Mutual stabilisation between $M^{II}4L_6$ tetrahedra and $M^{II}X_4^{2-}$ metallate guests

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A complex host–guest equilibrium employing metal ions incorporated into both the host and guest is discussed. $M^{II}X_4^{2-}$ metallate guests are shown to provide a good size and shape match for encapsulation within the $M_4L_6$ tetrahedral capsules, facilitating the generation of previously unreported Zn$_4$L$_6$ complexes. Displacement of the initial, primary template anion (ZnBr$_4^{2-}$) by a secondary template anion (ClO$_4^{-}$) is shown to result in the formation of a pentagonal-prismatic Zn$_{10}$L$_{15}$ structure that incorporates both Br$^{-}$ and ClO$_4^{-}$. Furthermore, the formation of heterometallic complexes provides direct evidence for metal exchange between the guest and host complex.

Introduction

A variety of complex three-dimensional supramolecular architectures have been templated using non-coordinating anions bound in internal binding pockets. Such anions may be present either as the counterion of the metal salt, or added externally. The dynamic nature of host–guest complexes, which enables their formation, can facilitate subsequent structural rearrangements driven by replacement of one anion for another. Building upon these foundations, here we show that complex $M^{II}X_n^{2-}$ anions may serve as templates for intricate three-dimensional structures. These anions are constructed from the same dicaticonic first-row transition-metal ions that form the vertices of the host framework, and they are generated during the same equilibration process that generates the host. Their use as structural templates provides an added layer of complexity in the synthesis of metal–organic capsules.

Subcomponent self-assembly, where ligands and complex come together during the same overall process, has proven useful in the preparation of a variety of new materials and functional structures. We have previously reported its use in the preparation of a diverse range of discrete three-dimensional architectures starting with $p$-toluidine and 6,6’-diformyl-3,3’-bipyridine subcomponents, and cobalt(ii) or zinc(ii) salts (NO$_3^{-}$, BF$_4^{-}$, ClO$_4^{-}$ or CF$_3$SO$_3^{-}$) as templates. In contrast, when cobalt(ii) or zinc(ii) bis(trifluoromethane)sulfonimide (triflimide, NTf$_2^{-}$) is employed, a dynamic library (DL) is observed because the triflimide anion is too large to serve as a template. Herein we discuss addition of halide or pseudohalide anions Cl$^{-}$, Br$^{-}$, I$^{-}$, OCN$^{-}$ or SCN$^{-}$ to Zn(NTf$_2$)$_2$- or Co(NTf$_2$)$_2$-containing DL’s. These reactions result in the in situ generation of complex metallate anions, which serve as optimal templates for $M_4L_6$ tetrahedra. Structural interconversion between a Zn$_4$L$_6$ tetrahedron, Zn-1, containing a centrally bound dianionic guest, and a larger Zn$_{10}$L$_{15}$ pentagonal architecture, Zn-2, templated by five mono-anionic ClO$_4^{-}$ anions was also demonstrated.

Results and discussion

Synthesis of $[MX_4]^{2-}$ $\subset M_4L_6$ complexes

When the subcomponents of $L$ (Scheme 1) were mixed with Co$^{II}$(NTf$_2$)$_2$ (6 equiv. $L$ per 4 equiv. Co$^{II}$), a broad $^1$H NMR spectrum characteristic of a dynamic library of different complexes showing both of the equilibration processes necessary for complex formation: (i) formation of the metallate template and; (ii) self-assembly of the ligand $L$. 

Scheme 1 Subcomponent self-assembly of $[MX_4]^{2-}$ $\subset M_4L_6$ complexes showing both of the equilibration processes necessary for complex formation; (i) formation of the metallate template and; (ii) self-assembly of the ligand $L$. 

$M^{II}$ = Co$^{II}$, Zn$^{II}$
$\times$ = Cl$^{-}$, Br$^{-}$, I$^{-}$, OCN$^{-}$, SCN$^{-}$
coordination complexes (Co-DL) was observed. Following the addition of KBr (4 equiv. per 5 equiv. Co) to Co-DL, the 1H NMR spectrum displayed a single sharp resonance per ligand proton environment (Fig. S1†). ESI-MS analysis provided evidence for a tetraboromocobaltate [CoBr4]2− adduct of 1 (Fig. S2†). No evidence for a simple bromide adduct was obtained when less KBr was employed, suggesting that [CoBr4]2− provides an optimal size, shape and Coulombic fit for the cavity of 1.

Further evidence for formation of [CoBr4]2− was obtained through UV-Vis spectroscopy, which displayed characteristic absorbance bands in the 500–800 nm window. The complex absorbance profile is indicative of the multiple accessible electronic excited states which are available to tetrahedral cobalt species,43 and corresponds well to the absorbance of the [CoIIBr4]2− dianion in the absence of the Co4L6 tetrahedral cage (Fig. S3†).

Unambiguous structural characterisation of the host–guest complex was obtained through single-crystal X-ray analysis (Fig. 1a and b). The structure of [CoBr4]2− ⊂ Co-I was solved in cubic space group F4,32 with one twelfth of the T-symmetric molecule in the asymmetric unit. The encapsulated [CoBr4]2− anion thus occupies a special position of tetrahedral symmetry. Its Co–Br bond length of 2.424(1) Å is in good agreement with the average value of 2.40(2) Å found for [CoBr4]2− anions reported in the Cambridge Structural Database.46

A variety of other dianions were previously investigated as potential guest molecules for this metal–ligand system, however, no evidence was obtained for the formation of a host–guest complex upon addition of Na2SO4, Na2SeO4, Na2TeO4, Na2MoO4, (NH4)2SnF6, or K2CrO4.31 The increased energy of desolvation associated with dianions relative to their monomeric counterparts generally makes their encapsulation less energetically favourable.34,37–39 Thus [CoBr4]2− is both the first dianionic guest to be bound within Co-I, and the largest guest to be accommodated within its cavity (Scheme 1).

The tight fit of [CoBr4]2− within the tetrahedral host is apparent from Fig. 1b, as is the complimentary shape of the host–guest pair.38 The centrally bound tetrahedral guest adopts a complementary orientation to that of the CoII4L6 tetrahedron, wherein each bromide points directly out of an open face of the capsule. In contrast, when the octahedral anion SbF6− was bound within Co-I, not all of the fluorine atoms can orient towards an open face of the tetrahedron (Fig. 1c). The shape match between [CoBr4]2− and 1 thus facilitates accommodation of this large (115 Å3) anion within Co-I. Further evidence of a good fit is provided by the observation that the Co–Co distances (9.763(1) Å of host 1 in [CoBr4]2− ⊂ Co-I were not observed to lengthen from the 9.681(3)–9.823(3) Å and 9.753(3)–9.863(3) Å Co–Co distances observed for the SbF6− and MeCN inclusion complexes, respectively. This observation indicates that the increased size of the dianionic template does not affect the size of the central binding pocket, leading us to hypothesise that other tetrahedral species, which have longer arms that are capable of protruding through the faces of the capsule,41–43 might prove suitable templates for Co-I.

Our hypothesis was tested using the pseudohalide OCN−, which displays similar reactivity to Br−, and is known to generate the [CoII(NCO)4]2− dianion.44,45 The addition of KOCN (0.4 equiv.) to Co-DL followed by sonication and heating at 323 K for 24 hours yielded a 1H NMR profile consistent with the formation of a tetrahedral capsule. The broad 1H NMR profile associated with the mixture of species present in the Co-DL complex was simplified to show one set of resonances per ligand proton, as was observed with KBr. ESI-MS also confirmed formation of the [CoII(NCO)4]2− ⊂ Co-I complex (Fig. S5†).

The ability to generate host–guest complexes such as [CoII(NCO)4]2− ⊂ Co-I, where the guest molecule occupies the centre of the capsule but extends through the open faces of the host expands the potential range of applications for these metal–organic cage molecules. Most importantly this property allows host–guest complexes to be generated using guest molecules that are larger than the volume of the internal binding pocket, a property that restricts potential application of these capsules.43

In common with cobalt(n) complexes, zinc(n) complexes can be either four- or six-coordinate. Furthermore, no crystal field stabilisation effects are observed for zinc(n) complexes due to their d10 electronic configuration, implying that a wider range of ligands might be employed to generate the tetrahalozincate

![Fig. 1](image-url)
inclusion complexes. The diamagnetic nature of this series of complexes also allows more subtle changes in the chemical environments of the host–guest complexes to be probed by $^1$H NMR (Fig. 2).

As with Co$^{II}$, when a range of halide and pseudohalide anions were added directly to the Zn-DL generated from the subcomponents of L and Zn(NTs)$_2$, the complex $^1$H NMR associated with the mixture of species in the dynamic library simplified to give a well defined $^1$H NMR profile displaying one resonance per ligand proton, consistent with the formation of Zn$_4$L$_6$ tetrahedral complexes (Fig. 2). The $^1$H NMR resonance corresponding to H$_E$ (Fig. 2), which is directed across the open face of the Zn$_4$L$_6$ complexes, showed the greatest variation in chemical shift upon guest binding. For the series of halide anions, Cl$^-$, Br$^-$ and I$^-$, a significant downfield shift of H$_E$ relative to SbF$_6^-$ < Zn-1 was observed. This observation is consistent with increasing contact between the larger centrally-bound anions and H$_E$. In contrast, the octahedral SbF$_6^-$ anion cannot align itself effectively with the open faces of the tetrahedron and therefore does not effect a substantial change in the chemical shift of H$_E$.

The $^1$H NMR profiles of the host–guest complexes containing [Zn(NCO)$_4$]$^{2-}$ and [Zn(NCS)$_4$]$^{2-}$ reflect the longer protruding arms of these guest anions, compared with the metalate anions generated from the halide precursors. The H$^E$ resonances for Zn-1 encapsulating [Zn(NCO)$_4$]$^{2-}$ and [Zn(NCS)$_4$]$^{2-}$ are less downfield-shifted compared with those of the Zn-1 complexes generated in the presence of Cl$^-$, Br$^-$ and I$^-$.

Furthermore, the 4,4$'$-bipyridine resonance, H$_I$, experiences a downfield shift which we attribute to interaction of this proton with the protruding isocyanato or isothiocyanato arm (Fig. 3). Formation of the isocyanato and isothiocyanato complex anions is expected to be preferred over the corresponding cyanato and thiocyanato isomers, in accordance with the position of these ligands in the spectrochemical series.$^{44}$

Dynamic rearrangement of [MX$_4$]$^{2-}$ < M$_4$L$_6$ complexes

The observation that Br$^-$ could induce formation of 1 when added to Co-DL is of interest for two reasons: firstly, the ability to selectively bind tetrahalometalate anions over their corresponding halide anions in organic media, particularly chloride, is of great interest in the hydrometallurgy industry.$^{48,49}$ Secondly, Br$^-$ binds within the centre of Co$^{II}$L$_{14}$ pentagonal prism, Co-2,$^{40}$ which self-assembles from p-toluidine, 6,6-diformyl-3,3'-bipyridine and Co$^{III}$ClO$_4$ in acetonitrile.$^{40}$ Similarly, addition of KBr (0.6 equiv.) to the reaction mixture generated from p-toluidine, 6,6-diformyl-3,3'-bipyridine and Zn(ClO$_4$)$_3$ generates a Zn$_{10}$L$_{15}$ complex with five peripherally bound ClO$_4^-$ anions and a centrally bound bromide anion, Br$^-$ < Zn-2. The single-crystal X-ray structure of Br$^-$ < Zn-2 (Fig. 4) indicates that the complex is isostructural to previously reported M$_{10}$L$_{15}$ complexes formed from the same ligand,$^{31,40}$ and to its chloride bound analogue (details of Cl$^-$ < Zn-2 are provided in the ESI†).

No significant differences are observed in the size of the perchlorate-containing peripheral binding pockets of the
the presence of both ClO₄⁻ solution (Fig. S20): Zn⁻⁻⁻⁻¹
no evidence was obtained for formation of [ZnBr₄]²⁻. Under identical conditions, KBr (0.6 equiv.) was added to two portions of Zn⁻⁻⁻⁻¹, one of which contained LiClO₄ (0.8 equiv.). In the absence of perchlorate, the tetrabromozincate anion was generated and acted as a template for tetrahedron Zn⁻⁻⁻⁻¹. In the presence of perchlorate, however, the larger pentagonal prismatic structure, Zn⁻⁻⁻⁻², was formed with a single bromide anion bound at its centre, and no evidence for formation of Zn⁻⁻⁻⁻¹ was observed (Scheme 2). The different pathways followed by the system highlight its finely balanced thermodynamics, and the differential influence of the tetraboromozincate and perchlorate template anions. Increasing the salt concentration of the reaction mixture led to significant precipitation, but after addition of more than 1.3 equiv. of LiClO₄, which is more than four times the amount required to template the M₃₀L₁₅ complex from Zn-DL, to a solution containing solely [ZnBr₄]²⁻, the ratio of Zn⁻⁻⁻⁻¹ : Zn⁻⁻⁻⁻² was 1 : 8. In contrast, increasing the concentration of Br⁻ in the reaction mixture of Br⁻ ⊂ Zn⁻⁻⁻⁻² generated precipitate, and no evidence was obtained for formation of [ZnBr₄]²⁻ ⊂ Zn⁻⁻⁻⁻¹ in solution (Fig. S20†). These results indicate that the pentagonal prismatic structure Zn⁻⁻⁻⁻² is favoured over the tetrahedron, Zn⁻⁻⁻⁻¹, in the presence of both ClO₄⁻ and Br⁻ in solution.

For all of the reactions described herein, no evidence was obtained for association of the M₄L₆ complexes with either simple halide anions, or more than one dianion. ESI-MS of each reaction mixture displayed peaks corresponding to [MX₄]²⁻ ⊂ [M₄L₆]³⁺, where the remaining charge was in all cases balanced by triflimide counterions. The same results were obtained when the preformed anions [CoBr₄]²⁻ and [CoCl₄]²⁻ were added directly to Co-DL.

The self-assembly of heterometallic species was also investigated. ¹H NMR analysis of reaction mixtures generated from addition of [CoBr₄]²⁻ or [CoCl₄]²⁻ anions to Zn-DL indicated several paramagnetically shifted resonances clustered across distinct spectral regions. In contrast to the complex ³¹P NMR, the ESI-MS spectra consisted of well-defined m/z peaks displaying an unusually wide distribution. Taken together these observations led us to postulate that a dynamic library of mixed metal species had been formed with overlapping m/z peaks (Fig. S21–26†). Observation of complexes containing multiple cobalt metal centres indicates that the [CoX₄]²⁻ salts are a suitable source of cobalt(II) for generation of Co₄L₆ tetrahedra in the presence of solubilising counterions.

Conclusions

In conclusion, a family of novel [MX₄]²⁻ ⊂ M₄L₆ host–guest complexes has been identified, extending the limited range of M₄L₆ tetrahedral complexes known to encapsulate dianionic guests. These new [MX₄]²⁻ ⊂ M₄L₆ complexes readily self-assemble from complex dynamic libraries upon addition of a range of halide or pseudohalide anions. Self-assembly of both the host and the guest occurs during the same equilibration process, and neither the host nor guest was observed to be stable in the absence of the other under these conditions. Further investigations into the properties of these capsules indicated that association of more than one dianion per tetrahedral capsule is not favoured for solubility reasons, an observation previously made by Ward and co-workers. An unprecedented symbiosis was thus observed, in which cobalt(II) or zinc(II) exist in a dynamic equilibrium occupying roles either within a tetrahedral metallic guest or as an octahedral metal centre defining the vertices of the host framework. A clear hierarchy was observed in the stability of these structures, whereby the pentagonal prism Zn⁻⁻⁻⁻² was favoured over such tetrahedral structures as [ZnBr₄]²⁻ ⊂ Zn⁻⁻⁻⁻¹. These structures and their dynamic interconversions may have future roles to play in novel hydrometallurgical processes.

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Notes and references
