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Review

Recent advances in crystalline oxidopolyborate complexes of *d*-block or *p*-block metals: structural aspects, syntheses and physical properties

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 Abstract:
 Crystalline materials containing hybrid inorganic-organic metal borates (complexes
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 with oxidoborate ligands) display a variety of novel framework building blocks.
 The structural
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 aspects of these hybrid metallaoxidoborates containing Cd^(II), Co^(II), Cu^(II), Ga^(III), In^(III), Mn^(II), Ni^(II) or
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 Zn^(II) metal centres are discussed in this review.
 The review describes synthetic approaches to these
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 hybrid materials, their physical properties, their spectroscopic properties and their potential applications.
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Keywords: Borates; Coordination compounds; Inorganic-organic hybrids; Oxidoborate ligands.

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1. Introduction

In borate chemistry (*n.b.* oxidoborate is the recommended IUPAC name for oxidized 20 oxygen containing borates [1]) the boron centres are bound to oxygen atoms as sp^2 hybrid-21 ized triangular $\{BO_3\}$ (Δ) or sp^3 hybridized $\{BO_4\}$ tetrahedral (T) structural units [2-4]. 22 These fundamental units can be aggregated, employing organic cations or transition-23 metal cations as templating agents [5], with condensation and oxygen atom corner sharing 24 into larger oxidoborate clusters. In these structures, terminal oxygen atoms (*i.e.* those not 25 corner shared) are generally also bound to hydrogen atoms as hydroxy groups [2-4]. Such 26 oxidoborates are often described as hydrated borates [6] and these compounds are readily 27 formed under relatively mild conditions [2-6]. Harsher conditions can lead to anhydrous 28 borates and although very rare, even to the possibility of edge sharing oxygen atoms [7]. 29 Descriptors have been developed by Christ and Clark [2] and by Burns and co-workers 30 [4] to designate reoccurring structural motifs as framework building blocks (FBBs) and 31 Christ and Clark's descriptors [2] are used in this review. Hydrated oxidiborates may 32 enter the primary coordination shell of metals, with formation of O-donor coordinate 33 bonds with the result of even more complex and diverse species. Coordination com-34 pounds containing oxidopolyborate ligands are therefore an important sub-class of syn-35 thetic oxidoborate compounds and recent progress in this area is the subject of this review. 36 Since insular oxidopolyborate anions partnered by cationic transition metal complexes do 37 not contain oxidoborates as *ligands* they are not within the scope of this review and are 38 excluded. Metal complexes often contain more than one ligand type and those that con-39 tain oxidoborate ligands and conventional organic ligands with typical donor atoms may 40 also be classified as inorganic-organic hybrid materials [9]. Such hybrid compounds, may 41 potentially have unique and useful properties as a result of combining and/or enhancing 42 properties associated with non-hybrid single materials. 43

This review is designed to be comprehensive within the defined topic and covers 44 recently reported (twenty-first century) literature. It reports on structural chemistry, syn-45 thetic methods, physical properties and possible applications and focusses on complexes 46 containing oxidoborate ligands. The following *d*-block and *p*-block metal ions form such 47 complexes: Cd^(II), Co^(II), Cu^(II), Ga^(III), In^(III), Mn^(II), Ni^(II) and Zn^(II) and sub-sections are dedi-48cated to each metal. Oxidoborate ligands are discussed by increasing boron number 49 within these metal borate subsections. Compounds containing oxidoborates ligands were 50 first reported in the twentieth century and for information on such compounds the reader 51 is referred to an earlier review [3] which surveys general structural aspects of oxidoborate 52 chemistry. 53

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2. Structural (XRD) studies

2.1. Cadmium(II) borates

Several oxidoborate coordination compounds of Cd(II) have been reported during the 57 review period: $[Cd(dab)_{0.5}(dab')_{0.5}(B_5O_7(OH)_3)]_n$ (1) (dab = 1,4-diaminobutane) [8], 58 [Cd(dap)0.5(dap')0.5{B5O8(OH)}]n·nH2O (1,3-diaminopropane) (2) [9], 59 $[Cd(tren){B_8O_{11}(OH)_4}]_n$ tris(2-aminoethyl)amine) [10], (3) (tren) 60 (4) $[Cd(pn){B_6O_7(OH)_6}]n \cdot nH_2O$ = 1,2-diaminopropane) [11], (pn 61 $[Cd_3(H_2O)_4(NO_3)_2]{B_6O_9(OH)_2}_n$ (5) [12] and four related compounds, exemplified by 62 $[pyH]_2[Cd(py)_2[B_{14}O_{20}(OH)_6]]$ (6) (py = pyridine) [13]. The FBB's in these compounds are 63 mostly complex with the pentaborate(2-) unit in 1 is based on a FBB of 4(4-1) with a pen-64 dant $-B(OH)_2$ unit (FBB = 1) replacing a terminal H atom on a T unit of the larger FBB and 65 can be designated a compounded descriptor $5:[4:(2\Delta+2T)+\Delta]$. The FBB for **2** is based on 66 the pentaborate(2-) unit $5:(3\Delta + 2T)$. The octaborate(2-) and the tetradecaborate(4-) FBBs in 67 3 and 6 are also best designated as compounded descriptors 8:[5:(4A+T)+3:(2A+T)] and 68 14:[$\{7:(5\Delta+2T)\}_2$] respectively, with the latter dimer based on a FBB unit of 7. The con-69 densed hexaborate(2-) units in 4 and 5 have a more standard descriptor based on a FBB 70 unit of 6 *i.e.* 6:(2∆+3T). 71

Compound **1** is a hybrid inorganic-organic 3-D coordination polymer [8]. The 22electron Cd^(II) centre adopts a distorted six-coordinate octahedral complex with two *trans N*-donor from two different 1,4-dab ligands (forming a 1-D chain) and four *O*-donor atoms from two oxidoborate network ligands (Figure 1a).



Figure 1. The building units of (a) $[Cd(dab)_{0.5}(dab')_{0.5}\{B_5O_7(OH)_3\}]_n$ (1), (b) $[Cd(tren)\{B_8O_{11}(OH)_4\}]_n$ (3), and (c) $[Cd(pn)\{B_6O_7(OH)_6\}]_n$ nH₂O (4).

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Compound **2** is also a hybrid inorganic-organic 3-D coordination polymer [9]. In **2** the 2-D layers of repeating neutral [Cd{ $B_5O_8(OH)$ }] units are linked by bridging dap units into a 3-D structure, with each 20-electron Cd^(II) centre being 5-coordinate and ligated by two monodentate *N*-donor dap ligands and three *O*-donors from the oxidoborate framework. 91

Compound **3** is a Cd^(II) complex comprised of an anionic oxidoborate ligand 92 $\{B_8O_{11}(OH)_4\}^2$ fragment coordinated to a supporting $\{Cd(tren)\}^{2+}$ fragment (Figure 1b) [10]. 93 The 20-electron Cd^(II) centre in **3** is five-coordinate and is coordinated by four *N*-donors 94 from the tren ligand and one *O*-donor from the bridging O atom from the $\{B_3O_6(OH)\}$ subunit of the octaborate(2-) anion. These $\{B_3O_4(OH)\}$ sub-units, with additional pendant 96 $\{B_5O_7(OH)_3\}$ sub-units, link together into a 1-D chain polymer. These 1-D chains are joined 97 together into a 3-D framework *via* extensive H-bond interactions. 98

Compound 4 is an octahedral 22-electron Cd^(II) complex with a *cis*-pn ligand and four 99 O-donors from two hexaborate(2-) ligands, forming a 1-D coordination polymer chain 100 [11]. One of the hexaborate ligands is coordinated fac to the Cd^(II) centre via the three OH 101 groups bound to tetrahedral boron atoms and the final coordination bond is formed from 102 a trigonal boron OH group from another hexaborate(2-) unit (Figure 1c). The structure of 103 this complex is described as [Cd(1,2-dap)][B₆O₁₁(OH)₂]·H₂O in [11] but is better formulated 104as $[Cd(pn){B_6O_7(OH)_6}]n nH_2O$ (4). The hexaborate(2-) units in 4 not only bridge two Cd^(II) 105 centres but are also further linked together to form an extended network which is inter-106 connected via a strong H-bonding. 107

Compound **5** has a unique structure and it is comprised of 2-D planes of condensed 108 $\{B_6O_9(OH)_2\}^{2-}$ units coordinated to the terminal 22-electron $Cd^{(II)}$ centres of a linear 109 $\{Cd_3(H_2O)_4(NO_3)_2\}^{4+}$ unit (Figure 2a) is such a way as to form a crosslinked 3-D network 110 [12]. The central 22-electron $Cd^{(II)}$ centre of the centrosymmetric $\{Cd_3(H_2O)_4(NO_3)_2\}^{4+}$ sub- 111 unit (marked Cd* in Figure 2a) has two *trans* NO₃⁻ ligands and all three Cd^(II) centres are 112 octahedral with remaining sites occupied by terminal or bridging H₂O ligands or ox- 113 idoborate *O*-donor centres. 114



Figure 2. (a) The cationic {Cd₃(H₂O)₄(NO₃)₂}⁴⁺ unit of [(Cd₃){B₆O₉(OH)₂}₂(NO₃)₂(H₂O)₄]_n (5). Donor atoms marked O and OH from oxidohexaborate clusters, and (b) the building unit of Cd^(II)-centered metallaoxidoborate cluster anion, *trans*-[Cd(py)₂{B₁₄O₂₀(OH)₆}]²⁻ (6).

Compound **6** is typical of a series of four hybrid Cd^(II) oxidoborates that exhibit 3-D 124 open-framework with novel topologies [13]. All the networks are comprised of a novel 125 Cd^(II) centered complex *trans*-[Cd(py)₂{B₁₄O₂₀(OH)₆}]²⁻ and there are further interionic links 126 *via* H-bonding interactions. As shown in Figure 2b, the $[B_{14}O_{20}(OH)_6]^4$ coordinates by four 127

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O-donors from the four boroxole {B₃O₃} rings to the Cd^(II) centre in a square planar ar-128 rangement, with two additional axial N-donor ligands, resulting in a 22-electron octahe-129 dral complex.

2.2. Cobalt(II) borates

The triborate(1-) ligand with the FBB unit of a 6-membered boroxole ring with two 133 $\{BO_3\}$ units and one $\{BO_4\}$ unit *i.e.* $3:(2\Delta + T)$ (Figure 3a) is present in the recently synthe-134 sized salt [Co(H2O)₆]2[NO₃]2[Co(H2O)₄{B₃O₃(OH)₄}2]·2H2O (7) [14]. The 19-electron Co^(II) 135 centres are both octahedral and the two [B₃O₃(OH)₄] ligands in the neutral 136 [Co(H2O)4[B3O3(OH)4]2] complex are trans and are coordinated by hydroxy O-donors 137 bound to tetrahedral boron atoms.



Figure 3. (a) The structure of $[Co(H_2O)_4[B_3O_3(OH)_4]_2]$ (7), and (b) the building unit of $[Co(dap)_{0.5}(dap')_{0.5}[B_4O_7]]_n$ (8).

A diagram of the structure of $[Co(dap)_{0.5}(dap')_{0.5}[B_4O_7]]_n$ (8) [8] (Figure 3b) shows a five-coordinate trigonal-bipyramidal coordination geometry at the 17-electron Co^(II) centre with two N-donor coordinate bonds, from two different dap ligands and forming a 1-D chain, and three O-donor atoms from three anionic $\{B_4O_7\}^{2-}$ tetraborate units. The Christ and Clark descriptor for these units with a FBB of 4 is $4-1:(2\varDelta+2T)$.

The basic building unit of [Co(tren){OB₅O₆(OH)₃] (9) is shown in Figure 4a [15]. The 153 $[OB_5O_6(OH)_3]^{2-}$ ligand can be described as $5:(4\Delta + T)$ with one of four trigonal units depro-154 tonated. The 17-electron Co^(II) centre exhibits a trigonal-bipyramidal coordination geome-155 try, with four N-donor atoms and one O-donor atom. Compound 9 forms 3-D supramo-156 lecular network through extensive H-bond interactions. 157



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Figure 4. Drawings of (**a**) the uncharged unit of [Co(tren) {OB₅O₆(OH)₃}] (**9**), (**b**) the dianion observed in [1-cyanopiperazinium][Co{B₆O₇(OH)₆}]·4H₂O (**11**), and (**c**) the structure of [Co{(NH₂CH₂O)₃B₆O₇(OH)₃] (**12**).

> The homoleptic bis(hexaborate(2-))cobalt(II) complexes [piperazine-1,4-dium] 166 [Co{ $B_6O_7(OH)_6$]_2].6H₂O (**10**) [16] and [1-cyanopiperazinium][Co{ $B_6O_7(OH)_6$]_2].4H₂O (**11**) 167 [17] have been recently synthesized and characterized crystallographically. The oxidoborate ligands in **10** and **11** are designated 6:(3_4 + 3_7) and it is the three hydroxyl O atoms on the three {BO4} centres that coordinate in *fac*- geometries to the octahedral **19**electron Co^(II) centres (Figure 4b). In both these structures there are strong templating interionic H-bond interactions. 172

> The derivatized hexaborate(2-) ligand observed in $[Co{(NH_2CH_2CH_2O)_3B_6O_7(OH)_3}]$ 173 (12) [18] has three 2-amino ethoxy groups in place of the three -OH groups on the tetrahedral boron centres of a hexaborate(2-) ion found in 10 and 11. It functions as a hexadentate 175 ligand through the three *O*-donors, bound to the aminoethyl substituents and the three amino *N*-donors to the 19-electron Co^(II) centre (Figure 4c). 177

2.3. Copper(II) borates

Oxidoborates coordinated to Cu^(II) centres are not uncommon and are available for 180 diborate(2-), pentaborate(1-), hexaborate(2-) and icosaborate(12-) anions. The diborate(2-) 181 anion is observed as part of the templated oxidoborate found in 182 [H₃O]₄[Cu₇(NH₃)₂(H₂O)₄{B₂₄O₃₉(OH)₁₂]¹3H₂O (13) [19] and this will be discussed later on 183 in this sub-section. 184

As shown in Figure 5a, $[Cu(pn)_2\{B_5O_6(OH)_4][B_5O_6(OH)_4]\cdot4H_2O$ (14) [20] is an ionic 185 compound comprised of a cationic hybrid $Cu^{(II)}$ complex containing a pentaborate(1-) ligand based on a FBB of $5:(4\varDelta+T)$. This +1 cation is partnered with an additional insular 187 $[B_5O_6(OH)_4]$ anion. The 19-electron $Cu^{(II)}$ ion in 14 has a distorted square based pyramidal 188 geometry with four *N*-donor atoms, and an axial *O*-donor pentaborate(1-) with a T^5 [21] 189 of 0.87. A sixth O-donor H₂O potential 'ligand' is axially *trans* to the pentaborate(1-) ligand but the Cu-O distance is not within normal (or even long) bonding distances. 191



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Figure 5. (a) The cation present in $[Cu(pn)_2\{B_5O_6(OH)_4\}][B_5O_6(OH)_4]\cdot 4H_2O$ (14), and (b) the uncharged building unit of $[Cu(deen)\{B_6O_7(OH)_6\}]\cdot 5H_2O$ (19).

Coordinated hexaborate ligands are well represented in Cu(II) coordination chemistry 201 as illustrated by the following examples: [Cu(NH₃)₂{B₆O₇(OH)₆}]_n·2nH₂O (15) [22], 202 $[Cu(en){B_6O_7(OH)_6}]n \cdot 3nH_2O$ (16) (en = 1,2-diaminoethane) [23], 203 $[Cu(dmen){B_6O_7(OH)_6}] \cdot 4H_2O$ (17) (dmen = N,N-dimethyl-1,2-diaminoethane) [20], 204 $[Cu(tmeda){B_6O_7(OH)_6}]$ ·6H₂O (tmeda = N, N, N', N'-tetramethyl-1,2-diaminoethane) 205 (18) [20], and $[Cu(deen){B_6O_7(OH)_6}] \cdot 5H_2O$ (19) (deen = N, N-diethyl-1,2,-diaminoethane) 206

[22]. The organic N-donor ligands in 17, 18 and 19 are relatively sterically demanding and 207 the resulting neutral coordination complexes have square based pyramidal 5-coordinate 208 19-electron Cu^(II) geometries. Each Cu^(II) centre is coordinated by two *N*-donors and three 209 O-donors from the oxidohexaborate(2-) ligands. This is illustrated in Figure 5b for 19. The 210 organic N-donor ligands in 15 and 16 are relatively small and this permits the Cu^(II) centres 211 to adopt 6-coordinate tetragonally distorted octahedral geometries with the formation of 212 additional Cu-O coordinate bonds from bridging oxidoborate ligands, in a similar way to 213 that observed in the Cd^(II) complex, **4**. These 21-electron Cu^(II) centers are coordinated by 214 two N-donors ligands, three O-donors from the fac-hexaborate(2-) ligand with a sixth site 215 from an O-donor of an 'adjacent' hexaborate(2-) by formation of a 1-D coordination poly-216 meric chain. 217

Three new examples of Cu^(II) complexes containing oxidoicosaborate(12-) ligands 218 [H₃O]₄[Cu₇(NH₃)₂(H₂O)₄{B₂₄O₃₉(OH)₁₂}]·13H₂O have been prepared: (13)[19], 219 $H_6[Cu_4O\{B_{20}O_{32}(OH)_8\}] \cdot 25H_2O$ (20) [24] and $H_6[Cu_4O\{B_{20}O_{32}(OH)_8\}] \cdot 34H_2O \cdot 8B(OH)_3$ (21) 220 [24]. These three compounds are fundamentally structurally identical to Cu^(II)/oxidoi-221 cosaborate(6-) complexes HM5[Cu4O{B20O32(OH)8}]·32H2O (M = Na, K) first synthesized by 222 Heller and described in his early (1986) borate structural chemistry review [3]. The struc-223 ture of the anion in 13, 20 and 21 is drawn in Figure 6. It is best described as comprised on 224 four square planar 17-electron Cu^(II) ions and a central μ_4 -O²⁻ ion supporting and sur-225 rounded by large oxidoicosaborate(12-) ring structure. This ring structure itself is 'tetram-226 eric' with four alternating FBB's of {B1} and {B4} sub-units linked into a larger 24-mem-227 bered ring with a compound designation of $20:\{4:(2\Delta+2T)+\Delta\}_4$. Compound **21** has 8 ad-228 ditional B(OH)3 molecules per oxidoicosaborate(12-) moiety and these are situated within 229 channels which are available within the giant structure formed by close-packing the large 230 multi-metallic oxidoborate anions [24]. 231



Figure 6. Drawing of (a) the structure of the (oxidotetracopper)oxidoicosaborate(6-) anion, [Cu₄O{B₂₀O₃₂(OH)₈]⁶⁻, found in 13, 20 and 21, and (b) the 'top layer' of 13 which is comprised of a linear {Cu₃}⁶⁺ chain and two coordinated oxidodiborate(2-), [B₂O₃(OH)₂]²⁻, anions. The four O* atoms also bridge to the four Cu atoms in the lower level of 13.

Compound **13** is unique and contains seven Cu^(II) centres. It can be considered as a large anion based on two 'layers': the 'lower layer' is as is illustrated in Figure 6a and this layer supports a linear {Cu₃}⁶⁺ unit in an 'upper layer'. The {Cu₃}⁶⁺ unit is further supported by two peripheral diborate(2-) anions, and all seven **19-electron** Cu^(II) centres are 5-coordinate square based pyramids. This 'top layer' of **13** is illustrated in Figure 6b. Structurally, compound **13** has another unusual feature: the presence of B-O groups. It is very 248

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rare for oxidiborates prepared and crystallized from aqueous solution to display B-O²⁴⁹ groups arising from trigonal B centres [2]. It is also interesting to note that all such B-O²⁵⁰ groups in the 'upper layer' of **13** are found bridging two (μ 2-) or three (μ 3-) Cu^(II) centres and that the central O²⁻ ion bridges five (μ 5-) Cu^(II) centres [19]. There are also four B-O²⁵² groups in the 'lower layer' of **13**, and in **20** and **21** that each bridge (μ 2-) two Cu^(II) centres. 253

2.4. Gallium(III) and indium(III) borates

Ga(III) and In(III) borates are conveniently considered together. The hybrid oxidoborate, 256 $[Ga(en)_2 \{B_5O_8(OH)_2\}]_n \cdot nH_2O$ (22) was first synthesized in 2012 [25]. The two en N-donor 257 bidentate ligands occupy four coordination sites around an octahedral 22-electron Ga(III) 258 centre and the two remaining cis- coordination sites are connected to two different 259 {B₅O₈(OH)₂}³ anions forming a 1-D chain structure (Figure 7a). Two recently reported 260 compounds $[Ga(teta){B_5O_8(OH)_2}]nnH_2O$ (23) (teta = tetraethylenetriamine) and 261 [In(teta){B5O8(OH)2}]n 1.5nH2O (24) have similar structures [11]. Compound 22 was also 262 reported in 2013 together with three other related compounds: [In(en)2{B5O8(OH)2}]·H2O 263 (25), [In(dap)2{B5O8(OH)2}]·H2O (26), and [In(dien){B5O8(OH)2}]n (27) [26]. The structures 264 of 25 and 26 are essentially the same as 22 with a change of metal (to In(III) in 25) or ligand 265 and metal (to In^{III} and dap in **26**). Compounds **22-27** contain $\{B_5O_8(OH)_2\}^{3-}$ anions and these 266 anions are based on the frequently observed $5:(4\Delta+T)$ pentaborate(1-) anion, $[B_5O_6(OH)_4]^-$ 267 , but is additionally double deprotonated. The related compound Rb2n[Ga{B5O10}]n-4nH2O 268 (28) has also been reported [25]. Here, the $5:(4\Delta + T)$ building block is deprotonated 4 269 times to form the $[B_5O_{10}]^{5-}$ anion with four of these anions coordinated to a tetrahedral 18-270 electron Ga^(III) centre in chain-like 2-D structures. 271

Compound **27** is unique and features octahedral $In^{(III)}$ centres coordinated by a tridentate dien ligand (*fac*-) and three monodentate {B₅O₈(OH)₂}³⁻ ligands which bridge to other $In^{(III)}$ centres in such a way as to produce a 'dimeric' 1-D chain with oxidoborate *O*bridges between two $In^{(III)}$ centres (Figure 7b). The H-bonding interactions between adjacent chains leads to a supramolecular H-bonded network. 276



Figure 7. The building unit of (**a**) [Ga(en)₂{B₅O₈(OH)₂}]_n·nH₂O (**22**), and (**b**) the dimeric In^(III) centres in [In(dien){B₅O₈(OH)₂}]_n (**27**).

2.5. Manganese(II) borates

One 15-electron five-coordinate trigonal-bipyramidal Mn^(II) complex with a coordinated oxidoborate ligand has been reported, K₇[(BO₃)Mn{B₁₂O₁₈(OH)₆)}]·H₂O (**29**) [27]. The structure of this will be discussed in section 2.7 since **29** is isostructural with an analogous **20-electron** Zn^(II) complex and forms part of a family of structurally related of Zn^(II) compounds.

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The recently synthesized salt $[Ni(H_2O)_6]_2[NO_3]_2[Ni(H_2O)_4]_3O_3(OH)_4]_2[2H_2O$ (30) 295 [14] is isostructural with 7. The two [B₃O₃(OH)₄]- ligands in the neutral 296 [Ni(H₂O)₄{B₃O₃(OH)₄}₂] complex are *trans* on octahedral 20-electron Ni^(II) centres and are 297 coordinated by hydroxy O-donors bound to tetrahedral boron atoms of the triborate(1-) 298 anion. 299

Two Ni^(II) hexaborate(6-) complexes have recently been synthesized during the re-300 period: $[Ni(en)(H_2O)_2\{B_6O_7(OH)_6\}]$ ·H_2O view (31)[28] and [Ni(dmen)(H₂O){B₆O₇(OH)₆}]5H₂O (32) [28]. These compounds are neutral molecules and contain the [B₆O₇(OH)₆]²⁻ ligand as described earlier for 4, 10, 11 and 15-19 (sections 2.1, 303 2.2 and 2.3). These ligands are also found in Zn^(II) chemistry (section 2.7). Both 31 and 32 304 are octahedral about the 20-electron Ni^(II) centres but differ in the denticity of the hexabo-305 rate(2-) ligands which are bidentate in **31** (Figure 8a) and tridentate in **32**. Both compounds 306 demonstrate numerous intermolecular and intramolecular solid-state structure directing H-bond interactions. 308



Figure 8. (a) The uncharged complex of $[Ni(en)(H_2O)_2[B_6O_7(OH)_6]]$ -H₂O (31) showing the bidentate nature of the $[B_6O_7(OH)_6]^2$ ligand, (b) the building unit of $[Zn{B_3O_4(OH)_3}]n$ (35), and (c) the building unit of $[Zn(pn){B_4O_6(OH)_2}]n$ (36).

> The derivatized hexaborate(2-) ligand, with three 2-amino ethoxy groups, in place of the three -OH groups on the tetrahedral boron centres, functions as a hexadentate ligand in the 20-electron Ni^(II) complex, [Ni(NH2CH2CH2O)3{B6O7(OH)3}] (33) [18]. Compound 33 is isostructural with 12.

2.7. Zinc(II) borates

There have been more oxidoborate complexes reported for Zn^(II) than for any other metal. The coordinated oxidoborate ligands range in size from triborate(1-) to dodecaborate(6-).

The recently synthesized salt [Zn(H2O)6]2[NO3]2[Zn(H2O)4{B3O3(OH)4}2]2H2O (34) 328 [14] is isostructural with 7 and 30. The two [B₃O₃(OH)₄] ligands in the neutral 329 [Zn(H₂O)₄{B₃O₃(OH)₄}] complex are *trans* on octahedral 22-electron Zn^(II) centres and are 330 coordinated by hydroxy O-donors bound to tetrahedral boron atoms. 331

Structural characterization of the commercially important Zn^(II) triborate, 332 $[Zn{B_3O_4(OH)_3}]n$ (35), revealed that it was a cross-linked 1-D coordination chain polymer 333 of Zn^(II) [29]. The oxidoborate forms a 1-D polymeric chain (Figure 8b) and each 18-electron 334 Zn^(II) centre is coordinated by two oxygen atoms from two adjacent monomeric unit of the 335 chain, and completes its tetrahedral arrangement by coordination from two hydroxide O-336

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donors from a neighboring chain to crosslink the structure into a 2-D network. The network also has numerous interchain H-bond interactions. Each triborate(2-) FBB unit is a 6-membered boroxole ring with one $\{BO_3\}$ and two $\{BO_4\}$ units, *i.e.* $3:(\varDelta+2T)$. 339

Tetraborate ligands based on the 4(4-1) FBB are represented by the following com-340 pounds $[Zn(pn){B_4O_6(OH)_2}]_n$ (36) [30], $[Zn(dap)_{0.5}(dap')_{0.5}{B_4O_6(OH)_2}]_n \cdot nH_2O$ (37) [30], 341 and [Zn(dab)0.5(dab')0.5{B4O6(OH)2}]n·nH2O (38) [31]. Compounds 36-38 have identical 342 Zn^(II)/ligand/borate stoichiometries with similar, but non-identical, structures. All three 343 contain tetrahedral 18-electron Zn^(II) centres with two N-donor amine and two O-donor 344 tetraborate ligands, with the latter condensed into 1-D tetraborate chains. Compound 36 345 contains a chelating pn ligand (Figure 8c) whereas 37 (and 38) has two dap (or dab) ligands 346 on each Zn^(II) centre both bridging other Zn^(II) centres and forming a 1-D coordination pol-347 ymer chains in cross-linked inorganic-organic 2-D layered structures. 348

The FBB of 5:(4 Δ +T) is present in [Zn(tren){B₅O₇(OH)₃}] (**39**) [32] and this ligand is 349 identical to that found in **9** (section 2.2). [Zn(dab)_{0.5}(dab')_{0.5}{B₅O₇(OH)₃}]_n (**40**) [8] and 350 [Zn(appip){B₄O₆(OH)(OB{OH}₂})]_n·3nH₂O (**41**) (appip = *trans*-1,4-bis(3-aminopropyl)pi-351 group on a tetrahedral boron centre of the tetraborate moiety with a 5:[4:(2Δ +2T)+ Δ] 353 framework and are isostructural with **1** (section 2.1).

The 6:(3/2+37) FBB, described in sections 2.1, 2.2, 2.3 and 2.6, is also observed as a 355 hexaborate(2-) ligand in Zn^(II) complexes. The following complexes have been prepared 356 $[Zn(dien)\{B_6O_7(OH)_6\}] \cdot 0.5H_2O$ (42) [33], $(NH_4)_2[Zn(H_2O)_2\{B_6O_7(OH)_6\}_2] \cdot 2H_2O$ (43) [33], 357 [Zn(en){B₆O₇(OH)₆]_n·2nH₂O (44) [34] and [Zn(pn){B₆O₇(OH)₆]_n·1.5nH₂O (45) [34]. The 22-358 electron Zn^(II) centres in 42-45 are all octahedral with N_3O_3 , O_6 , N_2O_4 and N_2O_4 donor sets, 359 respectively. The hexaborate(2-) ligands in 42, 44 and 45 are tridendate but are bis(biden-360 tate) and *trans* in the centrosymmetric anion in **43** (Figure 9a). Compounds **44** and **45** are 361 1-D coordination polymers with a -OH group of a trigonal boron of the coordinated hex-362 aborate(2-) anion bridging onto another Zn^(II) centre. This configuration has also been ob-363 served in Cd^(II) (4, section 2.1) and Cu^(II) (15 and 16, section 2.3) chemistry. 364



Figure 9. Drawings of (a) the insular anion in $(NH_4)_2[Zn(H_2O)_2\{B_6O_7(OH)_6\}_2]_2H_2O$ (43), and (b) the repeating unit of $[\{Zn(en)_2\{B_7O_{10}(OH)_3\}_2]_n$ (46).

 $\{\{Zn(en)_2B_7O_{10}(OH)_3\}_2\}_n (46), \text{ also formulated as } [Zn(en)_2\{B_7O_{12}(OH)\}\} \text{ in Ref. [9], is a } 374 \\ 3-D \text{ coordination polymer comprised of an octahedrally coordinated } 22-electron Zn^{(II)} \text{ cen-} \\ 375 \\ \text{tre based on square planar } [Zn(en)_2] \text{ units axially coordinated by a } O-\text{donor condensed } 376 \\ \text{oxidoheptaborate framework. The oxidoborate FBB in 46, } [B_7O_{10}(OH)_3]_n^{2n-}, \text{ can be de-} \\ \text{scribed by a compounded descriptor } 7:[(3:2\varDelta+T)+(3:2\varDelta+T)+\varDelta] \text{ with triangular BO}_2(OH) \\ 378 \\ \text{cross-linking units (Figure 9b).} \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 379 \\ 370 \\ 370 \\ 379 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\ 370 \\$

[Zn(en)₂{B₈O₁₁(OH)₄}]_n (47) [10] was reported in 2017 and has a similar stoichiometry 380 to **3** but is isomeric, with a different oxidoborate condensation mode and a 1-D polymer 381 chain. (Figure 10a). Nevertheless, the octaborate(2-) anion is best designated by the com-382 pounded descriptor 8:[5: $(4\Delta + T)$ +3: $(2\Delta + T)$].



Figure 10. The repeating unit of $[Zn(en)_2\{B_8O_{11}(OH)_4]]_n$ (47), and (b) a drawing of the $[(Hdap)Zn\{B_{12}O_{18}(OH)_6\}]^3$ - anion in (H2dap)3[(Hdap)Zn{B12O18(OH)6}]2.14H2O (51).

 $[Zn_2(dap)(dap'){B_8O_{13}(OH)_2}]n$ (48) [30] has a similar structure to 37 and 38 with the 393 18-electron Zn^(II) centres tetrahedral with 2 monodentate bridging dap *N*-donor ligands. 394 The octaborate(4-) O-donor ligands are based on two condensed tetraborate(2-) building blocks *via* an O bridge, $8:\{4(2\Delta+2T)\}_2$. However, this now results in 2-D layers rather than 1-D chains and the 2-D layers are further crosslinked into 3-D networks, by the amines.

The large insular anion [B12O18(OH)6]6been reported has in 398 $K_7[(BO_3)Zn\{B_{12}O_{18}(OH)_6\}]$].H₂O (49) [35], [(Hen)Zn\{B_{12}O_{18}(OH)_6\}Zn(en)(Hen)]·8H₂O (50) 399 [36] and (H2dap)3[(Hdap)Zn{B12O18(OH)6}]2-14H2O (51) [33]. The dodecaborate(6-) anion is 400 based on a hexameric FBB of 3 *i.e* 12; $\{3:(\Delta+2T)\}_6$, with all tetrahedral boron centres linking 401 boroxole {B₃O₃} rings, generating a larger inner 12-membered B/O alternating B₆O₆ ring. 402 Stereochemically, three of these potential O-donor atoms point to one side of the large 403 ring and three face towards the other side. The anion in 51, $[(Hdap)Zn\{B_{12}O_{18}(OH)_6\}]^3$ -404 which is typical of the oxidoborate building blocks contained within these structures, has 405 a tetrahedral 18-electron Zn^(II) centre coordinated by fac O-donors from the dodecabo-406 rate(6-) ligand and a monodentate N-donor from a protonated 1,3-diamminopropane lig-407 and (Figure 10b). There is extensive H-bonding between anions forming a supramolecular 408 3-D H-bonded lattice.

3. Synthetic methods

3.1 Slow crystallization by solvent evaporation

This method involves slow crystallization from aqueous or a miscible aqueous/or-415 ganic solution which originally contained B(OH)₃ and metal precursor complexes. The 416 oxidoborate compounds are templated by the transition-metal/p-block metal complexes 417 present in the dynamic combinatorial library (DCL) of oxidoborate anions that are present 418in solution in equilibrium concentrations [37]. Products are generally under thermody-419 namic control especially when metal-ligand exchange equilibria (and hydroxyox-420 idoborate-H2O equilibria) are fast. The solvents used for evaporation in the boric acid 421

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solution are often H2O or H2O/EtOH or H2O/MeOH. Crystallization of the product may 422 take a few hours to several weeks. This is illustrated for the preparation of 423 [Cu(en){B₆O₇(OH)₆]_n3nH₂O (16) from [Cu(en)₂]SO₄ and B(OH)₃ [20]. [Cu(en)₂]SO₄ (3.6 424 mmol) and BaSO48H2O (3.6 mmol) were dissolved in H2O (20 ml). The solution was 425 stirred for 20 mins at room temperature and the precipitate that formed (BaSO4) was re-426 moved by filtration. B(OH)₃ (25 mmol) was added to the filtrate which was then stirred 427 for 40 mins. The resulting solution was left in several small vials to crystallize. After stand-428 ing for 35 days the product 16 was collected by filtration as blue crystals in 31% yield. 429 Compounds 11, 13-19, 31, 32, 35, 42-45, 50 and 51 were prepared by a method similar to 430 this. Oxidopolyborates prepared by this method are often insular salts and are often less 431 condensed than those prepared by the methods described below, which generally use 432 more forcing conditions. 433

3.2 Solvothermal/hydrothermal methods

Many organic-inorganic oxidoborates described in this article are synthesized by sol-436 vothermal (or hydrothermal) methods. In this method the non-aqueous solvent (or H₂O) 437 is placed in a sealed reaction vessel together with boron containing materials, transition-438 metal/p-block metal salts and organic ligands and heated to a specific temperature (usu-439 ally 100-250 °C) for a set time period. Solvothermal reactions can lead to metastable, ki-440netically controlled, products. This method is illustrated by the preparation of 441 [Cd(dap)0.5(dap')0.5{B5O8(OH)}]n·nH2O (2) [9]. H3BO3 (5 mmol) and Cd(NO3)2·4H2O (1 442 mmol) were slowly added to a N,N-dimethylformamide (2 mL) and 1,3-dap (1 mL) solu-443 tion with stirring. The white emulsion was then sealed in a Teflon-lined 25 mL autoclave 444 reactor and heated at 180 °C for 7 days. After cooling to room temperature, crystals of 2 445 $(36\% \text{ yield based on H}_3BO_3)$ were collected by filtration, washed with H₂O, and dried in 446 air. Compounds 1-6, 8, 9, 12, 20-29, 33, 36-41 and 46-49 were prepared by this general 447 method. This relatively low temperature hydrothermal method is popular amongst many 448 researchers since the necessary autoclave reactors are widely available in both academic 449 and industrial laboratories. 450

3.3 Molten salt methods

In this method a molten salt (120-250 °C) is used as both solvent and reactant and the 453 reaction vessel is charged with B(OH)₃ and any other necessary reaction materials. This is 454 illustrated by the preparation of $[Co(H_2O)_6]_2[NO_3]_2 [Co(H_2O)_4 \{B_3O_3(OH)_4\}_2]_2 H_2O$ (7) [14]. 455 Co(NO₃)₂·6H₂O (1 mmol) was placed in a 25 mL beaker and heated in an oil bath at 120 °C 456 until fully molten (ca. 1 min). H₃BO₃ (0.8 mmol) was then added and the mixture was 457 stirred until the H₃BO₃ was completely dissolved (ca. 30 mins). The solution was then al-458 lowed to cool to room temperature and left to crystallize. After 14 days, transparent col-459 orless crystals of 7 (30% based on H_3BO_3) were deposited on the bottom of the beaker. Compound 7 was isolated by filtration. Compounds 30 and 34 were also prepared by this 461 method. 462

4. Physical, spectroscopic properties and potential applications

The oxidoborate complexes described in this manuscript are all crystalline solids 465 with high melting (or decomposition) points. Techniques used to characterize the ox-466 idoborates and to study their possible applications include structural studies (single-crys-467 tal XRD, powder-XRD, TEM), optical properties (diffuse reflectance spectroscopy, nonlin-468 ear optical studies (NLO), photoluminescence), magnetic properties, vibrational spectros-469 copy, thermal studies and catalytic investigations. 470

Oxidoborate complexes derived from transition-metals are generally coloured but 471 the *d*¹⁰ ions (Cd^(II), Zn^(II), Ga^(III) and In^(III)) are colorless. UV/Vis absorption properties of a 472 number of oxidopolyborate complexes (1, 4, 5-8, 22-24, 28, 30, 34, 36, 37, 40, 41) have been 473 studied by diffuse reflectance spectroscopy and band-gaps with energies of ranging from 474

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The band gaps of $[Cd(pn){B_6O_7(OH)_6}]_n:nH_2O$ (4), $[Ga(teta){B_5O_8(OH)_2}]_n:nH_2O$ (23) 477 and $[In(teta){B_5O_8(OH)_2}]n 1.5nH_2O$ (24) show interesting but different temperature ef-478 fects. Compounds 23 and 24 exhibited blue maximum luminescence at 446 and 472 nm, 479 respectively, when excited at 352 and 342 nm, respectively. The luminescence intensity of 480 these two compounds increased with a decreasing temperature and was at its most in-481 tense at 80 K. Compound 4 exhibited the maximum luminescence at 444 nm with a 360 482 nm excitation light source, and the luminescence intensity increased with the decrease of 483 temperature to 230 K where it was at its most intense. The luminescence intensity of 4 484 decreased when the temperature was further lowered to 80 K [11]. Compounds 485 $[M(dab)_{0.5}(dab')_{0.5}[B_5O_7(OH)_3]]_n$ (M = Cd, (1); M = Zn (40)) have been shown to display blue 486 luminescence with maximum fluorescent emission at 423 and 412 nm, when excited at 487 and 356 or 384 nm, with lifetimes of 4.67 and 4.09 ns, for 1 and 40, respectively [8]. It is 488 reported that the blue luminescence of two compounds originates from the their inorganic 489 oxidoborate frameworks. 490

Nonlinear optical (NLO) effects are also displayed by the following hybrid metal ox-491 idoborates $Rb_{2n}[Ga\{B_5O_{10}\}]n^4nH_2O$ (28) [25], $K_7[(BO_3)Mn\{B_{12}O_{18}(OH)_6)\}]H_2O$ (29) [27], 492 [Zn(pn){B4O6(OH)2}]n (36) [30] and [Zn(appip){B4O6(OH)(OB{OH}2)]n 3nH2O (41) [30]. For 493 example, compound **41** belongs to the space group *P2*₁, and since this point group is non-494 centrosymmetric 41 might be expected to exhibit second order nonlinear optical effects; 495 indeed, powdered **41** does exhibit SHG behavior with a response of 29 mV using a laser 496 of beam energy with 2.83 mJ/pulse [30]. Compound 36 had a response of 24 mV whilst the 497 response of the SHG standard, KH2PO4 (KDP), was 175 mV under similar conditions [30]. 498 Compound 28 crystallizes in chiral space group C2221. SHG measurements on a Q-499 switched Nd:YAG laser with sieved powdered samples (70-100 mesh) revealed that com-500 pound 28 displays a moderately strong SHG response approximately equal to that KDP 501 [25]. The SHG response of 29 was again moderate and equal to that of KDP [27]. 502

Many of the compounds described within this review are diamagnetic but those con-503 taining Cu^(II) (13 - 21) and Ni^(II) (31 - 32) centres are paramagnetic. The magnetic properties 504 of the Ni^(II) complex 30 have not been reported. Magnetic properties of the Co^(II) (7 - 12) 505 and the Mn^(II) (29) oxidoborate complexes were also not reported, although these would 506 be expected to be paramagnetic. The µeff values (per Cu^(II) atom) of the multi-metallic com-507 plexes [H₃O]₄[Cu₇(NH₃)₂(H₂O)₄{B₂4O₃₉(OH)₁₂]·13H₂O (13), H₆[Cu₄O{B₂₀O₃₂(OH)₈]·25H₂O 508 (20) and $H_6[Cu_4O[B_{20}O_{32}(OH)_8]]_34H_2O_8B(OH)_3$ (21) are much lower than expected by the 509 spin-only formula and 20 and 21 are temperature dependent and display anti-ferromag-510 netic behavior [24]. 511

A complex borate, [Ni(en)₃]n[Hen]n[B₉O₁₃(OH)₄]n·nH₂O, containing isolated cations 512 and a partially condensed anionic a one-dimensional oxidoborate chain has ferroelectric 513 properties at room temperature [39]. The electrical hysteresis loop was observed when an 514 electric field between -28 and +28 kV was applied to the sample. Spontaneous polarization 515 (P_s of about 52 nCcm⁻²) also occurred during the measurement with remnant polarization 516 (P_r of about 30 nCcm⁻²) and a coercive field (E_c) of about 28 kVcm⁻¹. The P_s of this com-517 pound is very close to that of KH2PO4 type ferroelectrics. The related compound, 518 [Ni(en)2(pip)][B5O6(OH)4]2, which contains insular oxidoborate anions rather than coordi-519 nated oxidoborate ligands, was reported as the first templated oxidoborate with ferroe-520 lectric properties [38]. The P_s value for [Ni(en)₂(pip)][B₅O₆(OH)₄]₂ of 20 nCcm⁻² for is sig-521 nificantly higher than that of typical organic ferroelectric compounds, *e.g.* β-quinol-meth-522 anol ($P_s = 6 \text{ nCcm}^{-2}$). A related pentaborate derivative, $[Zn(dab)_{0.5}(B_5O_7(OH)_3)]_n$ 523 (40), has been prepared more recently, and exhibits a wide Ec value of ca. 6 kV cm⁻¹ to 11 524 kV cm⁻¹ [8]. 525

Vibrational (IR) data are commonly reported for oxidoborate complexes. In addition 526 to diagnostic bands associated with organic ligands (O-H, N-H, C-C, C-N, C-C *etc.* 527 stretches and bends) strong absorptions associated with ligand-M and B-O stretches (1400-528

1000 cm⁻¹) are often observed but usually are not diagnostic since they are in the IR fin-529 gerprint region. In general B-O stretching modes can be subdivided into four specific re-530 gions (asymmetric stretches: Btrig-O, 1450-1300 cm⁻¹; Btet-O, 1150-1000 cm⁻¹; symmetric 531 stretches: Btrig-O, 960-890 cm⁻¹, Btet-O, 890-740 cm⁻¹) with bending and deformation modes 532 at lower energy [40]. Nevertheless, Li and co-workers have tabulated diagnostic wave-533 numbers for specific smaller oxidoborate anions [40]. Hexaborate(2-) derivatives, with a 534 FBB of $6:(3\Delta+3T)$, are fairly common as oxidoborate ligands (e.g. compounds 4, 5, 10-12, 535 15-20, 42-45) and a diagnostic bands at 955 and 808 cm⁻¹ have been proposed for this anion 536 [20, 40]. Tentative diagnostic bands (1047, 952, 902 and 857 cm⁻¹) have also been reported 537 for icosaborate(6-) derivatives (49-51) [33]. 538

The most studied property of inorganic-organic hybrid borates is their thermal be-539 havior. Indeed, all compounds described within this review, except 10, 12, 33 (which are 540reported in crystallographic journals [16, 18]), have their thermal properties documented. 541 Many of the thermal studies have been undertaken in air from room temperature up to 542 800 °C, although a few (4, 23, 24) report data obtained through heating under N2 to a sim-543 ilar temperature. Generally, thermal decomposition is a multistage process which in-544 cludes a low temperature loss of interstitial molecules (where present), a moderate tem-545 perature dehydration and cross-linking of oxidoborate hydroxyl groups, and a high tem-546 perature oxidation process (or removal of volatile organics) to generate a glassy borate 547 residual solid. This solid can be formulated as an anhydrous metal borate (*i.e.* a mixed 548 metal/boron oxide) with a M:B ratio that maintains the M:B ratio of the original ox-549 idoborate complex. These principles are illustrated compounds for 550 $[Cd(pn){B_{0}(OH)_{12}]nnH_{2}O$ (4), $[H_{3}O]_{4}[Cu_{7}(NH_{3})_{2}(H_{2}O)_{4}[B_{24}O_{39}(OH)_{12}]^{-1}3H_{2}O$ (13) and 551 [Zn(en){B₆O₇(OH)₆]_n·2nH₂O (44). Compound 44 decomposes thermally (in air) by the loss 552 of 2 interstitial H2O (100-180 °C), condensation of hydroxyoxidoborate with loss of 3H2O 553 (180-320 °C) and oxidation of the en ligand (320-470 °C) to leave as a residue ZnB₆O₁₀ 554 (=ZnO3B2O3) [34]. Compound13 thermally decomposes in air by the loss of 21 (interstitial 555 and coordinated) H2O and 4 (coordinated) NH3 molecules (at 70-160 °C) and condensation 556 of hydroxyoxidoborate groups with loss of 8H2O (at 160-300 °C) to afford Cu7B24O43 (= 557 7CuO12B2O3) [19]. Occasionally, the individual processes overlap but the endpoint is the 558 same. Thus, 4 losses weight continuously in one step upon heating, under N₂, from room 559 temperature to 800 °C [11]. In this process the guest water molecules are removed together 560 with pn ligands and the oxidopolyborate fully condenses to afford CdB₆O₁₀ (= CdO3B₂O₃) 561 [11]. Non-metal cation polyborates have been considered as thermal precursors to porous 562 materials [5]. To date this has not been successful [41 - 43] but recent studies on 563 H6[Cu4O{B20O32(OH)8}]25H2O (20) and H6[Cu4O{B20O32(OH)8}]34H2O8B(OH)3 (21), which 564 contain large 3-D intersecting channel systems with PLATON calculated solvent accessi-565 ble voids of ca. 60%, suggest that mesoporous materials may be available from hybrid 566 organic oxidoborates [24]. 567

Fire retardancy is often linked to a compounds thermal properties and $[Zn{B_3O_4(OH)_3}]_n$ (**35**) [31] is well known for this property and has been commercially exploited. Many zinc borates also possess such properties [44] and the recently prepared hybrid oxidoborate complex $[Zn(H_2O)_6]_2[NO_3]_2$ · $[Zn(H_2O)_4{B_3O_3(OH)_4}_2]$ · $2H_2O$ (**34**) [14] has also been reported as a promising flame retardant, at 15 wt% loading, for ABS (acrylonitrile butadiene styrene), an important engineering thermoplastic material.

Hybrid metal-organic oxidoborates have been used as precursors to thermally prepared catalysts [9] *e.g.* carbon material catalysts derived from thermal treatment of [Cd(dap) $_{0.5}$ (dap') $_{0.5}$ {B₅O₈(OH)}]_m·nH₂O (**2**) and [{Zn(en)}{B₇O₁₀(OH)}]₂]_n(**46**) have been used 576 to electrochemically reduce CO₂ to CO. Initial results indicate that the oxidoborate is a useful catalyst precursor since at 1.4V CO formation is 51% higher than that of RHE. This 578 may be a promising development as a low cost alternative to traditional noble metal catalysts. 580

5. Conclusions.

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Since metals generally have multiple coordination sites they often have the ability to 583 form coordination linkages with organic (N-donor) and oxidoborate (O-donor) ligands. 584These organic/inorganic oxidopolyborate hybrid complexes are structurally diverse and 585 often have polymeric structures based on unique frameworks. The hybrid organic ox-586 idoborate materials in such integrated structures can afford materials with unique prop-587 erties and which may overcome possible deficiencies inherently present in related single 588 component systems. Hybrid inorganic-organic oxidoborates have attracted considerable 589 recent research attention, and interesting properties such as photoluminescence, ferroe-590 lectric and NLO and catalytic properties have been discovered and these are summarized 591 within the review. Initial results indicate that these properties can be tailored and ma-592 nipulated by adjusting the organic ligands, the oxidoborate motif and/or the and central 593 metallic ion, and that such compounds are a promising area for future research. 594

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Abbreviations: appip = *trans*-1,4-bis(3-aminopropyl)piperazine); dab = 1,4-diaminobutane; dap =6041,3-diaminopropane; deen = N,N-diethyl-1,2,-diaminoethane); dmen = N,N-dimethyl-1,2-dia-605minoethane; en = 1,2-diaminoethane; pip = piperidine; pn = 1,2-diaminopropane; py = pyridine; teta606= tetraethylenetriamine; tren = tris(2-aminoethyl)amine); tmeda = N,N,N',N'-tetramethyl-1,2-dia-607minoethane.608

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 $NH(CH_2CH_2NH_2)_2), (NH_4)_2[Zn\{\kappa^2-O-B_6O_7(OH)_6\} (H_2O)_2]\cdot 2H_2O \text{ and } (1,3-pn)_3[(\{\kappa^1-NH_3\{CH_2\}_3NH_2)Zn\{\kappa^3-O-B_{12}O_{18}(OH)_6\}]_2\cdot 14H_2O.$ 686 Inorganics, 2019, 7, 44. DOI: 10.3390/inorganics7040044. 687

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