** Shedding Light on the Protonation States and Location of Protonated N Atoms of Adenine in Metal–Organic Frameworks **

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ABSTRACT: We report the syntheses and structures of five metal–organic frameworks (MOFs) based on transition metals (NiII, CuII, and ZnII), adenine, and di-, tri-, and tetra-carboxylate ligands. Adenine, with multiple N donor sites, was found to coordinate to the metal centers in different binding modes including bidentate (through N7 and N9, or N3 and N9) and tridentate (through N3, N7, and N9). Systematic investigations of the protonation states of adenine in each MOF structure via X-ray photoelectron spectroscopy revealed that adenine can be selectively protonated through N1, N3, or N7. The positions of H atoms connected to the N atoms were found from the electron density maps, and further supported by the study of C–N–C bond angles compared to the literature reports. DFT calculations were performed to geometrically optimize and energetically assess the structures simulated with different protonation modes. The present study highlights the rich coordination chemistry of adenine and provides a method for the determination of its protonation states and the location of protonated N atoms of adenine within MOFs, a task that would be challenging in complicated adenine-based MOF structures.

INTRODUCTION

Metal–organic frameworks (MOFs) are crystalline materials consisting of an infinite network of metal ions or clusters bridged by organic ligands through coordination bonds into one-, two-, or three-dimensional extended structures. Highly porous MOFs have various topologies, compositions, and properties have been reported, including MOF materials with high internal surface areas and void volumes, and remarkable properties have been reported, including MOF materials with various topologies, compositions, and applications in areas such as magnetism, sensing, gas separation, and catalysis. Canonical nucleobases are often classified into two subgroups based on their structures: purines (adenine and guanine) and pyrimidines (cytosine, thymine, and uracil). With the higher number of heteroatoms, the purine nucleobases are the better bridging ligands than the pyrimidine nucleobases, especially with adenine being extensively used for forming complexes and networks with transition metals.

Adenine (9HAde, Figure 1) is a rigid ligand with five potential coordination sites for metal binding, i.e., two imidazolate, two pyrimidinate N atoms, and an –NH2 group. The basicity order of the N atoms is N9 > N1 > N7 > N3 ≫ N10, and the pKₐ of 9HAde is 9.8. Due to the 9HAde’s imidazolate functionality with the coordination bridging mode of μ₂-N7,N9, several MOFs with zeolitic topologies have been reported. However, due to the high versatility of adenine, which displays a variety of monodentate and bridging bidentate or tridentate metal coordination modes, an accurate prediction of Ade-based MOFs structures is often impossible. This, in fact, has been the impetus for the discovery of a large number of both porous and nonporous Ade-based MOFs with diverse structures and topologies that have been reported in the literature over the past few years.

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CH4 selectivity,28 tunable porosity,29,31,35 and properties such as mesoporosity with low densities, high surface areas, and large pore volumes.32 Application in controlled drug delivery has also been explored.36 Despite the high utility of adenine in MOF synthesis and due to the different possible protonation states of adenine (H2Ade+, HADE, and Ade−), assigning the protonated N atoms and finding the idealized rotating groups.37 Since the rule is mainly valid for noncoordinated aromatic H atoms and to localize the protonated N atoms of adenine in different possible binding modes of adenine ligands within these materials will be discussed based on their crystal structures, whereas their protonation states and the location of the protonated N atoms are reviewed through a detailed analysis of the XPS data and C−N−C angles.

**EXPERIMENTAL SECTION**

**Synthesis.** The reagents, including adenine (9H Ade), 1,3,5-benzenetribacrylic acid (Hbtc), isophthalic acid (H2ipa), 3,5-pyrazoledicarboxylic acid (Hpzdc), 1,2,4,5-benzenetetraacrylic acid (H4btec), NiCO3, CuCO3, and ZnCO3, were purchased from Sigma-Aldrich and used as received.

**Synthesis of [Ni2(btec)(7HAde)2(H2O)2]** (SION-35). A mixture of NiCO3 (71 mg, 0.40 mmol), Hbtc (63 mg, 0.30 mmol), 9H Ade (40 mg, 0.30 mmol), and water (6 mL) was placed in a Teflon reactor. The mixture was heated at 160 °C for 4 days, and then gradually cooled to room temperature at a rate of 0.1 °C/min. Green block-type crystals of SION-35 were obtained in 43% yield (based on NiCO3). Anal. Calcld for ([Ni2(C9H3O6)(C5H4N5)(H2O)2]·(H2O)0.2): C 43.26, H 2.60, N 19.41; found: C 43.11, H 2.92, N 19.36.

**Synthesis of [Ni2(btc)(Ade)(H2O)5]** (SION-31). A mixture of NiCO3 (54 mg, 0.40 mmol), Hbtc (84 mg, 0.40 mmol), 9H Ade (54 mg, 0.40 mmol), and water (6 mL) was placed in a Teflon reactor. The mixture was heated at 150 °C for 72 h, and then gradually cooled to room temperature at a rate of 0.3 °C/min. Colorless needle crystals of SION-35 were obtained in 22% yield (based on ZnCO3). Anal. Calcld for ([Cu2(C9H3O6)(C5H4N5)2(H2O)2]·(H2O)0.7): C 32.45, H 3.39, N 13.52; found: C 32.68, H 3.42, N 13.33.

**Single-Crystal X-ray Diffraction (SCXRD).** SCXRD data of SION-31 were collected on a Bruker AXS SMART Apex diffractometer equipped with a CCD detector. The X-ray beam was generated using graphite monochromated Mo Kα radiation from a molybdenum X-ray tube operating at 40 kV and 30 mA. A single crystal was immersed in silicone oil, and picked with a polymide loop which in turn was mounted on a goniometer. Diffraction data were collected at 100.0 K. Single crystals of SION-32, SION-33, SION-34, and SION-35 were measured using the synchrotron radiation at the European Synchrotron Radiation Facility, Beamline BM01. Suitable single crystals were selected and mounted onto the diffractometer equipped with a CCD area detector. The crystals were kept at 100.0 K during data collection. Using Olex2,38 their structures were solved with the ShelXT structure solution program using Intrinsic Phasing and refined with the ShelXL refinement package using Least Squares minimization.40 Atomic positions were found from the difference-Fourier maps and refined anisotropically for all non-H atoms. Positions of aromatic H atoms were refined using a riding model, while H atoms in methyl and hydroxy groups were refined as fixed with eq of atoms in terminating groups.

**Powder X-ray Diffraction (PXRD).** PXRD data of SION-31, SION-32, SION-33, SION-34, and SION-35 were collected with Cu Kα radiation at 298 K. Their phase purity was confirmed by the comparison of the simulated PXRD patterns (derived from the Mercury software) to the experimental ones.

**X-ray Photoelectron Spectroscopy (XPS).** XPS spectra of SION-31, SION-32, SION-33, SION-34, and SION-35 were recorded on a Scienta ESCA 300 spectrometer located at the NCESS facility at Daresbury Laboratory, U.K., which incorporated a rotating anode Al Kα (hv = 1486.6 eV) X-ray source and had an effective instrument resolution of 400 meV. The spectrometer was...
Figure 2. SION-31: (a) Depiction of the Ni-Ade secondary building unit (SBU) \([\text{Ni}_2(\text{Aden})(\text{H}_2\text{O})_4(\mu_2-\text{H}_2\text{O})_2]\)^{2+}, with Ade^{-} acting as the bridging ligand via N3, N7, and N9, linking the Ni-Ade SBU with Ni2. (b) Coordination environment of Ni2; the carboxylate group of btec is not involved in coordination to Ni^{2+} and is enclosed in a red ellipse. (c) Mono- and bidentate coordination modes of the btec ligand. (d) The Ni-Ade SBU (colored in green) connects the btc-based chains (orange zigzag line), resulting in a two-dimensional layered structure. Color scheme: Ni, green; C, gray; O, red; N, blue; and H, yellow.

RESULTS AND DISCUSSION

Synthesis and Characterization. The syntheses of SION-31, SION-32, SION-33, SION-34, and SION-35 were all performed under hydrothermal reaction conditions, with the reactants being carbonate metal salts, adenine, and different carboxylate ligands in pure water. The molar ratios of the reactants were screened to obtain phase-pure products. Green block type crystals of SION-31 were formed as a pure phase, with the formula of \([\text{Ni}_2(\text{btc})(\text{Aden})(\mu_2-\text{H}_2\text{O})(\text{H}_2\text{O})_4]\)_2·3\(\text{H}_2\text{O}\) as determined by SCXRD (vide infra). This is well in agreement with the molar ratio of the reactants NiCO_3:H_2btc:9HAde of 2:1:1 in the reaction. When H_2ipa, a dicarboxylic acid, is used instead of H_3btc, SION-32 was formed, with the formula of \([\text{Ni}(\text{ipa})(\text{3Aden})]\). It is worth noting that the use of a slight excess NiCO_3 and 9HAde (molar ratio of NiCO_3:H_2ipa:9HAde of 3:2:3) is crucial to obtain the pure phase of SION-32. Similarly, in the case of SION-33, \([\text{Ni}_2(\text{btc})(7\text{Aden})(\text{H}_2\text{O})_2]_2\cdot4\text{H}_2\text{O}\), the slight excess of the tetracarboxylic acid H_4btc and adenine (molar ratio of NiCO_3:H_4btc:9HAde of 2:3:3) is the key for the formation of the product.

Since different MOF structures were obtained with different carboxylate ligands, the metal ion and/or the carboxylate ligands were changed to further explore the versatility of this reaction scheme. SION-34, \([\text{Cu}_3(\text{pzdc})(\text{1Aden})(\text{H}_2\text{O})_2]_2\), formed when CuCO_3 and H_3pzdc were used, while SION-35, \([\text{Zn}_2(\text{btc})(\text{Aden})(\text{H}_2\text{O})_2]_2\), was the product when ZnCO_3 was combined with H_2btc.

The bulk phase purity of SION-31, SION-32, SION-33, SION-34, and SION-35 was confirmed by PXRD, with both the theoretical and the experimental X-ray powder diffraction patterns well in agreement (Figure S1). Their purity was further confirmed through elemental analysis (see the Experimental Section) and SEM images (Figures S2–S4).

The thermal stability of SION-31, SION-32, SION-33, SION-34 and SION-35 was also investigated through TGA analysis. As illustrated in Figure S5, the TGA profiles of SION-31, SION-32, SION-33, SION-34, and SION-35 follow the same trend, showing an initial weight loss corresponding to the removal of the lattice H_2O molecules, followed by the release of the metal bound H_2O molecules at higher temperature (Table S2). The weight loss in SION-31 occurred in two steps: i. the first step corresponds to the loss of the guest H_2O molecules in the temperature range of 30–130 °C; ii. the second step is attributed to the loss of the coordinated H_2O molecules in the temperature range of 130–250 °C. A total weight loss of 22.5% is in agreement with the loss calculated from elemental analysis (23.4%). Decomposition of the framework starts at 400 °C. SION-32 shows no weight loss as there are no bound H_2O molecules to the metal center and the cavities are too small to accommodate any H_2O molecules. This is consistent with the.
elemental analysis as it was found that there are 0.2 H2O guest molecules per formula unit comprising 0.4% of the total weight of the structure. The TGA profile of SION-32 shows its excellent stability up to 450 °C. The same trend with the profile for SION-31 was also observed for SION-33. The final weight loss of 14.4% is slightly larger than the one calculated from elemental analysis (11.3%). TGA for SION-34 shows that guest and coordinated H2O molecules were removed in the temperature range of 30–280 °C with the total weight loss being 8.1%. This weight loss observed in TGA is slightly lower than that calculated from elemental analysis (10.0%). Decomposition of the framework starts at 295 °C. Finally, the TGA profile of SION-35 shows that it is stable up to 210 °C, and in the temperature range of 210–370 °C, the coordinated H2O molecules are removed. The weight loss of 8.4% is in agreement with the calculated proportion (9.1%) of the H2O molecules bound to ZnII from the elemental analysis. 8.4% is in agreement with the calculated proportion (9.1%) of the H2O molecules bound to ZnII from the elemental analysis. The same trend with the profile for SION-31 was also observed for SION-33. The final weight loss of 14.4% is slightly larger than the one calculated from elemental analysis (11.3%). TGA for SION-34 shows that guest and coordinated H2O molecules were removed in the temperature range of 30–280 °C with the total weight loss being 8.1%. This weight loss observed in TGA is slightly lower than that calculated from elemental analysis (10.0%). Decomposition of the framework starts at 295 °C. Finally, the TGA profile of SION-35 shows that it is stable up to 210 °C, and in the temperature range of 210–370 °C, the coordinated H2O molecules are removed. The weight loss of 8.4% is in agreement with the calculated proportion (9.1%) of the H2O molecules bound to ZnII from the elemental analysis. 

**Single-Crystal X-ray Diffraction Analysis.** SION-31 crystallizes in the monoclinic space group P21/n. The asymmetric unit (ASU) contains two symmetrically independent NiII environments (Ni1 and Ni2) which are linked via the btc and adenine ligands. Two carboxylate groups of the btc ligand coordinate to the NiII ions, while the third site is free of any metal binding. Since the bond distances between the C and O atoms, C16–O9 and C16–O10, are comparable (1.263(3) Å and 1.273(3) Å, respectively) and intermediate between the values of 1.35 and 1.21 Å characteristic for, respectively, single and double C(sp2)–O bonds, this site is a deprotonated carboxylate group; i.e., the btc ligand is btc3−. The charge is balanced with the adenine ligand in its anionic form Ade−. Two NiII ions are part of a dimeric unit, which is bridged by two Ade− ligands via N3 and N9 and two μ2-O2H4 molecules to form a Ni-Ade secondary building unit (SBU) [Ni2(Aden)(H2O)2(μ2-O2H4)]2+ (Figure 2a). The octahedral coordination sphere of each Ni1 ion is filled by two terminal H2O ligands. The bond lengths of the Ni–Oterminal H2O bonds are 2.085(2) Å (Ni1–O2), 2.020(2) Å (Ni1–O3), while the bond lengths of the Ni–Obridging H2O are 2.1793(17) Å and 2.1234(16) Å (Ni1–O1). These bond distances are consistent with those reported in the literature ([Ni2(O2CFcCO2)2(2,2′-bpy)(μ2-OH2)2]·CH3OH·2H2O:52 Ni–OH, 2.107(9)/2.138(3) Å, [Ni2(H2O)bdptz](OTs)2·2CH3OH·H2O:53 Ni–OH 2.159(1)/2.170(4) Å), and are strikingly different from the bonds reported for bridging hydroxyl ligands (bis(1,5-mesityl-3-nitroformazanato)-bis-μ-hydroxonickel:54 Ni–OH 1.890(7)/1.884(4) Å). The Ni2 ion also adopts an octahedral coordination geometry, with one chelating and one monodentate btc3−, one monodentate Ade− via N7, and two terminal H2O molecules as ligands (Figure 2b). SION-31 consists of two-dimensional sheets constructed by connecting the [Ni2(Aden)2(H2O)4(μ2-O2H4)]2+ units augmented at two extremities with Ni2 atoms (Figure 2a) via two deprotonated carboxylate groups of the btc3− ligands (Figure 2c,d). The third carboxylate group of btc3− forms H-bonds with guest and coordinated H2O molecules. The noncoordinated H2O molecules (O1W, O2W, and O3W) are hydrogen bonded with the N1 atom of Ade− and carboxylate O9, O11, and O12 atoms of btc− (O9, O11, and O12) ligands, (O1W···N1 2.712(3) Å, O1W···O11 3.025(3) Å, O2W···O9 2.801(3) Å, O3W···O11 2.618(3) Å), as well as with coordinated water molecules. The structure is close packed and exhibits no accessible voids. 

![Image](image-url)
The coordination of the ligand is disordered as it assumes two equivalent orientations related to one another by a mirror plane. The coordination environment of NiII is of distorted octahedral geometry, with four O atoms from two chelating ipa2⁻ ligands and two N atoms from two disordered 3HAde ligands completing its coordination geometry (Figure 3a). Each of the 3HAde and ipa2⁻ ligands behaves as a bridging linker between the NiII ions, giving rise to a two-dimensional layered structure (Figure 3b,c).

The packing of the two-dimensional layered structure of SION-32 (Figure 3c) reveals that there is no accessible volume, as confirmed by PLATON.56,57 Topological analysis reveals that the two-dimensional layers in SION-32 form a square-lattice net (sql), which possesses single 4-connected nodes centered on the NiII atoms, while the ipa⁺ and the 3HAde ligands represent the edges (Figure S7).

SION-33 crystallizes in the triclinic space group P\(^{\overline{1}}\). The ASU of SION-33 consists of one NiII atom, one neutral 7Hade ligand (protonated via N7, vide infra), half the btec4⁻ ligand, one coordinated and two guest H2O molecules. The 7Hade ligand binds to two chemically equivalent NiII centers through N3 and N9. Two bridging 7Hade molecules coordinate on opposite sides of the NiII unit, forming a “blade” dimer, [Ni\(_2\)(7Hade)\(_2\)\((H_2O)_2\)]\(^{4+}\) (Figure 4a). Two symmetrically independent carboxylate groups of the btec4⁻ ligand bind to NiII in two distinct ways: the first one coordinates to both NiII atoms within the Ni\(_2\) unit in a monodentate mode through O1, whereas the neighboring carboxylate binds solely to one NiII atom through O4 (Figure 4b). A terminal coordinated H2O molecule completes the octahedral coordination environment of NiII. The [Ni\(_2\)(7Hade)\(_2\)\((H_2O)_2\)]\(^{4+}\) dimers connected into one-dimensional chains extend along the b-axis (Figure 4c). The O3W water molecule links these chains along the [001] direction, through O3W···O2 (2.962(13) Å) and O3W···N10 (2.930(13) Å) hydrogen bonds to one such chain, and through the N10···O3W (2.937(15) Å) H-bond to the parallel one. Along [100], there are hydrogen bonds linking subsequent NiII chains directly (O1W···O4, 2.806(8) Å), and by means of the O2W water molecule (N7···O2W, 2.781(10) Å, and O2W···N1, 2.943(9) Å). Therefore, the array of hydrogen bonds combines 1D chains of [Ni\(_2\)(7Hade)\(_2\)\((H_2O)_2\)]\(^{4+}\) units into a three-dimensional supramolecular assembly. The packing and space filling representations of SION-33 show that it is not porous, which is further confirmed by PLATON.56

SION-34 crystallizes in the monoclinic space group P2\(_1\)/c. The ASU consists of three symmetrically independent CuII centers (Cu1, Cu2, and Cu3), two fully deprotonated pzdc3⁻ ligands, one neutral 1HAde ligand (protonated via N1, vide infra), and four coordinated H2O molecules. Cu1 has a square planar geometry and is coordinated to one N9 atom from 1HAde, one pyrazolate N2 and one carboxylate O1 atom from
the same pzdc$^{3-}$ ligand, while the fourth position is occupied by one H$_2$O, with the Cu1–O1W bond distance of 1.920(2) Å (Figure 5a). Cu2 is five coordinated with a tetragonal pyramidal geometry. The equatorial positions of Cu2 are occupied by two carboxylate O atoms (O3 and O5) and two N atoms (N4 and N6) from two adjacent pzdc$^{3-}$ ligands. The axial position is occupied by the O2W atom from a H$_2$O molecule with a bond distance of 2.323(2) Å (Figure 5a). Cu3 is also coordinated by five donor atoms, giving rise to a trigonal bipyramidal coordination geometry. The coordination environment of Cu3 is provided by N12 and O7 atoms from the pzdc$^{3-}$ ligand, one N7 from 1HAde, and two O atoms, O3W and O4W, from two distinct H$_2$O molecules (bond distances of 2.281(2) and 1.969(2) Å, respectively). As shown in the Figure 5a, the connection of Cu1–Cu2–Cu3 via the pzdc$^{3-}$ and 1HAde affords a one-dimensional chain extended along the crystallographic [01] direction. The adjacent one-dimensional chains are interlocked by the π–π stacking interactions between the pyrimidine rings of 1HAde (distance between two 1HAde ligands: 3.231(4) Å) (Figure 5b), and by a system of hydrogen bonds to form a three-dimensional supramolecular array. Water molecules O2W, O3W, and O4W, which protrude from the surface of a nearly planar 1D chain, serve as donors in a range of hydrogen bonds to the carboxylate O atoms of the pzdc$^{3-}$ ligands situated above and below the chain: O2W···O1, 3.028(3) Å; O2W···O7, 2.789(3) Å; O3W···O2, 2.719(3) Å; O3W···O6, 2.705(4) Å; O4W···O4, 2.677(3) Å. In addition, the pyrimidine N1 and amino N10 atoms of the 1HAde ligand provide the chain with linkage to the neighboring chains in the perpendicular direction (N1···O8, 2.785(3) Å; N10···O8, 2.911(3) Å). The dense packing of SION-35 indicates that it is nonporous.

SION-35 crystallizes in the monoclinic space group P2$_1$/c. The ASU contains two independent Zn$^{II}$ ions (Zn1 and Zn2), one btc$^{3-}$, one bidentate Ade$^-$, and two terminal H$_2$O ligands bound to Zn2. Zn1 ions adopt a trigonal bipyramidal geometry, whereas Zn2 ions adopt a distorted tetrahedral coordination geometry. Zn1 and Zn2 are connected via the imidazolate N atoms, N7 and N9, respectively, of the Ade$^-$ ligand (Figure 6a). The completion of the coordination sphere of Zn1 is provided by four carboxylate O atoms from four adjacent btc$^{3-}$ ligands with monodentate binding mode (Figure 6a). Two terminal H$_2$O molecules are coordinated to Zn2, and the fourth position is occupied by a monodentate carboxylate O atom from the btc$^{3-}$ ligand. These two specific coordination geometries of Zn$^{II}$ present in SION-35 are rarely observed in a single MOF. Each btc$^{3-}$ ligand is bound to five Zn$^{II}$ atoms: four Zn1 and one Zn2 (Figure 6b). One carboxylate group of btc$^{3-}$ is coordinated solely to Zn1 in a monodentate mode, whereas each of the other two groups is coordinated through a syn–anti coordination mode to two Zn$^{II}$—to two Zn1 and to both Zn1 and Zn2 (Figure 6b). Both btc$^{3-}$ and Ade$^-$ are packed in a zigzag fashion along the c-axis and as in a flat two-dimensional layer along the b- and a-axes (Figure 6c,d). Through the coordination of the carboxylate O atoms of the btc$^{3-}$ ligand with Zn (Zn1–O4: 2.373(3) Å and Zn1–O1: 2.113(3) Å) complemented by the system of hydrogen bonds, the Zn–Ade–btc layers are linked to each other propagating the structure into 3 dimensions. PLATON software reveals that there is no solvent accessible volume.

To determine the topology of SION-35, small rings consisting of no more than 8 covalent or dative bonds were identified in the material. This includes an 8-membered ring consisting of both Zn1 and Zn2 ions coordinated to two carboxylate groups from separate btc$^{3-}$ ligands. This complex was reduced to a single node connected to 8 neighbors, including a 3-connected node representing the btc$^{3-}$ ligand. The resulting 3,3,8-connected net has been reported in six previous structures as the “3,3,8T25” topology (Figure S8).

Investigation of the Protonation States of Adenine and Location of the Protonated N Atoms. As previously mentioned, adenine can afford different protonation states, and it is often very challenging to accurately observe the presence of H atoms by elemental analysis or SCXRD. The protonation states of the adenine ligand within a MOF structure can sometimes be assigned based on the charge balance of the structure, but the location of the protonated N atoms is not straightforward. XPS has been widely utilized as a reliable technique to assign protonation states of heterocyclic compounds. This technique is based on the general observation that individual chemical environments/functional groups often exhibit similar ranges of binding energy values where the bonding or electronegativity is similar. For example, the neutral 9HAde is expected to exhibit three photoemission N 1s peaks corresponding to three different nitrogen environments, i.e., the one of the N1, N3, N7 atoms, as all of them have the same C=N=C environment, and therefore should have very similar binding energies, which are different from the ones of N9 (C=NH–C environment) and N10 (C=NH$_2$ environment). This prediction, in fact, was experimentally demonstrated by Feyer et al. when they studied the XPS spectra of 9HAde in the gas phase. The N 1s peaks

Figure 6. SION-35: (a) Coordination environments of Zn1 and Zn2; Zn1 holds a trigonal bipyramidal geometry and is coordinated via N7 of Ade$^-$ and four O atoms from four adjacent btc$^{3-}$ ligands; Zn2 is coordinated to N9 from Ade$^-$, one O atom from btc$^{3-}$ and two terminal H$_2$O molecules, affording a distorted tetrahedral geometry. (b) Each btc$^{3-}$ ligand is coordinated to five Zn$^{II}$ ions. (c, d) The orientation of btc$^{3-}$ and Ade$^-$ (in a zigzag fashion along c-axis and in a flat two-dimensional layer along a- and b-axes) around Zn1 and Zn2 affords a three-dimensional compact structure. Color scheme: pink, Zn; gray, C; red, O; blue, N; and white, H.
from each N atom of adenine in the gas phase appear in the range of 398−402 eV. The three peaks correspond to N1, N3, and N7, while the other two peaks represent N9 or N10. The difference in the binding energies of the N atoms becomes less pronounced when the XPS spectrum of adenine is measured in the condensed phase, i.e., in the solid state, probably due to the influences of intermolecular interactions. Furukawa et al. recorded the XPS spectrum from a thick film (≫1 monolayer) of 9HAde on Cu(110); they fitted the N 1s XPS spectrum with two components with binding energies of 399.3 eV (attributed to the imine nitrogen: N1, N3, and N7) and 400.6 eV (attributed to both protonated NH or NH$_2$ groups: N9 and N10). However, the ratio of these two components was found to be 75:25, significantly different from the expected imine:amine ratio of 60:40. XPS measurement for 9HAde was repeated in the solid state, and we observed the same spectrum as reported by Furukawa et al. (Figure 7a), with a peak close to 399 eV and a high energy shoulder, and adapted the three component model that has been successful for the gas phase. The three components represent N environments with different numbers of N−H bonds. Component A represents imine nitrogen with no N−H bonds: N1, N3, and N7. Component B represents N9 which has one N−H bond, and component C represents N10 in the −NH$_2$ groups. Each
component was modeled as a Guassian–Lorentzian peak. The area ratio of the components A:B:C was constrained according to the number of N atoms in 9H Ade they comprised, i.e., to the ratio of 3:1:1. The full width at half-maximum (FWHM) of all components was constrained to be equal in order to give a more physically realistic model. The neutral 9H Ade compound was fitted with the model described, with the components A, B, and C to be at binding energies of 398.49, 399.14, and 400.06 eV, respectively (Figure 7a and Table 1). A difference in

| binding energy (eV) of component | A    | B     | C     |
|----------------------------------|------|-------|-------|
| 9H Ade                           | 398.49 | 399.14 (+0.65) | 400.06 (+1.57) |
| SION-31                          | 399.10 | 399.10 (+0.00) | 400.59 (+1.49) |
| SION-32                          | 399.17 | 399.49 (+0.32) | 400.58 (+1.41) |
| SION-33                          | 399.12 | 400.08 (+0.96) | 400.94 (+1.82) |
| SION-34                          | 399.04b | 400.69 (+1.65) | 400.69 (+1.65) |
| SION-35                          | 399.07 | 399.08 (+0.01) | 400.32 (+1.25) |

“Values in brackets are the differences of binding energy from component A. For SION-34, the component A includes the N atoms in the pzdc⁻ ligand. Components A, B, and C were constrained to have the same FWHM, and the peak areas were constrained in the ratio A:B:C of 3:1:1.

binding energies of 0.65 eV was observed between components A and B, and 1.57 eV between A and C. In the gas phase, the corresponding differences in binding energies were 1.3 and 2.3 eV. The model described, based on the chemical states expected in 9H Ade, fits the experimental data well.

The XPS spectra of SION-31, SION-32, SION-33, SION-34, and SION-35 were subsequently collected, with 9H Ade in the solid phase used to model the XPS spectra collected for each of these materials (Figure 7b–f). It should be noted that, since the pzdc⁻ ligand in SION-34 contains different N atoms (two pzdc⁻ ligands per Ade⁻ ligand, therefore four nonprotonated N atoms per Ade⁻), these must also be included in the model for that material. The binding energy of each of the fitted components for all materials was allowed to vary, and the refined values are tabulated in Table 1. It can be seen that, in two cases, in SION-31 and SION-35, the components A and B appear at very similar binding energies (<0.1 eV difference), whereas, in SION-32, SION-33, and SION-34, the separation was at least in the range of 0.3 eV. The appearance of these two components at the same binding energy was interpreted as the deprotonation of N9, bringing its binding energy down to coincide with the other deprotonated nitrogen environments N1, N3, and N7. On the other hand, separation between the components A and B suggests a loss of equivalence of chemical environments between one out of four following atoms: N1, N3, N7, or N9, and the remaining three. Therefore, the XPS data suggest that adenine is fully deprotonated and acts as an anionic ligand in SION-31 and SION-35, whereas, in SION-32, SION-33, and SION-34, adenine is protonated, acting as a neutral ligand.

At this point, the structural questions for SION-31 and SION-35 are fully resolved: Ade⁻ is a fully deprotonated ligand, which is in agreement with the formulas provided with SCXRD, assuring the charge balance. However, the information gained by the XPS that SION-32, SION-33, and SION-34 structures contain a neutral HAde ligand is inconclusive, since the protonation of adenine may take place on either N1 or N3 for SION-32 and SION-34, while the sites available in SION-33 are N1 or N7.

First, the difference-Fourier maps of SION-32, SION-33, and SION-34 were examined and probed for additional electron density maxima. Since these structures are solved using a synchrotron radiation diffraction data, the probability for peaks originating from H atoms appearing in the Fourier map was relatively high. Indeed, in the case of SION-33 and SION-34, the Fourier maps revealed the location of H atoms. In SION-33, two electron density maxima were identified in the proximity of N7 (0.55 e/Å³ 0.887 Å and 0.54 e/Å³ 1.104 Å apart from this atom), and no maxima near N1. At the same time, the peak of 0.75 e/Å³ as far as 0.850 Å from N1 was found in the difference-Fourier map of SION-34, while again no peaks were found in the vicinity of N3. In both cases, N1 and N3 act as acceptors of H bonds from neighboring guest water molecules, whose H atoms have also been localized in a similar way from the electron density map. However, the information obtained from the difference-Fourier map of SION-32 did not allow for an unequivocal assignment of H atom positions, which might be due to the disorder of adenine over two sites. Thus, an attentive consideration of adenine ligand geometry turned out to be necessary to determine the H atom position in SION-32.

In order to substantiate our observations of different protonation states present in the adenine-based SION-31, SION-32, SION-33, SION-34, and SION-35, and to resolve the problem of localization of the protonated N sites in SION-32, SION-33, and SION-34, we systematically studied the bond angles of adenine in these structures. As a reference, the observation made by Singh was used that, in 6-membered rings of purine derivatives, the C–N–C angles fall in the range of 125 ± 3° for the protonated N atoms, and of 116 ± 3° when the H atom is absent. For 5-membered rings, a similar relation states that, if N7 is protonated, then this 5-membered ring is more symmetrical, whereas, in the case of deprotonation of N7, the C5–N7–C8 angle increases by 3.8–5°. The respective bond angles for SION-31, SION-32, SION-33, SION-34, and SION-35 are listed in Table 2.

| Table 2. Bond Angles (in Degrees) Calculated from the Crystal Structures of SION-31, SION-32, SION-33, SION-34, and SION-35 |
|-------------------|----------|----------|----------|
| C6–N1–C2         | 119.8(2) | 112.9(2)a | 103.1(2)a | 103.2(2)a |
| C2–N3–C4         | 117.4(14) | 110.2(1b) | 106.1(1b) |
| C5–N7–C8         | 118.3(7) | 111.5(8)a | 106.5(7) | 104.6(7)a |
| C4–N9–C8         | 124.6(3) | 113.7(3) | 102.9(3)a | 103.2(3)a |
| C11–C15–C6–N1–C2 | 119.5(4) | 112.0(4) | 104.3(3)a | 104.8(3)a |

“Metal coordinated to the N atom. aAngles modeled as part of an idealized 5-membered ring.

A survey of the Cambridge Structural Database (CSD version 5.38) was performed, and from the 61 adenine-based MOF crystal structures deposited with the CSD (Table S23), the bond angle values were extracted and are presented in the form of histograms (Figure 8). In many cases, the protonation of sites of adenine is excluded, as they bind to the neighboring metal centers by means of coordination bonds. In all other cases, the N sites of adenine were classified as protonated or nonprotonated, and marked, respectively, as red and blue bars.
in Figure 8. For the N1 atom, the span of the C6–N1–C2 angle was found to equal 119 ± 3° for the deprotonated sites, and the only structure with this site protonated, [Cd(1H Ade)-(SO4)],65 was reported with the bond angle of 124.095° (Figure 8a). Considering the bond angle values for SION-31, SION-32, SION-33, SION-34, and SION-35, we observe that the C6–N1–C2 angle has similar values for SION-31, SION-32, SION-33, and SION-35, while such an angle in SION-34 is noticeably larger. Since the former group of angles fall in the range characteristic for deprotonated N1 sites, and the latter one is closer to the literature account on the N1 protonated adenine MOF, the data suggest that, in SION-31, SION-32, SION-33, and SION-35, the adenine ligand is N1 deprotonated, while, in SION-34, the adenine ligand is protonated through N1 (Table 2). No published structures were found to be protonated at the N3, and all C2–N3–C4 angles fall in the range of 112 ± 2° (Figure 8b). Respective angles for SION-31, SION-33, SION-34, and SION-35 also fall in this range, which confirms the deprotonation of adenine on N3 (Table 2). The respective angle for SION-32, however, is visibly larger (Table 2). No literature reference is available, so we resorted to ab initio calculations to confirm the location of protonation. For SION-32 and SION-34, i.e., the two MOFs where the protonation may take place on N1 or N3, we simulated and DFT optimized the respective N1 and N3 protonated structures, thus exploring all possibilities of protonation.

Figure 8. Distribution of the (a) C6–N1–C2, (b) C2–N3–C4, (c) C5–N7–C8, and (d) C4–N9–C8 bond angle values in the adenine-based MOF structures found in the Cambridge Structural Database.

Table 3. Bond Angles and Relative Energies of DFT Optimized Structures of SION-32, SION-33, and SION-34 Simulated with All Probable Protonation Schemes

|          | C6–N1–C2 (deg) | C2–N3–C4 (deg) | C5–N7–C8 (deg) | ΔE (kcal/mol) |
|----------|----------------|----------------|----------------|---------------|
| SION-32, N1-protonated | 123.50          | 113.97         |                | 15.44 (N3)    |
| SION-32, N3-protonated | 120.74          | 117.03         |                |               |
| SION-33, N1-protonated | 123.21          | 102.32         | 12.90 (N7)     |
| SION-33, N7-protonated | 119.55          | 107.84         |                | 12.90 (N7)    |
| SION-34, N1-protonated | 124.45          | 114.46         |                | 13.29 (N1)    |
| SION-34, N3-protonated | 120.96          | 117.52         |                |               |

The energetically favored site of N protonation is given in each ΔE column entry in parentheses.
angles found in these DFT relaxed structures are listed in Table 3. Comparison between these data and the angles found experimentally (Table 2) confirms the N3 deprotonation of adenine in SION-34 (DFT: C2−N3−C4 of 114.46° in the N3 deprotonated and 117.52° in the N3 protonated structure; SCXRD: C2−N3−C4 of 113.7°(3)), and strongly suggests the protonation of SION-32 on N3 (DFT: C2−N3−C4 of 113.97° in the N3 deprotonated and 117.03° in the N3 protonated structure; SCXRD: C2−N3−C4 of 117.7°(3)). In addition, the calculations report an energy difference of 15.4 kcal per mol of HAdv residues in favor of N3 protonation for SION-32. Meanwhile, in SION-34, the N1 protonation is favored by 13.3 kcal/mol, and its DFT-optimized angle distributions for N1 protonated (124.45°) and N3 deprotonated (114.46°) are closer to those reported for the structure in Table 2. Thus, based on insurmountable evidence, the adenine ligand in SION-34 is N1 protonated, while in SION-32 is clearly N3 protonated. We note that SION-32 was previously reported as an N1 protonated structure; however, in that study, the complicated disorder of adenine was not fully resolved (probably due to the low quality of data recorded at room temperature), thus not allowing for the precise geometrical analysis. In terms of the 5-membered ring present in the adenine ligand of the CSD reported MOFs, two maxima in the distribution of the C5−N7−C8 angle were observed: the first one at 103 ± 1° for the nonprotonated, and the second one at 107 ± 2° for the protonated N7 sites (Figure 8c). In the first case, an unexpectedly high value of this angle found at the nonprotonated N7 site, 107.964°, originated from [Co₃(H₂O)₆](μ₃−OH)₆(sip)₂][μ(CH₃)₆] as an example of a structure with the adenine ligand disordered over two sites. In the second case, the structures with extremely high (114.52°) and low (101.109°) coordination geometries as well as different coordination binding modes of the ligands used (H₃btc, H₂ipa, H₄btc, H₃pzdc, HAdv) resulted in a range of structural topologies. The phase purity of each material was confirmed by PXRD and elemental analysis, and their thermal stability was checked by TGA.

Particular interest of our study has been drawn upon the coordination, protonation, and location of protonated N atoms of adenine present in all materials presented herein. XPS has successfully been used to determine the protonation of adenine in SION-31, SION-32, SION-33, SION-34, and SION-35. The N 1s photoelectron peaks were deconvoluted to three components, and the binding energy shifts between them showed significantly different values in the spectra of fully deprotonated SION-31 and SION-35, and those of protonated SION-32, SION-33, and SION-34. The protonation sites of adenine, derived from the difference-Fourier maps, were further confirmed by the study of C−N−C bond angles supported by the literature survey. As previously observed by Singh, the protonation of a given N site gives rise to an increase of the adjacent C−N−C bond angle, which is further supported by the DFT calculations. This strategy can be utilized to other MOF materials with very complex formulas where the balance of the charge of the formula is proven to be challenging.

| N site nonprotonated | N site protonated |
|---------------------|------------------|
| C2−N1−C6            | 119 ± 4°         |
| C2−N3−C4            | 112 ± 2°         |
| C5−N7−C8            | 103 ± 1°         |
| C4−N9−C8            | 103 ± 1°         |

Table 4. Regularities in the C−N−C Angle Values of Adenine Ligand within the Adenine-MOF Crystal Structures Reported in the Literature

![Protonation states and coordination modes of adenine](image-url)

Figure 9. Protonation states and coordination modes of adenine within SION-31, SION-32, SION-33, SION-34, and SION-35.

**Conclusions**

By varying the synthesis conditions, five adenine-based MOFs were isolated, and their crystal structures were solved using SCXRD. SION-31 and SION-32 were found to be two-dimensional layered structures, SION-33 and SION-34 are based on one-dimensional chains (or three-dimensional supramolecular assemblies), while SION-35 is a three-dimensional compact framework. A variety of metal (Ni, Cu, and Zn) coordination geometries as well as different coordination binding modes of the ligands used (H₃btc, H₂ipa, H₄btc, H₃pzdc, Ad) resulted in a range of structural topologies. The phase purity of each material was confirmed by PXRD and elemental analysis, and their thermal stability was checked by TGA.

Additional details and experimental procedures are provided in the supporting information.
ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.inorg-chem.7b02761.

PXRD patterns, SEM images, TGA curves, crystallographic data, literature survey details, and topological analysis (PDF).

Accession Codes

CCDC 1574033 and 1576594–1576597 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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Notes

The authors declare no competing financial interest.

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