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Article Oxidoborates templated by cationic nickel(II) complexes and self-assembled from B(OH)₃

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Abstract: Several oxidoborates, self-assembled from B(OH)3 and templated by cationic Ni^(II) coordi-11 nation compounds, have been synthesized by crystallization from aqueous solution. These include 12 the ionic compounds trans-[Ni(NH₃)₄(H₂O)₂][B₄O₅(OH)₄]·H₂O (1), s-[Ni(dien)₂][B₅O₆(OH)₄]₂ (dien = 13 N-(2-aminoethyl)-1,2-ethanediamine (2), trans-[Ni(dmen)₂(H₂O)₂] [B₅O₆(OH)₄]₂2H₂O (dmen = N,N-14 dimethyl-1,2-diaminoethane) (3), $[Ni(HEen)_2][B_5O_6(OH)_4]_2$ (HEen = N-(2-hydroxyethyl)-1,2-dia-15 minoethane) (4), [Ni(AEN)][B5O6(OH)4]H2O (AEN = 1-(3-azapropyl) -2,4-dimethyl-1,5,8-triazaocta-16 2,4-dienato(1-)) (5), trans-[Ni(dach)2(H2O)2][Ni(dach)2] [B7O9(OH)5]24H2O (dach = 1,2-diaminocyclo-17 hexane) (6), and the neutral species $trans-[Ni(en)(H_2O)_2[B_6O_7(OH)_6]]\cdot H_2O$ (7) (en = 1,2-dia-18 minoethane), and [Ni(dmen)(H2O){B₆O₇(OH)₆}]5H2O (8). Compounds 1 - 8 have been character-19 ized by single-crystal XRD studies and by IR spectroscopy and 2, 4 - 7 were also characterized by 20 thermal (TGA/DSC) methods and powder XDR studies. The solid-state structures of all compounds 21 show extensive stabilizing H-bond interactions, important for their formation, and also display a 22 range of gross structural features: 1 has an insular tetraborate(2-) anion, 2 - 5 have insular pentabo-23 rate(1-) anions, 6 has an insular heptaborate(2-) anion ('O+' isomer), whilst 7 and 8 have hexabo-24 rate(2-) anions directly coordinated to their Ni(II) centers, as bidentate or tridentate ligands, respec-25 tively. The Ni^(II) centers are either octahedral (1 - 4, 7, 8) or square-planar (5), and compound 6 has 26 both octahedral and square-planar metal geometries present within the structure as a double salt. 27 Magnetic susceptibility measurements have been undertaken on all compounds. 28

Keywords:Borate anions; H-bonding; Heptaborate(2-); Hexaborate(2-); Nickel(II) complex; Ox-29idoborate; Pentaborate(1-); Self-assembly; Templated synthesis, Tetraborate(2-); X-ray structures.30

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1. Introduction

Synthetic borates (oxidoborates is the IUPAC recommended name for this class of 34 compounds [1]) show great structural diversity and their structures complement the variety of structural motifs observed in borate minerals [2-5]. Borate minerals have large 36 scale economic importance [6-8] (*e.g.* as vitreous, agriculture, fire retardant products) and 37 some of these minerals, and their synthetic counterparts, have also found more specialist 38 uses [9-18] (*e.g.* as fluorescent, optical SHG and wide band-gap semi-conductor materials). 39

Structurally borates salts consist of metallic or non-metallic cationic centers and hydroxyoxidoborate units, with variable degrees of condensation, as either insular anions or as anionic 1-D chains, 2-D layers or 3-D networks [2-5]. Oxidoborate materials can be synthesized from aqueous solution or from solid-state or solvothermal methods and the latter non-aqueous methods often lead to the formation of the more highly condensed 44

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Copyright: © 2021 by the authors. Submitted for possible open access publication under the terms and conditions of the Creative Commons Attribution (CC BY) license (https://creativecommons.org/license s/by/4.0/). oxidoborate structures [5]. In aqueous solution oxidoborate speciation is pH and boron-45concentration dependent [19,20], and a dynamic combinatorial library (DCL) [21] of ox-46idoborate anions co-exist in rapidly attained aqueous equilibria.47

Many oxidoborate anions have been successfully crystallized from aqueous solution 48 using $B(OH)_3$ and an appropriate templating cation *e.g.* triborate(1-), tetraborate(2-), pen-49 taborate(1-), hexaborate(2-), heptaborate(2-) (two isomers), heptaborate(3-), octabo-50 rate(2-) (two isomers), nonaborate(3-), tetradecaborate(4-), and pentadecaborate(3-) ani-51 ons [5]. These salts are selectively templated from the solution by the cations present in 52 self-assembly processes [22-24]. However, pentaborate(1-) salts often crystallize out of 53 such solutions and this is particularly so if the cation is relatively small and has a charge 54 of +1 [5]. This process primarily occurs since the $[B_5O_6(OH)_4]^-$ anion is ideally shaped to 55 form giant H-bonded lattices through anion-anion interactions with 'cavities' large 56 enough to be occupied by the small cations [25-28]. 57

We are interested in expanding the structural diversity of isolated and coordinated 58 oxidoborate chemistry and have developed a strategy of incorporating relatively highly 59 charged (>+1) and/or cations, with the potential of forming multiple H-bond interactions, 60 into the aqueous DCL of borate anions so that they can template and crystal engineer 61 novel structures [29, 30]. Using labile cationic transition-metal complexes [31] introduces 62 a further DCL of potential cations into reaction medium and leads to the possibility of 63 oxidoborate anions entering the primary coordination shell of the metal, as O-donor lig-64 ands. 65

With this in the forefront of our mind, we have recently investigated the reactions 66 of labile complexes of Zn^(II) [32-35] and Cu^(II) [36-40] and have prepared novel species con-67 taining coordinated [B2O3(OH)2]²⁻ [39], [B5O6(OH)4]⁻ [37], $[B_6O_7(OH)_6]^2$ [34-38], 68 [B₁₂O₁₈(OH)₆]⁶⁻ [32, 35] and [B₂₀O₃₂(OH)₈]¹²⁻ [39] ligands. Ni^(II) complexes and their reac-69 tions with B(OH)₃ have received little attention to date and reported oxidoborate species 70 are restricted to an unusual hexadentate tris(aminoethoxy)hexaborate(2-) anion complex, 71 $[Ni{(H_2NCH_2CH_2O)_3B_6O_7(OH)_3}]$ [41], and a series of salts containing the insular pentabo-72 rate(1-) [42-47] or heptaborate(2-) [33, 48] anions. 73

Ni^(II) complexes have a d^8 electronic configuration and should also be relatively labile74[31]. In this manuscript we report on the use coordination complexes of Ni^(II) containing75N-donor ligands NH₃, en, dien, dmen, HEen, AEN, dach (See Figure 1 for ligand structures and abbreviations) to synthesize eight new oxidoborate compounds. Their single-76crystal structures and their spectroscopic and physical properties are reported.78



Figure 1. Drawings of the organic ligand structures found in compounds **1** - **8** with abbreviations as follows: (a) en, 1,2diaminoethane), (b) dien, *N*-(2-aminoethyl)-1,2-ethanediamine, (c) HEen, *N*-(2-hydroxyethyl)-1,2-diaminoethane, (d) dmen, *N*,*N*dimethyl-1,2-diaminoethane, (e) dach, 1,2-diaminocyclohexane, (f) AEN, 1-(3-azapropyl)-2,4-dimethyl-1,5,8-triazaocta-2,4dienato(1-).

2. Results and Discussion

2.1. Synthesis and general discussion

The Ni^(II) complex oxidoborates **1** - **8** were prepared in acceptable yields as a crystalline solids from the reaction of a Ni^(II) complex cation hydroxide salt with boric acid in a ratio of either 1:5 or 1:10 (Scheme 1). The Ni^(II) complex cation hydroxides were obtained *in situ* from the corresponding salts containing either chloride or sulfate anions. Anion exchange (from Cl⁻) was used for **2**, **4**, **5**, **6**, and **8** and stoichiometric reactions with Ag₂O (from Cl⁻) was used for **1** and **3** and a stoichiometric reaction with Ba(OH)₂·8H₂O (from SO₄²⁻) was used for **7**.

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 Scheme 1. Reagents and stoichiometry of reactants involved in self-assembly of Ni^(II) oxidoborates from cationic Ni^(II) complexes
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 with B(OH)₃ in aqueous or methanolic aqueous solution.
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The new compounds were identified by XRD single-crystal studies (Section 2.2), IR 102 spectroscopic analysis and accompanied by thermal decomposition analysis (TGA and 103 DSC) and powder XRD studies for 2, 4 - 7 (Section 2.3). Six of these eight compounds 104 contain insular oxidoborate anions and two compounds contain coordinated hexabo-105 rate(2-) anions; the oxidoborate anions found in these new compounds are drawn out in 106 Figure 2. The coordination geometry around the Ni^(II) centers are either octahedral $(1 - 4_r)$ 107 7, 8) or square-planar (5) and compound 6 has both octahedral and square-planar metal 108 geometries present within the structure as a double salt. The new Ni^(II) oxidoborate com-109 plexes are coloured with those with square-planar centres (5 and 6) are orange/red with 110 the others having blueish hues. All compounds were stable in the solid state were insolu-111 ble in organic solvents and decomposed by aqueous solution (see ¹¹B NMR, section 2.3). 112 Since the products arise through self-assembly processes that we believe are associated 113 with H-bonded structure directing effects induced by the cations present [21-24] these in-114 teractions are discussed in detail in Section 2.2.2. 115

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Figure 2. Drawings of the oxidoborate structures found in (a) 1, (b) 2 - 5, (c) 7, 8 and (d) 6.

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2.2 Crystallographic studies on 1 - 8 and their solid-state H-bonding interactions.

2.2.1 General considerations for 1 - 6 and the structures of 7 and 8.

Selected crystallographic data for 1 - 8 can be found in Experimental Methods (Sec-123 tion 3) and full crystallographic details are available as supplementary information. Com-124 pounds 1 - 6 are all anionic compounds containing discrete insular cationic Ni^(II) coordi-125 nation complexes with N-donor or O-donor ligands, partnered with anionic oxidopoly-126 borate anions. The cations in 2 - 6 are as found in their precursor complexes whilst 1 con-127 tains the partially aquated trans-[Ni(NH₃)₄(H₂O)₂]²⁺ cation. The cationic charges are neu-128 tralized by the insular oxiodopolyborate anions $[B_4O_5(OH)_4]^2$, $[B_5O_6(OH)_4]^2$, or 129 [B₇O₉(OH)₅]²⁻ for 1, 2 - 5 and 6, respectively. Compounds 1, 3, 5 and 6 also contain one, 130 two, one and four interstitial H₂O molecules per formula unit, respectively. The dien lig-131 and in 2, and the HEen ligand in 4 are both disordered over two positions. Compounds 7 132 and 8 are uncharged coordination compounds with $[B_6O_7(OH)_6]^{2-}$ coordinated as O-donor 133 ligands to octahedral Ni^(II) centres either as a bidentate or a tridentate ligand in 7 or 8, 134 respectively. Compounds 7 and 8 have one and five interstitial H₂O molecules, respec-135 tively, and are also partially aquated. There is some disorder of the interstitial H₂O mole-136 cules in both 7 and 8 and some additional disorder in the en ligand and the dmen ligands 137 of 7 and 8. Compound 6 is unusual and is best formulated as a double salt. It contains both 138 octahedral trans-[Ni(dach)2(H2O)2]2+ and square-planar [Ni(dach)2]2+ cations partnered by 139 two crystallographically equivalent [B7O9(OH)5]²⁻ anions and four interstitial H2O mole-140 cules. The dach ligands are also disordered in 6 but have chair conformations with their 141 amino groups equatorial and with one dach ligand in each cation having R,R stereochem-142 istry and the other S,S. Interestingly, two of these interstitial H₂O molecules (O22) in 6 are 143 in 'axial' positions in the square-planar complex at 3.1517(13) Å (T = 0.61 [49]). In the 144 octahedral complex these two H₂O ligands (O21) are at 2.1422(15) Å (T = 0.98). A structur-145 ally related (but d9) Cu(II) compound, prepared by an identical method [37], contains a 146 square-planar (also with very long axial H₂O 'bonds', T = 0.70) and a Jahn-Teller distorted 147 axially elongated octahedral complexes (T = 0.80). Structurally, the insular [B₇O₉(OH)₅]²⁻ 148dianion is previously known to adopt either a 'chain' or 'O+' isomeric forms [28, 50-52] 149 and the 'O^{+'} isomer is present in 6. The new Ni^(II) oxidoborates complex salts 1 - 6 all con-150 tain cations and anions that are known in other salts and the gross structures of the com-151 ponent ions, bond distances and internuclear angles for the component ions in 1 - 6 are as 152 expected and need no further comment [5, 53-55]. The uncharged complexes 7 and 8 are 153 structurally novel although similar compounds have been observed in Co(II), Cu^(II) and 154 Zn^(II) oxidoborate chemistry [34-38, 56-58]. Compounds 7 and 8 contain isolated species 155 [35, 37, 38] rather than polymer 1-D coordination chains [34, 36, 38] and these two struc-156 tures will be discussed in more detail below. 157

A drawing of the structure of *trans*-[Ni(en)(H₂O)₂{B₆O₇(OH)₆}]-H₂O (7), with selected 158 crystallographic numbering, can be found in Figure 3. Compound 7 is a distorted neutral 159 octahedral Ni^(II) complex with *trans* H₂O ligands (containing O21 and O22), a disordered 160 1,2-diaminoethane ligand and a bidentate $[B_6O_7(OH)_6]^2$ -ligand that is coordinated through 161 two hydroxy groups (containing O8 and O9). A single disordered water of crystallization 162 containing O31 (s.o.f. 0.47(2)) or O31B (s.o.f. 0.53(2)) is also shown. The O21-Ni1-O22 angle 163 is 171.67(12)° and the average Ni-(OH2) distance is 2.089(3) Å. This average distance is 164 slightly longer than the average oxidoborate O8/O9-Ni1 distance of 2.065(3) Å. The che-165 lating en ligand significantly distorts the Ni^(II) centre where the Ni1-N1 and Ni1-N2 dis-166 tances are 2.086(4) and 2.085(4) Å, respectively. The N1-Ni1-N2 angle is 82.80(16)° and the 167 oxidoborate/en ligand O8-Ni1-N2 and O9-Ni1-N1 angles are both significantly <180°, 168 whilst the O8-Ni1-O9 angle is 90.48(14)°. These Ni-N and Ni-(OH2) bond lengths are 169 within the expected ranges for Ni^(II) complexes [53-55] but this is the first example of a Ni-170 O bondlength arising from an oxidoborate ligand and it is not significantly different 171 from other N-O bondlengths. The B-O bondlengths and OBO and BOB angles of the hex-172 aborate(2-) ligand are in accord with previously reported data for this ligand including 173 three very long B-O bondlengths (1.498(6)-1.538(6)°; av. 1.517(6)°) to the three-coordinate 174 O atom (O1) [28, 29, 48, 51, 52]. There is an intramolecular H-bond from an oxidoborate 175 hydroxy group (containing O10) and the coordinated aqua ligand (containing O21), 176 O10H10-O21, with a O10-O21 distance of 2.704(5) Å and a O10-H10-O21 angle of 144.9° 177 (Figure 3). This H-bond is part of two R_1 ⁽¹⁾(8) rings [59] and both rings involve the Ni^(II) 178 centre and two Ni-O coordinate bonds. There is a similar H-bond interaction in the only 179 other known example bidentate hexaborate(2-) of а complex, 180 $(NH_4)_2[Zn(H_2O)_2(B_6O_7(OH)_6)_2]$, but here, interestingly, the H-bond originates from the 181 aqua ligand and the uncoordinated hydroxy site of the oxidoborate is the acceptor [35]. 182 183



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Figure 3. Drawing of structure *trans-*[Ni(en)(H₂O)₂{B₆O₇(OH)₆}]·H₂O (7), showing selected crystallographic numbering.

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A drawing of the structure of $[Ni(dmen)(H_2O)\{B_6O_7(OH)_6\}]$:5H₂O (8), with selected 187 crystallographic numbering, is shown in Figure 4. Compound 8 is a distorted neutral octahedral Ni^(II) complex with disordered aqua ligand (containing O21) and dmen ligands 189 (0.496(5) s.o.f shown) and a tridentate $[B_6O_7(OH)_6]^{2-}$ ligand coordinated through three 190 hydroxy groups (containing O11, O12 and O13). The five waters of crystallization contain 191 O31-O35. The Ni1-O21 distance for the aqua ligand (2.131(11) Å) is significantly longer 192 than the Ni-O distances associated with the three hydroxy groups of the tridentate ox-193 idoborate ligand (2.0415(17)-2.0978(17) Å; av. 2.078(2) Å) but this difference was not sig-194 nificant for 7. The N1-Ni1-N2 angle associated with the chelating dmen ligand is 84.5(12)° 195 and angles between the three O-donor atoms of the fac-hexaborate ligand range from 196 84.26(7)-90.66(6)°; av. 86.94(8)°. The aqua ligand with O21 is trans to oxidoborate O11 with 197 an angle of 173.2(3)°, and the other trans angles are significantly <180.0°. The Ni1-N dis-198 tances of 2.031(11) Å (N1) and 2.13(2) Å (N2) are significantly different but there is more 199 steric demand at N2 as it is substituted by the two methyl groups. Overall, the 200 bondlengths and angles associated with the Ni^(II) centre and those within the hexabo-201 rate(2-) ligand in 8 are in accord with comparable data for 7. 202



Figure 4. Drawing of structure of [Ni(dmen)(H2O){B6O7(OH)6}]5H2O (8), showing (selected) crystallographic numbering.

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2.2.2 Solid-state H-bonding interactions in 1 - 8.

Compounds 1 - 8 are self-assembled and crystallized from the various oxidoborate 208 and the Ni^(II) amine complexes that are each in equilibrium in the reacting aqueous solu-209 tion [19,20,31]. There is a strong preference for the oxidoborate anion to enter the primary 210 coordination shell in related Cu^(II) and Zn^(II) chemistry with the formation of energetically 211 favourable O-donor coordinate bonds [32, 34-40, 58]. Interestingly, Ni^(II) salts are described 212 as labile, although in practice they are often considerably less labile than corresponding 213 Cu^(II) and Zn^(II) complexes [31]. In accord with this, the Ni^(II) cations used in this study 214 generally remain more or less intact in salts 1 - 6 after prolonged crystallization from aque-215 ous solution, with formation of oxidoborate O-donor bonds only observed in compounds 216 7 and 8. H-bond interactions also appear to be of paramount importance in driving this 217 crystallization process and these interactions are described in detail for each compound 218 in the following paragraphs. 219

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A drawing of the components of the formula unit of 1, showing important atomic 220 numbering, is shown in Figure 5. The cations, anions and interstitial H₂O molecules have 221 numerous H-bond donor and/or acceptor sites and all potential H-bond donor sites are 222 used in the solid-state structure. The dotted red lines in Figure 5 illustrate H-bond inter-223 actions which link cation to anion (O11H11...O6), cation to interstitial H₂O (N11H11...O21), 224 and anion to interstitial H₂O (O6H6···O21) forming a R₂²(8) ring incorporating the Ni^(II) 225 centre. These interactions as well as anion-anion interactions template the formation of 226 the tetraborate salt. Full details of all H-bond interactions are available in the supplemen-227 tary information. Siveav and co-workers [46] have analysed H-bond interactions in tetra-228 borate(2-) systems using the terminology of Zolatarev et al. [60] but we prefer the com-229 plimentary and more widely known Etter symbolism [59] in this manuscript. R₂²(8) inter-230 actions are very important at stabilizing hydroxyoxidopolyborate anions and three of the 231 four boron based hydroxy centres are involved in such interactions: observed in $R_{2}^{2}(8)$ 232 interactions are very important at stabilizing hydroxyoxidopolyborate anions and three 233 of the four boron based hydroxy centres are involved in such interactions with O8-H8-O5' 234 and O9'-H9'...O3 paired in a R22(8) ring and O7-H7...O1' and O7'-H7'...O1 forming a recip-235 rocal pair in another $R_{2}^{2}(8)$ ring. The fourth boron based hydroxy group O6H6 is a donor 236 to a H₂O molecule, O21 (Figure 5) forming an unusual R₃(8) ring {Ni1-O11A-H11A···O6-237 H6-O21-H11E-N11} and is also part of a larger R44(12) ring {O6-H6-O21-H21A-O3'-B1'-238 O6'-H6'-O21'-H21A'-O3-B1-} (Figure 6). Since the cation is involved in templating the 239 structure its H-bond interactions are very important and these include O11-H11A-O6, 240 O11-H11B...O8, O12-H12A...O4, O12-H12B...O2, N14-H14A...O8, N12-H12D...O7 and N12-241 H12C. O7 which originate from the aqua or ammine ligands. Further patterns are difficult 242 to visualize although R2²(8) {Ni1-N12-H12C····O7-B2-O2···H12B-O12-} and a C2²(10) {Ni1-243 O11-H11A-O6-B1-O3-B3-O8-H14A'-N14'-} are discernable. 244



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Figure 5. Drawing of structure of *trans*-[Ni(NH₃)₄(H₂O)₂][B₄O₅(OH)₄]·H₂O [Ni(dmen)(H₂O){B₆O₇(OH)₆}]·5H₂O (**1**), showing (selected) crystallographic numbering and the R₃³(8) ring system: {Ni1-O11A-H11A···O6-H6···O21···H11E-N11}.



Figure 6. Drawing of the R4⁴(12) ring, {O6-H6…O21-H21A…O3'-B1'-O6'-H6'…O21'-H21A'…O3-B1-}, in *trans*-[Ni(NH₃)4(H₂O)₂][B₄O₅(OH)4]·H₂O (**1**).

Compounds 2 - 5 are all pentaborate(1-) salts and anion-anion H-bond interactions 254 are well known to strongly influence their formation [5, 25-28] since each pentaborate(1-) 255 anion has four donor H-bond sites that are sterically arranged to form 3D H-bonded ani-256 onic giant structures, in which the cations occupy vacant 'open-sites' in the anionic lattice. 257 The anion-anion interactions are conveniently described using a combination of 'acceptor-258 sites nomenclature', as defined by Schubert [26], and Etter terminology [59]. Compounds 259 2 - 4 have very similar anion-anion interactions and the four acceptor sites can be de-260 scribed as $\alpha, \alpha, \alpha, \beta$ with the α -site interactions reciprocal R₂²(8) and the β -interactions re-261 ciprocal $R_2^2(12)$; both types of interactions are common in pentaborate(1-) structures [5, 262 27]. The anion interactions in 5 are different with three pentaborate(1-) acceptor sites 263 (α, α, γ) , all reciprocal R₂²(8), and an interstitial H₂O acceptor site. The cations in **2** - **5** are 264 situated within the anionic lattices and have the potential to be H-bond donors (and ac-265 ceptors) which can further stabilize the anionic lattices and help to template their self-266 assembly. Examples of such interactions, which involve novel ring systems that incorpo-267 rate the Ni^(II) centres, ligands interstitial molecules and pentaborate(1-) anions, are $R_{44}(14)$ 268 {Ni1-N3B-H3B··O10-O1OH··O9-O9H··O4-B1-O1-B2-O7··H2B-N2B-} (2) (includes three 269 pentaborate(1-) anions), R2²(8) {Ni1-O11-H11B-07-B2-O2-H12A-N12-} (3) (one pentabo-270 rate(1-) anion), R₂²(8) {Ni1-N11-H11D··O11-B2-O7··H11-O11-} (4) (one pentaborate(1-) an-271 ion) and R₂²(8) {Ni1-N11-H11D-O1-B2-O7-H11-O11-} (4) (one pentaborate(1-) anion), and 272 R4⁴(16) {Ni1-N1-H1C···O5-B4-O4-B1-O3···H1A-O11···H7-O7-B2-O1···H4A-N4-} (5) (two 273 pentaborate(1-) anions and one interstitial H_2O with O11). The $R_2^2(8)$ interaction for **3** is 274illustrated in Figure 7. Full details of all these interactions (and associated atomic num-275 bering systems) are available in the supplementary information. 276

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 Figure 7. Drawing of the R2²(8) {Ni1-O11-H11B···O7-B2-O2···H12A-N12-} interaction involving the Ni^(II) in *trans*-[Ni(dmen)2(H2O)2]
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 [B5O6(OH)4]2·2H2O (3).
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Compound 6 is a heptaborate(2-) salt with numerous opportunities for interionic H-282 bonding interactions with the each anion possessing 14 potential acceptor sites and 5 do-283 nor sites and the two cations potentially having 20 donor sites and 4 acceptor sites; there 284 are also 4 interstitial H₂O molecules. Potentially templating cation-anion interactions in-285 clude the R₂²(8) {Ni1-N12-H12A···O10-B4-O3···H11A-N11-}, the R₃³(8) {Ni2-N12-286 H12D-023-H23B-011-H11D-N11-}, and the R2²(10) {Ni1-O21-H21B-01-B2-287 O6...H2B-N2-}. The familiar $R_2^2(8)$ anion-anion interaction is present in {O12-H12...O9'-B7'-288 O13'-H13'-O7-B6-} and the reciprocal {O11-H11--O5'-B5'-O11'-H11'--O5-B5-} and there is 289 also a R₂²(12) interaction, {O10-H10···O8'-B2'-O2'-B4'-O10'-H10'···O8-B2-O2-B4-}. The fifth 290 heptaborate hydroxy group (containing O14) does not form a donor bond but O14 has a 291 $R_{2^{2}}(10)$ acceptor interaction with a cation, as described above. 292

Intermolecular H-bonding, in addition to the intramolecular interaction found in 7 293 (Section 2.2.1) can be found in 7 and 8. Such interactions for 7 include $R_2^2(6)$ {Ni1-O8-294 H8--O2--H1AA-N1-} and {Ni1-O9-H9--O3--H2BD-N2-}, R44(8) {O21-H21B--O10H10--O21'-295 H21B'...O10'-H10'...} and a reciprocal R22(12) {O12-H12...O6'-B3'-O5'-B5'-O12'-H12'...O6-296 B3-O5-B5-}. There are 5 interstitial H₂O's per Ni(II) centre in 8 and their main roles are to 297 act as 'spacers' [28] by forming H-bond bridges between neutral Ni^(II) oxidoborate com-298 plexes. Templating interactions occur when the oxidoborate ligand of one complex en-299 ters the second coordination sphere of another. This occurs in reciprocal oxidoborate lig-300 and R₂²(12) interactions, {O11-H11...O10'-B6'-O7'-B1'-O11'-H11'...O10-B6-O7-B1-} and 301 {O12-H12...O9'-B5'-O4'-B2'-O12'-H12'...O9-B5-O4-B2-}; these are that similar to one ob-302 served in 7 and in another heptaborate(2-) system [52]. A R₂²(8) {O8-H8-O2'-B4'-O8'-303 H8'-O2-B4-} interaction also links oxidoborate sections of neighbouring complexes. 304

2.3. Magnetic, spectroscopic, thermal and p-XRD characterization of compounds 1-8

Magnetic susceptibility measurements on 1 - 8 confirmed that all, excepting 5, are paramagnetic with χ_m values ranging from 2500 x10⁻⁶ to 3600 x10⁻⁶ cm³mol⁻¹ (μ_{eff} of 2.6 - 3.0 308

BM) corresponding to 2 unpaired electrons per formula unit. Compound **5** is diamagnetic 309 $(\chi_m = -170 \times 10^{-6} \text{ cm}^3 \text{mol}^{-1})$, and this value is in accord with the Ni^(II) centre being squareplanar. The χ_m values of **1** - **4**, **7** and **8** are typical of octahedral Ni^(II) complexes [61] and 311 have similar values to those obtained for the starting octahedral Ni^(II) complexes. Compound **6** is unusual in that it has a μ_{eff} corresponding to two unpaired electrons but there are *two* Ni^(II) centres in the formula unit; the XRD study confirmed that one Ni^(II) centre is 314 square planar and the other is octahedral. 315

Compound 5, in D₂O solvent, gave ¹H and ¹³C NMR signals for the AEN ligand at 316 similar intensities and chemical shifts to those obtained for [Ni(AEN)]Cl-H2O and other 317 reported data [62]. ¹H and ¹³C NMR signals for the organic ligands (in D₂O) were not ob-318 servable for the other Ni^(II) oxidoborates despite their apparent solubility, due to their par-319 amagnetic properties. Despite the possible lability of the organic ligands the complexes 320 the organic ligands are not sufficiently labile to be remote from the Ni^(II) centres during the 321 lifetime of the NMR experiment. ¹¹B NMR spectra for all D₂O solutions arising from the 322 Ni^(II) oxidoborates were observed and a probable explanation for this being two-fold: (i) 323 compounds with insular oxidoborate anions (1 - 6) are not coordinated to the Ni^(II) centres 324 and hence are not in close proximity and unaffected by the paramagnetic centres, and (ii) 325 the coordinated hexaborate(2-) ligands are significantly more labile than the N-donor lig-326 ands in 7 and 8. The ¹¹B spectra were not particularly characteristic of the specific ox-327 idopolyborate anions in 1 - 8 since it is known that dissolution of oxidopolyborate anions 328 in aqueous solution yields solutions containing equilibrium mixtures of various ox-329 idopolyborate anions whose concentrations are pH are boron concentration dependent 330 [19-20]. However, pentaborate(1-) salts often afford spectra in a characteristic pattern with 331 three signals at ca. +17, +13 and +1 ppm which have been assigned to B(OH)₃/[B(OH)₄]⁻, 332 $[B_3O_3(OH)_4]^-$ and $[B_5O_6(OH)_4]^-$ species [5, 27, 28]; compounds 2 - 4 all showed this charac-333 teristic pattern. Dilute solutions often give just one signal, assigned to B(OH)₃/[B(OH)₄]⁻, 334 at a chemical shift which is variable [19] but can be calculated assuming fast exchange of 335 $B(OH)_3/[B(OH)_4]^{-1}$ according to their relative proportions present in the oxidopolyborate 336 anion originally present [27]. Unsurprisingly therefore, compounds 1 and 5 - 8 all gave 337 single signals at +11.1, +16.3, +15.8, +17.7 and +15.8 ppm respectively. The observed chem-338 ical shifts for the tetraborate(2-) (1) and the pentaborate(-) (5) salts are as would be ex-339 pected from this calculation (+11.0 and +16.1, respectively) but chemical shifts for the hex-340 aborate(2-) (7 and 8) and the heptaborate(2-) (6) salts are ca. 2 ppm more downfield than 341 expected (13.8 and 14.6 ppm calc., respectively), indicating that more B(OH)3 might be 342 present than expected. Similar behaviour has been observed before for hexaborate [34-38] 343 and heptaborate [33] complexes. 344

Vibration spectroscopy is often used to help characterize oxidopolyborate species 345 since strong, and often diagnostic, B-O stretches can be observed by IR spectroscopy be-346 tween 1600 and 600 cm⁻¹[63]. In particular diagnostic bands have been reported for pen-347 taborate(1-) [5, 27,28, 64], hexaborate (2-) [35, 37, 63], and heptaborate(2-) [28, 33, 52] ani-348 Compounds 1 - 8 all displayed such diagnostic bands with the diagnostic band for ons. 349 the pentaborate(1-) anion (ca. 1925 cm⁻¹) clearly present in 1-5. Likewise the diagnostic 350 bands for hexaborate(2-) anion (ca. 960 and 810 cm⁻¹) and the heptaborate(2-) anion (ca. 351 860 cm⁻¹) were observed in IR spectra of 6 and 7, and 8, respectively. 352

Selected Ni^(II) oxidoborates (2, 4 - 7) were thermally decomposed air and insights 353 into these decompositions were obtained through TGA/DSC studies. The Ni^(II) ox-354 idoborates all yielded green glassy residues by 700 °C with masses consistent with anhy-355 drous Ni^(II) oxidoborates being produced with Ni:B stochiometries in the elemental ratios 356 of their precursors. Thus, the pentaborates 2 and 4 were decomposed to NiB10O16 whilst 5 357 gave Ni2B10O17, the heptaborate (6) gave Ni2B14O23 and the hexaborate (7) yielded NiB6O10. 358 These decomposition reactions are generally two-step processes with the first steps being 359 a lower temperature (<280 °C) endothermic dehydration (loss of interstitial H2O and con-360 densation reactions of the hydrated oxidoborates) followed by higher temperature (280 – 361 700 °C) exothermic oxidations of the organic ligands. Details for individual compounds 362 (2, 4 - 7) are given in the experimental section. This type of decomposition behaviour has 363 been observed previously for other hydrated Ni^(II) oxidoborates [33, 42-48] and more gen-364 erally applies to the thermolysis of many hydrated transition-metal oxidopolyborates [30, 365 33, 35, 37]. Powder XRD data were also obtained for this selection of Ni^(II) oxidoborates 366 (viz. 2, 4 - 7). The data obtained were good matches with data calculated from the single-367 crystal XRD data indicating that these samples were crystalline and homogeneous. 368

3. Experimental Methods

3.1. General

The Ni^(II) complexes *s-fac*-[Ni(dien)₂]Cl₂·H₂O [65] [Ni(dmen)₂]Cl₂·4H₂O[66], 372 [Ni(HEen)2]Cl2 [67], [Ni(AEN)]ClH2O [62], trans-[Ni(dach)2(H2O)2]Cl2 [68] were prepared 373 by modified literature procedures. FTIR spectra were obtained as KBr pellets on a Perkin-374 Elmer 100FTIR spectrometer (Perkin-Elmer, Seer Green, UK). ¹¹B NMR spectra were ob-375 tained on a Bruker Avance-400 spectrometer (Bruker, Coventry, UK) on samples dis-376 solved in D₂O at 128 MHz. TGA and DSC were performed on an SDT Q600 instrument 377 (TA Instruments, New Castle DE, USA) using Al₂O₃ crucibles with a ramp rate of 10 °C 378 per minute (RT to 1000 °C in air). X-ray crystallography was performed at the EPSRC na-379 tional crystallography service centre at Southampton University. Magnetic susceptibility 380 measurements were performed on a Johnson-Matthey magnetic susceptibility balance. 381 (Johnson-Matthey, UK). CHN analyses were obtained from OEA Laboratories (Calling-382 ham, Cornwall). 383

3.2 X-ray crystallography.

Crystallographic data for 1 - 8 are given in the experimental section. The crystallo-386 graphic data collection of compounds were performed on a Rigaku FR-E+ diffractometer 387 (1, 3 - 8) equipped with an AFC12 goniometer, an HG Saturn 724+ detector and either HF 388 varimax confocal mirrors (1, 3, 4, 6 - 8) or VHF varimax confocal mirrors (5). Data for 2 389 were obtained on DLS synchronous beamtime 119 equipped with a crystallographic go-390 niometer and a Rigaku HG Saturn 724 detector. Data for all compounds were obtained at 391 100(2)K. Cell determinations and data collections were carried out using either CrysA-392 lisPro [69] (1) or CrystalClear [70] (2 - 8). Data reduction, cell refinement and absorption 393 correction were carried out using either CrysAlisPro [69] (1 - 3, 5 - 8) or CrystalClear [70] 394 (4). Using Olex2 [71] the structures were solved using SHELXT [72] and models or refined 395 with SHELXL [73]. Structure refinement details are given in the supplementary material. 396 CCDC 2101911-2101918 (1 - 8 respectively), contains the supplementary crystallographic 397 data for this paper. These data can be obtained free of charge via 398 http://www.ccdc.cam.ac.uk/conts/retrieving.html (or from CCDC, 12 Union Road, Cam-399 bridge, CB2 1EZ. Fax: +44 1223 336033; E-mail: deposit@ccdc.cam.ac.uk) 400

3.3 Preparation of trans- $[Ni(NH_3)_4(H_2O)_2][B_4O_5(OH)_4]H_2O(1)$.

NiCl₂6H₂O (1.00 g, 4.0 mmol) dissolved in ethanol (25 mL) and gave a clear green 403 solution. Aqueous NH3 solution (2.2 mL, 25%, 24 mmol) was added dropwise to the green 404 solution with stirring and then was left to stir for a further 5 mins to give a dark blue 405solution. Ag₂O (0.926 g, 4 mmol) was rapidly added and stirred at room temperature for 406 5 mins and the precipitate which had formed (AgCl) was removed by filtration. B(OH)³ 407 (1.23 g, 20 mmol) was added to the dark blue filtrate and left to stir for a further 10 mins. 408 The reaction mixture was filtered into vials and left to allow for slow evaporation of the 409 solvent. After 2 weeks a light blue solid (1) had formed in the bottom of a vial and this 410was collected by filtration. (0.6 g, 40 %). M.p. = >300 °C. χ_m = 2500×10⁻⁶ cm³mol⁻¹. 411 H₂₂B₄N₄NiO₁₂. Anal. Calc.: H = 6.0 %, N = 15.1 %. Found: H = 5.7 %, N = 15.3. ¹¹B/ppm: 11.1. 412 IR (KBr/cm⁻¹): 3375(s), 3245(s), 1634(m), 1450(s), 1402(s), 1345(s), 1230(m), 1152(m), 413 1059(m), 1002(s), 936(m), 828(m). sc-XRD crystal data: Mr = 372.16, monoclinic, P21/c (No. 414

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14), a = 8.27730(10) Å, b = 20.5277(3) Å, c = 8.01330(10) Å, $\beta = 91.2830(10)^{\circ}$, $\alpha = \gamma = 90^{\circ}$, V = 4151361.23(3) Å³, T = 100(2) K, Z = 4, Z' = 1, $\mu(MoK_{\alpha}) = 1.492$ mm⁻¹, 15477 reflections measured, 416 3128 unique ($R_{int} = 0.0329$) which were used in all calculations. The final wR_2 was 0.0528 417 (all data) and R_1 was 0.0216 (I > 2 σ (I)). 418

3.4 Preparation of s-fac- $[Ni(dien)_2][B_5O_6(OH)_4]_2$ (2).

s-fac-[Ni(dien)2]Cl2H2O (1.05 g, 3.0 mmol) was dissolved in H2O (10 mL) and added 421 to a suspension of excess DOWEX® monosphere 550A OH- activated anion exchange resin 422 (40 g) in H₂O (40 mL). The mixture was left to stir for 30 h at RT and then filtered to 423 remove the resin. The filtrate, containing s-fac-[Ni(dien)2](OH)2, was reduced in volume 424 to 15 mL and MeOH (15 mL) was added. B(OH)3 (1.83 g, 29.7 mmol) was added to the 425 H2O/MeOH solution which was then gently warmed with stirring for 3 hours. The solu-426 tion was concentrated to 5 mL under reduced pressure and the concentrated solution was 427 left for a few days for crystallization. Purple crystals of **2** were carefully recovered by 428 filtration and dried in air (0.70 g, 34 %). M.p. = 287-289 °C (dec.). χ_m = 3516×10⁻⁶ cm³ mol⁻¹. 429 CsH34B10N6NiO20. Anal. Calc.: C = 13.7 %, H = 4.9 %, N = 12.0 %. Found: C = 14.0 %, H = 5.4 430 %, N = 12.1 %. ¹¹B/ppm: 1.5 (1 %), 13.3 (13 %), 17.4 (86 %). IR (KBr/cm⁻¹): 3362(s), 3333(s), 431 3297(s), 3263(s), 1412(s), 1330(s), 1303(s), 1135(m), 1059(m), 1042(m), 1028(m), 964(m), 432 931(m), 915(s), 774(s), 772(m), 707(s). TGA: 100-200 °C, condensation of polyborate which 433 loss of four H2O 14.9 % (10.2 % calc.); 200-650 °C, oxidation of organic content 40.3 % (39.7 434 % calc.); residual NiB₈O₁₃ 59.7 % (60.3 % calc.). p-XRD d-spacing/Å (% rel. int.): 6.22 (24), 435 5.88 (100), 5.57 (94), 4.27 (15), 3.79 (19), 3.71 (28), 2.94 (17). sc-XRD crystal data: Mr = 701.22, 436 437 89.108(5)°, $\gamma = 89.110(5)°$, V = 728.75(8) Å³, T = 100(2) K, $Z = 1, Z' = 0.5, \mu$ (Synchrotron_{0.6889Å}) = 438 0.702 mm^{-1} , 6342 reflections measured, 3152 unique ($R_{int} = 0.0476$) which were used in all 439 calculations. The final wR_2 was 0.2109 (all data) and R_1 was 0.0767 (I > 2 σ (I)). 440

3.5 Preparation of trans-[Ni(dmen)₂(H₂O)₂][B₅O₆(OH)₄]₂·2H₂O (3).

[Ni(dmen)2]Cl24H2O (0.97 g, 3.0 mmol), and Ag2O (0.75 g, 3.2 mmol) were rapidly 443 stirred in H₂O (30 mL) at room temperature for 40 mins and the precipitate which had 444 formed (AgCl) was removed by filtration. B(OH)3 (1.85 g, 30 mmol) was added to the dark 445 blue filtrate and left to stir for a further 40 mins. The reaction mixture was filtered into 446 vials and then left to allow for slow evaporation of solvent. After 3 weeks a light blue solid 447 (3) had formed in the bottom of a vial and this was collected by filtration. (0.9 g, 40 %). 448 M.p. = > 300 °C. χ_m = 3600×10⁻⁶ cm³mol⁻¹. C₈H₄₀B₁₀N₄NiO₂₄. Anal. Calc.: C = 12.9 %, H = 5.4 449 %, N = 7.5 %. Found: C = 13.0 %, H = 5.5 %, N = 7.5. ¹¹B/ppm 16.6 (85%), 13.9 (14%), 1.2 450 (1%). IR (KBr/cm⁻¹): 3510 (m), 3348(s), 3285(s), 1608(m), 1415(s), 1310(s), 1129(m), 1043(m), 451 1015(m), 920(s), 780(s), 707(m), 486(m). sc-XRD crystal data: Mr = 743.25, triclinic, P-1 452 (No. 2), a = 8.6570(3) Å, b = 9.7443(3) Å, c = 10.2101(3) Å, $\alpha = 107.087(3)^{\circ}$, $\beta = 102.747(3)^{\circ}$, $\gamma = 102.747(3)^{\circ}$ 453 95.002(3)°, V = 792.23(5) Å³, T = 100(2) K, Z = 1, Z' = 0.5, μ (MoK_{α}) = 0.708 mm⁻¹, 12978 re-454 flections measured, 3628 unique ($R_{int} = 0.0163$) which were used in all calculations. The 455 final wR_2 was 0.0731 (all data) and R_1 was 0.0270 (I > 2 σ (I)). 456

3.6 Preparation of [Ni(HEen)₂][B₅O₆(OH)₄]₂ (4).

The preparation of [Ni(HEen)₂](OH)₂ was carried out as described previously for 2 459 from [Ni(HEen)2]Cl2 (1.00 g, 2.95 mmol) and Dowex (40 g). B(OH)3 (1.83 g, 29.6 mmol) was 460 added with stirring. The reaction mixture was stirred for a further 2 h at room tempera-461 ture. The solvent was then evaporated to 5 mL using a rotary evaporator, then the result-462 ing solution was left for 10 d in several NMR tubes for crystallization to yield purple crys-463 tals of 4 (0.76 g, 37 %). M.p. = 288-289 °C (dec.). $\chi_m = 3430 \times 10^{-6} \text{ cm}^3 \text{ mol}^{-1} \text{ C}_8\text{H}_{32}\text{B}_{10}\text{N}_4\text{NiO}_{22}$. 464 Anal. Calc.: C = 13.7 %, H = 4.6 %, N = 8.0 %. Found: C = 13.7 %, H = 4.7 %, N = 8.0 %. 465 ¹¹B/ppm: 1.2 (3 %), 13.4 (23 %), 18.0 (74 %). IR (KBr/cm⁻¹): 3375(s), 3288(s), 3322(s), 1410(s), 466 1321(s), 1196(m), 1141(s), 1022(s), 919(s), 774(s), 706(s). TGA: 230-290 °C, condensation of polyborate which loss of four H₂O 10.3 % (10.2 % calc.); 290-700 °C, oxidation of organic content 37.9 % (39.9 % calc.); residual NiB₁₀O₁₆ 62.1 % (60.1 calc.). p-XRD d-spacing/Å (% rel. int.): 6.18 (40), 5.88 (75), 5.50 (100), 3.94 (26), 3.78 (38), 3.66 (25). sc-XRD crystal data: M_r = 703.18, Monoclinic, P_{21}/c , a = 8.3891(2) Å, b = 11.6666(3) Å, c = 14.3092(4) Å, α = γ = 90°, β = 90.035(2)°, V =1400.47(6) Å³, T = 100(2) K, Z = 2, Z' = 0.5, μ (MoK_α) = 0.791 mm⁻¹, 6239 reflections measured, 6239 unique (R_{int} = 0) which were used in all calculations. The final

3.7 Preparation of $[Ni(AEN)][B_5O_6(OH)_4]H_2O$ (5).

 wR_2 was 0.1124 (all data) and R_1 was 0.0499 (I > 2 σ (I)).

The preparation of [Ni(AEN)](OH) was carried out as described previously for 2 477 from [Ni(AEN)]Cl·H2O (2.10 g, 7.0 mmol) and Dowex (25 g). B(OH)3 (2.22 g, 36 mmol) was 478 added to the aqueous solution which was gently warmed with stirring for 3 h, and then 479 its volume was reduced to 5 mL under reduced pressure. The solution was then distrib-480 uted in several NMR tubes for crystallization and left for 10 days to yield dark red crystals 481of 5 (1.30 g, 38 %). M.p. = 279-280 °C (dec.). χ_m = -170×10⁻⁶ cm³ mol⁻¹. C₉H₂₅B₅N₄NiO₁₁. Anal. 482 Calc.: C = 22.6 %, H = 5.3 %, N = 11.7 %. Found: C = 22.7 %, H = 5.4 %, N = 11.7 %. ¹H/ppm: 483 2.0 (s, 6H, CH₃), 2.5 (t, 4H, J = 6 Hz, CH₂ of en), 3.2 (t, 4H, J = 6 Hz, CH₂ of en), 4.8 (s, 11H, 484 NH2, H2O, OH, CH). ¹³C/ppm: 19.9, 43.0, 53.6, 160.5. ¹¹B/ppm: 16.3. IR (KBr/cm⁻¹): 3377(s), 485 3326(s), 3288(s), 3246(s), 3169(s), 1615(m), 1566(m), 1536(m), 1476(s), 1442(s), 1416(s), 486 1393(s), 1333(s), 1295(s), 1129(m), 1094(s), 1072(m), 1019(m), 924(s), 778(m). TGA: 100-180 487 °C, loss of one interstitial H2O 2.5 % (3.7 % calc.); 180-280 °C, condensation of polyborate 488 which loss of two further H2O 10.0 % (11.3 % calc.); 280-700 °C, oxidation of organic con-489 tent 46.1 % (48.0 % calc.); residual Ni2B10O17 53.9 % (52.0 % calc.). p-XRD d-spacing/Å (% 490 rel. int.): 5.65 (69), 5.38 (100), 4.14 (46), 3.89 (33), 3.81 (53), 3.65 (58), 2.03 (41), 1.43 (33). sc-491 XRD crystal data: M_r = 478.09, Triclinic, P-1, a = 8.5386(2) Å, b = 11.3328(3) Å, c = 11.8717(3) 492 Å, $\alpha = 115.914(3)^\circ$, $\beta = 101.626(2)^\circ$, $\gamma = 99.670(2)^\circ$, V = 968.39(5)Å³, T = 100(2) K, Z = 2, Z' = 100(2) K, Z =493 1, μ (MoK_a) = 1.065 mm⁻¹, 16639 reflections measured, 4433 unique ($R_{int} = 0.0216$) which 494 were used in all calculations. The final wR_2 was 0.0765 (all data) and R_1 was 0.0278 (I > 495 $2\sigma(I)$). 496

3.8 Preparation of trans-[Ni(dach)₂(H₂O)₂] [Ni(dach)₂] [B₇O₉(OH)₅]₂4H₂O (6).

The preparation of [Ni(dach)2(H2O)2](OH)2 was carried out as previously described 499 for 2 from trans-[Ni(dach)2(H2O)2]Cl2 (1.00 g, 2.53 mmol) and Dowex (40 g). B(OH)3 (1.56 500 g, 25.2 mmol) was added to the solution which was then gently warmed with stirring for 501 3 h. The solution volume was reduced to 5 mL under reduced pressure. The product (6) 502 was isolated by filtration, and then allowed to dry in an oven at 40 °C for 3 h to afford 503 orange prismatic crystals (0.78 g, 47 %). M.p. = 270-272 °C (dec.). χ_m = 3030×10⁻⁶ cm³ mol⁻¹. 504 C₂₄H₇₈B₁₄N₈Ni₂O₃₄. Anal. Calc.: C = 22.3 %, H = 6.1 %, N = 8.7 %. Found: C = 21.8 %, H = 6.3 505 %, N = 8.4 %. ¹¹B/ppm: 15.8. IR (KBr/cm⁻¹): 3663(s), 3271(s), 2929(m), 2863(m), 1468(m), 506 1453(s), 1350(s), 1180(s), 1125(m), 1066(s), 986(w), 854(m), 807(m). TGA: 100-190 °C, loss of 507 four interstitial and two coordinated H2O 7.5 % (8.4 % calc.); 190-270 °C, condensation of 508 polyborate which loss of five further H2O 15.0 % (15.3 % calc.); 270-720 °C, oxidation of 509 organic content 48.2 % (50.7 % calc.); residual Ni2B14O23 51.8 % (49.3 % calc.). p-XRD d-510 spacing/Å (% rel. int.): 10.67 (100), 9.81 (89), 6.25 (58), 5.35 (70), 4.26 (48), 3.74 (55), 2.03(59). 511 sc-XRD crystal data: M_r = 645.85, Monoclinic, C2/c, a = 22.3539(4) Å, b = 11.0192(2) Å, c = 512 22.8834(4) Å, $\alpha = \gamma = 90^{\circ}$, $\beta = 107.630(2)^{\circ}$, V = 5371.94(18) Å³, T = 100(2) K, Z = 8, Z' = 1, 513 μ (MoK_a) = 0.806 mm⁻¹, 33628 reflections measured, 6154 unique ($R_{int} = 0.0281$) which were 514 used in all calculations. The final wR_2 was 0.1087 (all data) and R_1 was 0.0384 (I > 2 σ (I)). 515

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3.9 Preparation of trans- $[Ni(en)(H_2O)_2\{B_6O_7(OH)_6\}]$ · H_2O (7).

An aqueous solution of en (2.30 g, 70 %, 26.79 mmol) was added to an aqueous solu-521 tion (10 mL) of NiSO46H2O (2.12 g, 8.06 mmol). The reaction mixture was stirred for 5 522 mins, and then an aqueous solution (10 mL) of Ba(OH)₂·8H₂O (2.54 g, 8.06 mmol) was 523 added. The reaction mixture was stirred for 30 mins and then filtered to remove the BaSO4. 524 A solution of B(OH)₃ (4.98 g, 80.6 mmol) in H₂O (10 mL) was added to the filtrate with 525 stirring. The reaction mixture was stirred at room temperature for 2 h. The solution was 526 then distributed in several vials and left for 10 d to yield faint blue crystals of 7 (1.20 g, 33 527 %). M.p. = 245-247 °C (dec.). χ_m = 2588×10⁻⁶ cm³ mol⁻¹. C₂H₂₀B₆N₂NiO₁₆. Anal. Calc.: C = 5.3 528 %, H = 4.5 %, N = 6.2 %. found: C = 5.4 %, H = 4.5 %, N = 6.2 %. ¹¹B/ppm: 17.7. IR (KBr/cm⁻ 529 ¹): 3400(s), 3350(s), 2924(w), 1420(m), 1380(s), 1360(s), 1133(s), 1095(s), 1044(s), 955(m), 530 908(m), 809(s). TGA: 100-180 °C, loss of one interstitial H2O and two coordinated water 531 molecules 0.8 % (12 % calc.); 180-280 °C, condensation of polyborate which loss of three 532 further H₂O 24.3 % (23.9 % calc.); 280-700 °C, oxidation of organic content 38.6 % (37.2 % 533 calc.); residual NiB₆O₁₀ 61.4 % (62.8 % calc.). p-XRD d-spacing/Å (% rel. int.): 9.78 (100), 534 8.26 (24), 6.77 (53), 4.88 (48), 3.99 (34), 3.38 (30), 2.74 (47). sc-XRD crystal data: Mr = 451.77, 535 triclinic, P-1 (No. 2), a = 8.5116(7) Å, b = 9.7946(5) Å, c = 9.8073(8) Å, $\alpha = 89.873(5)^{\circ}$, $\beta =$ 536 82.901(7)°, $\gamma = 74.372(6)°$, V = 780.94(10) Å³, T = 100(2) K, Z = 2, Z' = 1, $\mu(MoK_{\alpha}) = 1.333$ mm⁻ 537 ¹, 5883 reflections measured, 5883 unique ($R_{int} = 0$) which were used in all calculations. The 538 final wR_2 was 0.1902 (all data) and R_1 was 0.0682 (I > 2 σ (I)). 539

3.10 Preparation of [Ni(dmen)(H₂O){B₆O₇(OH)₆}]5H₂O (8).

The preparation of [Ni(dmen)₂](OH)₂ was carried out as described for 2 from 542 [Ni(dmen)2]Cl24H2O (1.00 g, 2.6 mmol) and Dowex (30 g). B(OH)3 (1.57 g, 25 mmol) was 543 added to the filtrate and left to stir for 30 mins. The turquoise solution was filtered into 544 vials and left to allow for slow evaporation of the solvents. After 2 weeks a blue-green 545 solid (8) had formed in the bottom of a vial and this was collected by filtration. Yield: (0.5 546 g, 36 %). M.p. = > 300 °C. χ_m = 3050×10⁻⁶ cm³mol⁻¹. C₄H₃₀B₆N₂NiO₁₉. Anal. Calc.: C = 9.0 %, 547 H = 5.7 %, N = 5.2 %. Found: C = 9.3 %, H = 5.5 %, N = 5.5. ¹¹B/ppm: 15.8. IR (KBr/cm⁻¹): 548 3399(s), 1642(m), 1416(s), 1355(s), 1247(s), 1120(s), 1052(m), 1009(m), 962(m), 859(m), 813(s), 549 700(m). sc-XRD crystal data: M_r = 533.87, triclinic, P-1 (No. 2), a = 9.1508(4) Å, b = 550 9.8599(4) Å, c = 13.5971(4) Å, $\alpha = 88.525(3)^{\circ}$, $\beta = 70.627(3)^{\circ}$, $\gamma = 65.245(4)^{\circ}$, V = 1041.79(8) Å³, 551 T = 100(2) K, Z = 2, Z' = 1, μ (MoK $_{\alpha}$) = 1.023 mm⁻¹, 15537 reflections measured, 4713 unique 552 $(R_{int} = 0.0461)$ which were used in all calculations. The final wR_2 was 0.1048 (all data) and 553 R_1 was 0.0403 (I > 2 σ (I)). 554

4. Conclusions

Eight new oxidoborate Ni(II) compounds have been prepared by self-assembly crys-556 tallization processes from aqueous solution starting from B(OH)₃ and selected Ni^(II) com-557 plexes. Six of these compounds are salts containing *insular* oxidoborate anions, but two 558 products are complexes containing *coordinated* oxidoborate(2-) anions, with energetically 559 favourable Ni-O bonds. The two products that contain coordinated hexaborate(2-) ligands 560 are neutral monomeric complexes, rather than the 1-D coordination polymers that we 561 have occasionally observed in related Cu^(II) and Zn^(II) chemistry. The new compounds are 562 templated in all cases by numerous strong structure-directing inter/intramolecular H-563 bonding interactions involving the oxidoborate ligands and many novel H-bonding mo-564 tifs are noted and described. This work further demonstrates that in general the inclusion 565 of labile metal complexes into aqueous solutions containing B(OH)³ can lead to the self-566 assembly of novel species including those with coordinated oxidoborate ligands and that 567 this synthetic strategy could be applied successfully to other systems.

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	Supplementary Materials: The following are available online at xxxxxxx: single-crystal XRD data., including H-bond interactions.	572 573
	Author Contributions: M.A.B. conceived the experiments; M.A.A. synthesized and characterized the complexes and grew the single-crystals; P.N.H. and S.J.C. solved the crystal structures; M.A.B. wrote the paper with contributions from all co-authors.	574 575 576
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	Conflicts of Interest: The authors declare no conflict of interest.	580
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