Hexaborate(2-) and Dodecaborate(6-) Anions as Ligands to Zinc(II) Centres: Self-assembly and Single-crystal XRD Characterization of [Zn\{3O-B6O7(OH)6}\{3N-dien\}]0.5H2O (dien = NH(CH2-CH2NH2)2), (NH4)2[Zn\{2O-B6O7(OH)6\}2 (H2O)2].2H2O and (1,3-pnH2)3[(1N-H3N(CH2)3NH2) Zn\{3O-B12O18(OH)6\}]2.14H2O (1,3-pn = 1,3-diaminopropane).

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Hexaborate(2−) and Dodecaborate(6−) Anions as Ligands to Zinc(II) Centres: Self-Assembly and Single-Crystal XRD Characterization of 

\[ \text{[Zn}\{\kappa^3O-B_6O_7(OH)_6}\}(\kappa^3N\text{-dien})\cdot0.5\text{H}_2\text{O} \text{ (dien = NH(CH}_2\text{CH}_2\text{NH}_2)\_2}, \text{(NH}_4\text{)}_2\text{[Zn}\{\kappa^2O-B_6O_7(OH)_6]\text{]}_2\text{(H}_2\text{O})_2]\cdot2\text{H}_2\text{O and (1,3-pnH}_2\text{)}_3[(\kappa^1N-H_3N\text{[CH}_2\text{]}_3\text{NH}_2)\text{Zn}\{\kappa^3O-B_{12}O_{18}(OH)_6\}]\cdot14\text{H}_2\text{O} \]

(1,3-pn = 1,3-diaminopropane)

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Abstract: Two zinc(II) hexaborate(2−) complexes, \([\text{Zn}\{\kappa^3O-B_6O_7(OH)_6}\}(\kappa^3N\text{-dien})\cdot0.5\text{H}_2\text{O} \text{ (dien = NH(CH}_2\text{CH}_2\text{NH}_2)\_2}, \text{(NH}_4\text{)}_2\text{[Zn}\{\kappa^2O-B_6O_7(OH)_6]\text{]}_2\text{(H}_2\text{O})_2]\cdot2\text{H}_2\text{O} \) (2), and a zinc(II) dodecaborate(6−) complex, \((1,3\text{-pnH}_2\text{)}_3[(\kappa^1N-H_3N\text{[CH}_2\text{]}_3\text{NH}_2)\text{Zn}\{\kappa^3O-B_{12}O_{18}(OH)_6\}]\cdot14\text{H}_2\text{O} \) (1,3-pn = 1,3-diaminopropane) (3), have been synthesized and characterized by single-crystal XRD studies. The complexes crystallized through self-assembly processes, from aqueous solutions containing 10:1 ratios of B(OH)_3 and appropriate Zn(II) amine complex: \([\text{Zn}(\text{dien})_2]\text{(OH)}_2\), \([\text{Zn}(\text{NH}_3)_4]\text{(OH)}_2\), and \([\text{Zn}(\text{pn})_3]\text{(OH)}_2\). The hexaborate(2−) anions in 1 and 2 are coordinated to octahedral Zn(II) centres as tridentate (1) or bidentate ligands (2) and the dodecaborate(6−) ligand in 3 is tridentate to a tetrahedral Zn(II) centre.

Keywords: dodecaborate(6−); hexaborate(2−); oxidoborate; polyborate; self-assembly; X-ray structure; zinc(II) complex

1. Introduction

There are more than two hundred known borate (polyborate) minerals, and many more known synthetic polyborates [1–3]. Borates are generally comprised of cationic moieties partnered with anionic units containing boron, oxygen, and in many cases hydroxyl hydrogen. Oxidoborates (or hydroxoyoxidoborates) are the more appropriate terms, but the term borate (or polyborate) has been used for many years and will be used in this manuscript. Borates are a class of compounds with rich structural diversity [4–7], and have been synthesized by solvothermal methods or from aqueous solution by the addition of B(OH)_3 to a solution containing the appropriate templating cation [7]. Polyborate salts obtained from aqueous solution usually contain discrete, isolated or insular hydroxyl anions, whilst polyborate salts prepared via solvothermal methods are often more condensed and contain anionic polymeric 1-D chains, 2-D layers or 3-D networks with a variety of framework building blocks [1,7]. Salts formed from aqueous solution often contain the pentaborate(1−)
[B₁₀O₂₆(OH)₁₆]⁻ anion since this anion is structurally well suited to forming crystalline supramolecular lattices, which are held together by strong H-bond interactions [8–11]. We have developed a strategy to overcome pentaborate(1−) salt formation by utilizing more highly charged (> (+1)) metal complex cations with ligands having the potential to form multiple H-bond interactions to template crystallization from aqueous solution of polyborate salts of unusual structures. In this context we have isolated two novel polyborate anions: [B₇O₉(OH)₁₀]³⁻ [12] and [B₈O₁₈(OH)₁₆]⁷⁻ [13]. We have also recently started to investigate Zn(II)/polyborate chemistry and have been able to isolate an insular bi-Zn(II) complex containing a rare dodecaborate(6−) anion [14] and two polymeric 1-D coordination chains with hexaborate(2−) ligands bridging Zn(II) centres [15]. There are a number of other structural reports on polyborate/Zn(II) chemistry [16–23], including the industrially important Zn[B₃O₄(OH)]₃ [24].

In this manuscript we describe the synthesis and XRD structures of two new Zn(II)/hexaborate(2−) complexes: [Zn[k⁵O-B₆O₇(OH)₆]k(N-dien)] 0.5H₂O (dien = NH(CH₂CH₂NH₂)₂) (1) and (NH₄)₂[Zn(k²O-B₆O₇(OH)₆)₂(H₂O)₂]-2H₂O (2). We also report a Zn(II)/dodecaborate(6−) complex (1,3-pnH₂)₃[k¹N-H₃N(CH₂)₃NH₂]Zn[k³O-B₁₂O₁₈(OH)₁₆]₂·14H₂O (1,3-pn = 1,3-diaminopropane) (3). All three complexes are insular and the hexaborate(2−) ligand is tridentate in 1, whereas in 2 it is bidentate to octahedral Zn(II) centres. The dodecaborate(6−) ligand in 3 is tridentate to a tetrahedral Zn(II) centre. The structures of these two anions are drawn schematically in Figure 1.

Figure 1. The (a) hexaborate(2−) anion, [B₆O₇(OH)₆]²−, observed in 1 and 2; and (b) dodecaborate(6−) anion, [B₁₂O₁₈(OH)₁₆]⁷−, observed in 3. These diagrams show the location of formal Lewis charges.

2. Results and Discussion

2.1. Synthesis and Characterization

Compounds 1, 2 and 3 were prepared in moderate yield through crystallization from aqueous solution initially containing B(OH)₃ and [Zn(dien)]²⁺[OH]₂, [Zn(NH₃)₄]⁺[OH]₂ or [Zn(pn)₃]⁺[OH]₂ for 1, 2 and 3, respectively. The hydroxide salts were prepared in situ from the corresponding sulphate salts by the addition of Ba(OH)₂ and removal of precipitated BaSO₄ (Scheme 1).
Compounds 1, 2 and 3 are formed through self-assembly processes. B(OH)₃, when dissolved in aqueous solution at moderate to high pH, exists not as boric acid but as a dynamic combinatorial library (DCL) [25,26] of a variety of polyborate anions which are in rapid equilibria [27,28]. Likewise, Zn(II) complexes are labile [29], and a DCL of Zn(II)/amine species are also present in the solution. The products crystallize from solution maximizing energetically favourable solid-state interactions, including coordination bonds, Coulombic attractions, H-bonding and steric effects [30,31].

Compounds 1, 2 and 3 were characterized spectroscopically (NMR and IR), by thermal DSC/TGA analysis and by single-crystal XRD studies (Section 2.2). They all gave satisfactory bulk elemental analysis.

The thermal TGA/DSC data obtained for 1–3 (see Supplementary Materials) were consistent with the structures determined by single-crystal X-ray diffraction studies (see below) and can be interpreted by multi-step decomposition processes. For 1 this involved loss of interstitial water (<190 °C), further loss of water with cross-condensation of hexaborate(2–) ligands (190–380 °C) and finally oxidation and/or evaporation of the organic dien ligand (380–650 °C) to leave an anhydrous zinc borate ZnB₁₂O₁₉ (= ZnO·3B₂O₃) as a glassy residue. Glassy solids with masses consistent with ZnB₁₂O₁₉ (= ZnO·6B₂O₃) were obtained as the final residues for both 2 and 3 since the initial starting Zn/B ratio was 1:12. The thermal decomposition of 3 followed a similar pattern to 1. Compound 2 had a TGA trace consistent with loss of initial interstitial water (<110 °C), loss of ammonia (110–250 °C), and final condensation of hexaborate(2–) anions (250–500 °C). Similar thermal behaviour has been observed in other metal polyborate species [12,13,24,32–35], including 1-D zinc hexaborate(1–) coordination polymers [Zn(en)][B₆O₇(OH)₆]·2H₂O and [Zn(pn)[B₆O₇(OH)₆]]·1.5H₂O [15]. Magnetic susceptibility χₘ data for 1–3 were ∼ −200 × 10⁻⁶ cm³·mol⁻¹ and typical for diamagnetic zinc(II) complexes.

IR spectra can be used to characterize polyborate species since characteristic B–O stretches are generally strong and often diagnostic [36]. Hexaborate(2–) ions, which are never “isolated” and usually found coordinated tridentate to metal centres, have been reported to show such bands at ∼953(m) cm⁻¹ and 808(s) cm⁻¹. Compound 1 displayed bands at 950(m), 861(m) and 806(s) whilst 2 showed bands at 953(m), 904 (s) and 857(m). Thus, the strong band usually observed at 808 cm⁻¹ was absent in 2 and replaced by a strong band at 904 cm⁻¹. This may be a reflection on the unusual centrosymmetric bidentate hexaborate(2–) coordination mode observed in 2. The IR spectrum of 3 showed peaks at 1047(s), 952(m), 902(s) and 855(m), and there were corresponding absorptions in the reported spectrum of [(H₃NCH₂CH₂NH₂)Zn[B₁₂O₁₈(OH)₆]Zn(en)(NH₂CH₂CH₂NH₃)]·8H₂O [14], which also contains a coordinated dodecaborate(6–) ion. Possible diagnostic absorption bands for this anion have not been described before.
Compounds 1–3 were all insoluble in organic solvents but “dissolved” with decomposition in aqueous solution. $^1$H, $^{11}$B spectra of these solutions were obtained in D$_2$O, as were the $^{13}$C spectra of 1 and 3. The $^1$H and $^{13}$C spectra showed peaks consistent with the organics present and the $^1$H spectra additionally displayed at H$_2$O/exchangeable hydrogen peak (H$_2$O, NH, BOH) at ~4.8 ppm. $^{11}$B spectra of 1–3 all showed a single signal at a +17.4, +15.9 and +14.0 ppm, respectively. These signals are all downfield of those calculated [10] (at infinite dilution) for the boron/charge ratio of three (+13.8) for a hexaborate(2$^-$) system, and two (+11.0) for the dodecaborate(6$^-$) ions. This assumes fast B(OH)$_3$/[B(OH)$_4$]$^-$ exchange [27,28] and is also associated with the pH of the solution. The influence of the zinc(II) ions may also be important here by reducing the effective charge at boron.

2.2. X-ray Diffraction Studies

The structures of 1, 2 and 3 were determined by single-crystal XRD methods. Crystal data are given in the experimental section and all XRD data are available as Supplementary Materials.

Compounds 1 and 2 both contained the hexaborate(2$^-$) anion coordinated to a Zn(II) centre and the structures of 1 and 2, showing their atomic numbering schemes, are shown in Figures 2 and 3, respectively. The anionic complex in 2 was centrosymmetric with the asymmetric unit comprising of half the anion with the zinc(2+) ion on the inversion centre. Compound 1 was a neutral zinc(II) complex with 0.5 waters of crystallization. The neutral Zn(II) complex, [Zn[B$_6$O$_{7}$OH$_6$](dien)], contained a tridentate ($\kappa^3$N) dien ligand and a tridentate ($\kappa^3$O) hexaborate(2$^-$) ligand. Compound 1 was disordered with two heavy atoms (O10, C4) of the ligand, and associated hydrogen atoms, split in a 1:1 ratio. One position also had an associated water of crystallization (O21). Compound 2 was a salt comprised of [NH$_4$]$^+$ cations, [Zn[B$_6$O$_7$(OH)$_6$](H$_2$O)$_2$]$^{2-}$ anions and interstitial H$_2$O molecules. Both hexaborate(2$^-$) ligands in 2 were bidentate ($\kappa^2$O) and the coordinated H$_2$O molecules were trans. The Zn–O (hexaborate) distances in 2 (2.0692(9) Å (O11) and 2.1208(9) Å (O12)) were within the range of distances observed for 1 (2.0612(11)–2.1864(10) Å) despite the change in coordination mode of the hexaborate(2$^-$) ligand. The Zn–O (H$_2$O) distance in 2 was 2.1292(9) Å (O21), and the three Zn–N (dien) distances in 1 ranged from 2.1283(14)–2.1473(15) Å. The angles about the Zn(II) centres were 82.56(5)–100.26(5)$^\circ$ and 166.45(5)–175.22(5)$^\circ$ for 1, and 87.90(3)–92.10(3)$^\circ$ and 180.00$^\circ$ for 2. These angles and distances were consistent with previous reported octahedral complexes of Zn(II) with O and N donor ligands [37]. Bond lengths (B–O) and OBO and BOB bond angles associated with the hexaborate(2$^-$) ligands in both 1 and 2 were very similar. For example, bond lengths to the central pyramidal O$^+$ (1.5154(18)–1.5231(18) Å, 1; 1.5053(15)–1.5247(16) Å, 2) > other bond lengths to four coordinate borons (1.4407(19)–1.4791(19) Å, 1; 1.4413(18)–1.4889(15) Å, 2) > bond-lengths to three coordinate borons (1.362(2)–1.418(4) Å, 1; 1.3570(17)–1.3793(17) Å, 2) and consistent with distances and angles previously reported specifically for hexaborate(2$^-$) complexes [15,32,38,39] and related polyborate systems [8–24,32–36,38–40].
Figure 2. Molecular structure of $[\text{Zn}\{\kappa^3\text{O-B}_6\text{O}_7(\text{OH})_6}\{\kappa^3\text{N-dien}\}] \cdot 0.5\text{H}_2\text{O}$ (dien = NH(CH$_2$CH$_2$NH$_2$)$_2$) (1) showing atomic labelling.

Figure 3. Molecular structure of the asymmetric unit of $(\text{NH}_4)_2[\text{Zn}\{\kappa^2\text{O-B}_6\text{O}_7(\text{OH})_6\}_2(\text{H}_2\text{O})_2] \cdot 2\text{H}_2\text{O}$ (2), showing atomic labelling.
H-bonding interactions are commonly observed in most polyborate solid-state structures. They were observed at many locations in the solid-state structures of 1 and 2 and must be partly responsible for the self-assembly of these structures from their constituents. Compound 1 showed H-bond interactions between the neutral complexes as well as these complexes and the water of crystallization. Compound 2 showed H-bond cation/anion and anion/H2O interactions. The energetically favourable reciprocal R2(8) (Etter [41] nomenclature) O8H8→O3*, O8*H8*→O3) linked hexaborate(2−) units in 1. There were also unusual R2(6) (O9H9→O12*H12*→O4) and R2(8) (N2H2→O8* and O13H13→O2*) arrangements between neighbouring hexaborate units in 1; the latter ring included Zn(1). Compound 2 also had two energetically favourable reciprocal R2(8) interactions between neighbouring hexaborate(2−) units (O13H13→O6*, O13*H13*→O6 and O8H8→O3*,O8*H8*→O3). There was also an unusual intramolecular H-bond in 2 between the coordinated H2O molecule and the hexaborate(2−) ligand (O21H21A→O13) as part of an intramolecular R1(8) system incorporating the Zn1 centre (Figure 4). The coordinated H2O also H-bonded to a neighbouring hexaborate O21H21B→O2*. O13 is the hexaborate hydroxyl oxygen atom that fulfilled the role as third coordination donor atom in 1 and in other tridentate hexaborate complexes. In this particular local environment of 2, the energetics of forming this H-bond and the H2O–Zn coordination bond must outweigh the energetics of a simple borate O–Zn coordinate bond. O13H13 also H-bonded to a neighbouring hexaborate (O13H13→O6*). Full details of these H-bond interactions are given in the Supplementary Materials.

Figure 4. The intramolecular O21H21A→O13 H-bond interaction in 2, d(O21–H21) 0.87 Å, d(H21–O13) 1.79 Å; d(O21–O13) 2.6446(13) Å; angle O21H21O13, 169.7°] which is part of two R1(8) rings, incorporating Zn–O coordinate bonds (symmetry i = 2 − x, 1 − y, 2 − z).

Compound 3 was an ionic compound comprised of [H3N(CH2)3NH3]2+ cations and [(H3N(CH2)3NH2)ZnB12O18(OH)6]3− anions, with the anions containing the dodecaborate(6−) ligand coordinated κ3O to a tetrahedral Zn(II) centre which also had a monoprotonated monodentate
k$^3$N-H$_3$N(CH$_2$)$_3$NH$_2$ ligand. There were also seven waters of crystallization per Zn(II) centre. A diagram of the structure is shown in Figure 5.

![Diagram of structure](image)

Figure 5. Diagram of $(1,3$-pn)$_2$[(k$^3$N-H$_3$N(CH$_2$)$_3$NH$_2$)Zn(k$^3$O-B$_{12}$O$_{18}$(OH)$_6$)]$_2$·4H$_2$O (1,3-pn = 1,3-diaminopropane) (3) showing atomic labelling.

The Zn–O (dodecaborate) distances in 3 [1.9592(18) Å (O3)–1.9717(18) Å (O1)] were shorter than those observed for 1 or 2, reflecting tetrahedral vs. octahedral coordination geometries. The Zn1N1 distance was 2.006(2) Å, and internuclear angles about Zn1 ranged from 103.43(7)–117.86(9)$^\circ$. These data are very similar to those of the closely related di-Zn(II) complex [(NH$_3$CH$_2$CH$_2$NH$_2$)Zn(B$_{12}$O$_{18}$(OH)$_6$)[Zn(en)(NH$_2$CH$_2$CH$_2$NH$_3$)] [14] that features 1,2-diaminoethane. The dodecaborate(6−) anion (Figure 1b) is comprised of six boroxole rings fused so as to produce a larger central [B$_6$O$_6$] ring, with each boron atom within this ring carrying a formal negative charge due to their four-coordinate nature. This anion was first reported in 1990 in the structure of Ag$_6$[B$_{12}$O$_{18}$(OH)$_6$]·3H$_2$O [42]. The dodecaborate(6−) anion in 3 is closely related to the deprotonated structures found in Na$_8$[B$_{12}$O$_{20}$(OH)$_4$] [43] and Zn$_6$[B$_{12}$O$_{24}$] [44]. The central ring oxygen atoms alternate up and down on different sides of the central ring and are ideally set-up to bind tridentate to a metal centre. The dodecaborate(6−) anion has been previously observed to coordinate in a tridentate mode in the following compounds: [(NH$_3$CH$_2$CH$_2$NH$_2$)Zn(B$_{12}$O$_{18}$(OH)$_6$)-Zn(en)(NH$_2$CH$_2$CH$_2$NH$_3$)] [14], Na$_2$Cs$_4$Ba$_2$[B$_{12}$O$_{18}$(OH)$_6$](OH)$_4$ [45], K$_7$[(BO$_3$)Mn[B$_{12}$I$_{18}$(OH)$_6$]]·H$_2$O [46] and K$_7$[(BO$_3$)Zn[B$_{12}$O$_{18}$(OH)$_6$]]·H$_2$O [47].
The six four-coordinate boron atoms had B–O distances ranging between 1.441(3)–1.506(3) Å and their O–B–O angles ranged from 106.3(2)–112.1(2)°. The remaining six borons of the anion were three-coordinate and had significantly shorter B–O distances 1.353(1)–1.386(3) Å and larger O–B–O angles 115.4(2)–123.1(2)°. These bond lengths are similar to those observed for \( ([\text{NH}_3\text{CH}_2\text{CH}_2\text{NH}_2]_2\text{Zn(B}_2\text{O}_1\text{OH}_6)_6\text{Zn(en)}(\text{NH}_2\text{CH}_2\text{CH}_2\text{NH}_3]) \) [14], other similarly fused boroxole systems [14,45–47] and the hexaborate(2–) complexes 1 and 2.

The hydroxyl hydrogen atom, the amino hydrogen atoms of the protonated 1,3-diaminopropane ions and ligands and the waters of crystallization form numerous H-bond interactions and they were presumably responsible—at least in part—for this remarkable self-assembly from mono-boron species. There are numerous cation/anion H-bond interactions, and three of the six potential dodecaborate hydroxyl interactions are \( \text{R}_2\text{O}(8) \): \( \text{O}2\text{H}2\text{O} \rightarrow \text{O}15\text{*}, \text{O}2\text{H}2\text{3} \rightarrow \text{O}9\text{*} \) and \( \text{O}2\text{H}2\text{4} \rightarrow \text{O}17\text{*} \), with only the latter reciprocal. “Simple” inter-borate H-bonds originate from \( \text{O}19\text{H}19 \) and \( \text{O}22\text{H}22 \) whilst \( \text{O}21\text{H}21 \) has a non-borate interaction and H-bonds to an \( \text{H}_2\text{O} \) (\( \text{O}31 \)). This configuration with only the latter reciprocal. “Simple” inter-borate H-bonds originate from \( \text{O}19\text{H}19 \) and \( \text{O}22\text{H}22 \) whilst \( \text{O}21\text{H}21 \) has a non-borate interaction and H-bonds to an \( \text{H}_2\text{O} \) (\( \text{O}31 \)). This configuration contrasts with that of \( ([\text{NH}_3\text{CH}_2\text{CH}_2\text{NH}_2]_2\text{Zn(B}_2\text{O}_1\text{OH}_6)_6\text{Zn(en)}(\text{NH}_2\text{CH}_2\text{CH}_2\text{NH}_3]) \) where all six were involved in \( \text{R}_2\text{O}(8) \) interactions. However, a structural motif that is similar to that found in \( ([\text{NH}_3\text{CH}_2\text{CH}_2\text{NH}_2]_2\text{Zn(B}_2\text{O}_1\text{OH}_6)_6\text{Zn(en)}(\text{NH}_2\text{CH}_2\text{CH}_2\text{NH}_3]) \) is that amino hydrogen atoms of the uncoordinated nitrogen (\( \text{N}2 \)) of the protonated 1,3-diaminopropane contrasts with that of \( ([\text{NH}_3\text{CH}_2\text{CH}_2\text{NH}_2]_2\text{Zn(B}_2\text{O}_1\text{OH}_6)_6\text{Zn(en)}(\text{NH}_2\text{CH}_2\text{CH}_2\text{NH}_3]) \) the hexaborate(2–) complexes 1 and 2. Full details of these H-bond interactions are in the Supplementary Materials.

### 3. Experimental

#### 3.1. General

All chemicals were obtained from commercial sources. Combustion analysis (CHN) were obtained from OEA laboratories Ltd. in Callington, Cornwall, UK. NMR spectra were obtained on a Bruker Avance spectrometer (Bruker, Coventry, UK) in \( \text{D}_2\text{O} \) at 400.1 MHz (\( ^1\text{H} \)), 100.6 MHz (\( ^13\text{C} \)) or 128.4 MHz (\( ^13\text{B} \)) with data reported as \( \delta \) (ppm) with positive chemical shifts to a high frequency of tetramethylsilane (TMS) (\( ^1\text{H} = 0.0, ^13\text{C} = 29.5, ^13\text{B} = 19.9 \)) and BF\(_3\)-OEt\(_2\) (\( ^13\text{B} = 19.9 \)). FTIR spectra were obtained on a PerkinElmer 100 FTIR spectrometer (PerkinElmer, Seer Green, UK) as KBr pellets. TGA/DSC analyses were undertaken in air on an SDT Q600 V4.1 Build 59 instrument (New Castle, DE, USA), using \( \text{Al}_2\text{O}_3 \) crucibles between 10–800 °C with a ramp temperature rate of 10 °C·min\(^{-1}\).

#### 3.2. Synthesis, Spectroscopic, Analytical and Crystallographic data for 1

A solution of \( \text{NH}((\text{CH}_3\text{CH}_2)\text{NH}_2)_2 \) (dien) (2.16 mL, 20 mmol) in \( \text{H}_2\text{O} \) (5 mL) was added to a solution of \( \text{ZnSO}_4 \cdot 7\text{H}_2\text{O} \) (1.79 g, 10 mmol) in \( \text{H}_2\text{O} \) (10 mL). The reaction mixture was stirred at room temperature for 60 min before the addition of \( \text{Ba(OH)}_2 \cdot 8\text{H}_2\text{O} \) (3.15 g, 10 mmol) in \( \text{H}_2\text{O} \) (25 mL). This mixture was rapidly stirred for a further 30 min. The white precipitate of \( \text{BaSO}_4 \) was removed by filtration and \( \text{B(OH)}_3 \) (6.18 g, 10 mmol) dissolved in \( \text{H}_2\text{O} \) (50 mL) was added to the filtrate, which was further stirred at room temperature for 3 h. The volume of this solution was reduced to 20 mL by gentle evaporation in a warm water bath. The concentrated solution was left for 10 days in NMR tubes for crystallization and yielded colourless crystals of \( [\text{Zn(dien)}]_2[\text{B}_6\text{O}_7(\text{OH})_6] \)·0.46\( \text{H}_2\text{O} \) (1) (1.9 g, 42%). Mp ≥ 300 °C. Anal. Calc.: C = 10.5%, H = 4.4%, N = 9.2%. Found: C = 10.7%, H = 4.1%, N = 9.3%. NMR. \( ^1\text{H} \)/ppm: 2.5 (m, 8H), 4.8 (s, 37H, \( \text{NH}_2 \), \( \text{H}_2\text{O} \), \( \text{OH} \)). \( ^13\text{C} \)/ppm: 38.10. \( ^13\text{B} \)/ppm: 17.4. IR (KBr/cm\(^{-1}\)): 3549(s), 3384(s), 1642(m), 1424(s), 1427(s), 1362(s), 1249(m), 1193(s) 1108(s), 1028(s), 951(m), 861(m), 808(m). TGA: 100–190 °C, loss of 0.46 interstitial \( \text{H}_2\text{O} \) 2.5 (1.8 calc.); 190–380 °C, condensation of polyborate with loss of three further \( \text{H}_2\text{O} \) 15.2% (13.7% calc.); 380–650 °C, oxidation of dien 38.5% (36.3% calc.); residual \( \text{ZnB}_6\text{O}_{10} \) 61.5% (63.4% calc.). Magnetic susceptibility: \( \chi_m = -210 \times 10^{-6} \text{cm}^3\text{mol}^{-1} \).

Crystal data: \( C_{11}\text{H}_{19}\text{B}_5\text{N}_3\text{Zn}_3 \), \( M_r = 456.46 \), monoclinic, \( C2/c \) (No. 15), \( a = 26.0212(3) \) Å, \( b = 9.15620(10) \) Å, \( c = 13.6318(2) \) Å, \( \beta = 99.5800(10)^\circ \), \( \alpha = \gamma = 90^\circ \), \( V = 3202.55(7) \) Å\(^3\), \( T = 100(2) \) K, \( Z = 8 \),
3.3. Synthesis, Spectroscopic, Analytical and Crystallographic Data for 2

A solution of NH₃ (35%, 2.4 mL, 36 mmol) was added dropwise to a solution of ZnSO₄·H₂O (1.08 g, 6 mmol) in H₂O (15 mL). The addition of Ba(OH)₂·8H₂O (1.89 g, 6 mmol) in H₂O (35 mL) followed by rapid stirring for 15 min resulted in a precipitate of BaSO₄ which was removed by filtration. B(OH)₃ (3.71 g, 60 mmol) dissolved in H₂O (30 mL) was added to the filtrate which was further stirred at room temperature for 30 min. The volume of this solution was reduced to 5 mL by gentle evaporation on a warm water bath and the concentrated solution was left for 3 days in NMR tubes for crystallization and yielded colourless crystals of [NH₄]₂[Zn(B₃O₆)(OH)₆]₂(H₂O)₂·2H₂O (2) (2.1 g, 48%). Mp ≥ 300 °C. Anal. Calc.: H = 3.8%, N = 3.8%. Found: H = 4.0%, N = 3.7%. NMR: 1H/ppm: 15.9, IR (KBr/cm⁻¹): 3212(s), 1400(s), 1357(s), 1048(s), 953(m), 904(m), 857(m). TGA: 100–110 °C, loss of 4 interstitial/coordinated H₂O 10.2% (9.9% calc.); 110–250 °C, loss of 2 NH₃ 15.5% (14.8% calc.); 250–500 °C, condensation of polyborate with loss of six further H₂O 31.1 (29.6 calc.); residual ZnB₆O₁₉ 68.9% (68.2% calc.). Magnetic susceptibility: χₘ = −290 × 10⁻⁶ cm³·mol⁻¹⁻¹.

Crystal data: B₄H₆Na₂O₃Zn, Mᵣ = 731.33, triclinic, P–1 (No. 2), a = 7.4831(2) Å, b = 7.8551(2) Å, c = 11.0111(3) Å, α = 108.065(2)°, β = 95.020(2)°, γ = 90.118(2)°, V = 612.68(3) Å³, T = 100(2) K, Z = 1, Z′ = 0.5, χ(Mo Kα) = 1.138 mm⁻¹, 16475 reflections measured, 2799 unique (Rint = 0.0314) which were used in all calculations. The final wR₂ was 0.0559 (all data) and R₁ was 0.0212 (I > 2σ(I)).

3.4. Synthesis, Spectroscopic, Analytical and Crystallographic Data for 3

A solution of NH₂CH₂CH₂CH₂NH₂ (1,3-pn) (2.52 mL, 30 mmol) in H₂O (10 mL) was added to a solution of ZnSO₄·H₂O (1.79 g, 10 mmol) in H₂O (10 mL). The reaction mixture was stirred at room temperature for 60 min before the addition of Ba(OH)₂·8H₂O (3.15 g, 10 mmol) in H₂O (25 mL). This mixture was rapidly stirred for a further 30 min. The white precipitate of BaSO₄ was removed by filtration and B(OH)₃ (6.18 g, 10 mmol) dissolved in H₂O (50 mL) was added to the filtrate, which was further stirred at room temperature for 30 min. The volume of this solution was reduced to 5 mL by gentle evaporation in a warm water bath. The product was collected by filtration and carefully washed with cold H₂O followed by CH₃COCH₃ and then dried at 40 °C for 1 h to yield colourless crystals of [H₃N(CH₂)₃N₃][H₃N(CH₂)₃N₃][H₂N(CH₂)₃N₂ZnB₁₂O₁₈(OH)₁₂]·4H₂O (3) (4.1 g, 46%). Mp ≥ 300 °C. Anal. Calc.: C = 10.0%, H = 5.9%, N = 7.8%. Found: C = 9.7%, H = 5.2%, N = 7.8%. NMR: 1H/ppm: 1.93 (p, 10H, CH₂), 3.01 (t, 20H, CH₂) 4.8 (s, 68H, NH₂), 6.9 (60 calc.). p-XRD: d-spacing (Å)/(% rel. int.): 9.98(36), 9.44 (100), 8.50 (54), 8.08 (35), 6.93 (43). Magnetic susceptibility: χₘ = −180 × 10⁻⁶ cm³·mol⁻¹⁻¹.

Crystal data: C₇H₄O₄B₁₂ZnO₃Zn, Mᵣ = 900.60, triclinic, P–1 (No. 2), a = 9.3681(2) Å, b = 10.6910(2) Å, c = 19.2746(4) Å, α = 82.954(2)°, β = 76.156(2)°, γ = 68.655(2)°, V = 1744.44(7) Å³, T = 100(2) K, Z = 2, Z′ = 1, χ(Mo Kα) = 0.821 mm⁻¹, 38,867 reflections measured, 7958 unique (Rint = 0.0389) which were used in all calculations. The final wR₂ was 0.1053 (all data) and R₁ was 0.0425 (I > 2σ(I)).

3.5. X-ray Crystallography

Single-crystal X-ray crystallography was undertaken at the Engineering and Physical Sciences Research Council (EPSRC) National Crystallography service at the University of Southampton, (Southampton, UK). Suitable crystals of 1, 2 and 3 were selected and mounted on a MITIGEN holder in perfluorooether oil on a Rigaku FRE+ equipped with HF Varimax confocal mirrors and an AFC12 goniometer and HG Saturn 724+ detector diffractometer. The crystals were kept at T = 100(2) K during data collection. Using Olex2 [48], the structures were solved with the ShelXT [49] structure solution.
program using the Intrinsic Phasing solution method. The models were then refined with ShelXL \cite{50} using least squares minimisation. Cambridge Crystallographic Data Centre (CCDC) 1898912 (1), 1898913 (2), 1898914 (3) contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via http://www.ccdc.cam.ac.uk/conts/retrieving.html (or from CCDC, 12 Union Road, Cambridge CB2 1EZ, UK; Fax: +44 1223 336033; email deposit@ccdc.cam.ac.uk).

4. Conclusions

The strategy of using more highly charged cationic labile transition-metal complexes to template self-assembly (by crystallization) of polyborate anions from alkaline aqueous solutions originally containing $\text{B(OH)}_3$ has resulted in the synthesis of three new zinc polyborate complexes in moderate yields (40–50%). These complexes contain either hexaborate($2^-$) or dodecaborate($6^-$) ligands and are stabilized by Zn–O coordinate bonds. The solid-state structures are further stabilized by multiple intramolecular and/or intermolecular H-bond interactions which are prevalent in polyborate structures.

Supplementary Materials: The following are available online at http://www.mdpi.com/2304-6740/7/4/44/s1. TGA and single-crystal XRD data. Cif and checkcif files.

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.

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