Perovskite-related ReO₃ structures: A diverse family, from oxides to metal-organic frameworks.

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Abstract

ReO₃-type structures can be described as ABX_3 perovskites in which the *A*-cation site is unoccupied. They therefore have the general composition BX_3 , where *B* is normally a cation and *X* is a bridging anion. The chemical diversity of such structures is very broad, ranging from simple oxides and fluorides, such as WO₃ and AlF₃, to more complex systems in which the bridging anion is polyatomic, as in the Prussian blue-related cyanides such as Fe(CN)₃ and CoPt(CN)₆. The same topology is also found in metal-organic frameworks, for example In(Im)₃ (Im = imidazolate), and even the well-known MOF-5 structure, where the *B*-site cation is itself polyatomic. This remarkable chemical diversity gives rise to a wide range of interesting and often unusual properties, including negative thermal expansion (ScF₃), photocatalysis (CoSn(OH)₆), thermoelectricity (CoAs₃), and even a report of superconductivity in a phase that is controversially described as SH₃ with a doubly interpenetrating ReO₃ structure. We present a comprehensive account of this exciting family of materials and discuss current challenges and future opportunities in the area.

1 Introduction

Recent developments in photovoltaic devices with active layers containing hybrid perovskite halides such as (MA)PbI₃ (MA = methylammonium)¹ have drawn our attention once more to the importance and versatility of perovskite-based materials. These iconic materials, of general formula AMX_3 [Figure 1(a)], have a long and distinguished his-

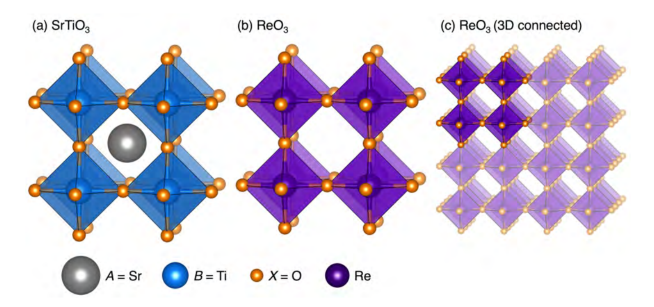


Figure 1: (a) The ABX_3 perovskite structure, exemplified by cubic SrTiO₃; (b) The ReO₃ structure, illustrated by ReO₃ itself, contains no cation on the *A*-site. It is therefore an array of corner-sharing BX_6 octahedra. These figures are drawn to scale to remind the reader that the ReO₃ unit cell is smaller due to the smaller size of the Re⁶⁺ cation; (c) The ReO₃ structure, drawn to show its extended connectivity. The VESTA software suite was used to create all visualizations of crystal structures in this manuscript.⁹

tory, with major milestones including the discoveries of ferroelectric BaTiO₃,² ferromagnetic SrRuO₃,³ superconducting YBa₂Cu₃O₇,⁴ magnetoresistive (La/Sr)MnO₃,⁵ and multiferroic BiMnO₃.⁶ Striking breakthroughs in the hybrid perovskite area include the discoveries of ferroelectric and multiferroic behaviour in formates such as (DMA)Mn(HCOO)₃, (DMA = dimethylammonium), where the *X* anion is an organic linker rather than a simple halide or oxide anion,⁷ and the ferroelectric, metal-free perovskites ((N-methyl-N'diazabicyclo[2.2.2]octonium)-ammonium triiodides) that were recently reported.⁸

It is often forgotten, however, that the ABX_3 perovskite structure is itself based on an even simpler structure type in which there are no cations in the *A*-site cavities. Such BX_3 -

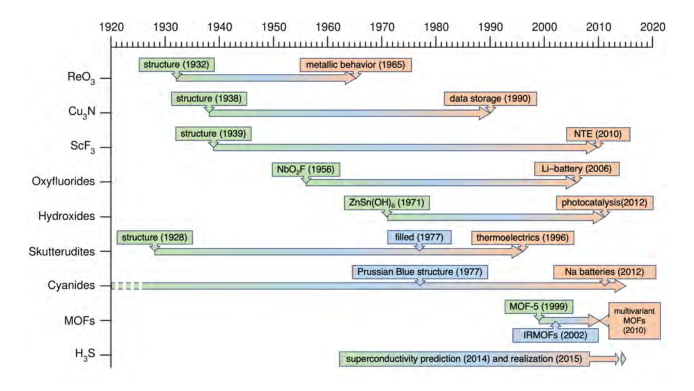


Figure 2: Timeline of ReO_3 -type compounds. It is interesting to note the long period between discoveries and reported properties, as well as the diverse chemical compositions and functionalities, of the ReO_3 -type materials.

type structures (these are typically MX_3 structures, where M is a metal) are most notably illustrated by the eponymous ReO_3 system, shown in Figure 1(b) and (c). It comprises a network of ReO₆ octahedra that share common corners in three dimensions, in this case with an Re–O–Re angle of 180° (though in general such linear linkages are unusual.) Recalling that the perovskite family now comprises an extensive range of systems with both simple and complex X anion linkers,¹⁰ we have begun to explore the extent to which this is also true for ReO₃-type materials. Our article addresses this question, drawing together the many examples of ReO₃-type phases, ranging from simple oxides and halides, such as WO₃ and ScF₃, to systems containing more complex linkers and nodes, such as the formates $M^{III}(HCOO)_3$, hypophosphites $M^{III}(H_2POO)_3$, imidazolates $M^{III}(C_3N_2H_3)_3$, and borohydrides Ln(BH₄)₃. We shall show that this family is far more extensive than has hitherto been recognized, and that they display a remarkable range of functionality that almost matches that of the perovskites. Figure 2 illustrates this point, showing the timelines of discoveries and properties of these ReO₃-type materials, as well as their considerable chemical scope. Furthermore, our survey of the literature in this area suggests that there are several additional systems that have yet to be explored.

One of the characteristics of the ReO_3 -type structures is their openness, which is immediately apparent when comparing ReO_3 itself with the perovskite structure in Figure 1. This feature is even more striking in expanded ReO_3 structures, such as those in which the *X*-site has a multi-atom linker. There are several aspects of the openness that can have an important influence on both the structures and properties of ReO_3 systems. In terms of the structures, the openness often results in tilting of the BX_6 octahedra, leading to more stable structures with bent B-X-B linkages.¹¹ Tilting of octahedra in perovskites has been extensively studied,¹²⁻¹⁴ including the analysis of group-subgroup relationships, and the concepts in this area have recently been extended to multi-atom X-site linkers.^{15,16} These well-established ideas can be equally applied to ReO₃-type structures but will not be discussed further here. However, we note that tilting, or the absence of it, can have a significant impact on electronic and magnetic properties, as will be seen in the case of ReO₃, itself, as well as in some of the cyanides. Turning to other manifestations of the openness of ReO₃-type structures, we note that cation insertion into the vacant A-site cavity is often facile, both in terms of the availability of space and the ease of ion transport; this has enabled a number of applications in the area of rechargeable batteries. The openness also amplifies the vibrational properties associated with the X-site linkers, which has led to some outstanding examples of negative thermal expansion. Finally, in the case of multiatom X-site linkers, where the openness is particularly high, we shall find many examples of systems that can accommodate guest molecules in the A-site cavities, leading in some cases to interesting adsorption-desorption behaviour.

Our article begins with a focus on simple inorganic systems, especially the oxides, fluorides and oxyfluorides, and then explores more complex phases with multi-atom linkers on the *X*-site. Note that one of the requirements for the adoption of the ReO₃-type structure or topology is that the system must be able to support octahedral connectivity at the *B*-sites. Bearing this in mind, we shall also consider some metal-organic frameworks

(MOFs) in which the *B* site node is itself polyatomic, the most striking of which is the metal-organic framework, MOF-5.¹⁷ The nomenclature for these materials in the MOF literature is quite varied and includes the use of the terms pcu (primitive cubic), α -plutonium and reo networks. For simplicity, we shall simply refer to them as ReO₃-type structures.

2 Inorganics

Oxides The oxide ReO₃, which is deep red in colour, was first reported in the early 1930's when its synthesis was described and the structure determined by powder X-ray diffraction.¹⁸ However, it was not until 1965 when Goodenough and co-workers reported that it was a metallic conductor that it attracted the attention of the scientific community.¹⁹ Oxide conductors were rarities at that time, and the behaviour of ReO₃ was particularly eye-catching because its conductivity was almost as high as that of copper metal. That finding led to subsequent studies of its electronic structure, which involves band formation from overlap between the rhenium 5d (t_{2g}) and O 2p ($p_{x/y}$) orbitals.²⁰ More than 50 years after the work of Goodenough, it remains the most striking example of a metallic transition metal oxide.

ReO₃ and WO₃ are the only binary oxides that crystallize with the ReO₃-type structure, but though structurally similar, they have properties that are quite different. For example, at room temperature, ReO₃ is primitive cubic and conductive, as described above, while WO₃ is monoclinic and insulating. The difference lies in their contrasting M–O–M bond angles, where ReO₃ has angles of 180°, and WO₃ has angles of between 165° and 179°.²¹ This distortion is due to the propensity for W atoms to off-center from the ideal cubic location location in the WO₆ octahedral environments, which density functional theory (DFT) calculations suggest is caused by non-populated antibonding orbitals in the conduction band. In ReO₃ it is the reverse, as the antibonding orbitals are populated and the off-centering is discouraged.²² This theory has been supported by experiment, since solid solutions between ReO₃ and WO₃ (Re_xW_{1-x}O₃) become cubic at an *x* value of 0.25; this change is also marked by an increase in conductivity.²³

ReO₃ and WO₃ also differ in that WO₃ undergoes multiple first-order phase transitions with temperature, whereas ReO₃ does not (the cubic ReO₃ structure is stable from liquid helium temperature to its 673 K melting point.)²⁴ However, ReO₃ does display negative thermal expansion (NTE).^{24,25} NTE is the phenomenon where a material contracts instead of expanding when heated. This property proves useful in composites, where the strain from the positive thermal expansion (PTE) of one phase can be offset by the NTE of another.²⁶ Studies by Rodriquez *et al.* described how for ReO₃ samples with high static disorder of oxygen, NTE was diminished.²⁷ The weakened NTE is due to the oxygen disorder, which alters the state of the antibonding band, reducing the Fermi pressure responsible for the structure's rigidity. In a more holistic sense, NTE is commonly seen in materials with corner connected *M*–*X* tetrahedra or octahedra which undergo concerted tilting or rocking motions when transverse vibrations (the movement of *X* species perpendicular to the *M*–*X*–*M* bond) are excited.²⁸ Famous NTE compounds such as ZrW₂O₈, Prussian blues and zeolites, all display some form of excited transverse vibrational modes related to their NTE behavior.²⁸ We shall see other striking examples of NTE in ReO₃-type systems in some of the following sections.

In addition to exhibiting NTE, ReO₃ displays pressure-induced first order phase transitions. These pressure dependent phase transitions (which involve octahedral rotations) occur as a stress relief,²⁹ as the ReO₆ octahedra do not distort easily because of the strong Re–O bonds.³⁰ The pressure induced phase transitions have been examined via diffraction experiments,^{31–33} but only recently with the implementation of high pressure Raman studies was a clear pressure phase diagram resolved.³⁴ Outside of physical pressure, chemical pressure in the form of lithiation can also induce phase transitions in ReO₃, as was first examined in the 1980's by Cava *et al.*,^{35,36} and more recently by Melot and co-workers,³⁷ where considerable correlated rotations of the corner-sharing ReO₆ octahedra are observed upon lithiation.

Lastly, though ReO₃ has generated a large amount of interest, it should not be overlooked that WO₃ is an attractive wide band gap semiconductor which can be tuned through intercalation, temperature, doping, and nano/microstructuring. Nano/microstructuring in particular, is quite attractive, as this can stabilize one of the three other WO₃ phases at room temperature,³⁸ or increase surface area for enhanced chemical functionality. For example, past work on WO₃ for its use as a photochromic,^{39,40} electrochromic,⁴¹ photocatalytic,⁴² superconductive,⁴³ ferroelectric,⁴⁴ as well as a gas-sensing material,^{45–48} is still quite relevant, with most areas still the focus of active research.⁴⁹ Current work on WO₃, for example, ranges from photoelectrochemical,^{50–54} photochromic,⁵⁵ conductive,⁵⁶ anodic,⁵⁷ photocatalytic,^{58–61} thermoelectric,^{62,63}, and gas-sensing^{64,65} applications, but there are some particularly interesting developments of late.

One recent work used WO₃ as an electron conductor in a core/shell nanowire BiVO₄ photoanode for photoelectrochemical water oxidation.⁶⁶ This was significant, as by utilizing this core/shell nanowire design the researchers overcame the low transport properties of BiVO₄ whilst still leveraging its high light absorption. Another recent example demonstrated how an apoferritin-encapsulated, nanoparticle coated, WO₃ material was capable of sensitive detection of certain molecules in exhaled breath, demonstrating great promise for reliable, non-invasive, diagnosis of diseases.⁶⁴ Lastly, recent work showed that WO₃ is able to tune the surface charge transfer of diamond.⁶⁷ In fact, the transfer doping efficiency with WO₃ was the highest per minimal surface acceptor coverage reported to date, marking a sizable advance in 2D diamond-based electronic devices. This recent work on WO₃, in combination with that of the past, attests to the wide variety of applications this material continues to have.

Nitrides The anti-ReO₃ crystal structure of Cu₃N was determined in 1938 by Juza and Hahn,⁶⁹ not long after the discovery of ReO₃ itself, and has been confirmed in more recent work.⁷⁰ Its crystal structure in space group $Pm\bar{3}m$ is the same as that of ReO₃, since the N–Cu–N bond angles are also 180°. Under high pressure, Cu₃N transforms to a simple

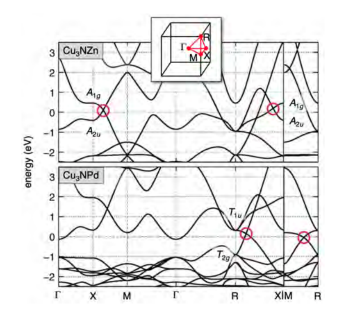


Figure 3: Electronic structures and Z_2 indices of (a) Cu_3NZn and (b) Cu_3NPd . Bands are drawn along the high-symmetry lines of the Brillouin zone (inset). The Dirac points are indicated by red circles. Parity eigenvalues are illustrated at the eight parity-invariant points in the first octant of the Brillouin zone. This figure and caption were reproduced with permission from Kim *et al.*⁶⁸

tetragonal structure, which is not surprising given the open nature of the structure.⁷¹ There has not been a detailed study of NTE in Cu₃N, but a recent report indicates that there is little, if any, expansion in the temperature range of 4 K - 100 K.⁷² More than fifty years from its discovery elapsed before the growth of thin films was reported,⁷³ and attention was drawn to the interesting semiconducting properties of Cu₃N. Shortly afterwards, in 1990, it was shown that such films, which are typically green, could be used as an optical data recording medium using infrared light.⁷⁴ Since copper does not react directly with nitrogen gas, the films were made by irradiating copper with nitrogen ions during film deposition onto a substrate (though bulk samples can be made from the reaction between CuF₂ and NH₃).⁷⁵ Since 1990, there has been extensive work on the preparation of high quality thin

films by using a variety of methods, especially magnetron sputtering and RF plasma jets. In addition to its potential for optical data storage, Cu₃N has also been proposed as a defect-tolerant semiconductor for PV applications,⁷⁶ while in condensed matter physics it has been suggested as a model for a new type of topological semi-metal exhibiting Dirac line nodes near the Fermi energy on doping with nonmagnetic transition metal atoms, such as Zn and Pd (Figure 3).⁶⁸ In the energy storage area, Cu₃N has been studied as a potential cathode material for lithium ion batteries, though the chemistry is complex and may well preclude any real applications in this field.⁷⁷ It has also been studied as a potential cathode for oxygen reduction in alkaline fuel cells.⁷⁸

Dark-blue sodium nitride, Na₃N, which is isostructural to Cu₃N, has been prepared by the reaction of metallic sodium or liquid Na-K alloy with plasma-activated nitrogen at low pressures.⁷⁹ Na₃N decomposes above 104° C into the elements, with ΔHf estimated at +64(2) kJ/mol.⁸⁰ The band gap of Na₃N is in the range 0.5 eV – 1.0 eV, which is similar to that of Cu₃N. It shows no phase change down to 20 K, but does exhibit strongly anisotropic effects.⁸¹ There is no tilting of the NNa₆ octahedra, nor is there any NTE, which is perhaps surprising. However, like Cu₃N, it does exhibit interesting topological semimetal behavior.⁸² It is also worth noting that neither Li₃N nor K₃N adopt the anti ReO₃-type structure.^{83,84}

Fluorides and Oxyfluorides ReO₃-type fluoride materials, of general formula MF_3 , are more common than their oxide counterparts, owing in part to the greater prevalence of

3+ oxidation state metals. The *M*-site metals for MF_3 compounds include M = Sc,⁸⁵ Co,⁸⁶ Al,⁸⁷ Cr,⁸⁸ V,⁸⁹ Ti,⁹⁰ and Fe.⁸⁸ It is worth noting that rare-earth cations, even the smallest example of lutetium (ionic radius of 0.861 Å)^{91,92}, appear to be too large to adopt the octahedral coordination required for the ReO₃-type structure. The majority of the single metal ReO₃-type fluoride compounds adopt a rhombohedral distorted variant at room temperature, with the exceptions being ScF₃, a notable NTE material, and MnF₃, which is Jahn-Teller distorted.⁸⁵ As will become apparent throughout this review, the majority of inorganic ReO₃-type materials have been studied for properties that are exhibited by ReO₃ itself: curious thermal expansion, as a host for guest cations, and unusual structural behaviour with pressure. This does not mean, however, that they do not, or cannot, display other interesting properties.

In terms of MF_3 compounds, ScF_3 is perhaps the most well known as it displays a remarkable temperature range of NTE (linked to the transverse vibrations of the F anions).⁹³ From 10 K to approximately 1100 K,⁹⁴ the material does not undergo any first-order phase transitions (although a possible quantum phase transition has been reported⁹⁵), and displays a coefficient of NTE near -8 ppm K^{-1} at room temperature. This is comparable to -9 ppm K^{-1} for the well known high performing NTE compound, ZrW_2O_8 .⁸⁵ Furthermore, it was recently shown that the degree of NTE in ScF_3 is particle size dependent,⁹⁶ and can actually be tuned between NTE, zero thermal expansion (ZTE), and PTE, by nanostructuring the product.⁹⁷ The tunability has been ascribed to local symmetry breaking as the particle size is decreased, i.e. the local structure is rhombodrally distorted (as evi-

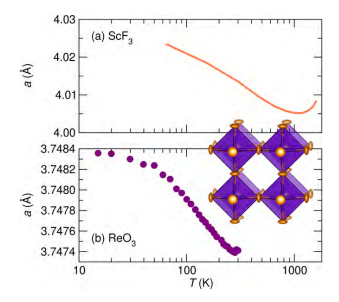


Figure 4: Lattice constant versus temperature (log scale). (a) The change in lattice constant for ScF₃; combination of synchrotron X-ray and neutron diffraction data.⁸⁵ (b) The change in lattice constant for ReO₃ based on neutron diffraction data.²⁷ The cubic ReO₃ structure type is shown as an overlay (95% thermal ellipsoids). Note the magnitude of the NTE is much greater for ScF₃ than for ReO₃. The Figure contains digitized data adapted with permission from Greve *et al.*⁸⁵ and Rodriguez *et al.*²⁷

denced by pair distribution studies) but appears cubic on average. The source of this local symmetry breaking is thought to be the increased surface pressure from the decreased particle size. This tuning can also be achieved by Zr doping.⁹⁸ It has also been found that NTE in ScF₃ can be essentially turned off by doping with a small amount of Fe and intercalating an equal amount of Li into the vacant *A*-sites.⁹⁹ The inclusion of Li ions limits the transverse vibrations of the fluoride ions, much the same way that guest molecules can dampen NTE in other porous NTE materials.^{100–104}

Redox active MF_3 materials that have been examined as intercalation materials for batteries include those where M = Ti, V, Mn, Co, and Fe.^{105,106} The predominant focus has been on FeF₃ as it displays the most well behaved charge/discharge profile, a viable voltage range (2V – 4V), and increased thermal stability upon lithiation.¹⁰⁷ However, due to the low electrical conductivity of FeF₃ (and fluorides in general), ball-milling with a conductive carbon to increase electron transport is always needed. In the case of FeF₃, a reversible capacity of 80 mAh g¹ is achievable after this process.¹⁰⁵ Aside from conductive carbon additives, nanostructuring the FeF₃ to increase the surface area for lithiation,^{108–112} or doping with cobalt, have been studied as ways to increase conductivity. However, theory suggests that FeF₃ becomes less stable with Co-doping¹¹³ and tends to form a tungsten-bronze related structure, a potential battery material in its own right.¹¹⁴ The tungsten-bronze structure type is also observed with iron fluoride hydrate, FeF₃·0.33 H₂O, and performs similarly to the Co doped variety.¹¹⁵ Lastly, there are reports that the perovskite NaFeF₃ can be completely (100%) desodiated/sodiated reversibly, and is of interest for future Na battery cathodes.¹¹⁶

Double fluoride ReO₃-type compounds, with the general formula *MM*^T₆, come in many varieties. Examples include the parent compound, $Sn^{2+}Sn^{4+}F_6$,¹¹⁷ $M^{2+}(Zr)F_6$ compounds (where M = Fe,^{118,119} Pd,¹²⁰ Co,¹²¹ Zn,¹²¹ Mg,¹²² Cu,¹²³ V,^{124,125} Ti,¹²⁴ Ca,^{119,126} Cr¹¹⁹), M^{2+} (Hf)F₆ compounds (where M = V,¹²⁴ Ti,¹²⁴), *M*MoF₆ compounds (where M= Na¹²⁷ and Cu¹²⁸), *M*NbF₆ compounds (where $M = Cr^{128}$ and V^{125,129}), and NaSbF₆.¹³⁰ There are examples within those compounds which display NTE,^{126,131,132} gas adsorption under pressure,¹²⁶ as well as Jahn-Teller distortions governed by the M^{2+} character.^{119,123,125} There are also the recent reports of the fluoride rich ReO₃-type compound, YbZrF₇,¹³³ which displays NTE below room temperature and ZTE at 300 K, and the Mg_{1x}Zr_{1+x}F_{6+2x} compounds in which the degree of thermal expansion can be tuned by varying x.¹³⁴

ReO₃-type oxyfluorides, with general formula MO_xF_u (M = Ti, ¹³⁵ V, ¹³⁶ Ta, ¹³⁷ Nb, ^{135, 137} Zr,¹³⁸ Mo¹³⁹) are also known, with the first example reported in the 1950's.¹³⁷ They are predominantly tertiary compounds, but there are examples of quarternary compounds, like the non-stoichiometric $Nb_{0.55}Zr_{0.45}O_{1.1}F_{0.8}$,¹³⁸ CrNb₂O₄F₅ and Fe₂Nb₃O₆F₉,¹⁴⁰ and MgNb₂O₃F₆ and MgTi₂OF₈.¹⁴¹ The ReO₃-type oxyfluorides have been primarily studied as intercalation materials for use in batteries, 135, 136 with the greater class of oxyfluorides viewed as a favorable middle ground between oxides and fluorides, leveraging the high voltage levels of the fluorides and increased stability of the oxides (due to the more covalent *M*–O bonds). As is the case with all battery materials, understanding the underlying crystal chemistry of each compound is important, as anion and cation order can have a substantial impact on material performance. Recent work in this respect was on NbO₂F and TaOF₂, where samples prepared via solution routes (as opposed to conventional solid state methods) were reported to have vacancies and hydroxide (OH⁻) ions introduced during preparation.¹⁴² In the same study, precise solid-state preparation of the materials, and subsequent diffraction studies, facilitated the development of a supercell model to describe the anion ordering motif. Another recent example was the application of VO₂F as a cathode material,¹³⁶ which demonstrated a reversible capacity of $250 \text{ mA} \text{ h g}^{-1}$ at 2.2 V (one equivalent of Li, C/50, with graphene additive). However, if the material is cycled past 2.2V (more than one equivalent of Li, C/50), it undergoes a conversion reaction which inhibits the cycling performance. This work was followed by a report where VO₂F was prepared via ball-

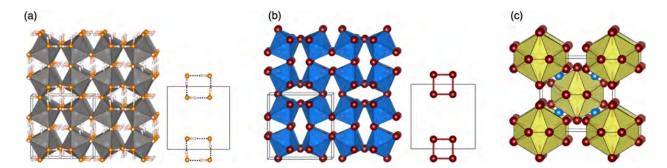


Figure 5: The highlighted structural similarity between (a) $In(OH)_3$ and (b) $CoAs_3$, drawn to scale. The bonding between As atoms follows the same motif as the "ice-rules" hydrogen bonding in $In(OH)_3$. (c) $CoAs_3$, emphasizing the 12 coordinate cavity, or cage, that can be filled with a cation. This cation "rattles" inside of the cavity, increasing thermoelectric performance via phonon scattering. Atom legend; indium = grey, oxygen = orange, hydrogen = white, cobalt = blue, arsenic = red, 12-coordinate site = yellow.

milling, as opposed to the high temperature solid-state synthesis of the original report,¹⁴³ and showed that the ball milled material had a reversable cyclability of 208 mA h g⁻¹ at 2.2 V, but at an improved cycle rate of C/20. This VO₂F material also underwent an irreversible phase change if cycled to lower voltages, which has been shown by multiple groups to be a disordered rock-salt phase.^{143–145} The most recent of those reports clarified the performance of the ball-milled VO₂F material,¹⁴⁵ and described that it delivers 276 mA h g⁻¹ (1.04 mol Li) and 206 mA h g⁻¹ (0.8 mol Li) at C/20 in its first and second cycles, respectively, with 97.5% capacity maintenance after 100 cycles. If taken to the irreversible rock-salt phase, 406 mA h g⁻¹ (1.54 mol Li, C/60) capacity was achieved, but with only 60% capacity maintenance after 50 cycles.

Hydroxides The hydroxide perovskites $[M(OH)_3, \text{ or } MM'(OH)_6]$ are a naturally occurring class of ReO₃-type materials well known to mineralogists,¹⁴⁶ with some synthetic examples joining the ranks. Members of this material class are showing great promise as future

oxygen evolution reaction (OER) catalysts, but there is still much to be understood about their fundamental crystal chemistry, as the hydrogen bonding of the –OH groups governs the final structure.

The single ReO₃-type perovskite, $M(OH)_3$, includes varieties where $M^{\text{III}} = \text{Al},^{147}$ Ga,¹⁴⁸ Fe,^{149,150} Sc,¹⁵¹ In,^{151,152} Lu,¹⁵³ and Yb.¹⁵⁴ The Ga, In, and Fe phases occur naturally, though all except for Fe (discussed below) can be prepared synthetically. The most common preparations are metathesis reactions, usually between a MCl₃ species and NaOH (to form a gel), which is then washed and heated with more base to crystallize the compound. Early reports of the Sc,¹⁵¹ In,^{151,152} and Lu¹⁵³ compounds described their structures, including a neutron diffraction study to establish the locations of the hydrogen atoms.¹⁵³ In the case of $In(OH)_3$ (as shown in Figure 5 (a)), we draw attention to how the hydrogen bonding follows a four-ring (square) pattern, but there are also "zigzag" and "crankshaft" hydrogen bonding patterns, which guide the structures of the hydroxide perovskites.¹⁴⁶ Pure Yb(OH)₃ has eluded synthetic efforts, although it is known to form solid solutions (up to 25% Yb) with both Lu(OH)₃ and In(OH)₃.¹⁵⁴ As stated above, Fe(OH)₃·nH₂O (bernalite is its mineral name),¹⁵⁵ is a curious case. There are no definitive reports of a laboratory prepared ReO₃-type Fe(OH)₃, but there are reports of a "Fe(OH)₃ powder" which is distinct from the $Fe(OH)_3$ gels that form in high pH Fe^{+3} cation containing solutions. This powder, we suspect, is the amorphous version of the ReO₃-type compound, with considerable disorder due to water molecules inside the cavities. This $Fe(OH)_3$ powder has been the subject of Mössbauer and magnetic characterization in order to differentiate it from the gels, albeit without definitive structural characterization.^{149,150} For the most part, these single metal materials have fewer applications than the double hydroxides discussed below, though $In(OH)_3$ can be synthesized with nanomorphology and has been converted to the more industrially relevant In_2O_3 whilst maintaining nanostructure.¹⁵⁶

There are numerous mineral examples of double perovskites, $MM'(OH)_6$, with a heavy predominance of stannates (the formal precursor is stannic acid, $Sn(OH)_4$). Some examples are $CaSn(OH)_6$ (burtite), $FeSn(OH)_6$ (natanite), $ZnSn(OH)_6$ (vismirnovite), and $MnSn(OH)_6$ (wickmanite). These stannate minerals crystallize in cubic space groups (except for $MnSn(OH)_6$, which is reported to be tetragonal) and exhibit cation ordering. There are also other minerals, such as $Fe^{3+}Sn^{4+}(OH)_5O$ (jeanbandyite), $NaSb^{5+}(OH)_6$ (mopungite), and $Fe^{2+}Ge^{4+}(OH)_6$ (stottite) which are not stannates. Synthetic varieties of double ReO₃ hydroxide compounds, such as $CuSn(OH)_6$,¹⁵⁷ CoSn(OH)6,¹⁵⁸ and SrSn(OH)6,¹⁵⁹ are also known, with the latter example by Woodward and co-workers illustrating that the ionic sizes and electronegativities of the *M*-site cations are important characteristics for ReO₃-type structure formation. This was made evident by demonstrating that both LiSb⁵⁺(OH)₆ and BaSn(OH)₆ both crystallize in other structural motifs.¹⁵⁹

Magnetic studies of double hydroxide perovskite materials have revealed that for systems with strong charge ordering, e.g. $MnSn(OH)_6$ and $CoSn(OH)_6$, the materials are paramagnetic down to 2 K, with $MnSn(OH)_6$ displaying weak antiferromagnetic interactions at low temperature.¹⁵⁸ FeSn(OH)₆ and FeSn(OH)₅O (where the former forms the

latter when left in air for a few hours) have also been examined,¹⁶⁰ with FeSn(OH)₆ exhibiting paramagnetism down to 1.4 K, and FeSn(OH)₅O ordering antiferromagnetically at 4 K. Additionally, the bulk Jahn-Teller distorted $CuSn(OH)_6^{157}$ displays weak antiferromagnetic behaviour and a spin-Peierls transition in strong magnetic fields at about 78 K.¹⁶¹

Most interestingly though, these compounds have found recent relevance as both photo and OER catalysts. Their photocatalytic performance is linked to how, when exposed to UV light, they generate OH^{-•} radicals, which can degrade organic molecules.¹⁶² The compounds $ZnSn(OH)_6^{163}$ and $CoSn(OH)_6^{164}$ have been of particular focus. In terms of OER performance, a recent example demonstrated the power of defect engineering by showing how an argon plasma treated sample of $Co_{0.90}Fe_{0.10}Sn(OH)_6$ performed better than a nontreated sample.¹⁶⁵ Due to the weaker Sn–OH bonds, Ar plasma treatment selectively removes Sn to expose more active Fe and Co sites, as well as increasing the micropore size of the material. When compared to the non Ar plasma treated material, the treated sample required a much lower overpotential for catalysis (270 mV at 10 mAcm⁻², with a Tafel slope of 42.3 mV dec^{-1}). Another recent example utilized a similar SnFeNi hydroxide system,¹⁶⁶ but implemented an electrochemical activation process to remove the Sn^{+4} cations. When compared to the well-known OER material, IrO_2 , the SnFeNi hydroxide material performed comparably. With the added benefit of being made from earth abundant metals, this is a promising field of research.

As the last examples indicated, defined control over the vacancies, and morphology,

of a double hydroxide perovskite proves essential to their use as functional materials. This kind of chemical control was previously exemplified by Nielson *et al.*¹⁵⁸, who demonstrated that if the hydrolysis of the Sn^{+4} ion can be kinetically controlled during the formation of $CoSn(OH)_6$ and $MnSn(OH)_6$ (by using a competing F⁻ anion), defined charge ordering of the metals can be promoted. This control is significant because if charge ordering can be maximized, materials like the OER catalysts should have the maximum amount of surface active sites available after Ar treatment.

Lastly, the ordering of the metals undoubtedly affects the hydrogen-bonding tendencies in certain systems, which as stated above, is a strong guide for the structure of the hydroxide perovskites. In future work, it would be worthwhile understanding how greater synthetic control can encourage certain hydrogen bonding arrangements (different polymorphs). Mineralogists have highlighted some of the significant, more fundamental, questions in this class of materials,¹⁴⁶ with the overarching problem being that the correct structures (e.g. space groups) of most hydroxide perovskites are not definitive. This is an area where better synthetic control can elucidate why certain polymorphs (tilts caused by hydrogen bonds) are favored in certain systems, with variable temperature and pressure¹⁶⁷ studies unraveling the poorly-understood phase transitions of the materials. Finally, with an average cavity diameter of 4.5 Å, many small molecules could be intercalated, but understanding how the hydroxide groups may play a role in guest-molecule intercalation is an open question.

Alloys with the ReO₃-type structure Skutterudite is a mineral that was named after the city of Skotterud, Norway, where it was first discovered in 1845. The naturally occurring version is a cobalt arsenide, CoAs₃, which also contains some nickel and iron on the cobalt site (Figure 5 (b)). The structure of the mineral form was first examined by X-ray diffraction in 1928 by Oftedahl,¹⁶⁸ who showed that it was cubic with space group $Im\bar{3}$, and a more precise determination was carried out by Mandel and Donohue in 1971.¹⁶⁹ The structure is formed from a network of corner-sharing $CoAs_6$ octahedra, as in ReO_3 , but these are sufficiently tilted that the arsenic atoms form approximately planar As₄ units (Figure 5). In order for the As₄ unit to be square, the x and y values of the As coordinates must obey the so-called Oftedal relationship, 2(y + z) = 1, but in the determination by Mandel and Donohue the values of x and y are such that the As₄ unit is rectangular with edges of 2.57 and 2.46 Å, respectively. There is clearly a significant amount of As-As bonding in CoAs₃, so the system can be regarded as a Zintl phase.¹⁷⁰ The structure is very similar to that of $In(OH)_3$, where the hydrogen bonding creates a square pattern (Figure 5), and indeed the two systems adopt the same $Im\bar{3}$ space group. Several closely-related families of naturally-occurring and synthetic alloys adopt the skutterudite structure, and this MX₃ family is known for M = Fe, Co, Ni, Ru, Rh, Ir, Os and X = As, Sb, P. However, we are not aware of any examples with M-site ordering, which is common for more ionic systems, as discussed previously.

The so-called "filled skutterudites" represent an exceptionally important extension of this family. The first examples were reported by Jeitschko¹⁷¹ and can be thought of as *A*-

site deficient perovskites, $A_x MX_3$, with x = 0.25 (i.e. their general composition is AM_4X_{12}). We have chosen to include them in this review because they are closer in composition to ReO₃ than perovskite and they have a wide range of very interesting electronic properties. Typical cases include systems such as LnM_4X_{12} , where Ln = rare-earth, M = Fe, Ru, Os, and X = P, As, Sb. Superconductivity below 4.08 K and 7.20 K was reported for $LaFe_4P_{12}$ and $LaRu_4P_{12}$, respectively, in 1981 by G. P. Meisner¹⁷² and magnetic ordering was found to occur at even lower temperatures when La is substituted by magnetic rare-earth elements such as Ce, Pr and Nd (superconductivity was not observed in the magnetic systems).

The initial discovery of superconductivity in LaFe₄P₁₂ and LaRu₄P₁₂ attracted the attention of the condensed matter physics community and lead to a large body of work on these materials in subsequent years. Beyond the studies on superconductivity, the most exciting development was the establishment by Sales *et al.* in 1996 that some of the filled skutterudites were excellent thermoelectric (TE) materials with, at that time, figures of merit, *ZT*, in excess of 1.0.¹⁷³ It was already recognized that conventional skutterudites had the potential to be good thermoelectrics,¹⁷⁴ and the filled skutterudites offer the opportunity for further enhancement. In particular, the rattling of the cations in the large *A*-site voids (Figure 5 (c)), along with the ease with which disorder can be introduced into these systems, both contribute to their excellent performance. These factors reduce the lattice thermal conductivity by scattering phonons and contribute to excellent TE performance, as discussed by Snyder and Toberer.¹⁷⁵ Subsequent developments in this area, including the use of filler cations with different charges, enabled the tuning of the carrier

density, leading to systems with TE performance that approach many of the better performing materials.¹⁷⁶ For example, $CoSb_3$ filled with various combinations of Ba, La and Yb have *ZT* values of up to 1.7 at 850 K, e.g. for $Ba_{0.08}La_{0.05}Yb_{0.04}Co_4Sb_{12}$.¹⁷⁷

While the bulk of the work on skutterudites for thermoelectric and other applications has focused on the filled systems, as described above, there has been a recent report of an excellent figure of merit for an unfilled system in which the thermal conductivity was minimized by controlling the nanostructure and microstructure of the material by means of careful annealing.¹⁷⁸ In this way, a *ZT* of 1.6 was reported for a synthetic alloy of composition $Co_{23.4}Sb_{69.1}Si_{1.5}Te_{6.0}$, despite the absence of rattling cations in the *A*-site cavity. When the nano/microstructure was not controlled by annealing, the thermal conductivity was approximately 4-5 times higher, leading to a 60% decrease in *ZT*.

Cyanides and other Prussian blue analogue materials In view of the extensive and historical literature on perovskite-related cyanides such as Prussian Blue, $Fe_4[Fe(CN)_6]_3 \cdot xH_2O$,¹⁷⁹ it is not surprising that similar ReO₃-type cyanides are known. These materials are generally referred to as Prussian blue analogues (PBAs), though the literature does not differentiate between those that have cations on the *A*-sites (i.e. they are perovskites) and those that do not (i.e. they are ReO₃-types). Several cyanide based examples with the ReO₃-type structure, however, are known, including a number of bimetallic systems such as $M^{II}Pt^{IV}(CN)_6$,¹⁸⁰ Ga^{III}Fe^{III}(CN)₆,¹⁸¹ and Fe₄[Ru(CN)₆]₃·18H₂O.¹⁸² The cavities are larger than those of the simple oxides and fluorides, so there is often solvent, typically water,

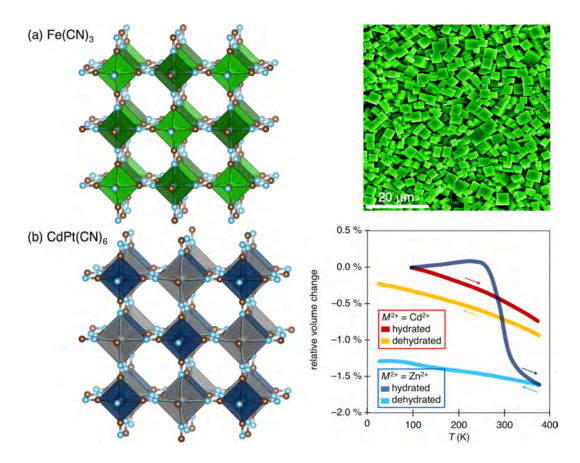


Figure 6: (a) Structure of Fe(CN)₃, with colorized SEM image of Berlin Green microcubes (SEM figure adapted with permission from Yang et al.¹⁸³ (b) The structure of dehydrated Cd^{II}Pt^{IV}(CN)₆ is drawn alongside a graph showing relative volume change as temperature is varied for M^{II} Pt^{IV}(CN)₆ (M = Zn, Cd). Intercalated water within the two compounds changes their NTE behavior by dampening the transverse vibrations of the –CN ligands. Data adapted with permissiom from from Goodwin *et al.*¹⁰⁰ Atom legend; Iron = light/dark green, carbon = brown, nitrogen = light blue, cadmium = grey, platinum = navy.

found within them. Anhydrous materials, solvated systems, and thiocyanates will be the focus of this section.

The simplest example is probably $Fe[Fe(CN)_6]$, i.e. $Fe(CN)_3$, which crystallizes in a doubled ReO_3 -type structure with alternate Fe^{III} ions coordinated by six carbons and six nitrogens, respectively, from the cyanide ions. It is sometimes known as Berlin Green be-

cause of its historical use as a pigment in paintings (Figure 6a).¹⁸³ The cation ordering, which gives rise to the doubling of the basic ReO₃ unit cell, is made possible by having the two iron sites in low-spin d^5 (FeC₆) and high-spin d^5 (FeN₆) states, respectively, as discussed below. The cavity in the as-synthesized material contains some water, but this can be removed under vacuum and samples of the anhydrous material have been studied as a function of temperature by synchrotron X-ray diffraction. Like many ReO₃-type structures, it exhibits pronounced NTE over a wide temperature range, in this instance from 100 K to 450 K.¹⁷⁹ As in the ReO₃-type oxides and fluorides, the NTE is associated with transverse vibrations of the carbon and nitrogen atoms of the cynaides; these are especially strong for the nitrogen atoms due to the longer Fe-N bonds. In an interesting variation on the above theme, a guest dependence of the NTE was reported for M^{II} Pt^{IV}(CN)₆ (M = Zn, Cd) by Goodwin *et al.*,¹⁰⁰ which showed that by including an intercalant species the transverse vibrations are dampened and NTE is suppressed (Figure 6b). As a point of interest, this dampening of NTE via intercalation is also observed for a related (non-ReO₃-type) porous rare earth cyanide compound,¹⁸⁴ and illustrates the similar structure-property relationship between the systems.

The magnetic properties of a hydrated form of $Fe[Fe(CN)_6]$ have been studied by low temperature neutron diffraction, Mössbauer spectroscopy and magnetic susceptibility methods, and the system is found to undergo a paramagnetic to ferromagnetic phase transition on cooling to 17.4 K.¹⁸⁵ This is consistent with the weak ferromagnetic exchange coupling that is expected between low-spin d⁵ Fe^{III} and high-spin d⁵ Fe^{III} via the cyanide

linker. There is also a long history of bimetallic PBAs that have been studied for their magnetic properties,¹⁸⁶ mainly in the context of molecular magnetism. This is a particularly fruitful area because the alternation of two different octahedrally coordinated cations that are linked by cyanide groups can give rise to a wide range of antiferromagnetic, ferromagnetic, and ferrimagnetic behavior. Most of the work in this area, however, has focused on systems with cations on the A-sites, so they are beyond the scope of this work, but there are a few examples of materials that adopt the ReO_3 -type structure. The most striking case is probably a phase with the approximate composition $VCr(CN)_6 xH_2O$, which has a ferrimagnetic ordering temperature of 315 K due to strong superexchange interactions between the vanadium and chromium (III) ions.¹⁸⁷ The system is more complex than it initially appears, however, because vanadium is present in both the V^{II} and V^{III} oxidation states, leading to a variety of possible magnetic interactions. Another example are the cobalt(II)-manganese(II)-chromium(III) compounds $(Co_x Mn_{1-x} [Cr(CN)_6]_{2/3} \cdot zH_2O,^{188})$ which depending on the compound of choice within the family, can be tuned between ferromagnetic and antiferromagnetic coupling depending on relatively humidity. The humidity-induced variations in the compounds are related to the coordination geometry of the Co ions, which in certain variants are undercoordinated due to $[Cr(CN)_6]$ vacancies, and can be coordinated to by water. Conversely, a counter example would be the ironruthenium PBA, $(Fe_4[Ru(CN)_6]_3 \cdot 18H_2O)$,¹⁸² which was examined in part for its potential magnetic behavior but found to display no ordering above 1.8 K. This material, however, is interesting as it demonstrates the effect of electron localization when compared to its

Prussian Blue parent and potassium intercalated sibling $(K_{1.2}Ru_{3.6}[Ru(CN)_6]_3)$, as the ironruthenium compound has decreased electrical conductivity and blue shifted optical intervalence charge transfer (IVCT) transitions.

Many PBA materials have also been explored for their potential applications in sodium ion batteries,¹⁸⁹ but work in this area has been hampered by low capacity utilization and poor cyclability. In the case of $Fe[Fe(CN)_6]$, the use of nanoparticles overcame these drawbacks, giving rise to a device with good kinetics, capacity and lifetime^{190,191} (we note that this strategy also worked for FeF_3 , as discussed earlier). There has also been work on both lithium¹⁹² and potassium¹⁹³ batteries with $Fe[Fe(CN)_6]$.

Finally, we would like to mention a very recent report of the first examples of thiocyanates with the ReO₃ structure.¹⁹⁴ The thiocyanate systems have the general composition M^{III} [Bi(SCN)₆] with M = Fe, Cr, Sc, and there is strict alternation of the Bi^{III} with the trivalent transition metal ions. This ordering is not too surprising, given the expected local bonding preference between sulfur and Bi^{III}. All the materials have band gaps in the visible or infrared regions, with the iron compound being as low as 1.20 eV, so they are strongly coloured and might be useful for applications that involve light harvesting. The *A*-site cavity would be expected to be even larger than those in the cyanides described above, but in fact the porosity is reduced by substantial monoclinic distortions of the hypothetical cubic structures. Nevertheless, the Cr^{III} compound exhibits reversible adsorption and desorption of water, forming a monohydrate. As in some of the other cyanides, the guest molecules have an impact on the thermal expansion, but there is no evidence of NTE in these systems.

Borohydrides M^{III} borohydrides, $M(BH_4)_3$, sometimes crystallize in the ReO₃ topology depending on synthetic route and the size of the metal ion. One well studied example is Y(BH₄)₃ (Figure 7) (a), a potential hydrogen storage material, which is attractive because it degrades to H₂ gas and solid YB₄ under 1–5 bar of partial hydrogen pressure (i.e. a typical operating condition of a H₂ storage system).^{195,198,199} Unlike other M^{III} borohydrides which have common by-products like elemental boron or diborane, YB₄ is more easily coverted back to Y(BH₄)₃, enabing a charge/discharge cycle.²⁰⁰ Crystallographically, Y(BH)₄ is reminiscent of the oxide ReO₃-type phases as it has two ReO₃-type polymorphs, α and β , which are both generated on synthesis by ball-milling and coexist at room temperature. However, the more dense and distorted α -Y(BH₄)₃ phase can be converted to the less dense and cubic β -Y(BH₄)₃ phase upon heating to 160°C - 180°C, ultimately decomposing past 190°C. Unlike the oxides, though, both phases show monotonic positive thermal expansion.

A more systematic study of the M^{III} –BH₄ reaction system with $M = \text{lanthanides}^{201}$ demonstrated the importance of the ionic radius of the M metal in determining whether the ReO₃-type structure type formed. Using ball-milling with a LiBH₄ precursor, the larger lanthanides (La, Ce, Pr, and Nd) formed only the cubic LiM(BH₄)₃Cl structure, while those closer in size to Y (Sm, Gd, Tb, Dy, Er, and Yb) all formed the α -Y(BH₄)₃ structure. The elements Sm, Er, and Yb are also stable in the β -Y(BH₄)₃ structure, under certain condi-

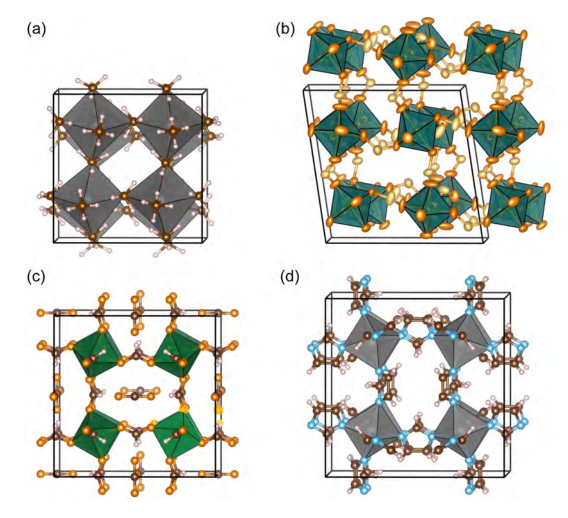


Figure 7: (a) Crystal structure of α -Y(BH₄)₃. Bonds are drawn between the Y and B atoms, to emphasize the ReO₃- type structure, though they are not present. It is in fact the bonding between Y and H atoms that guides the structure to be ReO₃-type.¹⁹⁵ Atom legend; yttrium = grey, boron = brown, hydrogen = beige. (b) Representative crystal structure with the ReO₃ topology of α -*M*(H₂PO₂)₃, shown for *M* = Ga in the *P*2₁/*n* space group (90% ellipsoids, H removed for clarity).¹⁹⁶ Atom legend; gallium = turquoise, oxygen = red, phosphorous = orange, hydrogen = beige. (c) Fe(HCOO)₃ with interstitial CO₂ (water removed for clarity).¹⁹⁷ Atom legend; iron = green, oxygen = orange, carbon = brown, hydrogen = beige. (d) In(im)₃. The imidazolate anions arrange in an alternating pattern in In(im)₃ to accommodate their size. Atom legend; indium = grey, nitrogen = blue, carbon = brown, hydrogen = beige.

tions. Further experimentation yielded the distorted ReO₃-type $M(BH_4)_3$ (M = La, Ce) through the reaction of MCl_3 and LiBH₄ in toluene at room temperature, followed by extraction of $M(BH_4)_3$ using dimethyl sufide [S(CH₃)₂] to form a solvated phase. Subsequent loss of S(CH₃)₂ from the adduct yields the $M(BH_4)_3$ phase.²⁰² The S(CH₃)₂ templating strategy combined with halide-free reagents (rare earth hydride and S(CH₃)₂·BH₃) was then applied generally to generate new structures, include the cubic $Pa\bar{3} \alpha$ -Pr(BH₄)₃ and α -Nd(BH₄)₃ structures,²⁰³ and α -Ce(BH₄)₃, β_2 -Pr(BH₄)₃, α -Lu(BH₄)₃, β -Pr(BH₄)₃.²⁰⁴ The latter paper also identified the structures of the S(CH₃)₂-adduct intermediates as layered phases, reminiscent of layered perovskites. Although the ReO₃ phases are not direct precursors, the related Li($M(BH_4)_3$)Cl phases have potential application as fast Li-ion conductors for solid-state battery electrolytes and gas storage,²⁰⁵ with some lithium borohydride phases having been tested in devices.²⁰⁶

Hypophosphites Until recently, the compound $V(H_2PO_2)_3$ was the only example of a hypophosphite containing ReO₃-type compound,²⁰⁷ but following work by the authors here, it is now understood to be a member of a polymorphic family of ReO₃-type materials, $M(H_2PO_2)_3$, where M = Al, Ga, and V [Figure 7 (b)]¹⁹⁶ There are four polymorphs seen in this family, (α , β , γ , δ), with each polymorph displaying different degrees of tilts and shifts of the *M* octahedra. Furthermore, each polymorph is preferred to varying degrees depending on the *M*, with each $M(H_2PO_2)_3$ forming at least two of the four polymorphs. The high pressure (above 1 GPa) δ phase has been observed for Ga(H₂PO₂)₃, which normally prefers the less dense α phase at STP. Synthetic control between each preferred polymorph is still

an open question, but to obtain at least one pure polymorph in each metal system, using an acid soluble oxide precursor is prefered (such as γ -Al₂O₃ instead of corundum for the Al system). Interestingly, the compound Al(H₂PO₂)₃, though its structure was unreported, is known to be an excellent bromine-free flame retardant in polymer blends.²⁰⁸ As the structure was only just reported, there are potentially advantages to using one polymorph as opposed to another if synthetic control can be leveraged.

3 Metal-Organic Frameworks

Systems with Organic Linkers We have seen in the previous section that there are several inorganic systems with the ReO₃-type structure in which the anionic linkers are polyatomic. Such compounds present an interesting variation on the perovskite topology because the vacant *A*-site may create a space that is large enough to accommodate adsorbed solvent or small gas molecules. This is seen in the cyanides and the thiocyanates, though the cavity in the hypophosphites appears to be too small for even monatomic neon.¹⁹⁶ Since the cavities defined by the larger molecular *X* sites may even be large enough to deliver permanent porosity, there is a significant similarity between such systems and metal-organic frameworks (MOFs). It is not surprising, therefore, to find that there are several examples of ReO₃-type materials in which the linker is an organic anion, and such materials can be properly regarded as MOFs. Examples of such systems, which are discussed below, include materials based on a wide variety of organic linkers, e.g. formates, azolates, guanidinates, and other, more complex ligands.

Formates, $M^{III}(\text{HCOO})_3 M = \text{Al}$, V, Fe, Ga, In ABX_3 perovskite-type phases where $X = (\text{HCOO})^-$, formate, have been extensively studied as they are easily prepared using solution synthesis and show a wide range of interesting ferroelectric, magnetic, and multiferroic properties.^{209–211} The classical Goldschmidt tolerance factor equation,²¹² which predicts the relative radii of cations and anions that are compatible with simple inorganic perovskite formation, has been extended to hybrid perovskites^{213,214} and can be used to predict the sizes of the *A* cations that will stabilize the perovskite structure in systems such as $AM^{II}(\text{HCOO})_3$. Because the cavity is quite large, it is not surprising that quite large protonated amines, such as dimethylammonium, $(\text{CH}_3)_2\text{NH}_2^+$, are required. As an extension of this observation, it is also not surprising to find that ReO₃-type systems of composition $M^{III}(\text{HCOO})_3$ have been reported, but only in the presence of neutral guest molecules that occupy the *A* site cavities.

A 1965 publication provided evidence that $V(HCOO)_3$ ·HCOOH may crystallize in the ReO₃ topology,²¹⁵ but a full structure determination for this compound has not been performed to date. However, the powder diffraction pattern²¹⁵ was indexed on a body-centred cubic structure that is consistent with M(III) formates reported more recently,¹⁹⁷ and other measures such as density are also close to the values reported for the Fe phase (V³⁺ and Fe³⁺ having similar molecular weights and radii). The neutral formic acid molecule, HCOOH, would be expected to occupy the *A* site.

The most recent work¹⁹⁷ on the $M^{\text{III}}(\text{CHOO})_3$ systems (M = Al, Fe, Ga, In) confirmed

that these phases, which can be readily formed by solvothermal or reflux methods, adopt an ReO₃-type structure with neutral *A*-site occupants assigned as combinations of H₂O, CO₂, and HCOOH [Figure 7 (c)]. All of these phases crystallize in the cubic $Im\bar{3}$ space group with a = 11.4Å– 12.2Å. Unlike many of the other carboxylate MOFs, the molecules occupying the *A*-sites in the M^{III} (CHOO)₃ systems could not be liberated while keeping the framework intact, suggesting that these molecules may have a structural role through strong hydrogen bonding or that the *M*–O bonds are insufficiently strong to stabilize the framework in the absence of guests. An alternate form of the Fe(CHOO)₃·guest ReO₃-type phase was also reported,²¹⁶ crystallizing in the trigonal $R\bar{3}c$ space group with a = 8.2Å, c = 22.6Å. It exhibits antiferromagnetic ordering at low temperatures.

The formate ReO₃-type compounds could perhaps be regarded as the simplest MOF systems and can be made from inexpensive, earth-abundant starting materials. What is particularly interesting is that they show a very strong preference for forming in the presence of CO₂, and Al(HCOO)₃·xCO₂/yH₂O can be obtained by bubbling CO₂ through an aqueous solution containing Al(OH)₃ and formic acid.¹⁹⁷ The fully stoichiometric phase, Al(HCOO)₃·CO₂, has yet to be achieved, but with a potential CO₂ loading of greater than 21 wt%, it is an interesting option for CO₂ storage.

Nitrogen-containing Organic Linkers In light of what has been reported for the formates, where the ReO₃-type MOFs contain metal-oxygen bonds, it would be reasonable to expect that MOFs with the ReO₃-type structure might also form with nitrogen-containing organic linkers that are capable of metal-nitrogen bonding. This has so far been realized for guanidinates, 1,2,4-triazolates, and imidazolates, though there are clearly other potential systems that might be feasible.

Only one guanidinate with the ReO₃-type structure is known to exist, Yb[C(NH₂)₃)₃.²¹⁷ This phase was synthesized in liquid ammonia and crystallizes in the cubic $Pn\bar{3}$ space group with a = 13.5 Å. Braced by the extensive hydrogen bonding between guanidinate anions, it has a relatively low density (1.9 g cm⁻³) and potential voids in the structure. Unlike many of the simple inorganic ReO₃-type systems, Yb(CN₃H₄)₃ shows positive thermal expansion rather than NTE. It is interesting to note that guanidine derivatives can play a role in perovskite-related materials both as an anionic linker (as in the present Yb phase) and as an *A*-site cation (as in the formate perovskites²¹⁸). Clearly the control of pH is very important in determining which of these roles it is able to play.

While azolates represent a well-known building block for MOFs (particularly ZIFs, where imidazolates are used as linkers), they are primarily used in tetrahedral coordination with M(II) cations. Nevertheless, there are a small number of ReO₃-type structures based azolates. Yb(tz)₃ (tz = 1,2,4-triazolate) was prepared from Yb metal and a melt of the azole, crystallizing in the cubic $Pm\bar{3}m$ space group with a = 6.8 Å.²¹⁹ Yb(tz)₃ is arguably not strictly an ReO₃-type as half of the tz ligands coordinate η_1 while the other half coordinate η_2 , giving ytterbium a coordination number of nine; however, the overall topology is ReO₃-like. This phase is well-packed and shows no potential porosity. Ga(trz)₃

(trz = 1,2,4-triazolate) was later reported by the same group,²²⁰ forming at the interface between the immiscible liquid metal and liquid triazole. This phase crystallizes in the cubic I_{23} space group with a = 12.3 Å, and gas sorption testing showed that Ga(tz)₃ is also a dense framework with no accessible porosity. We are not aware of any ReO₃-type structure based on 1,2,3-triazolate linkers.

Most recently, imidazole (im) was successfully used to form $In(im)_3 xG$ (G = guest) ReO₃-type phases via multiple pathways including ionothermal, solvothermal, and solventless syntheses.²²¹ Three phases were obtained - I, II, and III - with structural parameters of $Im\bar{3}, a = 13.0 \text{ Å}; R\bar{3}, a = 18.9 \text{ Å}, c = 10.5 \text{ Å}; \text{ and } R\bar{3}, a = 9.1 \text{ Å}, c = 22.3 \text{ Å}, \text{ respectively}$ [Figure 7 (d)]. While gas sorption showed no accessible porosity, varying amounts of unbound neutral molecules appeared to be trapped inside the framework cavities of phases I and II. Phase III was reported as a completely dense, guest-free structure. All structures were solved from PXRD, and the identities and quantities of neutral guest molecules in I and II were assigned by NMR to a range of molecules including solvent and solvent decomposition products.²²¹ The structures were distinguished by different octahedral tilting patterns. Phases I and II showed a reversible, discontinuous, hysteretic phase transition at ca. 90°C. The synthesis method affected the transition temperature, and it was suggested that this was due to the different guest molecules trapped within the framework. It would seem unlikely that In^{III} is the only trivalent cation capable of forming ReO₃-type networks with imidazolate, so this seems to be a very interesting area for future exploration.

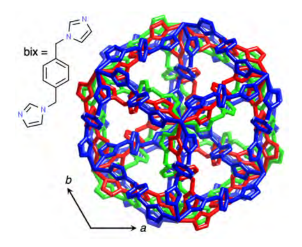


Figure 8: Triply interpenetrating MOF based on a *bis*-imidazole ligand. Figure was adapted with permision from Abrahams *et al.*.²²²

Systems with More Complex Linkers and Interpenetration There are several examples of positively charged ReO₃-type MOF frameworks with large organic linkers and chargecompensating anions in the *A*-site cavities,²²³ but these are more like perovkites than ReO₃ systems (albeit with inverse polarity). They are sometimes referred to in the MOF literature as having the α -polonium structure type (α -Po being the only simple cubic metal structure), irrespective of the framework being charged and the presence of anions in the *A*-site cavities. An important example is provided by the large, neutral *bis*-imidazole ligands (Figure 8), which forms ReO₃-type frameworks with Cd²⁺ nodes.²²² What is particularly interesting is that the frameworks are triply interpenetrating, presumably to reduce the unoccupied space in the channels. Such behaviour is also seen in some of the MOF-5 related materials discussed below. The positive charges of the frameworks in these structures are balanced by nitrate or tetrafluoroborate anions which sit in the small cavities created by the interpenetrating ReO₃ nets. One would imagine that neutral frameworks of this type could be obtained by using a trivalent cation in combination with anionic ligands.

MOFs with Polyatomic *B*-site Cations In this final section on MOFs with the ReO₃-type structure, we shall focus on an even more complex group of materials in which the cations on the *B*-site are polyatomic clusters, again linked by organic ligands. The most iconic example is MOF-5, shown in Figure 9, where tetrahedral $Zn_4O_6^+$ clusters are connected by 1,4-benzenedicarboxylate (1,4-bdc) anions to form a 3-D network of composition $[Zn_4O](1,4-bdc)_3$ [Figure 9 (a)].¹⁷ This important discovery from Yaghi's group lead to another concept known as reticular synthesis,^{224,225} whereby structures with even larger cavities could be obtained by using longer linkers, as shown in Figure 9 (b). By so doing, it is possible to tune the porosity of the MOF system for specific applications. This family of MOFs is, therefore, justifiably famous for being a model for the optimization of porous frameworks for applications in separation and catalytic processes.²²⁶ We note that interpenetration is often found in structures based on the longer linkers used to synthesize some of the IRMOF materials, though steps can be taken to mitigate this outcome.

There can be endless variations on the MOF-5 theme if we allow for the possibility that the 1,4-bdc linker might bear substituents on the benzene rings. Typical examples include $-NH_2$, -Br, -Cl, $-NO_2$.²²⁸ Yaghi refers to such systems as multivariate MOFs on account of the range of compositional variations that are possible, such as combining eight distinct functionalities into a single phase. The chemical variations are not of solely academic interest, since it is possible to tune the adsorption behaviour of the MOF for partic-

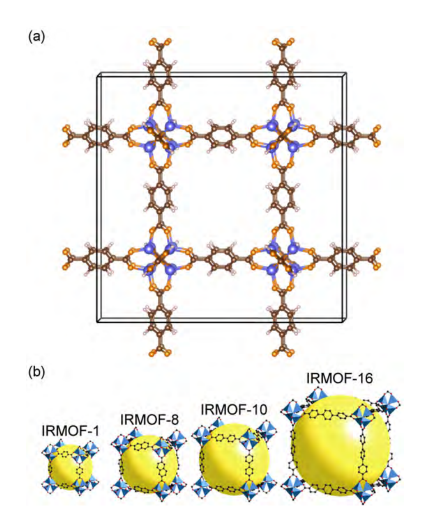


Figure 9: (a) The structure of the metal-organic framework, MOF-5, with a Zn_4O unit on the *M*-site, 1,4-bezenedicarboxylate anions on the *X*-site, of an ReO₃ topology.¹⁷ This structure type is referred to as pcu (primitive cubic) in the MOF community. Atom legend; zinc = light blue, oxygen = orange, carbon = brown, hydrogen = white. (b) Isoreticular series of MOFs, where IRMOF-1 = MOF-5. Panel b was from Rosi *et al.*²²⁷

ular applications. In one instance, the selectivity for CO_2 adsorption compared with CO could be enhanced by 400% by fine-tuning the composition.²²⁸

There are symmetry constraints on the types of systems that can form an ReO₃-type network, since the building unit at the *B*-site node must be compatible with 6-fold connectivity, as discussed in a review by Zaworotko *et al.*²²⁹ This is achieved in MOF-5 by using the two oxygens from one end of the 1,4-bdc carboxylate linker to bond to two different zinc ions in the cluster. In spite of this constraint, there are many other examples of systems that adopt the ReO₃-type structure (which is often referred to as the pcu - primitive cubic - topology in the recent MOF literature, though this appellation is sometimes interpreted more loosely than we have done in this review). Eye-catching examples include a system in which the nodes use $Cu_2(CO_2)_4$ paddlewheel clusters as vertices.²³⁰. To give a sense of the potential scope of this area, we note that Snurr and Hupp explored the methane storage and delivery capacity of 122,835 hypothetical pcu frameworks²³¹ from the MOF database of Wilmer *et al.*,²³² though the vast majority of these have never been synthesized.

There are many other MOFs in the literature that are described as PCU-type that we have not included here because they do not preserve the cubic or pseudo-cubic architecture of the ReO₃-type structures. Typically they have different linkages along different axes, such as the pillared-layered examples^{233,234} and other structures based on paddlewheel nodes.²³⁵

4 Concluding Remarks

The chemical diversity of materials with the ReO_3 -type structure is much greater than we had imagined when we began to think about this article in the summer of 2018. Furthermore, their properties and applications, both actual and potential, have also surpassed our expectations. The timeline shown in Figure 2 highlights some of the important systems that have been developed over the last eighty-plus years, from their initial discovery or structural characterization to the demonstration of their most striking properties. In addition to the remarkable range of their behaviour, we are struck by the long gaps between the initial discoveries of several of the materials and the establishment of their functionalities. In the case of ScF₃, for example, seventy years elapsed between the first structural characterization in 1939²³⁶ and the recognition of its outstanding NTE behaviour in 2010! At the same time, several new ReO₃-type families of been reported in the last decade, including the borohydrides, hypophosphites, guanidinates and imidazolates, though the properties of these new families have yet to be explored.

One of the most exciting discoveries in the very recent past concerns a 2015 report that hydrogen sulfide, H_2S , transforms to H_3S under high pressure and that this new phase is superconducting with a Tc of 203 K.²³⁷ This possibility was predicted in 2014²³⁸ and subsequent work in 2016 has indicated that the superconducting H_3S phase adopts a doubly interpenetrated ReO₃ structure.²³⁹ However, others have claimed that the superconducting phase is (SH⁻)(SH³⁺) and adopts the perovskite structure with (SH⁻) on the *A*-site,²⁴⁰ and Table 1: Compilation of cases where the ReO_3 analogue of a known perovskite family is unknown, or *vice versa*. The unknown phases are labled with "NR" (not reported). In some cases, e.g. with 1,2,3-triazole, both are unknown but might be expected to form.

Perovskite family	ReO_3 analogue
(AmineH)Pb X_3	BiX_3 (NR)
$A^{I}B^{II}(OH)_{3}$ (NR)	$M^{\rm III}({\rm OH})_3$
$(AmineH)M^{II}(N_3)_3$	$M^{III}(N_3)_3$ (NR)
(AmineH) <i>M</i> ^{II} (Gua) ₃ (NR)	<i>M</i> ^{III} (Gua) ₃
(AmineH) <i>M</i> ^{II} (Im) ₃ (NR)	$M^{ m III}(m Im)_3$
$(AmineH)M^{II}(1,2,4-triazole)_3$ (NR)	M^{III} (1,2,4-triazole) $_3$
$(AmineH)M^{II}(1,2,3-triazole)_3$ (NR)	M^{III} (1,2,3-triazole) ₃ (NR)
(AmineH)M ^{II} (tetrazole) ₃ (NR)	M^{III} (tetrazole) $_3$ (NR)

recent calculations broadly support the idea of such a disproportionation of H_2S .²⁴¹

We noted at the beginning of the article that most of the ABX_3 perovskite families can form ReO₃ analogues, and *vice versa*, so it is appropriate at this point to revisit this issue and identify the systems where this has not been established. Table 4 shows a number of cases where the analogues are not known, or even where neither form is known but could potentially exist. In some cases, such as the ReO₃ form of BiI₃, the compound prefers a layered structure, which is presumably stabilized by strong van der Waals interactions between the iodide ions. In the case of the hypothetical $A^{1}B^{11}(OH)_3$, it might be that the protons that protrude into the *A*-site cavity render it unsuitable for hosting a metallic cation. However, there are several nitrogen-containing linkers that might lend themselves to the formation of new ReO₃-type or perovskite families, though we note that some of these may be high energy materials, such as the azides and the tetrazoles. Future work in these areas is certainly needed and has the potential to produce new phases with interesting and unusual properties.

Conflicts of interest

There are no conflicts to declare.

Acknowledgments

HAE thanks the National Research Council for financial support through the Research Associate Program (RAP). AKC thanks the Ras al Khaimah Centre for Advanced Materials for financial support.

Competing Interests The authors declare that they have no competing financial interests.

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