Forum

Structures and Properties of Functional Metal Selenites and Tellurites

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Metal selenites and tellurites are a class of very important compounds. In this paper, the structures and properties of metal selenites or tellurites combining with transition-metal (TM) ions with the d⁰ electronic configuration or tetrahedral MO₄ building blocks of post-transition main-group elements were reviewed. Most compounds in the alkali or alkaline-earth-d⁰ TM-Se^{IV} (or Te^{IV})-O systems exhibit extended anionic architectures composed of distorted octahedra of (d⁰) TM cations and tellurite or selenite groups. The distortion of the octahedron is always away from the lone-pair cation, and some of them exhibit excellent second-order nonlinear optical properties due to the adductive effects of two types of bond polarizations. Because of the high coordination number of Ln^{III} ions, most of compounds in the Ln-d⁰ TM-Se^{IV} (or Te^{IV})-O systems are not second-harmonic-generation active; however, they are able to emit strong luminescence in the visible or near-IR region; also in most cases, the d⁰ TM cations are in tetrahedral geometry and are well separated from selenite or tellurite groups. It is also interesting to note that the selenite group is normally "isolated", whereas the TeO_x (x = 3-5) can be polymerized into a variety of discrete polynuclear anionic clusters or extended architectures via Te-O-Te bridges.

Introduction

Noncentrosymmetric (NCS) compounds exhibit many interesting and useful properties such as ferroelectricity, piezoelectricity, and second-order nonlinear optical (NLO) behavior.¹ With inorganic materials, the macroscopic acentricity is often a manifestation of the asymmetric coordination environments of the cations. One of the important classes of NCS compounds is based on cations susceptible to secondorder Jahn-Teller (SOJT) distortion.² Such a type of distortion can occur in two different types of cations, d⁰ transition metals (TMs; Ti⁴⁺, V⁵⁺, Nb⁵⁺, Mo⁶⁺, W⁶⁺, etc.) and cations with stereoactive lone pairs (Se⁴⁺, Te⁴⁺, Sb³⁺, etc.), both in asymmetric coordination environments. With the octahedrally coordinated d⁰ TMs, SOJT effects occur when the empty d orbitals of the metal mix with the filled p orbitals of the ligands. In extended structures, this mixing results in a host of nearly degenerate electronic configurations that can be removed through the spontaneous distortion of

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the d⁰ TM.² The d⁰ TM cation can be distorted toward either a face (local C_3 direction), an edge (local C_2 direction), or a corner (local C_4 direction) of the MO₆ octahedron. The situation with the lone-pair cations is more complex. The structural distortion and polarization was thought to be through the mixing of the metal cation s and p orbitals. Recently, it is believed that the oxide anion also plays an important role in the lone-pair formation. The interaction of the s and p orbitals of the metal cation with the oxide anion p states is critical for lone-pair formation. No matter how the lone pair is created, its structural consequences are profound because the lone pair "pushes" the oxide ligands toward one side of the cation, resulting in a highly asymmetric coordination environment.³ Studies have shown that compounds containing both types of cations may have a large second-harmonic-generation (SHG) efficiency due to the "constructive" addition of these polarizations, which will be discussed in more detail later.

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Tetrahedral groups such as silicates, phosphates, and germanates are very important building units in zeolites and other porous materials that have been widely used as

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catalysts.⁴ Some of phosphates such as KDP (KH₂PO₄) and KTP (KTiOPO₄) are also very important second-order NLO materials.⁵ Therefore, the combination of these tetrahedral groups with the lone pair containing a Te^{IV} or Se^{IV} cation may also result in new inorganic compounds with novel structural topologies and SHG properties.

It should also pointed out that the transition-metal oxyhalides of Se^{IV} or Te^{IV} display many types of novel structures and interesting magnetic properties such as low-dimensional magnets, etc. Transition-metal tellurium(IV) or selenium(IV) oxyhalides can be regarded as "chemical scissors". The later TM cations form bonds to both oxygen and halide, while lone-pair cations tend to form bonds only to oxygen anions. This chemical difference can be utilized to reduce the dimensionality of the TM arrangements with interesting magnetic properties.⁶ It is amazing that many compounds in above systems can be generally formulated as $M_{n+1}(QO_3)_nX_2$, where M, Q, and X represent the TM, Se (or Te), and the halide anion, respectively. Our exploration of the missing members in the nickel system afforded four new compounds, namely, $Ni_{n+1}(QO_3)_nX_2$ (Q = Se, X = Cl, Br, n = 4; Q = Te, X = Cl, n = 6 and 10).⁷ We also prepared two new cadmium(II) tellurium(IV) oxychlorides, namely, $\{Cd_2(Te_6O_{13})\}\{Cd_2Cl_6\}$ and $Cd_7Cl_8(Te_7O_{17})$, which display several unusual structural building units: two new types of 1D tellurium(IV) oxide anions (Te₆O₁₃²⁻ and $Te_7O_{17}^{6-}$) (Scheme 1), 1D Cd₂Cl₆ double chains, and 2D Cd₇Cl₈ sheets.⁸ We also obtained a series of lanthanide transition-metal tellurium(IV) oxyhalides with three different types of structures, namely, DyCuTe₂O₆Cl, ErCuTe₂O₆Cl, ErCuTe₂O₆Br, Sm₂Mn(Te₅O₁₃)Cl₂, Dy₂Cu(Te₅O₁₃)Br₂, and Nd₄Cu(TeO₃)₅Cl₃.⁹ The halide anion may be involved in metal coordination or remain isolated.

In this paper, our discussion will be focused on only two important systems: combination of Te^{IV} or Se^{IV} with d^0 TMs or with the tetrahedral groups of main-group elements such as SiO₄, GeO₄, BO₄, PO₄, etc. Our group and many other groups have been exploring new NCS compounds in these two systems during the past few years (Table 1). Of course, other Te^{IV} or Se^{IV} compounds are also very important; for **Scheme 1.** Selected Examples of Tellurium(IV) Oxide Anions with Polynuclear Cluster Units or Extended Structures



example, several good SHG materials have been discovered in compounds with two types of lone-pair cations.³

Part I. Combination of d⁰ TM Ions with Se^{IV} or Te^{IV}

1. Alkali (or Alkaline Earth) $-d^{0}$ TM $-Se^{IV}$ (or Te^{IV})-OSystem. This system contains a large number of compounds, some of which possess good SHG properties (Table 1). The TM ions with the d⁰ electronic configuration are mostly Mo⁶⁺ and W⁶⁺, but there are also some examples involving V⁵⁺, Nb⁵⁺, and Ta⁵⁺.

Five compounds with four different structural types were reported in alkali-metal-molybdenum(VI) selenium(IV) oxide systems A₂MoSeO₆ (A = Na⁺, K⁺, Rb⁺) and A₂(MoO₃)₃(SeO₃) (A = NH₄⁺, Cs⁺).¹⁰ The first three compounds feature a 3D anionic network, a 0D anionic dimer, and a 1D anionic chain, respectively. The 3D anionic structure of Na₂MoSeO₆ (NCS space group *P*2₁3) is based on alternative linkages of MoO₆ octahedra (distorted toward a face) and SeO₃ groups, forming tunnels of Mo₂Se₂ fourmembered rings and Mo₄Se₄ eight-membered rings; the sodium cations occupy the larger tunnels. Na₂MoSeO₆ revealed a weak SHG intensity of about 10 × SiO₂. The structure of K₂MoSeO₆ contains two edge-sharing MoO₆ octahedra, each of which is further chelated by a SeO₃ group, forming [Mo₂Se₂O₁₂]⁴⁻ clusters that are separated by K⁺ ions.

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Table 1. TeIV and SeIV Compounds Showing NCS Structures and SHG Properties

compound	space group	structural feature	SHG efficiency	ref
TeO ₂	P41212	3D network of corner-sharing TeO ₄ groups	$5 \times SiO_2$	3b
Te_2O_5	$P2_1$	3D network of corner-sharing Te ⁴⁺ O ₃ groups and Te ⁶⁺ O ₆ octahedra	$400 \times SiO_2$	3b
TeSeO ₄	Ia	3D network of corner-sharing Se ⁴⁺ O ₃ and Te ⁴⁺ O ₅ groups	$400 \times SiO_2$	3a, 3b
Te ₂ SeO ₇	$Pmn2_1$	2D layer composed of $Se^{6+}O_4$ tetrahedra and $Te^{4+}O_4$ groups	$200 \times SiO_2$	3b
Bi ₂ TeO ₅	Abm2	3D network of corner-, edge-, or face-sharing BiO ₃ , BiO ₅ , and TeO ₃ groups	$300 \times SiO_2$	3c
Na ₂ MoSeO ₆	P2 ₁ 3	3D anionic structure based on alternative linkages of MoO ₆ octahedra and SeO ₃ groups	$10 \times SiO_2$	10a
Na2Mo3Te3O16	12	1D chain consisting of Mo_3O_{14} trimers that are connected $Te_3O_8^{4-}$ anions	$500 \times SiO_2$	11a
(NH ₄) ₂ WTe ₂ O ₈	$P2_1$	2D layer consisting of WO ₆ octahedra connected to TeO ₄ polyhedra	$250 \times SiO_2$	12a
Na ₂ W ₂ TeO ₉	Ia	3D structure built from a corrugated 2D tungsten oxide layer further interconnected by TeO ₃ groups	$500 \times SiO_2$	12b
Rb ₂ W ₃ TeO ₁₂	P31c	2D tungsten oxide layer capped by TeO ₃ groups on one side	$200 \times SiO_2$	13b
Cs ₂ W ₃ TeO ₁₂	$P6_3$	2D tungsten oxide layer capped by TeO ₃ groups on one side	$400 \times SiO_2$	13b
BaMo ₂ TeO ₉	$P2_1$	2D layer built by dimers of edge-sharing MoO_6 octahedra that are interconnected by TeO_3 groups	$600 \times SiO_2$	14b
BaW ₂ TeO ₉	$P2_1$	2D layer built by dimers of edge-sharing WO_6 octahedra that are interconnected by TeO_3 groups	$500 \times SiO_2$	14b
$Cd_4V_2Te_3O_{15}$	<i>P</i> 2 ₁ 2 ₁ 2 ₁	3D network in which the cadmium tellurite layers are further interconnected by both "isolated" VO ₄ tetrahedra and 1D vanadium-oxide helical chains	$1.4 \times \text{KDP}$	23
$B_2Se_2O_7$	$P2_{1}2_{1}2_{1}$	3D network composed of $B_2 O_7{}^{8-}$ anions interconnected by Se^{IV} atoms	$2.2 \times \text{KDP}$	27

In Rb₂MoSeO₆, the MoO₆ octahedra are interconnected into a 1D chain via corner-sharing, the SeO₃ groups are hanging on both sides of the chain, and each of them shares an edge with a MoO₆ octahedron.^{10a} The last two compounds (space group *P*6₃) are isostructural and feature a 2D hexagonal tungsten bronzelike anionic layer of MoO₆ octahedra capped on one side by pyramidally coordinated Se^{IV};^{10b} such a layer was also reported in Cs₂(MoO₃)₃(TeO₃) and A₂(WO₃)₃(TeO₃) (A = Rb, Cs).^{11b,12c}

Compounds with three different structural types have been reported in alkali-Mo⁶⁺-Te⁴⁺-O systems.¹¹ Na₂Mo₃Te₃O₁₆ in NCS space group I2 exhibits a quasi-1D crystal structure, with each chain consisting of Mo₃O₁₄ trimers (composed of three edge-shared MoO₆ octahedra) that are connected to Te₃O₈⁴⁻ anions (formed by one TeO₄ group corner-sharing with two TeO₃ groups). Both of the Mo^{6+} and Te^{4+} cations are in the asymmetric coordination environments attributable to SOJT effects. The Mo⁶⁺ cations are distorted toward an edge of the MoO_6 octahedron (local C_2 direction). The SHG efficiency of Na₂Mo₃Te₃O₁₆ is approximately $500 \times \alpha$ -SiO₂ and is phase-matchable. The strong SHG efficiency is maintained up to the melting temperature (around 450 °C).^{11a} The use of other alkali-metal ions led to two other structural types.^{11b} $A_2Mo_3TeO_{12}$ (A = NH₄, Cs) with noncentrosymmetric space group P63 contain 2D hexagonal tungsten oxide related $(Mo_3TeO_{12})^{2-}$ anionic layers interleaved with $NH_4^+/$ Cs^+ ions. The TeO₃ groups cap on the same side of the anionic layer. $A_4Mo_6Te_2O_{24} \cdot 6H_2O$ (A = Rb, K) is composed of discrete centrosymmetric (Mo₆Te₂O₂₄)⁴⁻ anionic aggregates and alkali-metal ions. In this hexamolybdoditellurite anion, the Mo₆O₂₄ flat hexagonal ring, formed by edgesharing of six MoO₆ octahedra, is capped by tellurium on both sides. Isolated Mo₆O₁₈ and Mo₅O₁₅ cyclic clusters capped by SeO₃ groups were also reported with amines as template cations.¹³ The Mo⁶⁺ ions are distorted toward a face of the MoO₆ octahedron in A₂Mo₃TeO₁₂ (A = NH₄⁺, Cs⁺), whereas they are distorted toward an edge in A₄Mo₆Te₂O₂₄•6H₂O (A = Rb⁺, K⁺). The Te⁴⁺ cations in both types of compounds adopt the same asymmetric coordination environments and are bonded to three O atoms.

Three types of compounds were found in the alkali-Te^{IV}-W^{VI}-O systems, namely, (NH₄)₂WTe₂O₈, $Na_2W_2TeO_9$, and $A_2W_3TeO_{12}$ (A = K⁺, Rb⁺, Cs⁺).¹² (NH₄)₂WTe₂O₈ crystallizes in the noncentrosymmetric (NCS) polar space group $P2_1$ (No. 4). It exhibits a 2D structure consisting of WO₆ octahedra connected to TeO₄ polyhedra; the ammonium cations are located at the interlayer region. Both W⁶⁺ and Te⁴⁺ cations are in asymmetric coordination environments attributable to SOJT effects. The WO₆ octahedron exhibits three short and three long W-O bonds. (NH₄)₂WTe₂O₈ revealed a moderate SHG efficiency of approximately $250 \times \alpha$ -SiO₂.^{12a} The noncentrosymmetric $Na_2W_2TeO_9$ (space group Ia) exhibits a 3D structure comprising distorted WO₆ octahedra linked to asymmetric TeO₃ groups. The WO₆ octahedra form a corrugated 2D tungsten oxide layer through corner-sharing; these layers are further interconnected by TeO₃ groups. Both Te^{IV} and W^{VI} cations are in local acentric environments attributable to SOJT effects. Powder SHG measurements on polycrystalline Na₂W₂TeO₉ indicated a strong SHG intensity of approximately $500 \times SiO_2$, and the material is phase-matchable (type I).^{12b} Cs₂W₃TeO₁₂ is isostrutural with Cs₂Mo₃TeO₁₂.^{11b} Although K₂W₃TeO₁₂ and Rb₂W₃TeO₁₂ have chemical formulas similar to that of Cs₂W₃TeO₁₂, their structures are somehow different. All three compounds contain the same

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2D tungsten oxide layer of corner-sharing WO₆ octahedra with W₃ and W₆ rings. The TeO₃ groups in K₂W₃TeO₁₂ act as interlayer linkers to form a 3D structure. In Rb and Cs compounds, the TeO₃ groups only cap on the same side of the W₃ rings; therefore, the compounds are 2D. The K compound is centrosymmetric, whereas the Rb and Cs compounds are acentric and revealed strong SHG efficiencies of 200 and 400 × SiO₂, respectively. These can be explained by the different radii of the alkali-metal ions; the larger one will have larger interlayer O···O separations, which allows a similar capping of TeO₃ groups and the formation of noncentrosymmetric structures.^{12c}

The $AE^{2+}-Mo^{6+}(W^{6+})-Se^{4+}(Te^{4+})-O$ systems are still less explored.14 Two compounds in the Ba-Mo-Se-O system were reported, BaMoO₃(SeO₃) and BaMo₂O₅(SeO₃)₂. BaMoO₃(SeO₃) exhibits a layered structure in which MoO₆ octahedra (distorted toward a face) are bridged by SeO₃ groups, forming Mo₂Se₂ four-membered rings and Mo₃Se₃ six-membered rings. BaMo₂O₅(SeO₃)₂ is acentric ($Cmc2_1$) and features a 3D network composed of pairs of cornersharing MoO₆ octahedra bridged by SeO₃ groups. The MoO₆ octahedron is distorted toward an edge (the local C_2 direction).14a BaMo2TeO9 and BaW2TeO9 are isostructural and crystallized in the polar space group $P2_1$; their structures feature an anionic layer composed of MO₆ octahedra linked to the asymmetric TeO₃ polyhedra. The MoO₆ octahedra in BaMo₂TeO₉ are distorted toward a face (along the local C_3 [111] direction) with three short and three long Mo-O distances. The WO₆ octahedra in BaW₂TeO₉ exhibit two types of distortion: toward a face as discussed above and toward an edge with two short, two normal, and two long W-O bonds. Two MO₆ octahedra form a dimer via a M-O-M bridge and such dimeric units are further interconnected by TeO₃ groups via M-O-Te bridges. BaMo₂TeO₉ and BaW2TeO9 revealed extremely strong SHG responses of 600 and 500 \times SiO₂, respectively.^{14b}

Several phases were reported in A-V5+-Se4+-O systems: AVSeO₅ (A = Rb, Cs), A(VO₂)₃(SeO₃)₂ (A = K^+ Rb^+ , Cs^+ , NH_4^+), and the V^{5+}/V^{4+} mixed-valent KV_2SeO_7 .¹⁵ $A(VO_2)_3(SeO_3)_2$ (A = K⁺, Rb⁺, Cs⁺, NH₄⁺) are isostructural with $A_2(MoO_3)_3(SeO_3)_2$ (A = NH₄⁺, Cs⁺),^{10b} featuring hexagonal layers of corner-sharing VO₆ octahedra, with the SeO₃ groups capping on one side of the vanadium(V) oxide layer; the interlayer distances are much shorter than those of the corresponding Mo compounds because of the much fewer cations needed to balance the charge.^{15a,b} The structure of the polar AVSeO₅ (A = Rb, Cs; space group $P2_1$) is a 3D anionic network formed by alternative linkages of VO₅ square pyramids and SeO₃ groups via corner-sharing, creating two types of helical tunnels along the b axis composed of four- and eight-membered rings. The alkali-metal ions occupy the large tunnels.^{15b} KV₂SeO₇ contains both V⁵⁺ and V^{4+} ions, in octahedral and tetrahedral coordination environments, respectively. Its structure features a double layer of $\{V_2SeO_7\}^-$ composed of corner-sharing VO₆ octahedra, VO₄ tetrahedra, and SeO₃ groups. 15c

Only one compound was reported for the alkaline earth– V^V – Se^{IV} –O family. $Ba_{2.5}(VO_2)_3(SeO_3)_4 \cdot H_2O$ exhibits a 2D-layered structure consisting of layers of VO₅ square pyramids linked to SeO₃ polyhedra, with the Ba^{2+} cations and water molecules occupying the interlayer region.¹⁶ Each layer consists of eclipsed 12-membered V₆Se₆ rings and smaller eight-membered V₄Se₄ rings. Each V⁵⁺ is bonded to five O atoms in a distorted square-pyramidal environment with two "short" [1.642(9) and 1.647(9) Å] and three "normal" bonds [1.952-(8)–2.006(9) Å]. Three of the five O atoms are further bonded to Se⁴⁺ cations, whereas the two "short" V–O bonds remain terminal. The selenite groups adopt two types of binding modes: tridentate bridging with three Se–O–V bridges and bidentate with only two Se–O–V bridges.

Reports on $A(Ae)-Nb^{5+}$ (or $Ta^{5+})-Se^{4+}$ (or $Te^{4+})-O$ systems are still rare.¹⁷ The structural backbone of Na14Nb3Te49O18 may be considered as two sets of cornershared NbO₆ octahedral chains. The "first" chain consists of one row of corner-shared NbO₆ octahedra that are linked by TeO₃ and TeO₄ polyhedra, whereas the "second" chain consists of two rows of corner-shared NbO₆ octahedra that are linked by TeO₃ groups. Each of these chains of octahedra runs parallel to the [010] direction. The "first" and "second" chains are linked by the TeO₃ and TeO₄ groups, along the [100] and [010] directions, resulting in the 3D topology. The Te⁴⁺ cation that connects the two chains is partially occupied. The Na⁺ cations reside in the spaces between the two chains. The second compound, NaNb₃Te₄O₁₆, also exhibits a 3D structure consisting of corner-linked NbO₆ octahedra that are connected to asymmetric TeO₃ and TeO₄ groups. Similar to Na_{1.4}Nb₃Te_{4.9}O₁₈, the structural backbone of NaNb₃Te₄O₁₆ may also be considered as an infinite chain of corner-shared NbO₆ octahedra. The chain consists of three corner-shared NbO₆ octahedra running along the [001] direction. These octahedra are infinite, through additional corner-sharing, in the [010] direction. The TeO₃ and TeO₄ groups serve to link these NbO₆ octahedra. The Na⁺ cations reside in the spaces between the NbO₆ octahedra. In both compounds, the distortion of the NbO₆ octahedron is along the local C_4 direction.^{17a} The crystal structures of $Ba_2M_6Te_2O_{21}$ (M = Nb, Ta) are dominated by the $[M_6O_{21}]^{12-}$ anionic network with incorporated Ba²⁺ and Te⁴⁺ ions. Tunnels based on M₄, M₃Te and M₆Te₂ rings are formed along the [010] direction; the Ba²⁺ ions occupy the tunnels of eight-membered rings.^{17b}

2. $Ln-d^0$ TM-Te^{IV} or Se^{IV}-O Systems. Only three compounds, namely, LaNbTeO₆ and La₄M₂Te₆O₂₃ (M = Nb, Ta), were reported by the Halasyamani group before our group's work on this research field.¹⁸ LaNbTeO₆ consists of 1D corner-linked chains of NbO₆ octahedra that are

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connected by TeO₃ polyhedra. La₄M₂Te₆O₂₃ (M = Nb, Ta) also has a 1D crystal structure, consisting of different types of chains that run parallel to the c axis. One chain consists of corner-linked MO₆ octahedra. Each MO₆ octahedron shares additional corners with TeO3 and TeO4 groups. The TeO₃ polyhedra are separated from each other, but the TeO₄ groups are connected through an O atom into the TeO4 "chain", which is broken at random intervals because of the defects on the bridging O atoms. The other chain is the isolated TeO₄ chain that is also broken at random intervals because of the defects. In all three compounds, the Nb⁵⁺ or Ta⁵⁺ cation is distorted toward an edge with two "short", two "normal", and two long M⁵⁺–O bonds; the short ones are terminal and the long ones are connected to a Te⁴⁺ cation, whereas the remaining two bonds are involved in the $M^{5+}-O-M^{5+}$ bridges. In all three compounds, Te^{4+} cation is in an asymmetric coordination environment attributable to its stereoactive lone pair.

We think that a slight change of the radius of the Ln^{III} ion may induce a completely different structure; in other words, the "lanthanide contraction" may play an important role in the structures formed. Furthermore, lanthanide compounds are capable of emitting strong luminescence in the visible and near-IR regions; both oxyanions of d⁰ TM and Te^{IV} (or Se^{IV}) are good activators for luminescence of the Ln^{III} ions. Like corresponding alkali or alkaline-earth compounds, Ln^{III} compounds with NCS structures may also be formed. Therefore, systematic investigation of the lanthanide selenium(IV) or tellurium(IV) oxides with additional TM ions with a d^0 electronic configuration may give rise to new lanthanide NLO materials or luminescent materials. Solid-state reactions of lanthanide(III) oxide (and/or lanthanide(III) oxychloride), MoO₃ (or WO₃) and TeO₂ at high temperature led to 12 new compounds with 8 different types of structures, namely, Nd2MoSe2O10, Gd2MoSe3O12, La2MoTe3O12, Nd2MoTe3O12, $Ln_2MoTe_4O_{14}$ (Ln = Pr, Nd), $La_2WTe_6O_{18}, Nd_2W_2Te_2O_{13}$, and $Ln_5MTe_7O_{23}Cl_3$ (Ln = Pr, Nd; M = Mo, W).¹⁹

Nd₂MoSe₂O₁₀ can also be formulated as Nd₂(MoO₄)-(SeO₃)₂. Its structure features a 3D network in which the Nd^{III} ions are interconnected by SeO₃²⁻ anions and MoO₄ tetrahedra (Figure 1). Both Nd1 and Nd2 are eightcoordinated by eight O atoms with Nd–O distances in the range of 2.407(7)–2.540(7) Å. The Mo^{VI} atom is in a slightly distorted tetrahedral coordination environment with Mo–O distances in the range of 1.740(7)–1.816(6) Å. The interconnection of Nd1 atoms via bridging selenite groups leads to a $\langle 001 \rangle$ layer, whereas Nd2 atoms are bridged by MoO₄ tetrahedra to form a $\langle 002 \rangle$ layer. The above two types of layers are further interconnected via Nd–O–Se bridges into a 3D network (Figure 1). The lone pairs of the Se^{IV} cations are oriented toward the cavities of the structure.

When the molar ratio of Se/Ln was increased to 3:2, $Gd_2MoSe_3O_{12}$ with a different structure was obtained.



Figure 1. View of the structure of $Nd_2MoSe_2O_{10}$ down the *a* axis. The MoO₄ tetrahedra are shaded in cyan. Nd, Se, and O atoms are drawn as green, yellow, and red circles, respectively.



Figure 2. View of the structure of $Gd_2MoSe_3O_{12}$ down the *b* axis. The MoO₆ tetrahedra are shaded in cyan. Gd, Se, and O atoms are drawn as green, yellow, and red circles, respectively.

Gd₂MoSe₃O₁₂ can also be formulated as Gd₂(MoO₃)(SeO₃)₃, which can be considered as one O²⁻ anion of Nd₂MoSe₂O₁₀ being replaced by the third selenite group. The structure of Gd₂MoSe₃O₁₂ features a 3D network of gadolinium(III) selenite, with the MoO₆ octahedra occupying the cavities of the network. Both Gd atoms in the asymmetric unit are eightcoordinated, with Gd-O distances ranging from 2.337(5) to 2.485(5) Å. Unlike that in Nd₂MoSe₂O₁₀, the Mo^{VI} atom in Gd₂MoSe₃O₁₂ is octahedrally coordinated by three O atoms from three selenite groups and three O2- anions (Figure 2). The MoO₆ octahedron is distorted toward a face (local C_3 direction), exhibiting three "long" and three "short" Mo–O bonds. The magnitude of the distortion (Δ_d) is 1.492. The interconnection of the Gd^{III} ions by selenite groups results in a 3D network with tunnels running along the baxis. The MoO₆ octahedra are located at the tunnels formed by gadolinium(III) selenite (Figure 2).

The structure of La₂MoTe₃O₁₂ is composed of two La³⁺ ions, one MoO₄²⁻ anion, and one Te₃O₈⁴⁻ anion. It can be considered as two selenite groups in Nd₂MoSe₂O₁₀ being replaced by a Te₃O₈⁴⁻ anion. The La³⁺ ion is nine-coordinated by seven O atoms from four Te₃O₈⁴⁻ anions and

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Figure 3. View of the structure of $La_2MoO_4(Te_3O_8)$ down the *a* axis. The MoO₄ tetrahedra are shaded in cyan. La, Te, and O atoms are drawn as green, yellow, and red circles, respectively.

two O atoms from two MoO_4^{2-} anions. The La–O distances range from 2.451(8) to 2.821(8) Å. The trinuclear $Te_3O_8^{4-}$ anion is formed by one TeO₄ group corner-sharing with two TeO₃ groups. The Mo^{VI} cation is in a slightly distorted tetrahedral coordination geometry, with the Mo–O distances ranging from 1.754(8) to 1.793(11) Å. The interconnection of the La³⁺ by chelating and bridging $Te_3O_8^{4-}$ anions led to a 3D network with tunnels along the *a* axis. These tunnels are formed by 10-membered rings (six La and four Te). The MoO₄ polyhedra are located at the above tunnels and connected with the La³⁺ ions via corner-sharing (Figure 3).

Nd₂MoTe₃O₁₂ can be formulated as Nd₂(MoO₄)(TeO₃)-(Te₂O₅); it can also be considered as the Te₃O₈⁴⁻ anion in La₂MoTe₃O₁₂ being replaced by a tellurite and a ditellurite group. Different from the La³⁺ ion in La₂MoO₄(Te₃O₈), the Nd^{III} ion in Nd₂(MoO₄)(TeO₃)(Te₂O₅) is eight-coordinated by eight O atoms, with the Nd–O distances in the range of 2.324(4)–2.603(4) Å. The Te^{IV} cations of the tellurite and ditellurite groups are coordinated by three O atoms in an asymmetric environment. The Te–O distances are in the range of 1.838(4)–1.987(2) Å. The interconnection of Nd^{III} ions by bridging tellurite and ditellurite groups afforded a layered architecture (Figure 4). The MoO₄ tetrahedron was capped on four Nd^{III} ions of the 2D layer with the terminal O1 atom toward the interlayer space (Figure 4). The interlayer distance is about 8.85 Å.^{19a}

The use of more TeO₂ in the syntheses led to $Ln_2MoTe_4O_{14}$ (Ln = Pr, Nd). Both compounds are isostructural and feature a 3D network in which the Ln^{III} ions are interconnected by 1D Te₄O₁₀⁴⁻ chains and MoO₄ tetrahedra (Figure 5). Therefore, Ln₂MoTe₄O₁₄ (Ln = Pr, Nd) can also be formulated as Ln₂(MoO₄)(Te₄O₁₀) (Ln = Pr, Nd). Among two unique Ln^{III} atoms in the asymmetric unit, one is eightcoordinated by eight tellurite O atoms whereas the other is eight-coordinated by six tellurite O atoms and two O atoms from two molybdate anions. The Mo^{VI} atom is in a slightly distorted tetrahedral coordination environment with Mo–O



Figure 4. View of the structure of $Nd_2(MoO_4)(TeO_3)(Te_2O_5)$ down the *c* axis. The MoO₄ tetrahedra are shaded in cyan. Nd, Te, and O atoms are drawn as green, yellow, and red circles, respectively.



Figure 5. View of the structure of $Ln_2MoTe_4O_{14}$ down *a*-axis. The MoO₄ tetrahedra are shaded in cyan. Ln, Te, and O atoms are drawn as green, yellow, and red circles, respectively.

distances in the range of 1.733(4) - 1.779(4) Å. The Te^{IV} atoms are three- or four-coordinated by O atoms in an asymmetric environment. The Te-O distances are in the range of 1.826(3) - 2.457(3) Å. The TeO₃ and TeO₄ groups are interconnected via corner-sharing into a novel 1D $Te_4O_{10}^{4-}$ anionic chain (Scheme 1). The interconnection of Nd1 and Nd2 atoms via bridging Te₄O₁₀⁴⁻ anionic chains led to neodymium(III) tellurium(IV) oxide layers, which are further interconnected by MoO₄ tetrahedra into a 3D network with apertures along the *a* axis. The aperture is formed by 10-membered rings containing six Ln^{III} ions, two MoO₄²⁻ anions, and two TeO₄ groups. The lone pairs of the Te^{IV} atoms are oriented toward the apertures of the structure (Figure 5). It is worth comparing the structures of $Ln_2MoTe_4O_{14}$ (Ln = Pr, Nd) with those of $La_2MoTe_3O_{12}$ and Nd₂MoTe₃O₁₂. All four compounds contain MoO₄ tetrahedra; however, their Te-O architectures are completely different: La₂MoTe₃O₁₂ features discrete Te₃O₈⁴⁻ anions, whereas Nd₂MoTe₃O₁₂ contains both TeO₃²⁻ and dimeric Te₂O₅²⁻ anions, and 1D Te₄O₁₀⁴⁻ anions are formed in $Ln_2MoTe_4O_{14}$ (Ln = Pr, Nd). Nd₂MoTe₃O₁₂ features a layered structure with MoO4 tetrahedra hanging between two



Figure 6. View of the structure of $La_2WTe_6O_{18}$ down the *a* axis. The WO₄ tetrahedra are shaded in cyan. La, Te, and O atoms are drawn as green, yellow, and red circles, respectively.

neighboring layers, whereas the MoO_4 tetrahedra in $La_2MoTe_3O_{12}$ are located at the tunnels of lanthanum(III) tellurium(IV) oxide. In $Ln_2MoTe_4O_{14}$ (Ln = Pr, Nd), the MoO_4 tetrahedron serves as a bridge between two lanthanide(III) tellurium(IV) oxide layers.

The above synthetic methods were also applied to the Ln^{III}-W^{VI}-Te^{IV}-O systems. La₂WTe₆O₁₈ was obtained when we attempted to prepare the $W^{\ensuremath{\text{VI}}}$ analogue of La2MoTe3O12. La2WTe6O18 can also be formulated as $La_2(WO_4)(Te_3O_7)_2$. The La1 atom is 10-coordinated by seven O atoms from two $Te_3O_7^{2-}$ anions and three O atoms from three WO₄ tetrahedra. The La-O distances range from 2.487(4) to 2.884(8) Å. The W^{VI} atom is tetrahedrally coordinated by four O atoms, with the W-O distances ranging from 1.724(6) to 1.786(1) Å. The Te^{IV} atom is coordinated by four O atoms with Te-O distances in the range of 1.850(4) - 2.193(1) Å. The TeO₄ groups are interconnected into a Te₃O₇²⁻ layer via corner-sharing (Scheme 1). Within the layer, three-membered rings and "pear-shaped" six-membered rings are found. Similar Te₃ and "pear-shaped" Te6 rings have been reported in Cd₇Cl₈(Te₇O₁₇).⁸ The interconnection of La³⁺ ions by bridging WO₄²⁻ anions led to a 2D [La₂WO₄]⁴⁺ layer along the *ab* plane. One $[La_2WO_4]^{4+}$ layer and two $Te_3O_7^{2-}$ layers are further interconnected into a thick layer in the *ab* plane via La-O-Te bridges (Figure 6). The thickness of the layer is about 7.65 Å, and the interlayer opening width is about 3.0 Å. This type of layer can also be viewed as the $[La_2WO_4]^{4+}$ layer being sandwiched between two $Te_3O_7^{2-}$ anionic layers. The lone-pair electrons of Te^{IV} atoms are oriented toward the interlayer space (Figure 6).

 $Nd_2W_2Te_2O_{13}$ with a different structure was obtained in our attempts to prepare the Nd^{III} analogue of La₂WTe₆O₁₈. $Nd_2W_2Te_2O_{13}$ features a 3D network structure in which the W_2O_{10} dimers occupy the large apertures formed by neodymium tellurite (Figure 7). Among two unique Nd³⁺ ions in the asymmetric unit, one is eight-coordinated whereas the other one is nine-coordinated. The Nd–O distances range from 2.333(5) to 2.681(5) Å. Unlike that in La₂WTe₆O₁₈, both of the W^{VI} cations in Nd₂W₂Te₂O₁₃ are octahedrally coordinated. W1 is coordinated by two O atoms from two tellurite groups and four terminal O atoms, whereas W2 is coordinated by one O atom from a tellurite group and five



Figure 7. View of the structure of $Nd_2W_2Te_2O_{13}$ down the *a* axis. The WO₄ tetrahedra are shaded in cyan. Nd, Te, and O atoms are drawn as green, yellow, and red circles, respectively.

terminal O atoms. A pair of WO₆ octahedra is interconnected via edge-sharing into a W₂O₁₀⁸⁻ dimer. Both of the WO₆ octahedra are severely distorted. The W1 atom is distorted toward a face (local C_3 direction) with three "short" [1.743(6)-1.834(5) Å] and three "long" [2.033(5)-2.300(5)]Å] W–O bonds, whereas the W2 atom is distorted toward an edge (C_2) with two "short" [1.747(5) and 1.767(5) Å], two "normal" [1.900(5) and 1.930(5) Å], and two "long" $[2.172(5) \text{ and } 2.232(5) \text{ Å}] \text{ W-O bonds. Again both W^{VI}}$ cations are distorted away from the O atoms that bonded to the Te^{4+} cations. The magnitudes of the distortion $(\Delta_{\! d})$ are 1.118 and 0.945 Å respectively for $W1O_6$ and $W2O_6$. The interconnection of the Nd^{III} ions by the tellurite groups resulted in a 3D network with two types of apertures along the a axis (Figure 7). The large apertures with a narrowlong shape are formed by 10-membered rings composed by four TeO₃ groups and six Nd^{III} ions, and the small ones are formed by four-membered rings composed of two Nd1 and two Te1O₃ groups. The W₂O₁₀ dimers are located at the large apertures (Figure 7). In Nd₂W₂Te₂O₁₃, each W₂O₁₀ dimer connects with two TeO₃ groups (one in a unidentate fashion, and the other in a bidentate bridging fashion) to form a $[W_2Te_2O_{13}]^{6-}$ anion. Therefore, the structure of Nd₂W₂Te₂O₁₃ also can be viewed as the Nd3+ ions being interconnected by [W2Te2O13]6- anions via Nd-O-Te and Nd-O-W bridges (Figure 7).

When chloride anion was introduced into the Ln–Mo (W)–Te–O system, isostructural Ln₅MTe₇O₂₃Cl₃ (Ln = Pr, Nd; M = Mo, W) were obtained. These compounds feature a 3D network of a lanthanide(III) molybdenum(VI) tellurium(IV) oxychloride with large apertures occupied by isolated Cl anions and the lone-pair electrons of Te^{IV} (Figure 8). Ln₅MTe₇O₂₃Cl₃ can also be formulated as Ln₅(MO₄)-(Te₅O₁₃)(TeO₃)₂Cl₃. The asymmetric unit of Ln₅MOTe₇O₂₃Cl₃ contains three unique Ln^{III} ions, one M^{VI} cation, and four Te atoms. Two Ln^{III} ions are 8-coordinated, whereas the third one is 10-coordinated. The Ln–Cl distances are significantly longer than those of the Ln–O bonds. The M^{VI} cation is in



Figure 8. View of the structure of $Nd_5MoTe_7O_{23}Cl_3$ down the *a* axis. The MoO₄ tetrahedra are shaded in cyan. Nd, Te, Cl, and O atoms are drawn as green, yellow, pink, and red circles, respectively.

a distorted tetrahedral coordination environment with Mo-O distances in the range of 1.714(10)-1.845(9) Å. Te1, Te3, and Te4 are three-coordinated by three O atoms, whereas Te2 is four-coordinated. Two Te1O₃ groups, two Te2O₄ groups, and a Te3O₃ group are corner-sharing to form a novel Te₅O₁₃⁶⁻ pentamer in a "semicycle" shape. The Te-O distances fall in the range of 1.845(8) - 2.162(6) Å. The interconnection of Nd^{III} ions via $Te_5O_{13}{}^{6-}$ and $TeO_3{}^{2-}$ anions resulted in a thick layer parallel to the *ab* plane. Neighboring lanthanide tellurium(IV) oxide layers are bridged by Cl2 atoms into a complicated 3D network, forming two different types of apertures (Figure 8). The MO₄ polyhedra are capped on both sides of the Ln₄O₂Cl₂ rings. The isolated Cl anions and the lone-pair electrons of Te^{IV} atoms of the Te₅O₁₃⁶⁻ groups occupy the large apertures formed by eight-membered rings (Figure 8).

The solid-state luminescent properties of Nd₂MoSe₂O₁₀, Nd₂MoTe₃O₁₂, Nd₂MoTe₄O₁₄, and Nd₂W₂Te₂O₁₃ were investigated at both room temperature and 10 K, and the solidstate luminescent properties of Nd₅MoTe₇O₂₃Cl₃ and Nd₅WTe₇O₂₃Cl₃ were studied at room temperature. Under excitation at 514 nm, the room temperature emission spectra for all six Nd^{III} compounds display three sets of characteristic emission bands for the Nd^{III} ion in the near-IR region: ⁴F_{3/2} \rightarrow ⁴I_{9/2}, ⁴F_{3/2} \rightarrow ⁴I_{11/2}, and ⁴F_{3/2} \rightarrow ⁴I_{13/2}. Nd₂MoTe₃O₁₂ contains only one independent Nd^{III} site, whereas $Nd_2MoSe_2O_{10}$, $Nd_2MoTe_4O_{14}$, and $Nd_2W_2Te_2O_{13}$ each contain two unique Nd^{III} sites with C_1 symmetry. Because of the crystal-field effect, each transition band is expected to be split into several subbands. The ⁴F_{3/2} is expected to split into two sublevels, whereas the complete degeneracy of ${}^{4}I_{9/}$ 2, ${}^{4}I_{11/2}$, and ${}^{4}I_{13/2}$ leads to five, six, and seven sublevels, respectively. Therefore, ${}^{4}F_{3/2} \rightarrow {}^{4}I_{9/2}$, ${}^{4}F_{3/2} \rightarrow {}^{4}I_{11/2}$, and ${}^{4}F_{3/2}$ \rightarrow ⁴I_{13/2} transitions will have a maximum of 10, 12, and 14 subbands if both the lower and upper levels of ${}^{4}F_{3/2}$ are populated as in the case of room temperature. When two and more unique Nd^{III} sites are present such as Nd₂MoSe₂O₁₀, Nd₂MoTe₄O₁₄, and Nd₂W₂Te₂O₁₃, the spectrum will be even more complicated. Because of the overlapping of some emission bands and resolution limit of the instruments, the observed emission peaks are usually fewer than that expected. At very low temperature such as 10 K, only the lower level of ${}^{4}F_{3/2}$ is populated; hence, the corresponding emission spectra are much simpler. Also the ${}^4F_{3/2} \rightarrow {}^4I_{11/2}$ transition, subbands are difficult to be resolved even at 10 K. The lowtemperature emission spectrum for Nd₂MoTe₃O₁₂ displays five and six subbands for the ${}^4F_{3/2} \rightarrow {}^4I_{9/2}$ and ${}^4F_{3/2} \rightarrow {}^4I_{11/2}$ transitions, respectively, which is in good agreement with the theoretical expectation. Assuming the energy of the lowest sublevel of ${}^{4}I_{9/2}$ is 0 cm⁻¹, the energies of the lower level of ${}^{4}F_{3/2}$ and the other four sublevels for ${}^{4}I_{9/2}$ are calculated. Likewise, we can calculate the energies for the six sublevels of ${}^{4}I_{11/2}$. Upon excitation at 580 nm, both Nd₂MoSe₂O₁₀ and Nd₂MoTe₄O₁₄ display 8 and 10 subbands for the ${}^4F_{3/2} \rightarrow {}^4I_{9/2}$ and ${}^4F_{3/2} \rightarrow {}^4I_{11/2}$ transitions at 10 K, whereas Nd₂W₂Te₂O₁₃ displays eight and nine subbands at 10 K upon excitation at 288 nm. The lifetimes for the ${}^{4}F_{3/2}$ \rightarrow ⁴I_{11/2} transition were measured to be 0.21 and 0.20 ms and 3.2 and 3.6 µs respectively for Nd₂MoSe₂O₁₀ and Nd₂MoTe₃O₁₂ and for Nd₂MoTe₄O₁₄ and Nd₂W₂Te₂O₁₃. The three PrIII compounds are capable of emitting luminescent light in the blue, green, and red regions. Pr5MoTe7O23Cl3 and Pr₅WTe₇O₂₃Cl₃ display four sets of emission bands at 491 nm (very strong, ${}^{3}P_{0} \rightarrow {}^{3}H_{4}$), 534 nm (moderate, ${}^{3}P_{0} \rightarrow$ ${}^{3}\text{H}_{5}$), 619 nm (moderate, ${}^{3}\text{P}_{0} \rightarrow {}^{3}\text{H}_{6}$), 649 nm (strong, ${}^{3}\text{P}_{0} \rightarrow$ ${}^{3}F_{2}$), and 735 nm (weak, ${}^{3}P_{0} \rightarrow {}^{3}F_{4}$) under $\lambda_{ex} = 448$ nm. Under the same experimental conditions, the corresponding emission bands for Pr₂MoTe₄O₁₄ are much weaker. Because of the so-called "crystal-field effect" as well as multiple Pr³⁺ sites, a few transition bands were split into several subbands.¹⁹

3. $\mathbf{TM}-\mathbf{d}^{0} \mathbf{TM}-\mathbf{Te}^{IV}$ or $\mathbf{Se}^{IV}-\mathbf{O}$ Systems. Reports on compounds in the TM $-\mathbf{Mo}^{V}-\mathbf{Te}^{IV}/\mathbf{Se}^{IV}-\mathbf{O}$ systems are much fewer than those of the alkali and alkaline-earth ones.²⁰ The structure of $\mathbf{Cd}(\mathbf{VO}_2)_4(\mathbf{SeO}_3)_3 \cdot \mathbf{H}_2\mathbf{O}$ can be described as a pillared layered architecture. The layer is composed of dimers of \mathbf{VO}_5 square pyramids and dimers of \mathbf{CdO}_7 decahedra connected to one another by sharing corner and/ or edge O atoms, whereas the pillars are composed of distorted \mathbf{VO}_6 octahedra connected to one another via corner-and edge-sharing with capping selenite groups; the lone-pair electrons of the \mathbf{Se}^{IV} atoms are oriented toward the long-narrow-shaped tunnels along the *b* axis.

We deem that the combination of the lone-pair electrons of Se^{IV} or Te^{IV} with the distorted MoO_6 octahedron may lead to new compounds in $TM-Mo^{VI}$ or $V^V-Se^{IV}/Te^{IV}-O$ systems with novel structures and unusual physical properties, such as magnetic or optical properties. So far, no nickel molybdenum(VI) selenite or tellurite has been structurally characterized. Our exploration of new phases in nickel

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Figure 9. $[Mo_4O_{16}]^{8-}$ tetranuclear cluster unit (a) and $[Ni_6O_{22}]^{32-}$ hexanulear cluster unit in $Ni_3(Mo_2O_8)(SeO_3)$ (b), and 1D $[Ni_3O_{11}]^{16-}$ corrugated chain along the *b* axis in $Ni_3(Mo_2O_8)(TeO_3)$ (c).

molybdenum(VI) selenite or tellurite systems led to three new compounds, namely, $Ni_3(Mo_2O_8)(XO_3)$ (X = Se, Te)²¹ and $Ni_3(MoO_4)(TeO_3)_2$.²²

 $Ni_3(Mo_2O_8)(XO_3)$ (X = Se, Te) display two different types of 3D structures containing [Mo₄O₁₆]⁸⁻ and [Ni₆O₂₂]³²⁻ clusters or 1D nickel oxide chains (Figure 9). The structure of Ni₃(Mo₂O₈)(SeO₃) features a 3D network in which $[Ni_6O_{22}]^{32-}$ cluster units are interconnected by $[Mo_4O_{16}]^{8-}$ clusters and SeO_3^{2-} anions. All three Ni^{II} ions in the asymmetric unit are octahedrally coordinated by six O atoms, with Ni–O distances ranging from 1.983(4) to 2.160(4) Å. Both Mo^{VI} cations are octahedrally coordinated by six O atoms. The Mo-O bond distances are in the range of 1.719(4)-2.304(4) Å. The Mo^{VI} cations are distorted toward an edge (local C_2 direction) with two "short", two "normal", and two "long" Mo⁶⁺-O bonds. The magnitudes of the distortions (Δ_d) were calculated to be 1.18 and 1.29 Å respectively for Mo1 and Mo2. The four Mo^{VI}O₆ octahedra are interconnected by edge-sharing to form a cyclic $[Mo_4O_{16}]^{8-}$ tetranuclear cluster unit (Figure 9a). Six NiO₆ octahedra are interconnected into a hexanuclear $[Ni_6O_{22}]^{32-}$ cluster unit through edge-sharing (Figure 9b). The intracluster Ni···Ni separations between a pair of edge-sharing NiO₆ octahedra are in the range of 3.034(1)-3.245(1) Å (Figure 9b). The $[Ni_6O_{22}]^{32-}$ cluster units are bridged by SeO_3^{2-} groups to form a 2D nickel selenite layer parallel to the ab plane. Neighboring nickel selenite layers are further interconnected by the [Mo₄O₁₆]⁸⁻ clusters via Mo-O-Ni bridges into a 3D network structure. The lone-pair electrons of the Se^{IV} atoms are oriented toward the cavities of the structure. The structure of Ni₃(Mo₂O₈)(TeO₃) is different from that of Ni₃(Mo₂O₈)(SeO₃), although their chemical formulas are comparable. It features a 3D structure in which the corrugated nickel oxide anionic chains are bridged by [Mo₄O₁₆]⁸⁻ cluster units and TeO₃²⁻ anions. As in Ni₃(Mo₂O₈)(SeO₃), the Ni^{II} and Mo^{VI} cations are octahedrally coordinated by six O atoms. Similar to that in Ni₃(Mo₂O₈)(SeO₃), the four Mo^{VI}O₆ octahedra in Ni₃(Mo₂O₈)(TeO₃) are interconnected via edgesharing to form a [Mo₄O₁₆]⁸⁻ tetranuclear cluster unit. Two Ni1O₆ and one Ni2O₆ octahedra are interconnected via edgesharing into a [Ni₃O₁₃]²⁰⁻ trinuclear unit. Such neighboring trinuclear units are further interconnected through edgesharing into a corrugated [Ni₃O₁₁]¹⁶⁻ anionic chain along the b axis with a Ni1-O2-Ni1 bond angle of 84.65(9)° (Figure 9c). This nickel oxide chain can also be viewed as Ni2O₆ octahedra being grafted onto the corrugated chain of Ni1O₆ through edge-sharing. It is interesting to note that [Ni₆O₂₂]³²⁻ clusters are formed in Ni₃(Mo₂O₈)(SeO₃), whereas corrugated [Ni₃O₁₁]¹⁶⁻ anionic chains are observed in Ni₃(Mo₂-O₈)(TeO₃). Both nickel oxide building units are based on Ni₃O triangles. The Ni₃O triangles in [Ni₆O₂₂]³²⁻ clusters are condensed via sharing Ni-Ni edges, whereas those in [Ni₃O₁₁]¹⁶⁻ anionic chains are interconnected through pairs of Ni-O-Ni bridges. These different nickel(II) oxide architectures may result from the different coordination modes of the selenite and tellurite groups as well as the different ionic radii of Se^{IV} and Te^{IV}. Neighboring corrugated nickel(II) oxide chains are bridged by TeO₃²⁻ anions to form a thick nickel(II) tellurite layer parallel to the ab plane. The

⁽²²⁾ Jiang, H. L.; Mao, J. G. Unpublished results.



(a)





Figure 10. 3D network of nickel(II) oxide in $Ni_3(MoO_4)(TeO_3)_2$ (a) and view of the structure of $Ni_3(MoO_4)(TeO_3)_2$ along the *a* axis. The NiO₆ and NiO₅ polyhedra are shaded in green, and MoO₄ tetrahedra are shaded in cyan. Te atoms are drawn as yellow circles.

thickness of the layer is about 12.1 Å. Such layers are further interconnected by the $[Mo_4O_{16}]^{8-}$ clusters via Mo–O–Ni bridges into a 3D network structure. The lone-pair electrons of the Te^{IV} atoms are oriented toward the tunnels of the structure.

When we increased the molar ratio of TeO₂/MoO₃ to 2:1, a new Ni–Mo–Te–O phase with NCS structure (space group $P2_12_12_1$), Ni₃(MoO₄)(TeO₃)₂, was obtained recently.²² Its structure features a novel 3D network of nickel(II) oxide with larger 1D tunnels along the *a* axis. The MoO₄ tetrahedra and TeO₃ groups cap the walls of the tunnels (Figure 10). Among three unique Ni^{II} ions in the asymmetric unit, two are octahedrally coordinated, whereas the third one is in a square-pyramidal geometry. NiO₆ octahedra are intercon-



Figure 11. View of the structure of $Cd_4V_2Te_3O_{15}$ down the *a* axis. The CdO_6 and VO_4 polyhedra are shaded in blue and green, respectively. Te and O atoms are drawn as yellow and red circles, respectively.



Figure 12. View of the structure of $Cd_6V_2Se_5O_{21}$ down the *b* axis. The CdO_6 , CdO_7 and CdO_8 polyhedra are shaded in cyan and VO_4 tetrahedra in green. Se and O atoms are drawn as pink and red circles, respectively.

nected by edge-sharing along the *a* axis into a 1D chain, whereas NiO₅ polyhedra are corner-sharing along the *a* axis into a 1D chain. These two types of chains are further interconnected via corner-sharing into a 3D open framework of nickel oxide, forming 1D helical tunnels along the *a* axis, which are based on Ni₁₂ rings (Figure 10a). Unlike those in Ni₃(Mo₂O₈)(SeO₃) and Ni₃(Mo₂O₈)(TeO₃), the Mo^{VI} cation is in a tetrahedral environment and there are no Mo–O–Mo or Mo–O–Te bridges. The MoO₄ tetrahedra and TeO₃ groups cap walls of the tunnels; the lone-pair electrons of Te^{IV} cations and terminal O atoms of the MoO₄ tetrahedra are pointing toward the center of the tunnel (Figure 10b). SHG measurements indicate that its SHG signal is very weak.

Both Ni₃(Mo₂O₈)(SeO₃) and Ni₃(Mo₂O₈)(TeO₃) obey the Curie–Weiss law above 60 K with Weiss constants (θ) of 15.4(2) and 5.5(1) K respectively for Se and Te compounds, indicating significant ferromagnetic interactions between



Figure 13. View of the structure of CdWTeO₆ down the *a* axis (a) and a 2D layer composed of Cd2O₇, WO₆, and Te1O₃ groups (b). The CdO₆ and CdO₇ polyhedra are shaded in green and the WO₆ octahedra in green. Te and O atoms are drawn as yellow and red circles, respectively.

magnetic centers. It is expected that the magnetic interactions should be dominated by the magnetic interactions between Ni^{II} ions within the hexanuclear $[Ni_6O_{22}]^{32}$ cluster in Ni₃(Mo₂O₈)(SeO₃) and the 1D $[Ni_3O_{11}]^{16-}$ anionic chain in Ni₃(Mo₂O₈)(TeO₃). Long-range magnetic ordering is observed below 28 and 56 K.

We also extended our studies to TM ions with a d¹⁰ electronic configuration such as Zn^{II} and Cd^{II} ions.²³ Zn^{II} or Cd^{II} compounds may display semiconducting properties, and they are also able to emit blue light. Solid-state reactions of zinc(II) or cadmium(II) oxide, V₂O₅ and TeO₂, at high temperature lead to two novel quaternary compounds, namely, Zn₃V₂TeO₁₀ and Cd₄V₂Te₃O₁₅.²³ Very recently Cd₆(V₂O₆)(SeO₃)₅ and CdWTeO₆ were also isolated.²⁴

The structure of $Zn_3V_2TeO_{10}$ is a complicated 3D network constructed by the interconnection of ZnO_5 , ZnO_6 , VO_4 , and



Figure 14. View of the structure of $Se_2B_2O_7$ down the *b* axis. BO_4 tetrahedra are shaded in cyan. Se, B, and O atoms are drawn as yellow, cyan, and red circles, respectively.

TeO₄ polyhedra via corner- and edge-sharing. Cd₄V₂Te₃O₁₅ with an acentric structure features a 3D network in which the cadmium tellurite layers are further interconnected by both "isolated" VO₄ tetrahedra and 1D vanadium oxide helical chains (Figure 11). Cd₄V₂Te₃O₁₅ displays a SHG efficiency of about 1.4 times that of KDP. It is stable up to 840 °C and shows little absorption in 0.6–10.0 μ m. Luminescent measurements indicate that both compounds exhibit broad emission bands in the blue-light region.²³

Cd₆V₂Se₅O₂₁ was obtained by solid-state reactions of cadmium(II) oxide, V₂O₅ and SeO₂. Its structure is composed of a 3D open framework of cadmium(II) selenite and 1D vanadium(V) oxide chains of corner-sharing VO₄ tetrahedra. The vanadium(V) oxide chains are inserted at the large tunnels of cadmium(II) selenite via V-O-Cd bridges. The Cd^{II} ions are coordinated by six, seven, or eight O atoms (Figure 12).^{24a} CdWTeO₆ crystallized in triclinic $P\overline{1}$; its structure features a complicated 3D network of corner/edge-sharing CdO₆ octahedra, CdO7 pentagonal bipyramids, WO6 octahedra, and TeO3 groups (Figure 13). There are three Cd^{II}, two W^{VI}, and two Te^{IV} sites in its asymmetric unit. Two Cd^{II} sites lying on the center of symmetry are octahedrally coordinated, and the third Cd^{II} ion in a general position is seven-coordinated in a pentagonalbipyramidal geometry. The CdO₆ octahedra form 1D chains along the b axis via edge-sharing; these 1D chains are capped by Te2O₃ groups. The WO₆ octahedra are distorted toward a face (local C_3 direction). Neighboring WO₆ octahedra are interconnected via corner-sharing into 1D chains also along the b axis. These tungsten oxide chains are further capped by Te2O₃ groups on one side, and they are further bridged by CdO₇ polyhedra via edge- and corner-sharing into a 2D layer parallel to the ab plane (Figure 13b). Two such layers are interconnected into a double layer via Cd-O-W bridges. The above two types of building units are alternating and interconnected along the c axis via corner-sharing (Te-O-Cd, Cd-O-W, and Cd-O-Cd bridges) into a 3D architecture (Figure 13a).^{24b}

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Part II. Metal Tellurites and Selenites Containing Tetrahedral Groups of Main-Group Elements

So far such compounds are still rare. With respect to tellurite phosphate oxides, a few materials have been reported, namely, Te₂O₃(HPO₄), Te₈O₁₀(PO₄)₄, Ba₂TeO(PO₄)₂, BaTeMO₄(PO₄) (M = Nb⁵⁺ or Ta⁵⁺), and A₂TeMo₂O₆(PO₄)₂ (A = K, Rb, Cs, Tl)²⁵ Structurally these materials are different; BaNbTeO₄(PO₄) contains 2D layers, whereas Te₂O₃(HPO₄) and Te₈O₁₀(PO₄)₄ exhibit 3D structures consisting of asymmetric TeO₄ and TeO₅ polyhedra that are linked through corner-sharing.^{25a-c} Ba₂TeO(PO₄)₂ displays a novel 1D chain structure that is composed of PO₄ tetrahedra and TeO₅ polyhedra. Te⁴⁺ cations are in asymmetric coordination environments attributable to their lone pairs. Two TeO₅ polyhedra form a dimeric unit via edge-sharing; two neighboring Te₂O₈ units are further interconnected via a pair of phosphate groups into a 1D chain along the b axis, and each Te^{IV} atom is also bonded to the third phosphate group through one Te-O-P bridge.^{25d} The isostructural mixedmetal phosphates, BaMTeO₄(PO₄) ($M = Nb^{5+}$, Ta⁵⁺), have novel layered crystal structures consisting of MVO6 cornersharing octahedral chains that are connected to Te^{IV}O₄ polyhedra and phosphate tetrahedra. The phosphate groups act as pendent groups between two neighboring layers. The Nb^{5+} cation distorts along the local C_4 direction of its octahedron with a "short-long-short-long" Nb-O-Nb bond motif. Furthermore, the Nb⁵⁺ cation displaces away from the oxide ligands that are bonded to Te^{4+} or P^{5+} cations, attributable to the structural rigidity of the TeO₄ and PO₄ polyhedra.^{25e} A₂Mo₂TeO₆(PO₄)₂ displays a pillared layered architecture composed of Mo₂P₂O₁₄ layers and TeO₄ groups as the pillars. Within the $Mo_2P_2O_{14}$ layer, the MoO_6 octahedra (distorted toward an edge) and PO₄ tetrahedra are interconnected via corner-sharing, forming Mo₂P₂ four-membered rings and Mo₄P₄ eight-membered rings.^{25f}

So far, there is no reports on metal selenites or tellurites containing additional SiO₄ or GeO₄ tetrahedra. Our exploration into the unexplored RE–Ge/Si–Te^{IV}–O systems afforded two novel lanthanum(III) tellurites with additional SiO₄ or GeO₄ tetrahedra, namely, La₄(Si_{5.2}Ge_{2.8}O₁₈)(TeO₃)₄ and La₂(Si₆O₁₃)(TeO₃)₂.²⁶

The structure of La₄(Si_{5.2}Ge_{2.8}O₁₈)(TeO₃)₄ features a 3D network composed of the $[(Si_{5.2}Ge_{2.8})O_{18}]^{4-}$ tetrahedral layers and the $[La_4(TeO_3)_4]^{4+}$ layers that alternate along the *b* axis. The germanate-silicate layer consists of corner-sharing XO₄ (X = Si, Ge) tetrahedra, forming four- and six-membered rings. The structure of La₂(Si₆O₁₃)(TeO₃)₂ is a 3D network composed of the $[Si_6O_{13}]^{2-}$ double layers and the $[La_2(TeO_3)_2]^{2+}$ layers that alternating along the *a* axis. The $[Si_6O_{13}]^{2-}$ double layer is built by corner-sharing silicate tetrahedra, forming four-, five-, and eight-membered rings.

The TeO_3^{2-} anions in both compounds are only involved in the coordination with the La^{3+} ions to form a lanthanum(III) tellurite layer.²⁶

We also developed a new SHG material, Se₂B₂O₇, by the combination of BO₄ tetrahedra with SeO₃²⁻ groups.²⁷ The structure of Se₂B₂O₇ features a 3D network composed of dimers of corner-sharing BO₄ tetrahedra and SeO₃ groups (Figure 14). The open framework of Se₂B₂O₇ can also be described as an interesting mixed (3,4)-connected net of the B and Se atoms (linked by -O- bridges). Right-handed helical tunnels along the *c* axis are formed. These tunnels are based on B₆Se₄ 10-membered rings. The lone pairs of the Se^{IV} cations are oriented toward the above tunnels. Se₂B₂O₇ is stable up to 300 °C. It exhibits a SHG efficiency of about 2.2 times that of KDP (KH₂PO₄). Se₂B₂O₇ is transparent in the range of 300–2000 nm, and it exhibits an optical band gap of 4.64 eV.²⁷

Conclusions and Outlook

Several factors have strong effects on the structural topologies of the resulting compounds. The slight change of the ionic radius of the cation such as the A^I or Ln^{III} ion could lead to a completely different structure such as, for example, $A_2W_3TeO_{12}$ (A = K⁺, Rb⁺, Cs⁺)^{12c} and $Ln_2MoTe_3O_{12}$ (Ln = La, Nd).^{19a} Also metal tellurites usually show structures different from the corresponding selenites because of the larger ionic radius of Te^{IV} than Se^{IV}, such as, for example, $Ni_3(Mo_2O_8)(QO_3)$ (Q = Se, Te).²¹ In some cases, isomerism exists for some metal tellurites. Recently, we obtained a cubic phase of $Ga_2Te_3O_9$ (I43d) and a hexagonal phase of $Ga_2Te_3O_9$ (P6₃/m) in the same hydrothermal reaction; the cubic phase is SHG active $(1 \times \text{KDP})$ and is densely packed, whereas the centrosymmetric hexagonal phase has a much lower density and features a 3D network with large tunnels along the c axis.²⁸ Molar ratios of the starting materials, synthetic methods, and reaction temperatures are also very important factors.

Several trends are obvious. The combination of the two types of cations susceptible to SOJT distortion, d⁰ TM and lone-pair cation (Te^{IV} or Se^{IV}), not only gives rise to a rich structural chemistry but also affords many NCS compounds with good SHG properties. As for the direction of the outof-center distortion, it was found that V⁵⁺ and Nb⁵⁺ usually displace toward an edge or corner, whereas Mo⁶⁺ and W⁶⁺ ions are more likely distorted toward an edge or face. The V⁵⁺ cation may adopt the square-pyramidal geometry besides a tetrahedron and a octahedron. Also it is not as stable as other d⁰ TM cations, and sometimes it is reduced to V⁴⁺ during the reaction. The bond polarization follows the following order: Te^{IV} > Se^{IV} and Mo⁶⁺ > V⁵⁺ > W⁶⁺ > $Nb^{5+} > Ta^{5+}$.² It is observed that Ln^{III} compounds are seldom acentric and the d⁰ TM cations are more likely tetrahedrally coordinated; this may be due to the higher coordination number for the Ln^{III} ions than the A^I and AE^{II} cations. It is

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worth mentioning that the Se^{IV} cation exists mostly as a SeO₃ group (in some cases as the diselinite group), whereas Te^{IV} can be three-, four-, or five-coordinated, and the most amazing aspect is that these TeO_x (x = 3-5) polyhedra can be polymerized into many types of polynuclear clusters or extended skeletons (Scheme 1) besides a ditellurite anion; such a polymerization is hardly observed in the organically templated or organically covalently bonded metal tellurites. Inorganic solids with various polymeric tellurium(IV) oxide anions include a Te₃O₈⁴⁻ trimer in La₂MoTe₃O₁₂, a Te₄O₁₁⁶⁻ tetramer in $\text{Er}_2\text{Te}_4\text{O}_{11}$,²⁹ a Te_5O_{13} ⁶⁻ pentamer in Ln_5MTe_7 - $O_{23}Cl_3$ (Ln = Pr, Nd; M = Mo, W), a 1D Te₄ O_{10}^{4-} in Ln₂Mo- Te_4O_{14} (Ln = Pr, Nd), a 1D $Te_6O_{13}^{2-}$ in {Cd₂(Te₆O₁₃)}- $\{Cd_2Cl_6\}$, a 1D Te₇O₁₇⁶⁻ in Cd₇Cl₈(Te₇O₁₇), layered Te₃O₇²⁻ in La₂WTe₆O₁₈, a 2D Te₄O₉²⁻ in K₂Te₄O₉ \cdot 3.2H₂O,^{30a} and a 2D Te₂O₅²⁻ in Ln(Te₂O₅)X (Ln = Nd, X = Cl, Br; Ln = Gd, X = Cl).^{30b,c} Certainly more examples will be discovered in the future, and extensive theoretical studies are needed to understand its origin.

There are also many fertile lands for our exploration in the field of metal selenites and tellurites. Systematic investigation on the $Ln^{III}-V^{5+}$ (Nb⁵⁺, Ta⁵⁺, Ti⁴⁺)-Te^{IV} (or Se^{VI})–O systems is expected to afford many compounds with new structural types as well as excellent luminescent materials. The mixing of two types of charge-balancing cations such as Ln-AE, AE-A, and Ln-A could change the overall charges of the anionic networks in addition to breaking the packing limitation for the single-type cation; thus, a variety of new structures can be obtained. The combination of lone-pair Te^{IV} or Se^{IV} with borate is also quite promising in the search for new SHG materials. In order to exhibit good second-order NLO properties, the above two types of bond polarizations should be "adductive", which is still a great challenge for a synthetic chemist. To fully address these problems, strong interactions between synthetic and theoretical chemists are necessary.

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