = 120^{\circ}$	
CoPS ₃	0.5	AP	C2/m	a = 5.844(1) Å	(Matsuoka et al., 2023)
				b = 10.127(1) Å	
				c = 6.562(4) Å	
				$\alpha = \beta = 90^{\circ}, \gamma = 107.04(2)^{\circ}$	
	17	HP-I	$P\overline{3}$	a = b = 5.570(5) Å	(Matsuoka et al., 2023)
				c = 5.13(2) Å	
				$\alpha = \beta = 90^{\circ}, \gamma = 120^{\circ}$	
NiPS ₃	0.7	AP	C2/m	a = 5.800(1) Å	(Harms et al., 2022a)
				b = 10.057(1)Å	
				c = 6.605(2) Å	
				$\alpha = \gamma = 90^{\circ}, \beta = 106.99(2)^{\circ}$	
	13	HP-I	P 3	a = b = 5.638(5) A	

Table 2 Crystal structures for the ambient pressure (AP) and high pressure (HP) phases of MPS₃.

				c = 5.88(2) Å	
	16	HP-II	$P\overline{3}m1$	$\alpha = \beta = 90^{\circ}, \gamma = 120^{\circ}$ $a = b = 9.720(1) \text{ Å}$	
				c = 4.94(1) Å $\alpha = \beta = 90^{\circ}, \gamma = 120^{\circ}$	
	24	HP-III	<i>P</i> 3 <i>m</i> 1	a = b = 5.582(3) Å c = 15.03(2) Å	
	27	HP-IV	Р3	$ \alpha = \beta = 90^{\circ}, \gamma = 120^{\circ} $ a = b = 5.488(1) Å	
				c = 9.92(4) Å $\alpha = \beta = 90^{\circ}, \gamma = 120^{\circ}$	
CdPS ₃	0	AP	C2/m	a = 6.218(1) Å	(Ouvrard et al., 1985a)
				b = 10.763(2) Å	
				c = 6.867(1) Å	
				$\alpha = \gamma = 90^{\circ}, \beta = 107.58(1)^{\circ}$	

	P (GPa)	Phase	Symmetry	Lattice constants	Ref.
MnPSe ₃	0	AP	$R\overline{3}$	a = b = 6.387(2)Å	(Wiedenmann et al., 1981)
				c = 19.996(6) Å	
				$\alpha = \beta = 90^{\circ}, \gamma = 120^{\circ}$	
	27	HP-I	C2/m	a = 5.266(3) Å	(Wang et al., 2016)
				b = 7.600(5) Å	
				c = 7.151(3) Å	
				$\alpha = \gamma = 90^{\circ}, \beta = 108.17^{\circ}$	
FePSe ₃	0	LP	R 3	a = b = 6.262(3) Å	(Wiedenmann et al., 1981)
				c = 6.716(2) A	(Liu et al., 2022)
			_	$\alpha = \beta = 90^{\circ}, \gamma = 120^{\circ}$	
	18	HP-I	R 3	a = b = 5.824 A,	(Wang et al., 2018b)
				c = 19.330 A,	
		4.D		$\alpha = \beta = 90^{\circ}, \gamma = 120^{\circ}$	
N1PSe ₃	2.2	AP	C2/m	a = 6.052(1) A	(Sun et al., 2023)
				b = 10.440(1) A	
				c = 0.705(1) A	
	63	UD I	C2/m	$\alpha = \gamma = 90$, $\beta = 108.45(2)$	(Sup at al. 2023)
	0.5	ПР-I	C2/m	u = 5.898(1)A b = 10.248(1) Å	(Suil et al., 2023)
				c = 6.303(1) Å	
				$\alpha = \gamma = 90^{\circ} \beta = 88.38(1)^{\circ}$	
	25.7	HP-II	$P\overline{3}1m$	a = b = 5.873(2) Å	(Sun et al., 2023)
	_0.7		2 9111	c = 4.274(4) Å	(~~~~~)
				$\alpha = \beta = 90^{\circ}, \gamma = 120^{\circ}$	

Table 3 Crystal structures for the ambient pressure (AP) and high pressure (HP) phases of MPSe₃.