Novel Cd (II) Coordination Polymers Afforded with EDTA or Trans-1,2-Cdta Chelators and Imidazole, Adenine, or 9-(2-Hydroxyethyl) Adenine Coligands

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Abstract: Three mixed-ligands of Cd(II) coordination polymers were unintentionally obtained: {[Cd(µ₃-EDTA)(Him)Cd(Him)(H₂O)₂]·H₂O}ₙ (1), {[Cd(µ₄-CDTA)(Hade)Cd(Hade)₂]}ₙ (2), and {[Cd(µ₃-EDTA)(H₂O)Cd(H⁹heade)(H₂O)]·2H₂O}ₙ (3), having imidazole (Him), adenine (Hade) or 9-(2-hydroxyethyl)adenine (9heade) as the N-heterocyclic coligands. Compounds 2 and 3 were obtained by working with an excess of corresponding N-heterocyclic coligands. The single-crystal X-ray diffraction structures and thermogravimetric analyses are reported. The chelate moieties in all three compounds exhibit hepta-coordinated Cd centers, whereas the non-chelated Cd center is five-coordinated in 1 and six-coordinated in 2 and 3. Him and Hade take part in the seven-coordinated chelate [Cd (EDTA) (H₂O)] moiety in 3. The thermogravimetric analysis (TGA) behavior of [Cd (H₂EDTA) (H₂O)]·2H₂O in 1 and 3 leads to a residue of CdO, whereas the N-rich compound 2 yields CdO·Cd(NO₃)₂ as a residue. Density functional theory (DFT) calculations along with molecular electrostatic potential (MEP) and quantum theory of atoms-in-molecules computations were performed in adenine (compound 2) and (2-hydroxyethyl)adenine (compound 3) to analyze how the strength of the H-bonding and π-stacking interactions, respectively, are affected by their coordination to the Cd-metal center.

Keywords: EDTA; CDTA; adenine; cadmium; polymers; crystal structure; DFT calculations

1. Introduction

Ethylenediaminotetraacetic acid (H₄EDTA) and its different anionic forms are the most investigated metal chelators among the amino-polycarboxylic/carboxylate ligands. Interest has been focused in both the chemical and technological fields because of the efficient chelating properties and the well-known ability of carboxylate groups to display broad metal binding modes. The ability of EDTA to form up to five-membered metal-N,N and metal-N,O chelate rings enables its recognized capability to act as a hexadentate chelator with many metal ions (M). Moreover, the conformational flexibility of the metal-N, N'-ethylenediamine ring seems to enhance the diversity of coordination modes.
We have paid attention to the bridging role in M(µ-EDTA)M entities, acting as two N-substituted iminodiacetate moieties linked together by the flexible ethylene spacer. A search in the Cambridge Structural Database (hereafter CSD) affords around forty different crystallographic results of these kinds of complexes with \(M = \text{Cu}^{II}\) (6 compounds), \(\text{Mo}^{V}\) (27 compounds), \(\text{W}^{V}\) (9 compounds), \(\text{Tc}^{V}\) (1 compound), or \(\text{Re}(I)\) (1 compound) and also including an Mo-W bimetallic example. Interestingly \(M-\text{(µ-EDTA)-M moieties}\) with first-row transition metal ions refer only to six compounds of Cu(II). Such results include the polymer \([\text{Cu}_2(\text{µ}_2\text{-EDTA})(\text{H}_2\text{O})_2]\cdot 2\text{H}_2\text{O}\) [CUEDTA01 and CUEDTA10 in CSD], the molecular compound \([\text{Cu}_2(\text{µ}_2\text{-EDTA})(\text{py})_2]\cdot 2\text{H}_2\text{O}\) (py = pyridine, NAMJOB in CSD) [1], the polymer \([\text{Cu}_2(\text{µ}_4\text{-EDTA})(\text{OH-py})_2]_n\) (3OH-py = 3-hydroxypiridine, PEZRES in CSD), and two sophisticated polymers having the \([\text{Cu}_2(\text{µ}_4\text{-EDTA})]\) moiety, silicate or phosphate, a poly-oxo-poly-wolframate anion, \(\text{Na}^+\) ions, and water molecules [2].

For a long time, we have been interested in the investigation of molecular recognition modes between copper(II) chelates and amino-purine bases (such as adenine, 2-aminopurine, 2,6-diaminopurine, and hypoxanthine) [3–13]. Our work also includes deaza- and aza-purines and closely related \(N\)-heterocycles, related to at least one \(N\) atom in each five- or six-membered rings of the purine skeleton [5]. The main aim of this work was to extend the knowledge of mixed-ligand dinuclear Cu (II) complexes of the type \((N\text{-ligand)}\text{M(µ-EDTA)}(N\text{-ligand})\) (such as NAMJOB [1] or PEZRES in CSD) to Cd (II), µ-EDTA, or µ-CDTA (where CDTA = trans-1,2-diaminocyclohexane-tetraacetate(4-) ion) as bridging-dinucleating chelators and imidazole (Him), adenine (Hade), or 9-(2-hydroxyethyl)adenine (9heade) as \(N\)-heterocyclic donor coligands. The following \(N\)-donor ligands were selected: the small heterocyclic fragment of adenine (Him), adenine itself (Hade), or the closely related synthetic nucleoside 9heade [13]. To this purpose, a series of syntheses were carried out maintaining a Cd (II):EDTA:CDTA ratio of 2:1:2 but different amounts of Him, Hade, or 9heade, from which the three reported polymers were crystallized. Hence, we describe the syntheses, thermal stability, and crystal structures of these novel compounds. For thermogravimetric analysis (TGA) comparison, \([\text{Cd(H}_2\text{EDTA)}(\text{H}_2\text{O})_2]\cdot 2\text{H}_2\text{O}\) [14] (ACAQOK in CSD) was also prepared. In the solid state of the polymer with adenine, an interesting network of H-bonds is formed. Moreover, in compound 3, antiparallel displaced \(\pi\)–stacking interactions are established. Both interactions were studied by density functional theory (DFT) calculations, focusing on the influence of the metal coordination on the relative strength of the \(\pi\)–stacking and H-bonds, which were estimated using the quantum theory of atoms-in-molecules (QTAIM) theory.

2. Materials and Methods

2.1. Reagents

\(\text{H}_4\text{EDTA}\) acid (TCI), trans-\(\text{H}_4\text{CDTA}\) (Aldrich), Him (Merck), adenine (Aldrich), 9heade (TCI, 98%), and \(\text{CdCO}_3\) (Alfa Aesar, 98%) were used as received. The synthetic purine-nucleoside 9heade can also be prepared as reported by T.Y. Shen et al. [15]. Its purity was tested by elemental CHN-analysis, FT-IR spectroscopy, and thermogravimetric analysis (TGA). Its recrystallization in various polar solvents gives crystals of 9heade consistent with the three reported crystalline forms [16]. Also, the purity of \(\text{CdCO}_3\) (98%) was tested by thermogravimetric analysis (TGA) by the experimental final residue (CdO) that results in agreement of the calculated value within 1% of assumed experimental error.

2.2. Crystallography

Colorless needle crystals of compounds 1–3 were mounted on a glass fiber and used for data collection. Crystal data were collected at 100(2) K, using a Bruker D8 VENTURE PHOTON III-14 diffractometer. Graphite monochromated MoK(α) radiation (\(\lambda = 0.71073\) Å) was used throughout. The data were processed with APEX2 [17] and corrected for absorption using SADABS (transmissions factors: 1.000–0.962) [18]. The structure was solved by direct methods using the program SHELXS-2013 [19] and refined by full-matrix least-squares techniques against \(F^2\) using SHELXL-2013 [19]. Positional and anisotropic atomic displacement parameters were refined for
all non-hydrogen atoms. Hydrogen atoms were located in difference maps and included as fixed contributions riding on attached atoms with isotropic thermal parameters 1.2/1.5 times those of their carrier atoms. Criteria of a satisfactory complete analysis were the ratios of rms shifts to standard deviations less than 0.001 and no significant features in the final difference maps. Atomic scattering factors were taken from the International Tables for Crystallography [20]. Molecular graphics were plotted with PLATON [21]. A summary of the crystal data, experimental details, and refinement results for compounds 1–3 are listed in Table S1; Table S2–S4 list the with cadmium (II) coordination bond lengths and angles and H-bonding information of 1–3. Crystallographic data for 1–3 were deposited in the Cambridge Crystallographic Data Center with the CCDC numbers 1995138-1995140.

2.3. Other Physical Measurements

Analytical data (CHN) were obtained in a Fisons–Carlo Erba EA 1108 elemental micro-analyzer. TGA was carried out (r.t. –950 °C) under an air flow (100 mL/min) by a Shimadzu Thermobalance TGA–DTG–50H instrument. To identify the evolved gases, during each TGA experiment, a series of 35 time-spaced FT-IR spectra were recorded with a coupled FT-IR Nicolet Magna 550 spectrometer.

2.4. Synthesis

In order to minimize the presence of undesired by-products, the strategies of the synthetic procedures described below used CdCO₃ (as metal ion source) and EDTA or CDTA chelating agents in their corresponding acid forms. These reactions were carried out in a Kitasato flask including its stopper but with its side outset open to maintain an open thermodynamic system that permitted gas flow. These syntheses yielded CO₂ (easily removed) as the main by-product and water (that was used as a solvent).

2.4.1. [[Cd(µ₃-EDTA)(Him)-Cd(Him)(H₂O)₂]·H₂O]n (1)

CdCO₃ (1 mmol, 0.17 g) and H₄EDTA (0.5 mmol, 0.15 g) were reacted in water (100 mL) in a Kitasato flask at 50–70 °C, with permanent stirring until a clear solution was obtained. The heating was ceased and then Him (1.1 mmol, 75 mg) was added at r.t. A clear reaction mixture was immediately obtained and then filtered without vacuum by a funnel provided with a G3-fritted glass disk (to remove any insoluble material) on a crystallization flask. The slow evaporation of the solution was controlled with the aid of a plastic film and produced the stable colorless crystals of 1 (two-three weeks at r.t.), which were removed and then dried in air at r.t. over several days. Yield: ~ 75%. Elemental analysis (%): Calc. for C₁₆H₂₆Cd₂N₆O₁₁: C 27.33, H 3.73, N 11.95; Found: C 27.21, H 3.64, N 11.95.

2.4.2. [[Cd(µ₄-CDTA)(Hade)-Cd(Hade)]ₙ (2)

CdCO₃ (1 mmol, 0.17 g) and H₄CDTA (0.5 mmol, 0.18 g) were reacted in water (100 mL) in a Kitasato flask at 50 °C, with permanent stirring for one day. A somewhat translucent solution was obtained. The heating was ceased and then Hade (2 mmol, 0.27 g) was added at r.t. in small portions. The reaction mixture was filtered without vacuum by a funnel provided with a G3-fritted glass disk (to remove some white material) on a crystallization flask. The slow evaporation of the solution was controlled with the aid of a plastic film and produced the stable colorless crystals of 1 (four weeks at r.t.), which were collected and then dried in air at r.t. for one week. Yield: ~60%. This procedure represents 100% Hade in excess. Similar results were recently obtained using 50% Hade (1.5 mmol) in excess. Elemental analysis (%): Calc. for C₂₉H₃₃Cd₂N₁₇O₈: C 35.82, H 3.42, N 24.49. Found: C 35.76, H 3.39, N 24.44.

2.4.3. [[Cd(µ₃-EDTA)(H₂O)-Cd(H₈heade)(H₂O)]·4H₂O]n (3)

CdCO₃ (1 mmol, 0.17 g) and H₄EDTA (0.5 mmol, 0.15 g) were reacted in water (100 mL) in a Kitasato flask at 50 °C, with permanent stirring until a clear solution was obtained. The heating
was ceased and then H9heade (2 mmol, 0.36 g) was added in small portions at r.t. A clear reaction mixture was obtained, left to cool, and then filtered without vacuum on a crystallization flask. The slow evaporation of the solution was controlled with the aid of a plastic film and produced the stable colorless crystals of 3 (three weeks at r.t.), which were collected and then dried in air at r.t. over several days. This procedure represents 100% H9heade in excess. These results were recently confirmed by repeating this procedure. Yield: ~65%. Elemental analysis (%): Calc. for C_{17}H_{33}Cd_{2}N_{7}O_{15}: C 25.51, H 4.16, N 12.25, Cd 28.09; Found: C 25.43, H 4.04, N 12.19.

2.4.4. [[Cd(H₂EDTA)(H₂O)]-2H₂O]₁ₙ

CdCO₃ (1 mmol, 0.17 g) and H₄EDTA (1 mmol, 0.29 g) were reacted in water (100 mL) in a Kitasato flask at 50–70 °C, with permanent stirring until a clear solution was obtained, which was left to cool at r.t. and then was filtered without vacuum (to remove any insoluble material) on a crystallization flask. The slow evaporation of the solution (controlled with the aid of a plastic film) produced the well-shaped colorless crystals of the desired product (two-three weeks at r.t.), which were removed and then dried in air at r.t. over several days. Yield: ~80%. Elemental analysis (%): Calc. for C_{10}H_{20}CdN_{2}O_{11}: C 26.30, H 4.41, N 6.13; Found: C 26.21, H 4.34, N 6.09. Polynova et al. [14] synthesized this compound by reaction between CdSO₄·xH₂O (x = 2.67) and Na₂H₂EDTA in water.

2.5. Theoretical Methods

All DFT calculations were carried out using the Gaussian-16 program [22] at the PBE1PBE-D3/def2-TZVP level of theory and using the crystallographic coordinates. The formation energies of the assemblies were evaluated by calculating the difference between the total energy of the assembly and the sum of the monomers that constitute the assembly, which were kept frozen. The molecular electrostatic potential was computed at the same level of theory and plotted onto the 0.001 a.u. isosurface. The quantum theory of atoms-in-molecules (QTAIM) [23] analysis was carried out at the same level of theory by means of the AIMAll program [24].

3. Results and Discussion

The following sections highlight the relevant structural features of compounds 1 to 3 and their thermal stability. Detailed tables with coordination bond lengths and angles as well as data concerning the H-bonds are supplied as Supporting Information (See Tables S2–S4). Tables summarizing TGA results are given as Supporting information (Tables S5–S8).

3.1. Synthetic Considerations

The utilization of cadmium carbonate and the acid form of the chelators for the syntheses of metal complexes reported herein. The synthesis [Cd(H₂EDTA)(H₂O)] (ACAQOK in CSD) is supported by the reaction:

\[
\text{CdCO}_3 + \text{H}_4\text{EDTA} \rightarrow [\text{Cd(H}_2\text{EDTA})(\text{H}_2\text{O})] + \text{CO}_2\uparrow 
\]

where two protons from the H₄EDTA acid react with carbonate anion yielding H₂O (the used solvent) and CO₂ (as an easily removable by-product).

Analogously, we can write:

\[
2 \text{CdCO}_3 + \text{H}_4\text{EDTA} \rightarrow [\text{Cd}_2(\mu\text{-EDTA})] + 2\text{CO}_2\uparrow + 2\text{H}_2\text{O}
\]

where

\[
2\text{CO}_3^{2-} + 4\text{H}^+ \rightarrow 2\text{CO}_2\uparrow + 2\text{H}_2\text{O}.
\]

This is a convenient synthetic strategy because with this simple process CO₂ and water are the unique by-products. The CO₂ is easy removed by the aid of simple physical actions, such as heating and/or stirring and/or moderate vacuum (or an appropriate combination of them). The stoichiometric
reaction that yields the formation of the polymer \([\text{[Cd}_2(\mu_7\text{-EDTA})]\cdot \text{H}_2\text{O}]_n\) (FAFJON in CSD) is given below:

\[
2n \text{ CdCO}_3 + n \text{ H}_4\text{EDTA} \rightarrow [\text{[Cd}_2(\mu_7\text{-EDTA})]\cdot \text{H}_2\text{O}]_n + 2n \text{ CO}_2\uparrow + n \text{ H}_2\text{O}.
\]

The appropriate utilization of \(\text{H}_4\text{EDTA}\) or \(\text{H}_4\text{CDTA}\) and N-heterocyclic coligands yields the polymeric compounds 1 to 3. For example, in compound 1 the stoichiometric reaction can be written as:

\[
2n \text{ CdCO}_3 + n \text{ H}_4\text{EDTA} + 2.2n \text{ Him} + n \text{ H}_2\text{O} (\text{solvent}) \rightarrow [\text{[Cd}(_3\text{-EDTA})(\text{Him})\cdot \text{Cd(Him)(H}_2\text{O)}_2]\cdot \text{H}_2\text{O}]_n (1) + 2n \text{ CO}_2\uparrow + 0.2n \text{ Him}.
\]

In this case, the crystallization of 1 needs a very small number of water molecules (from the solvent) and, simultaneously, a small amount of Him (used in 10% of excess) remains in solution.

3.2. Thermal Stability of \([\text{Cd(H}_2\text{EDTA)(H}_2\text{O)}]\cdot 2\text{H}_2\text{O}\) and the Polymeric Compounds 1 to 3

The following sections highlight the relevant structural features of compounds 1 to 3 and their thermal stability.

Under air-dry flow, the weight loss versus temperature in complex \([\text{Cd(H}_2\text{EDTA)(H}_2\text{O)}]\cdot 2\text{H}_2\text{O}\) consists of five steps (Figure 1a). The experimental results and assignations are summarized in Table S5.

![Figure 1](image)

**Figure 1.** Weight loss versus temperature in the TGA of \([\text{Cd(H}_2\text{EDTA)(H}_2\text{O)}]\cdot 2\text{H}_2\text{O}\) (sample: 9.99 mg; r.t. to 950 °C) (a) and compound 1 (10.95 mg; r.t. to 950 °C) (b).

The first step (25–185 °C) in the TGA of \([\text{Cd(H}_2\text{EDTA)(H}_2\text{O)}]\cdot 2\text{H}_2\text{O}\) agrees with the loss of non-coordinated water, because most of this lost weight occurs below 100 °C. The experimental data in the second step is higher than the calculated value, considering the loss of all aqua ligands as a consequence of the partial burning of the protonated organic ligand that could start before the end this step (275 °C). The formation of ammonia was not observed from the burning of \(\text{H}_2\text{EDTA}^2-\). The third step yields C-oxides and water. The last two steps produce methane, trace amounts of ethylene, and the three commonly observed N-oxides (\(\text{N}_2\text{O}, \text{NO}, \text{and NO}_2\)). The final residue seems to be non-pure \(\text{CdO}\).

The TGA plot for compound 1 is shown in Figure 1b and the corresponding results are summarized in Table S6. Compound 1 essentially loses the uncoordinated water and both aqua ligands in the first step, from room temperature (r.t.) to 185 °C. Organic ligands burn in the remaining steps, but mainly between 300 and 560 °C (steps 3–5). On the basis of the above for the molecular compound \([\text{Cd(H}_2\text{EDTA)(H}_2\text{O)}]\cdot 2\text{H}_2\text{O}\), the gases evolved in the third step suggest a partial overlap of these processes. Indeed, the weight loss in the last step (12.55%) is lower than expected for the loss of 2 Him ligands (19.36%). The observed weight for the final residue at 560 and 950 °C is consistent with the calculated weight for 2 \(\text{CdO}\).

The TGA plot for compound 2 is shown in Figure 2a and the corresponding results are summarized in the Table S7. The crystal structure of compound 2 (see below) revealed that there is no water in this compound and hence a relevant thermal stability should be expected for it. However, the first step from room temperature to 260 °C essentially shows water loss. Most of the weight loss occurs below 150 °C. We have assumed that the fresh sample used was not completely dry. On this basis, the
observed behavior agrees to a formula $[\text{Cd}(\mu_4\text{-CDTA})(\text{Hade})\cdot\text{Cd}(\text{Hade})_2]\cdot4\text{H}_2\text{O}]_n$. The calculated value to remove such water content is 6.90%, in good agreement to the experimental value (7.00%). No other hypothesis seems reasonable, because in this step, only water is lost (with small amounts of CO$_2$). Above 560 °C a stable residue of ~33.4% is formed. The calculated value to 2 CdO (24.59%) is too low, however, it is well known that the burning of this N-rich polymer can change it to a (not necessarily stoichiometric) cadmium oxy-nitrate [25]. Indeed, an estimation for CdO-Cd(NO$_3$)$_2$ as a final residue leads to a quite reasonable calculated value (~35%); therefore, we tentatively assigned the residue to cadmium oxy-nitrate.

![Figure 2](image)

**Figure 2.** (a) Weight loss versus temperature (r.t. to 950 °C) in the TGA of compound 2 (sample: 10.97 mg). (b) Weight loss versus temperature (r.t. to 950 °C) in the TGA of compound 3 (sample: 11.04 mg).

The TGA plot for compound 3 is shown in Figure 2b and the corresponding results are summarized in Table S8. Compound 3 shows a multi-step TGA behavior where the four first steps cannot be assigned (partial water-loss processes). However, all these steps and the corresponding range of temperatures match well to the amount of water and aqua ligands present in the compound. The remaining steps correspond to the burn of organic ligands. The final residue matches cadmium oxide.

### 3.3. Structural Description

#### 3.3.1. $[\text{Cd}(\mu_3\text{-EDTA})(\text{Him})\cdot\text{Cd}(\text{Him})(\text{H}_2\text{O})_2]\cdot\text{H}_2\text{O}]_n$ (1)

This section highlights the relevant structural features of compounds 1 to 3. The Supporting Information contains detailed data of coordination bond lengths and angles (Table S2) as well as H-bonds (Table S3) and π-stacking interactions (only found in 2 and 3, Table S3).

Compound 1 consists of 1D-polymeric chains running parallel to the a axis of the crystal. The asymmetric unit shows two non-equivalent metallic centers, Cd1 and Cd2 (Figure 3 and Table S2).

![Figure 3](image)

**Figure 3.** Perspective view of $[\text{Cd}(\mu_3\text{-EDTA})(\text{Him})\cdot\text{Cd}(\text{Him})(\text{H}_2\text{O})_2]\cdot\text{H}_2\text{O}]_n$ (1) showing the asymmetric unit, with the atom-numbering scheme and some H-bonds (orange dashed lines).
The hepta-coordinated Cd1 atom is chelated by the $\mu_3$-EDTA ligand and is also linked to one Him ligand, defining a distorted mono-caped octahedral coordination. That is now recognized as a rather common coordination for this [Kr]4d$^{10}$ soft Pearson’s acid metal ion. Because all donor atoms of EDTA behave as hard Pearson’s bases and imidazole is a borderline base with moderate steric relevance, the Cd1-N1(Him) bond (2.224(2) Å) is the shortest one in this coordination polyhedron.

Another interesting feature is the unequal metal binding pattern featured by the two half-EDTA moieties in the polymeric chain of 1. Cd1-N20 (2.406(2) Å, Cd1- O21 (2.389(2) Å) and Cd1-O23 (2.556(2) Å) bonds confer to the first half-EDTA the role of an N-substituted-iminodiacetate N,O,O’-tridentate chelator, with Cd1-O21 and C1-N20 bonds (2.40 Å) being shorter than the Cd1-O23 bond. Indeed, O23 seems to play the “mono-caped” role in the hepta-coordination of Cd1. Thus the -C22,O21,O22 and -C23,O23,O24 carboxylate groups act as monodentates. This role probably relaxes the steric constraints related to the EDTA chelator and the implication of the O24-carboxylate atom as a triply-acceptor for the H-bonds (see Table S3).

In clear contrast, the second half-EDTA moiety assumes the $\mu_3$-EDTA bridging role in the polymeric chain by means of its two nearly anti,syn-carboxylate groups. The torsion angles involved are: Cd1-O13-C14-O14 -178.9° and O13-C14-O14-Cd2 -6.0°, Cd1-O11-C12-O12 -159.4° and O11-C12-O12-Cd2 17.5°. The N10, O11, and O13 atoms give slightly dissimilar bond distances, Cd1-O13 (2.337(2) Å) < Cd1-N10 (2.395(2) Å) < Cd2-O11 (2.437(2) Å), with an averaged value of 2.39 Å. Hence, both carboxylate groups -C12,O11(anti),O12(syn) and -C14, O14(anti),O13(syn) are bidentate. That increases the denticity of the $\mu_3$-EDTA to eight.

The Cd2 center exhibits a roughly penta-coordination, with three rather short bond distances (~2.20 Å): Cd2-N3 (Him) (2.205 (2) Å), Cd2- O12 (2.213 (2) Å), and Cd2-O14 (2.194 (2) Å), from the two anti,syn-carboxylate groups of adjacent Cd(EDTA) chelated moieties. Two aqua ligands fulfil this center, with Cd2-O1 and Cd2-O2 distances averaging ~2.32 Å. The angle O1-Cd2-O2 (164.0 (1) °) is the closest to 180° in the Cd2 coordination. A detailed inspection of this center locates O11 and O13 carboxylate donors of Cd1 at 3.011 and 3.163 Å from Cd2 respectively. However, such weak contacts (also depicted in Figure 4) should not be considered as true coordination bonds for two main reasons. First, the Cd2--O11 and Cd2--O13 distances remarkably approach or even exceed the sum of the van der Waals radii (1.52 (O) + 1.58 (Cd) = 3.10 Å). Second, such “contacts” seem clearly influenced by two aqua-mediated intra-polymeric chain H-bonds (see Table S3), namely O2-H2W1···O21 (2.759 (3) Å, 174.3°) (also shown in Figure 4) and O2-H2W2···O24b (2.777(3) Å, 169.0°, see Table S3 for symmetry code). Such H-bonds and other inter-chain aqua-mediated bonds (involving OH$_2$ ligand) build a $R_3^2$(10) graph-set motif (Figure 5). The H$_2$O$_3^{4\text{d}}$ water molecule is also involved in one (Him) N4--H4···O3d interaction (2.717 (3) Å, 175.7°, see Table S3 for symmetry code) as shown in Figure 4.

Figure 4. Plot of 1 showing some H-bonds (orange dashed lines) and the $R_3^2$(10) graph-set motif involving both aqua ligands of the Cd2 center. For appropriate symmetry codes, see Table S3.
Figure 5. 2D H-bonded networks of 1 parallel to the ac plane are further pillared along the b axis and connected by additional (Him)N-H···O(carboxylate) and (aqua)OH···O(carboxylate) interactions. H-bonds are depicted as orange dashed lines.

All N-H (imidazole) and O-H (aqua ligands or water) of 1 are involved in N-H···O or O-H···O interactions, with rather short (H-donor)···(acceptor) distances (2.63-2.78 Å) and open angles (169-176°). Polymeric chains are H-bonded in layers parallel to the ac plane and in turn are H-bonded and pillared along the b axis of the 3D-crystal (Figure 5). In this array, the O24-carboxylate atom acts as a triply H-acceptor. The H2O (3) water molecule is involved in two O3-H···O(carboxylate) and one (Him)N-H···O interactions.

3.3.2. [Cd(µ4-CDTA)(Hade)-Cd(Hade)]n (2)

The crystal of compound [Cd(µ4-CDTA)(Hade)-Cd(Hade)]n (2) consists of fused metallacycles (Figure 6) as graphite-like layers generating a 2D-polymer framework, parallel to the ab plane (Figure 7) and pillared along the c axis of the crystal. This configuration is because all carboxylate groups of the µ4-CDTA ligand display an anti, syn-bridging role.

Figure 6. Asymmetric unit of [Cd(µ4-CDTA)(Hade)-Cd(Hade)]2(H2O)2·H2O)n (2) showing the atom-numbering scheme.
Figure 7. Crystal packing of 2 projected onto the ac plane. The polymeric layers parallel to the ab plane leave cyclohexane rings of CDTA chelators and Hade ligands oriented toward their external faces.

In the polymer, there are three crystallographically independent metallic centers. The Cd2 center of the fragment Cd(μ4-CDTA)(Hade) exhibits a distorted mono-capped octahedral coordination, where, interestingly, both N-donors of the trans-1,2-CDTA ligand form the largest coordination bonds (Cd2-N1 2.457(5) and Cd2-N2 2.469(5) Å). Two factors contribute to such polyhedron distortion: the fulfilled 4d\(^{10}\) electronic configuration of the Cd(II) center and the remarkable steric constraints of the trans-N(e),N(e)-CDTA conformer, which favors its hexadentate chelating role against the trans-N(a),N(a)-CDTA. The other four Cd2-O(carboxylate) bond distances of this polyhedron fall within the range 2.277(5)-2.364(5) Å (see Table S2). The hepta-coordination of Cd2 is accomplished by the Cd2-N3(Hade) bond, namely Cd2-N (23) 2.332(5) Å. It is assumed that the N3 atom is the N-heterocyclic atom with the lowest proton affinity [5]. However, the formation of this coordination bond is favored by the use of the most stable tautomer of the adenine, the (H(N9)ade), which in turn enables its cooperation with the intramolecular interligand interaction N29-29A···O42(CDTA) (2.713(8) Å, 137.2°). Moreover, this interaction permits the exocyclic amino group –N(26)H\(_2\) (as H-donor) and the N1 and N7 heterocyclic atoms (as H-acceptors) to participate in N-H···N interactions, which contribute to the crystal packing (see below).

Each half-Cd2(μ4-CDTA)(Hade) unit connects with two non-equivalent but rather similar fragments, Cd1(Hade)\(_2\) and Cd3(Hade)\(_2\). Four O-carboxylate donors from Cd2(μ4-CDTA)(Hade) units build a centro-symmetric octahedral coordination to the Cd1 or Cd3 centers. Again, their trans-H(N9)ade ligands bind the corresponding metal centers by means of a cooperation of the Cd1-N13 (2.366(5) Å) or Cd3-N33 (2.380(5) Å) bonds with the N19-H19A···O11=1 (coordinated carboxylate, 2.969(7) Å, 132.0°) or N39-H39A···O31=3 (non-coordinated carboxylate) interligand interactions (\(≠1=+x+2,–y,z; ≠3 = x,y+1,z\)). It is just the O11-coordinated or O31-non-coordinated nature of these H-acceptor atoms that introduce the above-mentioned dissimilarity between the Cd1 or Cd3 centers. The 2D-polymeric layers leave the cyclohexane moieties of CDTA and the adenine ligands oriented toward their external faces (Figures 7 and 8).

The layers of this polymer pillared along the c axis are held together by pairs of rather linear N6-H···N1 and N6-H···N7 interactions that build synthons of the \(R^2_5(9)\) graph-set motif (Figure 8).
Figure 8. Synthons of the $R_2^2(9)$ graph-set motif with rather linear N6-H···N1 and N6-H···N7 interactions connect the external faces of the polymeric layers of compound 2. This plot also shows (for Cd1 center) the cooperation between the Cd-N3(Hade) bond and the N9-H···O(coordinated carboxylate) interligand interaction. Data of these H-bonds (orange dashed lines) are reported in Table S3.

3.3.3. [[Cd(µ₃-EDTA)(H₂O)Cd(9heade)(H₂O)]·4H₂O]ₙ (3)

The asymmetric unit of this compound (see Figure 9) revealed its polymeric nature with two nonequivalent metallic centers. The first one corresponds to the Cd1(µ₄-EDTA)(H₂O) cheated moiety, where the distorted hepta-coordination is fulfilled by the hexadentate chelating role of EDTA and the O₂-aqua ligand. This unit implies that the synthetic 9heade ligand is unable to remove the O₂-aqua ligand from the Cd1 center, in spite of the use a molar ratio of 2:1:2 of reagents CdCO₃:H₄EDTA:9heade in the synthesis (which finally represents an excess of 100% of the N-heterocyclic coligand). The Cd1-O₂(aqua) is the shortest bond (2.232(4) Å) in this coordination polyhedron, where all other Cd1-N or Cd1-O bonds are in the rather narrow range of 2.34-2.46 Å (see Table S2). The differences in these last bond distances seem difficult to rationalize due to the sophisticated µ₄-EDTA bridging role. Again, the two half moieties of EDTA display clearly distinct bridging functions. A half-(N32)-EDTA uses the monodentate –C38,O39,O40 group (Cd1-O39 2.332(3) Å) along with the µ₂-η¹,η¹ - anti,anti-C33,O35,O36 bridging group (C1-N32 2.383(3), Cd1-O35 2.344(3), and Cd2-O36b 2.268(3) Å, b = -x+3/2,y+1/2,-z+3/2). The other half-(N21)-EDTA features the bidentate µ₂-η¹,η¹ - anti,syn-C23,O24,O25 bridging group CdN21(Cd1-N21 2.425(3), Cd1-O24 2.341(3), and Cd2-O25 2.229(3) along with the tridentate µ₂-η²,η¹-C27O28,O29 bridging group (Cd1-O28 2.418(3), Cd2-O28a 2.411(3), and Cd2-O29a 2.445(3) Å, a = -x+1/2,y+1/2,-z+3/2). In addition to the above mentioned Cd2-O25, C2-O28 and C2-O29 bonds, the Cd2 center fulfills its hexa-coordination with the Cd2-O1(aqua, 2.305(3) Å) bond and the Cd2-N27(9hedae, 2.267(4) Å) bonds. Two aspects are worth commenting on in this regard: (i) the Cd2-O1(aqua) bond is not the shortest one around the Cd2 center and (ii) the N9-(2-hydroxyethyl) arm of 9heade favors the formation of the Cd2-N7 bond against Cd-N3 (as found with Hade in compound 2), because this coordination mode also implies the cooperation with the interligand H-bond N6-H6B···O29a (3.067(5) Å, 153.6°). Note that the N1-9heade donor results are disfavored for metal binding (against to N7-9heade) because of the great steric factor displayed by the adjacent exocyclic-(N6) amino group.
The crystal of (3) consists of fused metallacycles (each having three Cd1 and three Cd2 centers, Figure 10) generating graphite-like 2D-polymeric frameworks, parallel to the ab plane (Figure 10b). These layers orientate 9heade towards both external faces. Such layers are pillared along the c axis, connected by many H-bonds (Figure 10 and Table S3) and a \( \pi \)-stacking interaction between the six-membered rings from the 9heade moieties (Figure 11 and Table S4).

3.3.4. CSD Search

Finally, we searched the Cambridge Structural Database (CSD) to investigate the structural features of Cd(II) complexes with chelators. Remarkably, only three Cd-CDTA derivatives were found in
the database (Table S9). For instance, in the polymer [(Cd(µ4-CDTA)·Mn(H2O)4)]·3H2O)n, refcode GAZPOM, the metal ion is also hepta-coordinated by the hexadentate CDTA plus an O-carboxylate from a neighboring CDTA ligand. Table S9 summarizes structural details featured by the thirteen structures with Cd(II) chelates. Remarkably, most of these compounds are polymers, in agreement with the structures of compounds 1–3. In addition, regardless of the chelating anion (H2EDTA2−, HEDTA3− or EDTA4−), eleven of these compounds also exhibit hepta-coordinated Cd (II), with the hexa-dentency of the chelator implemented with an O−(aqua or carboxylate) donor. The two exceptions for that (LOFKAT and IFELIP) correspond to structures where the HEDTA3− chelator plays a penta- or tetra-dentate role in hexa-coordinated-Cd compounds, also having one S-donor (LOFKAT) or two N-(heterocyclic) donor coligands (IFELIP).

3.4. DFT Calculations

The DFT study was focused on analyzing the interesting assemblies described above for compounds 2 and 3, in particular the influence of the coordination to Cd on the H-bonding and π–π interaction strength. First of all, the molecular electrostatic potential (MEP) of the surfaces of adenine and adenine coordinated to Cd were computed (see Figure 12) in order to analyze how the MEP values at the H-bond acceptor and donor groups of adenine change upon complexation. It can be observed that the N3 atom of adenine is the most nucleophilic/basic in agreement with the X-ray structure of compound 2 where the nucleobase is coordinated via an N3-atom. The MEP value at N1 is more negative than in N7. The most positive NH group corresponds to N9–H (+50 kcal/mol). Both H-atoms of the exocyclic NH2 group exhibit identical MEP values (+38 kcal/mol). We used a monomeric model of compound 2 (see Figure 12b) where two carboxylate groups were protonated to keep the neutrality of the system. The effect of the coordination to the Cd (CDTA) system is an increase of the nucleophilicity at N1 and N7 and a slight reduction of the MEP values at the exocyclic NH2 group. This is due to the effect of the formation of two strong and intramolecular H-bonds between the carboxylate ligands and the N9–H and C2–H groups with concomitant charge transference from the anion to the adenine ring. Therefore, the coordination of adenine to a Cd (CDTA) moiety increases the H-bond acceptor ability of adenine.

![Figure 12. Molecular electrostatic potential (MEP) surfaces for adenine (a) and a model of compound 2 (b), highlighting the electropositive (blue) and electronegative (red) regions of each molecule. The values at selected points of the surface are given in kcal/mol. For the model of compound 2, the surface around the CDTA ligand has been omitted for clarity.](image)

We selected the H-bonding network commented on in Figure 8 to analyze the energetic features of the H-bonds in 2. Figure 13a shows the H-bonded dimer of adenine [R2(9) synthon] observed in the solid state of 2. It presents a moderately strong dimerization energy (ΔE1 = −9.5 kcal/mol) due to the formation of two N–H···N H-bonds. Interestingly, the dimerization energy is the same as the dimer when the adenine is more favorable (ΔE2 = −11.1 kcal/mol, see Figure 13b), in agreement with the MEP analysis, evidencing that the coordination to Cd reinforces the H-bonds. For compound 3,
we performed a similar analysis for the peculiar π-stacking motif observed in 3, where the two H-atoms of the 2-hydroxyethyl arms are pointing to the aromatic rings (see Figure 13c,d), so a combination of C–H···π and π–π interactions are formed. For the study, a simplified model of 3 was used due to its polymeric nature. The carboxylate groups of EDTA were replaced by formate ligands in order to generate monomeric species (see small arrows in Figure 13d). The dimerization energy of the (2-hydroxyethyl)adenine is modest ($\Delta E_3 = -6.8$ kcal/mol, see Figure 13c), however, it becomes more favorable, $\Delta E_4 = -8.0$ kcal/mol, upon coordination to Cd, thus indicating that the antiparallel π-stacking is reinforced, likely due to an increase of the dipole···dipole interaction, since the dipole of the (2-hydroxyethyl)adenine molecule significantly increases upon coordination of Cd.

![Figure 13.](image-url)

Figure 13. (a) Dimer of adenine retrieved from the X-ray solid state of 2, distances in Å. (b) Dimer of a model of compound 2, distances in Å. (c) Isolated π,π-stacked dimer of two 9heade molecules based on the X-ray structure of 3. The distances are given in Å. (d) π,π-stacked dimer of a theoretical model of compound 3.

Finally, we used the QTAIM method to further analyze the noncovalent interactions highlighted in Figure 13 for compounds 2 and 3. A bond path and bond critical point (CP) interconnecting two atoms can be used as an unambiguous indication of interaction [26]. Moreover, this type of analysis has been recently used in similar systems [27–31]. The distribution of bond CPs and bond paths of the two motifs analyzed above are shown in Figure 14. The QTAIM analysis of the H-bonded dimer of 2 confirms the existence of the intramolecular H-bonds between the carboxylate groups and the C2–H and N9–H bonds. Moreover, each intermolecular N–H···N bond is characterized by a bond CP and bond path interconnecting the H and N-bonds. Moreover, the H-bonded dimer is further characterized by a ring CP as a consequence of the formation of the supramolecular $R^2_2(9)$ ring. The π-stacked dimer is characterized by four bond CPs, two of them interconnecting the N-atoms of the adenine rings (see Figure 14b), thus confirming the antiparallel displaced π-stacking interaction. The other two CPs characterize the two symmetrically equivalent C–H···π interactions involving the C–H bonds of the $sp^3$ C-atoms of the side arms and the six-membered ring. The bond path connects the H-atom to one C-atom of the ring. This combination of interactions agrees with the high dimerization energy obtained for this dimer of compound 3.
4. Concluding Remarks

The main objective of the present work was the synthesis of molecular compounds of the general formula (N-ligand)Cd(µ-chelator)Cd(N-ligand) with EDTA (1,3) or trans-1,2-CDTA (2) chelating agents and closely related N-heterocyclic coligands, however, this aim was not accomplished. Instead of that, three novel coordination polymers were obtained and are reported on herein. In such compounds, the chelated Cd(II) metal centers are hepta-coordinated. It should be emphasized that only three Cd-CDTA derivatives were found in CSD database and in only one of them (GAZPOM) is the metal center also hepta-coordinated (hexadentate CDTA plus an O-carboxylate from an adjacent CDTA).

This work demonstrated that the cadmium(II) hepta-coordination of the Cd(EDTA or CDTA) chelate moieties in these kinds of polymers can be preserved with relatively small N-heterocyclic coligands (imidazole or adenine) but cannot use larger ones such as 9heade. Our findings also open a promising window for further investigations on this matter with additional experiments.

Finally, the MEP analysis and DFT calculations show that the H-bonding and π-stacking interactions involving adenine or 9heade rings are enhanced upon coordination to Cd.

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