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# **Inorganic Chemistry**

# Carbonate-Templated Self-Assembly of an Alkylthiolate-Bridged Cadmium Macrocycle

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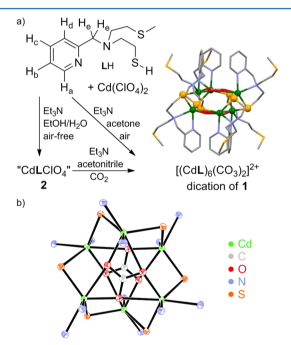
**Supporting Information** 

**ABSTRACT:** In the presence of  $Cd(ClO_4)_2$  and a base, a new mixed N,S-donor alkylthiolate ligand supported both carbonate formation from atmospheric  $CO_2$  and the selfassembly of a novel bicapped puckered  $(CdS)_6$  molecular wheel. The remarkable stability of the complex was demonstrated by slow intermolecular ligand exchange on the <sup>2</sup>*J*(HH) and *J*(<sup>111/113</sup>Cd<sup>1</sup>H) time scales at elevated temperature. Both  $CO_2$  and the base were required to convert amorphous "CdLClO<sub>4</sub>" precipitated in the absence of air to the carbonate complex. The complex shares structural features with the  $\zeta$ -carbonic anhydrase class associating cadmium(II) with the biogeochemical cycling of carbon and is the first structurally characterized carbonate complex of any metal involving an alkylthiolate ligand.

C arbonate is a versatile bridging species found in twodimensional and mono-, di-, tri-, tetra-, hexa-, and enneanuclear and higher complexes with at least 14 different coordination motifs.<sup>1</sup> Several hundred carbonate complexes involving all of the physiologically essential metals and a wide range of nonessential transition metals are structurally characterized.<sup>2</sup> In many cases, precursors to these carbonate complexes are able to hydroxylate atmospheric CO<sub>2</sub>, forming an intermediary species that spontaneously effects CO<sub>2</sub> fixation through combinatorial self-assembly and selective crystallization.<sup>3</sup> Global environmental problems associated with rising greenhouse gas concentrations justify continued investigation of these intriguing processes.

In biological systems, hydration of carbon dioxide  $(CO_2)$  is catalyzed by carbonic anhydrase (CA), a possibly ubiquitous metalloprotein with five zinc(II)-dependent forms across taxonomic kingdoms sharing no significant similarity in their primary sequence or overall structure.<sup>4</sup> Zinc(II) was viewed as an essential cofactor for this enzyme activity until the seemingly anomalous nutrient-like vertical oceanic profile of cadmium was linked to a highly active cadmium(II) carbonic anhydrase (CdCA) variant from the marine diatom *Thalassiosira weissflogii* grown under zinc(II)-limiting conditions, thus establishing the  $\zeta$ form of CA.<sup>5</sup> All six CA forms use invariant histidine residues for metal ligation and related aromatic N donors are quite common in synthetic complexes. Both the  $\beta$  and  $\zeta$  forms of these enzymes also have two invariant cysteine residues bound to the metal, but alkylthiolates have not yet been reported as metal ligands in synthetic carbonate complexes.  $^{\rm 2}$ 

Our interest in multidentate ligands providing simple models of amino acid side-chain donors for zinc triad coordination studies led to the preparation of LH.<sup>6</sup> Herein LH is reported to react with 1 equiv of  $Cd(ClO_4)_2 \cdot 6H_2O$  and excess Et<sub>3</sub>N in airsaturated acetone to form  $[(CdL)_6(\mu_3-CO_3)_2](ClO_4)_2$  (1), the first synthetic carbonate complex with alkylthiolate metal ligation (Figure 1a). The isolated carbonate complex provides a rare



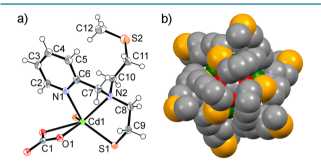
**Figure 1.** (a) Synthesis of **1** showing a stick structure of its dication. Atoms of the  $(CdS)_6$  macrocycle are highlighted as balls. (b) Perspective view of carbonate binding to the  $(CdS)_6$  macrocyclic core of **1**.

example of a bicapped molecular wheel (Figure 1b).<sup>7</sup> Alternatively, a noncrystalline "CdLClO<sub>4</sub>" (**2**) precipitate could be isolated from an EtOH/H<sub>2</sub>O solution of these reactants in the absence of air. <sup>1</sup>H NMR spectroscopy and electrospray ionization mass spectrometry (ESI-MS) are used to document the stability of **1** in solution and to show that both the base and CO<sub>2</sub> are

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required for its formation from **2**. Furthermore, this is the first study to provide experimental evidence for the fixation of  $CO_2$  by a cadmium(II) complex on a time scale of minutes or less.

Complex 1 has six crystallographically identical CdL units (Figure 2a). Each  $Cd^{II}$  atom has a bicapped tetrahedral NN'S<sub>2</sub>O<sub>2</sub>.



**Figure 2.** (a) ORTEP diagram of cadmium(II) coordination in 1 with an atomic numbering scheme and thermal ellipsoids at the 50% level. (b) Space-filling diagram of the  $1^{2+}$  cation looking down the  $C_3$  axis.

metal coordination environment (Figure S1 in the Supporting Information, SI). Each ligand uses an NN'S donor set to form a pair of fused five-membered chelate rings to one metal ion and a thiolate bridge to an adjacent Cd<sup>II</sup> atom. The thioether sulfur of L is pendant in **1**, separated from the closest Cd<sup>II</sup> atom by nearly 6 Å. The O donors are derived from asymmetric  $\eta^2$ -carbonate coordination with Cd–O distances of 2.230(2) and 2.690(3) Å, well within the sum of the van der Waals radii for Cd<sup>II</sup> (1.58 Å) and O (1.52 Å).<sup>8</sup> The asymmetric unit contains one-.

The novel obloid core of 1 is comprised of two trinuclear  $[(CdL)_3(\mu_3-CO_3)]$  units with 3-fold molecular symmetry (Figure 1b). Two trinuclear units with opposite chirality at N2 (all *R* vs all *S*) and joined to each other by six bridging thiolato S atoms form a puckered bicapped 12-membered (CdS)<sub>6</sub> macrocycle. The distance between the symmetry-equivalent Cd atoms in individual Cd<sub>3</sub>( $\mu_3$ -CO<sub>3</sub>) units is 4.882(4) Å. Sulfur-bridged Cd<sup>II</sup> atoms are separated by 3.679(3) Å. The asymmetric  $\eta^2$ -carbonate O atoms are bound 0.2628(17) Å above the coordinated plane of three alternate macrocycle Cd<sup>II</sup> atoms (Figure S2a in the SI). The carbonate C separation of 2.906(8) Å, less than the sum of the van der Waals radii (3.40 Å)<sup>8</sup> (Figure S2b in the SI), suggests assembly stabilization by  $\pi$ - $\pi$  interaction.

The bicapped (MetalS)<sub>6</sub> macrocycle is unique to 1. The dodecanuclear complex ([(cyclam)Mn<sup>IV</sup>(µ-O)<sub>2</sub>Mn<sup>III</sup>(H<sub>2</sub>O)(µ-OH)]<sub>6</sub>( $\mu_3$ -CO<sub>3</sub>)<sub>2</sub>)Cl<sub>8</sub>·24H<sub>2</sub>O<sup>9</sup> provides the only precedent for  $CO_2$  fixation in a puckered bicapped (MetalX)<sub>6</sub> macrocycle.<sup>2</sup> To the best of our knowledge, 1,3,5-triazine is the only other established 3-fold-symmetric templating species for a puckered  $(MetalX)_6$  macrocycle.<sup>10</sup> One hexanuclear dodecacarbonate of cadmium(II) is known,<sup>11</sup> but there are no bis(carbonates) of just zinc(II) or cadmium(II).<sup>2</sup> The five reported cadmium(II) carbonate complexes have varied metal coordination environments, including N<sub>4</sub>O<sub>2</sub>, N<sub>2</sub>O<sub>6</sub>, and N<sub>4</sub>(SR<sub>2</sub>)<sub>2</sub>O<sub>2</sub> (Figure S3 in the SI).<sup>12</sup> The published cadmium(II) carbonate complexes and 1 share perchlorate counterions and ligands with multiple N-donor groups. Both  $\mu_2$  and  $\mu_3$  binding modes are observed for carbonate in these cadmium(II) complexes, with hapticities ranging from nearly symmetrical  $\eta^2$  to highly asymmetrical  $\eta^1$ . As observed for 1, each Cd center in the  $\mu_3$ -carbonato complexes<sup>12b-d</sup> has two carbonate O atoms within the sum of the van der Waals radii for Cd<sup>II</sup> and O.<sup>8</sup> Although there is considerable ligand diversity among known metal carbonato complexes, the S donors have

been limited to a handful of thiolates bound to  $sp^2$ -hybridized C atoms (Figure S4 in the SI) and thioethers.<sup>2</sup>

In the context of extensive metal thiolate coordination studies, the paucity of synthetic carbonate complexes with alkylthiolate ligands is intriguing given their importance to biological CO<sub>2</sub> chemistry. Adventitious crystallization of multinuclear carbonate complexes requires kinetically efficient  $CO_2$  (<0.04% of air by volume) hydroxylation and deprotonation, as well as thermodynamic redistribution of the molecular assembly to exceed the saturation limit. Structural features likely to enhance carbonatedriven dynamic combinatorial self-assembly and prevent intermolecular interactions are evident. The staggered carbonates are close enough for  $\pi - \pi$  stacking (Figure S2b in the SI) and the sterically demanding pyridyl rings circumscribing the macrocycle faces have approximate edge-to-face orientations, severely limiting carbonate solvent exposure (Figure 1b). Thermodynamic and kinetics aspects of the formation of 1 were further investigated by <sup>1</sup>H NMR and ESI-MS.

 $^{1}$ H NMR comparisons of 1 and 2 in CD<sub>3</sub>CN supported the thermodynamic stability of 1 in solution (Figure 3). The  $^{1}$ H

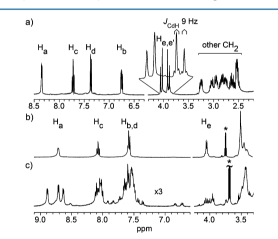
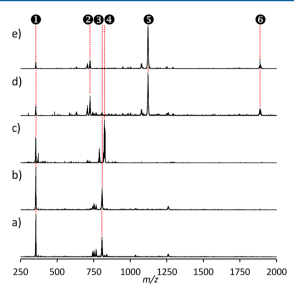


Figure 3. <sup>1</sup>H NMR spectra of (a) 1 at 80 °C and 2 at (b) +80 and (c) -40 °C (CD<sub>3</sub>CN; \* indicate CH<sub>2</sub> resonances of coprecipitated HNEt<sub>3</sub>ClO<sub>4</sub>).

NMR spectrum of 1 had a single set of ligand resonances from -40 to +80 °C and was stable to extended periods at elevated temperature. Strong geminal coupling between all of the methylene and ethylene <sup>1</sup>H atoms at elevated temperature is consistent with a well-defined structure. In addition, one of the methylene H<sub>e</sub> atoms had 9 Hz  $J(^{111/113}Cd^{1}H)$  satellites, which is to our knowledge the first observation of this interaction in a thiolate-ligated cadmium(II) complex (Figure 3a) and comparable in magnitude to couplings observed for related complexes. In contrast, the <sup>1</sup>H NMR spectrum of 2 had three major ligand environments in a 1:1:1 ratio and two minor ligand environments in a 1:1 ratio at -40 °C, suggesting at least two components (Figure 3c). A single exchange-averaged ligand environment was observed for 2 at 80 °C (Figure 3b) with geminal coupling of the methylene protons but not the ethylene protons. There was no evidence for any of the components of 2 in the <sup>1</sup>H NMR spectra of **1**.

Similarly, acetonitrile solutions of 1 and 2 had distinct ESI-MS speciation. The base peak for 1 was centered on m/z 1121 (Figure 4e) and corresponded to a combination of  $[Cd_3L_3CO_3]^+$  and  $[Cd_6L_6(CO_3)]^{2+}$  (Figure S5a in the SI). In the high mass spectrum (m/z 200–4000), a peak centered on m/z 2343 was



**Figure 4.** ESI-MS spectra for 0.2 mg/mL CH<sub>3</sub>CN solutions of 2 (bottom to top) (a) originally, (b) bubbled with CO<sub>2</sub> for 5 min, (c) with 1  $\mu$ L of NEt<sub>3</sub> added, (d) with 1  $\mu$ L of NEt<sub>3</sub> added and then bubbled with CO<sub>2</sub> for 5 min (qualitatively similar to the reverse order of the addition), and (e) **1**. Selected assignments with m/z of the isotope distribution maximum are (1) 355 ([CdL]<sup>+</sup>), (2) 725 ([Cd<sub>2</sub>L<sub>2</sub>OH]<sup>+</sup>), (3) 807 ([Cd<sub>2</sub>L<sub>2</sub>(ClO<sub>4</sub>)]<sup>+</sup>), (4) unassigned (atypical isotope distribution), (5) 1121 ([Cd<sub>3</sub>L<sub>3</sub>(CO<sub>3</sub>)]<sup>+</sup> and [Cd<sub>6</sub>L<sub>6</sub>(CO<sub>3</sub>)<sub>2</sub>]<sup>2+</sup>), and (6) 1887 ([Cd<sub>2</sub>L<sub>2</sub>(ClO<sub>4</sub>)<sub>2</sub>]<sup>+</sup>).

observed for  $[Cd_6L_6(CO_3)_2ClO_4]^+$  (Figure S5b in the SI). A modest number of additional ions with low relative abundance were observed. None of the carbonate-containing ions were detected in the ESI-MS spectra for dilute acetonitrile solutions of **2** prepared using stringent air-free conditions (Figure 4a). The base peak for **2** was  $[CdL]^+$  (Figure 4a). The other major component was  $[Cd_2L_2ClO_4]^+$  (m/z 807). Interestingly, the ESI-MS speciation of **2** was fairly limited compared to that observed for Hg(ClO<sub>4</sub>)<sub>2</sub> with *N*-(2-pyridylmethyl)-*N*-(2-(ethylthiolato)-amine.<sup>13</sup> The thoiether-group-found pendant in **1** may help to limit the oligomeric speciation in the absence of carbonate.

Finally, in situ generation of 1 by the addition of  $CO_2$  and  $NEt_3$  to 2 in acetonitrile was explored. Individually, neither of these additives produced solutions with ESI-MS (Figure 4b,c) or <sup>1</sup>H NMR (Figure S6b,c in the SI) peaks matching those for 1. The treatment of 2 with both additives resulted in ESI-MS ion patterns (Figure 4d) and <sup>1</sup>H NMR (Figure S6d in the SI) spectra that were qualitatively similar to those of samples prepared directly from 1. Furthermore, this transformation was complete within minutes, providing the first documentation of rapid  $CO_2$  fixation by a cadmium(II) complex.<sup>12</sup>

In summary, in the presence of  $Cd(ClO_4)_2$  and a base, the new potentially tetradentate thiol ligand LH was found to limit complex oligomerization, support efficient carbonate formation from atmospheric  $CO_2$ , and effectively sequester carbonate by the assembly of a novel puckered  $(CdS)_6$  macrocycle. Parallels between I and the metal binding sites of two CA forms, including the recently discovered CdCA, suggest that additional mixed N,S-donor alkylthiolate ligands may provide favorable electronics and sterics for carbonate formation and capture by cadmium(II) as well as possibly by zinc(II). We are actively investigating this possibility.

## ASSOCIATED CONTENT

#### **S** Supporting Information

X-ray crystallographic data in CIF format for 1 (CDC 902277), syntheses, crystal structure, and ESI-MS and NMR details. This material is available free of charge via the Internet at http://pubs. acs.org. The atomic coordinates for 1 have also been deposited with the Cambridge Crystallographic Data Centre. The coordinates can be obtained, upon request, from the Director, Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, U.K.

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#### Notes

The authors declare no competing financial interest.

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