Two New Adenine-based Co(II) Coordination Polymers: Synthesis, Crystal Structure, Coordination Modes and Reversible Hydrochromic Behavior

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ABSTRACT: We report the synthesis of two new 3D coordination polymers (CPs) based on Co(II), adenine and aromatic tetracarboxylate linkers. Adenine exhibits bidentate binding modes in both CPs, coordinating through the N1 and N6 sites in a first compact CP and through the more rare N1 and N7 sites in a second open, flexible and H2O-responsive CP. These differences together with an analysis of the extended coordination structures made of adenine reported in Cambridge Structural Database illustrate the rich coordination versatility of adenine as a building block for CPs. Although the latter CP is non-porous to N2 or CO2, it shows a reversible and detectable colour change from pink to purple, and vice versa, upon hydration and dehydration, respectively.

INTRODUCTION

Metal-Organic Frameworks (MOFs) or porous Coordination Polymers (CPs) are among the most attractive porous materials today, owing to their potential for applications such as gas storage; separation; catalysis; molecular sensing; adsorption for heat-pump processes; contaminant removal; contrast agents; and drug delivery systems. The infinite possibilities for connecting functional organic ligands to metal ions and clusters enables the formation of countless CPs with a range of pore sizes and shapes, extremely large surface areas, tailored internal surfaces, and flexible structures that give rise to novel phenomena such as gate-opening or sponge-like behavior. Bio-related linkers such as amino acids, peptides and nucleobases are very attracting building blocks that can be used for the formation of CPs due to their large number of oxygen and nitrogen atoms with different basicity that are available for metal coordination. In particular, adenine is a rigid connector with five potential coordination sites for metal binding: two imidazolate, two pyrimidinate N atoms and a NH2 group with basicity order of N6 > N1 > N2 > N1 > N6 (Scheme 1). It provides the widest range of neutral tautomers and protonated forms, and the non-coordinated nitrogen donor sites—especially, Watson-Crick (N1, N6) and Hoogsteen (N7, N6) faces—can interact through hydrogen bonding. Its rich metal binding and H-bonding capabilities, together with the rigidity of its molecular structure, make adenine an ideal bio-linker for constructing topologically diverse CPs. For instance, adenine has been used to build up robust porous CPs with applications for example in drug encapsulation and in the selective adsorption of greenhouse gases such as CO2.

Herein, we report the synthesis of two new 3D CPs composed of Co(II), adenine (ade) and aromatic tetracarboxylic acids. The selected tetracarboxylic acids are 1,2,4,5-tetrakis(4-carboxyphenyl)benzene (H4TCPB) and 1,2,4,5-benzene tetracarboxylic acid (H4BTeC). [Co3(μ4-O-TCPB)(μ-H4TCPB)(ade)]·4H2O (1) is based on a trinuclear Co(II) cluster, in which the three Co(II) ions are bridged by two bidentate adenine ligands through the N1 and N6 sites. [Co3(ade)(μ6-BTeC)(μ-H2O)(H2O)2]·4(H2O) (2) comprises binuclear Co(II) clusters, which are linked by bidentate adenine linkers (through the N1 and N7 sites) forming 1D Co(II)-adenine chains. The orientation of TCPB in 1 results in a compact network, whereas BTeC+ links the Co(II)-adenine

![Scheme 1. Illustration of adenine showing the labels for the N binding sites.](Image)
chains forming a 3D open network in 2. The structure 2 shows reversible structural transformations upon dehydration and hydration, which are accompanied by a detectable colour change from pink to purple, respectively. This colour change results from the dehydration process, which usually implies removal of coordinated H₂O molecules and therefore, conversion from octahedral (2, hydrated, pink) to tetrahedral (2′, dehydrated, purple) Co(II) coordination geometry (or vice versa, for the hydration process). It is found that 2′ is non-porous to N₂ or CO₂. Contrariwise, H₂O can diffuse within its framework and organize the well-ordered structure of 2.

EXPERIMENTAL SECTION

Materials and Characterization. Adenine (C₉H₇N₃), 1,2,4,5-benzenetetracarboxylic acid (H₂BTeC, C₁₀H₅O₄), and 1,2,4,5-tetakis(4-carboxyphenyl)benzene (H₂TCPB, C₂₅H₁₆O₈) were purchased from Sigma-Aldrich, and cobalt(II) carbonate (CoCO₃) was procured from Alfa Aesar. The reagents were used as received. IR spectra were recorded using an AutosorbIQ (Quantachrome Instruments). Field-Emission Scanning Electron Microscopy (FE-SEM) images were collected using a Quanta 650F Environmental Scanning Electron Microscopy (Field Emission Inc, USA). Aluminium was used as support. Powder X-ray Diffraction (PXRD) patterns were recorded at room temperature on an X’Pert PRO MPD diffractometer (PanAnalytical) using Cu Kα (λ = 1.7902 Å) emission line.

Synthesis of [Co₂(μ₂-H₂TCPB)(μ₂-H₂TCPB)(ade)₂]·H₂O (1). An aqueous solution (6 mL) of adenine (5 mg, 0.04 mmol), H₂BTeC (10 mg, 0.02 mmol) and CoCO₃ (10 mg, 0.08 mmol) was placed in a 12 mL scintillation vial and heated at 100 °C for 48 h. After slow cooling to room temperature during 6 h, 1 was obtained as pink crystals, which were filtered and air-dried (25% yield). Elemental analysis, calculated (%): C: 59.8, H 3.2 and N: 8.9; found: C: 59.0, H: 3.1 and N: 8.7. FT-IR (KBr, cm⁻¹): 3355(m); 1680(m); 1606(s); 1583(s); 1406(s) (br). 

Synthesis of [Co₄(μ₄-TCPB)(μ₄-CH₂TCPB)(μ₄-Sbr)₈·4(H₂O)] (2). A mixture of adenine (5 mg, 0.04 mmol), H₂BTeC (10 mg, 0.04 mmol) and CoCO₃ (10 mg, 0.08 mmol) in water (6 mL) was placed in a 12 mL scintillation vial and heated at 100 °C for 48 h. After slow cooling to room temperature during 6 h, pink crystals of 2 were obtained, filtered and air-dried (25% yield). Elemental analysis, calculated (%): C: 28.6, H: 3.4; N: 11.1; found: C: 28.5, H: 3.3, N: 11.3. FT-IR (KBr, cm⁻¹): 3300(sbr); 1682(m); 1583(s); 1400(s); 1379(sbr).

Crystallography. Crystallographic data for 1 was collected at 100K at XALOC beamline at ALBA synchrotron (λ = 0.79474 Å). Data were indexed, integrated and scaled using the XDS program and iMOSFLM programs. Absorption corrections were not applied. Single crystal X-Ray diffraction for 2 was collected at 298 K on a Bruker AXS SMART Apex diffractometer using graphite monochromated Mo-Kα radiation (λ = 0.71073 Å) and were corrected for Lorentz and polarisation effects. The frames were integrated with the Bruker SAINT software package. Absorption corrections were applied using the program SADABS giving max./min. transmission factors of 1.000/0.635. Both structures were solved by direct methods and subsequently refined by correction of F² against all reflections, using SHELXS2013 and SHELXL2013 within the WinGX package. All non-hydrogen atoms were refined with anisotropic thermal parameters by full-matrix least-squares calculations on F² using the program SHELXL2013. Hydrogen atoms were inserted at calculated positions and constrained with isotropic thermal parameters, except for the hydrogen atoms of N-H and –NH₂ groups and the water molecules in 2. The hydrogen atoms of the adenine moiety and the water molecules were located from difference Fourier maps, fixed and isotropically refined. By contrary, in the crystal structure of 1, the disorder associated with the oxygen atoms of the four crystallization water molecules hindered the localization of their hydrogen atoms. In 1, seven distorted atoms were found (one Co, five oxygen carboxylic atoms and one carbon) and the position of the atoms in adenine moiety was fixed and refined using DELU, EADP and SIMU instructions. Several reflections, for which I(obs) and I(calc) were differing more than 10 times SigmaW, were removed from the refinement. On the other hand, the alerts observed during the check cif can be explained by the quality of the crystal. The best crystal was chosen and attempts to obtain better data were unsuccessful. Although the diffraction experiments proceeded at 100 K under synchrotron radiation, the disorder of the structure could not be reduced and the values of the weighted R factor and wR2 remained high. Moreover, the short Inter D...A contact between the O5A...O6B and O5A...O6A atoms can be explained by the disorder associated with these O5 and O6. Crystal data of 1 and 2 and structure refinement data are listed in Table 1. Main bond distances and angles, and H-bond parameters of 1 and 2 are listed in Tables S1-S4 (ESI†).

| Table 1. Crystal and Structure Refinement data. |
|------------------------------ | ----------------- | ----------------- |
| Compound | 1 | 2 |
| Empirical formula | C₉H₇N₃O₃Co₂ | C₁₀H₅N₃O₄Co₂ |
| Formula weight | 1686.05 | 629.23 |
| Crystal system | Monoclinic | Monoclinic |
| Space group | P2₁/n | P2₁/n |
| CCDC ref | 1043962 | 1043963 |
| Unit cell dimensions | | |
| a (Å) | 9.77 | 9.47 |
| b (Å) | 17.33 | 12.58 |
| c (Å) | 22.71 | 18.98 |
| β (deg) | 100.9 | 93.5 |
| V (Å³) | 3775.8 | 2256.0 |
| Z | 2 | 4 |
| F (000) | 1710 | 1280 |
| θ range (°) | 1.66-53.82 | 1.94-28.39 |
| Ind refln (Rint) | 0.01349 | 0.10653 |
| Final R indices | R₁ = 0.01272 | R₁ = 0.0573 |
| [I > 2σ(I)] | wR² = 0.4590 | wR² = 0.1220 |

| Crystallography. Crystallographic data for 1 was collected at 100K at XALOC beamline at ALBA synchrotron (λ = 0.79474 Å). Data were indexed, integrated and scaled using the XDS and iMOSFLM programs. Absorption corrections were not applied. Single crystal X-Ray diffraction for 2 was collected at 298 K on a Bruker AXS SMART Apex diffractometer using graphite monochromated Mo-Kα radiation (λ = 0.71073 Å) and were corrected for Lorentz and polarisation effects. The frames were integrated with the Bruker SAINT software package. Absorption corrections were applied using the program SADABS giving max./min. transmission factors of 1.000/0.635. Both structures were solved by direct methods and subsequently refined by correction of F² against all reflections, using SHELXS2013 and SHELXL2013 within the WinGX package. All non-hydrogen atoms were refined with anisotropic thermal parameters by full-matrix least-squares calculations on F² using the program SHELXL2013. Hydrogen atoms were inserted at calculated positions and constrained with isotropic thermal parameters, except for the hydrogen atoms of N-H and –NH₂ groups and the water molecules in 2. The hydrogen atoms of the adenine moiety and the water molecules were located from difference Fourier maps, fixed and isotropically refined. By contrary, in the crystal structure of 1, the disorder associated with the oxygen atoms of the four crystallization water molecules hindered the localization of their hydrogen atoms. In 1, seven distorted atoms were found (one Co, five oxygen carboxylic atoms and one carbon) and the position of the atoms in adenine moiety was fixed and refined using DELU, EADP and SIMU instructions. Several reflections, for which I(obs) and I(calc) were differing more than 10 times SigmaW, were removed from the refinement. On the other hand, the alerts observed during the check cif can be explained by the quality of the crystal. The best crystal was chosen and attempts to obtain better data were unsuccessful. Although the diffraction experiments proceeded at 100 K under synchrotron radiation, the disorder of the structure could not be reduced and the values of the weighted R factor and wR2 remained high. Moreover, the short Inter D...A contact between the O5A...O6B and O5A...O6A atoms can be explained by the disorder associated with these O5 and O6. Crystal data of 1 and 2 and structure refinement data are listed in Table 1. Main bond distances and angles, and H-bond parameters of 1 and 2 are listed in Tables S1-S4 (ESI†). |
RESULT AND DISCUSSION

Synthesis and Characterization of 1 and 2. Hydrothermal reactions between CoCO$_3$, a basic metal salt source, adenine and H$_4$TCPB or H$_4$BTeC in water at 100°C for 48 h produced pink block-type crystals of 1 and 2, respectively. The simulated (derived from the single crystal structure of 1 and 2, vide infra) and experimental powder X-ray diffraction (PXRD) patterns were consistent (Figure S1, ESI†); confirming that 1 and 2 can be obtained as pure phases (as also evidenced by elemental analysis and SEM; Figure S2, ESI†). Thermogravimetric analysis (TGA) of 1 revealed multiple weight losses in the range of 315-500 ºC, corresponding to its decomposition (Figure S4; ESI†). In 2, the TGA profile indicates a weight loss of 17.5% (from 25 to 150 ºC), which we attributed to the loss of the remaining µ-H$_2$O molecule (essential to maintain the dinuclear cluster, vide infra), and occurs in a single step, forming CoO.

Crystal Structure Analysis of 1. In 1, there are two crystallographically independent Co(II) ions, which are both octahedrally coordinated forming a trinuclear cluster in the order of Co#1–Co#2–Co#1 (Figure 1a). The Co#1-Co#2 distance within this cluster is 3.62 Å. Co#1 is (NO$_5$)-octahedrally coordinated to the imidazolate N$_9$ atom of adenine and five carboxylate O atoms from three adjacent TCPB linkers. Co#2 is (N$_2$O$_4$)-octahedrally coordinated to the pyrimidinate N$_3$ atoms of two adjacent adenine ligands and to four neighboring TCPB linkers. Here, adenine is acting as a bridging ligand, linking the two Co#1 and Co#2 ions within the trinuclear cluster through the N$_3$ and N$_9$ sites (Figure 1a). The three-dimensional structure of 1 is constructed by the connection of these clusters through TCPB linkers. Within the asymmetric unit of 1, there are two distinct TCPB linkers: i) a TCPB$^-$ linker fully deprotonated that binds to six Co(II) ions with κ$^2$O,O′ and μ-κ$^2$O,O′:κO′ coordination modes (Figure 1b; Figure S5, ESI†); and ii) a H$_2$TCPB$^-$ linker partially protonated, in which two para-benzoate groups coordinate to four Co(II) ions with a μ-κO:κO′ coordination mode (Figure 1c; Figure S5, ESI†) and the other two protonated para-benzoate groups are hydrogen bonded with H$_2$O molecules, affording a compact 3D structure (Figure 1d).

In order to better understand the crystal structure of 1, a topological analysis was performed using TOPOS$^3$ program. As a result, 1 shows a 3D binodal 4,6-c net where each Co#1–Co#2–Co#1 cluster is connected to ten other clusters through 6 TCPB ligands with distances in the range of 15.77-26.63 Å. Overall, this MOF shows a new topology with the point symbol {4$^3$·6·10·8} [4$^3$·6$^2$] (Figure 2a).
Crystal Structure Analysis of 2. The basic unit of 2 is a binuclear Co#1-Co#2 cluster, in which both Co(II) ions exhibit (NO$_3$)-octahedral geometries, although with different coordination environments (Figure 3a). Co#1 is coordinated to the pyrimidinate N$_3$ atom of adenine, four carboxylate O atoms from four adjacent BTeC$_4$-ligands, and one μ-H$_2$O molecule bridging both Co(II) ions. In contrast, Co#2 is coordinated to the imidazolate N$_7$ atom of adenine, two carboxylate O atoms from BTeC$^+$, two terminal H$_2$O molecules and the μ-H$_2$O bridge (Figure 3a). The Co#1-Co#2 distance within this cluster unit is 3.63 Å. Comparison of the bond lengths for Co#1-μ-H$_2$O and Co#2-μ-H$_2$O (2.15 Å and 2.14 Å, respectively), and the angle between Co#1-H$_2$O-Co#2 (115.8º), with corresponding literature values for structures...
containing the same metal M-O-M unit (CSD codes: ZALXAL, UMAUGK).28-31 confirmed the presence of the H2O bridge (Table S4, ESI†). Adenine linkers bridge the Co#1-Co#2 clusters involving the N3 and N6 sites, thus forming 1D chains running along the c-axis (Figure 3b).

The use of the tetracarboxylate BTEC+ linkers was found to be prolific for the expansion of 2 in three dimensions, as each BTEC+ links four neighboring chains (one from each carboxylate site) (Figure 3c; Figure S6, ESI†). In particular, each BTEC+ binds to six Co(II) centers in two coordination modes: a monodentate χO mode, in binding Co#1 from the carboxylate positions 1 and 4; and a bidentate μ-χO:χO’ mode, in bridging Co#1 and Co#2 from the carboxylate positions 2 and 5. Framework 2 generates regular 1D channels (5.3 x 4.8 Å; 609.1 Å³; 27% of void space in unit cell)34 along the b-axis (Figure S7, ESI†). Figure 3d shows that the channels accommodate four guest H2O molecules per formula unit. Importantly, the amino -NH2 and the imidazolate -NH of adenine that reside within the channel establish H-bond networks with water molecules and oxygen atoms of the carboxylic groups. Moreover, the N4 pyridiminate nitrogen atom acts as acceptor of H-bond interactions established with water molecules (see all H-bonds in Table S5, ESI†). The theoretical pore volume of 2 calculated from the crystallographic structure after removing the guest H2O molecules is 0.176 cm³/g. In this case, the topological analysis showed that 2 presents the known fsc net (Figure 2b).

Adenine in CPs. An analysis of extended structures containing adenine and transition metal ions obtained from the Cambridge Structural Database (CSD; Allen, 2002) illustrates the rich coordination modes that can adopt adenine for constructing CPs. From this study, we found that adenine can act as a simple monodentate ligand, mainly through the N3 or N6 site.35-37 Examples of these coordination modes were observed by García-Terán et al. in 1D Zn(II)- and Cu(II)-oxalate chains, in which adenine coordinates these metal ions only through the N3 and N6 site, respectively (Figures 4a and 4b).

Adenine can also act as a bidentate ligand; having here a wide variety of combinations depending on the N sites that participate in the metal binding. Overall, we have detected that adenine can coordinate to two metal ions through the: i) [N3, N6];38,39 ii) [N7, N8];39,42 iii) [N5, N3];43 iv) [N5, N6];39 and v) [N5, N7] (in 2) sites. An example of the first coordination mode was reported by Stylianou et al. in [Ni(PZDC)2(ade)2(H2O)4][H2O]3 (where PZDC = 3,5-pyrazoledicarboxylate).38 In this structure, adenine participates in the formation of Ni(II) dimers by bridging two Ni(II) ions through the N3 and N6 sites (Figure 4c). These Ni(II) units are then connected through PZDC linkers, forming chains that are 3D hydrogen-bonded. The second coordination mode was observed in [Zn(ade)(INT)]·DMF (where INT = isonicotinate; DMF = dimethylformamide).40 In this compound, adenine connects Zn(II) ions through the N7 and N6 sites (Figure 4d), creating infinite 1D helices that are 3D linked by the INT linkers. Sanjib Das et al. described the third coordination mode in a 1D Cu(II) CP with formula [Cu(glycine)(ade)(NO3)(H2O)]. Here, adenine connects two Cu(II) ions through the N4 and N5 sites (Figure 4e), creating zig-zag chains that are 3D hydrogen bonded.40 An example of the last reported [N6, N7] coordination mode (Figure 4f) can be found in [Cd(ade)(IPA)(H2O)] (where IPA = isophthalate). In this 2D CP, adenine forms chains binding Cd(II) ions through the N4 and N5 sites, which are further 2D connected by the IPA linkers.39

In our reported extended structures 1 and 2, adenine also acts as a bidentate ligand coordinating to two Co(II) ions. In 1, adenine bridges two Co(II) ions through the N5 and N8 sites forming a trinuclear Co(II) cluster (Figure 4c). This coordination mode is very similar to the one observed by Stylianou et al.48 In contrast, adenine exhibits a rare coordination mode through the N5 and N7 sites in 2 (Figure 4g). To the best of our knowledge, this coordination mode was only observed in two previous discrete Cu(II)-based complexes.44-45 Through this [N5, N7] coordination mode, adenine participates in the formation of infinite Co(II)-adenine chains that are 3D linked by BTEC+ linkers.

Besides of acting as a bidentate ligand, adenine can also coordinate via more than two N sites. In fact, the most common coordination mode found in CPs is [N5, N7, N9].38,46-54 For example, adenine coordinates to three Cu(II) ions via the N5, N7, and N9 sites (Figure 4h) in a porous CP with formula [Cu3(ade)2(H2O)5][Cu(oxalate)(H2O)]2·14H2O.46 In this CP, four adenine ligands participate in the formation of a paddle wheel unit by bridging two Cu(II) ions through the N3 and N5 sites. These units are then connected through mononuclear entities ([Cu(oxalate)(H2O)] via the N2 site. More unusual coordination modes for adenine are [N6, N9, N9] and [N5, N5, N5]. An example of the [N6, N9, N9] coordination mode can be found in [(CH3Hg)2(ade)·EtOH (Figure 4i).54 In this structure, two CH3Hg+ ions are coordinated through the N6 and N9 sites, whereas another CH3Hg+ ion is coordinated via the N6 site forming 1D-chains. The other coordination mode was observed in [(CH3Hg)2(ade)(NO3)].55 In this case, four CH3Hg+ ions are linked by the N5, N6, and N9 sites (Figure 4i), creating a 2D structure.

Finally, adenine can also coordinate four metal ions via the N5, N9, N7, and N9 sites.56,57 An example of this coordination mode was reported by An et al. (Figure 4k) in [Zn(ade)4(BPDC)2O-2MeNH2]·8DMF·nH2O (where BPDC = biphenyl-edicarboxylate; bio-MOF-1).7 In this structure, Zn(II) ions and adenine coordinate forming chains composed of apex-sharing octahedral cages. These chains are interconnected via multiple BPDC linkers, expanding the structure to a 3D porous network.
Hydrochromic Behavior of 2. We observed that when 2 was heated at 150 ºC, a new dehydrated phase, $[\text{Co}_2(\text{ade})(\text{BTeC})(\text{H}_2\text{O})] \ (2')$, was formed, as evidenced by comparing the TGA profiles of 2 and 2' (Figures 5b). During this transformation, the crystals of 2 were cracked and changed color from pink to purple (Figure 5a). This is thought to be due to the change in coordination of Co#2, from octahedral to tetrahedral, which resulted from the loss of the coordinated H$_2$O molecules. The UV/Vis spectra revealed a red shift in the d-d transition of the Co(II), from $\sim 515$ nm (2) to $\sim 529$ nm (2'), and the absorption band broadened (Figure 5a). The sensitivity of these metal-based transitions to the geometry change, which occurs upon conversion of the Co(II) centers from six to mixed six/four coordination, is reminiscent of the color variations observed for other frameworks in which open transition metal sites can be generated.$^{58-60}$ Unfortunately, it was not possible to obtain more structural details of this transformation by determining the structure of 2' with single crystal X-ray diffraction, but its PXRD pattern showed loss of crystallinity to a high degree (Figure 5c). This is indicative that structural re-arrangements occurred upon desolvation and that 2' is highly disordered.

In order to assess the porosity of 2' and check the influence of the structural re-arrangements upon desolvation, N$_2$ and CO$_2$ isotherms collected at 77 K and 195 K, respectively, up to 1 bar (Figures S5o, S5u, ESI†). 2' adsorbs very small amounts of CO$_2$ (4.18 cm$^3$/g at 1 bar) with a BET surface area of 8.11 m$^2$/g, and the N$_2$ isotherm shows a characteristic type II shape confirming the non-porous behavior of 2'. This observation suggests that upon activation, dramatic structural changes occurred.

The structural changes observed between hydrated and dehydrated materials encouraged us to deeply analyze this hydrochromic behavior. For this, we studied how 2' responds to the presence of liquid H$_2$O and organic solvents. 2' was initially immersed in liquid H$_2$O for different periods of time, and the evolution of the structure was monitored by PXRD and UV/Vis spectroscopy. Notably, after 24 hours, the purple color of 2' had changed to pink and the corresponding absorption band had blue-shifted to 515 nm (Figure 5a; Figure S8, ESI†). The PXRD patterns showed an increase in crystallinity with time, which had stabilized after 2 days (Figure 5c). After 3 days, both the main reflection at $2\theta = 12.0^\circ$ and the absorption band at 515 nm, were essentially recovered compared with those observed for the newly synthesized 2 (Figures 5a-c).

**Figure 4.** Representation of the coordination modes of adenine observed in extended structures.
This recovery confirmed a large degree of reversibility for the 
H$_2$O-induced structural changes: the H$_2$O molecules can
coordinate to the Co#2 center, regenerating its octahedral
coordination geometry, and H$_2$O molecules can
diffuse within the structure, re-establishing the H-bonds between
them and the functional groups (mainly of the adenine), as
evidenced by elemental analysis (ESI†). The consistency an
d reversibility of these H$_2$O-induced structural changes were
confirmed by exposing a sample of 2 to three cycles of
alternate dehydration (heating the sample at 150 °C for
30 minutes) and rehydration (soaking in H$_2$O for 48 hours).
Figure 5d illustrates the alternating dehydration/hydration
behavior measured in 2, with reversible absorption band
shifts from 515 nm (2), to 529 (2'). Based on this observation
(H$_2$O importance for the recovery of 2), we checked if 2 could
be recovered through the formation of these specific H-
bonding interactions between the framework with other
polar and apolar solvents. Thus, the violet solid (2') was
soaked in liquid methanol, ethanol, dimethylformamide,
chloroform and hexane (Figure Sg, ESI†). It is found that the
violet color of 2' was retained after three days of treatment
with all the solvents suggesting the well-organized structure
of 2 was not formed. The PXRD of the methanol loaded
sample confirmed that 2 cannot be recovered, indicating that
the specific interactions between H$_2$O molecules and host
framework are key factors for retaining the structural
integrity of 2.\textsuperscript{13}

**Conclusions**

In conclusion, we have prepared two new 3D CPs assembled
from Co(II), adenine and aromatic tetracarboxylic acids.
From a structural point of view, 1 shows a compact 3D
binodal 4,6-c net with a new topology with the point symbol
{4^4.6^4.10^8}{4^4.6^2}. 2 shows a fsc net, in which adenine acts
as a bidentate linker through a rare coordination mode; that
is, through the N$_3$ and N$_7$ sites. In addition, 2 exhibits a
clearly observable and reversible color change from pink (2)
to purple (2'), and vice versa, upon dehydration and
hydration, respectively, indicative that structural re-
arangements occurred upon activation. This has a direct
impact on its sorption behavior as it is non porous to CO$_2$
or N$_2$. Contrariwise, we found that H$_2$O molecules can diffuse
within the framework and organize the 3D open structure of
2. Selective structural recovery is likely driven by the re-
establishment of the H-bonds occurred between H$_2$O
molecules and adenine in the host framework.

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**Figure 5.** (a) UV/vis spectra of 2 (pink), 2' (purple) and regeneration of 2 after the immersion of 2' in liquid H$_2$O (green).
Inset: color change from pink (2) to purple (2') upon dehydration and hydration, (b) TGA diagrams of 2 (pink) and 2' (purple), (c) PXRD study showing the re-generation of 2 upon immersion of 2' in liquid H$_2$O after three days: 2 (pink), 2' (purple) and 2' immersed in water for 1 day (orange), 2 days (grey) and 3 days (green), and (d) 2 can be activated and regenerated for at least three times as revealed by UV/vis spectroscopy retaining the comparable maximum absorption
wavenumber (and also followed by the characteristic color change).
Ramos, J. D.; Covelo, B.; Carballo, R.; Castiñeiras, A.; Niclós-Gutiérrez, J., *Inorg. Chim. Acta* 2002, 339, 160-170.

(46) García-Terán, J. P.; Castillo, O.; Luque, A.; García-Couceiro, U.; Román, P.; Lezama, L., *Inorg. Chem. 2004*, 43, 4549-4551.

(47) Li, T.; Chen, D.-L.; Sullivan, J. E.; Kozlowski, M. T.; Johnson, J. K.; Rosi, N. L., *Chem. Sci. 2013*, 4, 1746-1755.

(48) Hubert, J.; Beauchamp, A. L., *Can. J. Chem. 1980*, 58, 1439-1443.

(49) Charland, J.-P.; Britten, J. F.; Beauchamp, A. L., *Inorg. Chim. Acta* 1986, 124, 161-167.

(50) Pérez-Yáñez, S.; Beobide, G.; Castillo, O.; Cepeda, J.; Luque, A.; Aguayo, A. T.; Román, P., *Inorg. Chem.* 2011, 50, 5330-5332.

(51) Pérez-Yáñez, S.; Beobide, G.; Castillo, O.; Cepeda, J.; Luque, A.; Román, P., *Cryst. Growth Des.* 2012, 12, 3324-3334.

(52) Yang, E.-C.; Zhao, H.-K.; Ding, B.; Wang, X.-G.; Zhao, X.-J., *New J. Chem.* 2007, 31, 1887-1890.

(53) Wang, F.; Kang, Y., *Inorg. Chem. Commun.* 2012, 20, 266-268.

(54) Charland, J.-P., *Inorg. Chim. Acta* 1987, 135, 191-196.

(55) Charland, J. P.; Beauchamp, A. L., *Inorg. Chem.* 1986, 25, 4870-4876.

(56) Yang, E.-C.; Zhao, H.-K.; Feng, Y.; Zhao, X.-J., *Inorg. Chem.* 2009, 48, 3511-3513.

(57) An, J.; Farha, O. K.; Hupp, J. T.; Pohl, E.; Yeh, J. I.; Rosi, N. L., *Nat Commun* 2012, 3, 604.

(58) Uemura, K.; Kitagawa, S.; Kondo, M.; Fukui, K.; Kitaura, R.; Chang H.-C.; Mizutani, T., *Chem. Eur. J. 2002*, 8, 3587-3600.

(59) Rueff, J.-M.; Pillet, S.; Bonaventure, G.; Souhassou, M.; Rabu, P., *Eur. J. Inorg. Chem.* 2003, 4173-4178.

(60) Chen, Q.; Chang, Z.; Song, W.-C.; Song, H.; Hu, T.-L.; Bu, X.-H., *Angew. Chem. Int. Ed.* 2013, 52, 11550-11553.
Title: Two New Adenine-based Co(II) Coordination Polymers: Synthesis, Crystal Structure, Coordination Modes and Reversible Hydrochromic Behavior

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Two new 3D coordination polymers (CPs) based on Co(II), adenine and aromatic tetracarboxylate linkers and an analysis of the coordination modes of adenine found in extended coordination structures are reported. Interestingly, one of these CPs shows a reversible hydrochromic behavior, exhibiting a detectable colour change from pink to purple, and vice versa, upon hydration and dehydration, respectively.