Impact of Coordinated Pseudohalide Ions and Picolinamide on Supramolecular Synthons in Selected Zinc and Cadmium Complexes

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Dedicated to the memory of Prof. Dr. Jurij V. Brenčič.

Abstract

Four new Zn and Cd picolinamide (pia) complexes with cyanate and azide ions, namely [Zn(NCO)2(pia)2] (1), [Cd(NCO)2(pia)]n (2), [Zn(N3)2(pia)] (3) and [Cd(N3)2(pia)] · nH2O (4), have been prepared and characterized by IR and NMR spectroscopy, and thermal (TG/DTA) methods. For the compounds 1 and 3 X-ray analysis revealed mononuclear molecular structure with octahedral N4O2 environment around the zinc(II) center with cis-arrangement of ligands. Only in 1 the amide head-to-head supramolecular motifs, represented by an R24(8) designator, are observed, while in 3 none of the typical amide motifs were found. While the IR data suggest the polymeric structures for 2 and 4 in the solid state, all the NMR spectral data support the monomeric structure of the compounds in the solution.

Keywords: Zinc(II) and cadmium(II) complexes; Picolinamide; Crystal structure; Supramolecular synthons; IR and NMR spectroscopy; Thermal analysis (TG/DTA).

1. Introduction

The reasons for combining metal ions and organic ligands into supramolecular coordination networks are countless; they give access to structural diversity and properties likewise adsorption of small molecules, targeted spectral and magnetic properties, conductivity and catalytic activity that are less common for organic solids.1–7 These applications depend on the material’s structure and control of the synthesis is the crucial factor in the formation of new structures. The most important factors susceptible to change in the rational design of metal-organic frameworks are the pH value, solvent type, reaction temperature and time.8 In this context and in continuation of our earlier investigations, we undertook this research to implement more information for molecular recognition processes and relevance of the inter-ligand interactions on structure-properties-function relationship. Therefore, the usage of self-assembly of weaker intermolecular forces such as hydrogen bonds in creating large complex units via desirable motifs is a practical approach.

Recently, we have found for the amide moiety that in picolinamide (pyridine-2-carboxamide, pia) complexes it does not display its usual supramolecular behaviour, e.g. to build robust and reliable R24(8) synthons as it is generally so in organic systems. Instead, it participates more often in the formation of tetrameric R24(8) motifs involving other accessible hydrogen-bonding accepting atoms. For the thiocyanate complexes of Co(II), Ni(II) and Cu(II) with pia, the R24(8) tetrameric motif was found as a synthon of choice, while for Zn(II) complexes both supramolecular motifs, R24(8) and R34(8), were observed. As a continuation of our attempts in determination of ‘supramolecular conditions’ at which the self-complementary amide motif is a reliable and robust supramolecular synthon,9–16 we opted for the other two pseudohalide ions, azide and cyanate, and in this paper we report on their impact on supramolecular synthon formation in picolinamide complexes of zinc and cadmium.
2. Experimental

2.1. Materials and Physical Measurements

All chemicals were purchased from Aldrich Chemical Co. and were used as received without further purification. The CHN analyses were performed in a Perkin-Elmer 2400 Series II CHNS analyser in the Chemical Analytical Service Laboratories of the Rudjer Bošković Institute, Zagreb, Croatia. The IR spectra were obtained from KBr pellets in the range 4000–650 cm–1 with a Perkin-Elmer Spectrum RXI FT-spectrometer. Thermal measurements were carried out on a Mettler-Toledo TGA/SDTA 850e module. Data collections and structure refinement for compounds 1 and 3 are given in Table 1. Single crystals were mounted in a random orientation on a glass fibre. Data collections were carried out on an Oxford Diffraction Xcalibur four-circle kappa geometry single-crystal diffractometer with Sapphire 3 CCD detector, using a graphite monochromated MoKα (λ = 0.71073 Å) radiation, and applying the CrysalisPro Software system at 296 K. The crystal–detector distance was 60 mm. Data reduction, including ab-

2.2. Preparation of Complexes

2.2.1. General procedure for (1)-(4)

A warm aqueous solution of picolinamide (25 mL) was added dropwise with stirring to an aqueous solution of the corresponding metal nitrate (10 mL). An aqueous solution (10 mL) of sodium cyanate (1 or 2) or sodium azide (3 and 4) was added into the resulting mixture. The obtained clear solution was then left at room temperature allowing slow evaporation until X-ray quality crystals were formed. The crystals were filtered off, washed with small portions of cold water and dried in air.

2.2.2. Preparation of [Zn(NCO)2(pia)] (1)

Used: Zn(NO3)2·6H2O (0.30 g; 1 mmol), picolinamide (0.24 g; 2 mmol), sodium cyanate (0.13 g; 2 mmol). Yield: 58% (0.23 g; 0.58 mmol). Anal. Calc. for C12H12ZnN10O2: C, 30.16; H, 3.07 N, 21.35. Found: C, 30.26; H, 2.39; N, 33.29. Calc. for C8H6CdN8O2: C, 22.45; H, 1.81; N, 33.44%. FT-IR (Fig. S4) (KBr, cm–1): 3240(s), 3294(s), 3259(s), 2056(vs), 1670(s-vs), 1650(vs), 1586(s), 1570(s), 1484(w-m), 1422(s), 1330(m-s), 1306(m), 1282(m), 1218(w-m), 1177(w-m), 1161(w-m), 1155(m), 1058(m), 1023(m), 982(wv), 905(wv-w), 858(w), 822(w-m), 759(s), 751(s), 656(s-v), 639(s), 626(s), 550(s), 504(m). NMR: δ(H) [δ13(C)] (ppm): [C2] [148.9 br]; H3[C3] 8.16d [122.9]; H4[C4] 8.05t [139.3]; H5[C5] 7.67tm [127.7]; H6[C6] 8.63d [149.1]; NH2 8.37br, 7.93 br; C=O [166.4].

Abbreviations: br broad signal, d doublet, dm doublet of multiplets, t triplet, tm triplet of multiplets.

Caution! The azido and cyanato complexes are potentially explosive. Therefore, they should be prepared in small quantities and handled with care.

2.2.3. Preparation of [Cd(NCO)2(pia)]n (2)

Used: Cd(NO3)2·4H2O (0.31 g; 1 mmol), picolinamide (0.24g; 2 mmol), sodium cyanate (0.13 g; 2 mmol). Yield: 59% (0.19 g; 0.59 mmol). Anal. Calc. for C6H6CdN8O2: C, 30.03; Cd, 11.59; N, 21.73%. FT-IR (Fig. S4) (KBr, cm–1): 3433(vs), 3299(s), 3206(s), 2203(vs), 2166(vs), 1679(vs), 1651(vs), 1588(vs), 1571(vs), 1484(m), 1423(vs), 1348(m), 1307(s), 1266(m), 1177(w-m), 1161(w-m), 1115(m), 1053(m), 1017(s), 982(wv), 905(wv-w), 858(w), 822(w-m), 759(s), 751(s), 656(s-v), 639(s), 626(s), 550(s), 504(m). NMR: δ(H) [δ13(C)] (ppm): [C2] [149.9br]; H3[C3] 8.07d [121.9]; H4[C4] 8.01tm [137.8]; H5[C5] 7.61tm [126.5]; H6[C6] 8.64dm [148.4]; NH2 8.17br, 7.71 br; C=O [165.9]; Cd-NCO [127.2 br].
sorption correction, was done by CrysAlis RED program. The structures were solved by direct methods implemented in the SHELXS-97 program. The coordinates and the anisotropic thermal parameters for all non-hydrogen atoms were refined by full-matrix least-squares methods based on $F^2$ using the SHELXL-97 program. The aromatic hydrogen atoms were generated geometrically using the riding model with the isotropic factor set at 1.2 $U_{eq}$ of the parent atom. The amide hydrogen atoms were placed in the positions indicated by difference electron-density maps and in 1 their positions were allowed to refine together with individual isotropic displacement parameters, while in 3 they were constrained to ride on their parent N atom at distance 0.86 Å. Graphical work has been performed by ORTEP-3 for Windows and Mercury 1.4.2. The thermal ellipsoids were drawn at the 50% probability level.

2.4. NMR Measurements and Chemical Shift Calculations

All NMR measurements were performed on a Bruker DRX-500 Avance spectrometer equipped with a 5 mm triple broadband inverse probe (TBI) with a z-gradient

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**Table 1.** Crystal data and structure refinement for complexes 1 and 3.

| Compounds | [Zn(NCO)$_2$(pia)$_2$] | [Zn(N$_3$)$_2$(pia)$_2$] |
|-----------|------------------------|------------------------|
| **Empirical formula** | C$_{14}$H$_{12}$ZnN$_6$O$_4$ | C$_{14}$H$_{12}$ZnN$_{10}$O$_2$ |
| **Formula weight** | 393.69 | 393.71 |
| **Color and habit** | colourless, block | colourless, block |
| **Crystal system, space group** | Triclinic, $P1$ | Orthorombic, $Pna2_1$ |
| **Crystal dimensions / mm** | 0.26 × 0.33 × 0.47 | 0.27 × 0.54 × 0.59 |
| **Unit cell parameters:** | | |
| a / Å | 6.7675(2) | 12.6741(3) |
| b / Å | 8.8083(3) | 10.2571(3) |
| c / Å | 13.9214(4) | 11.8972(3) |
| $\alpha$ / ° | 78.832(3) | 90 |
| $\beta$ / ° | 88.1664(4) | 90 |
| $\gamma$ / ° | 80.833(3) | 90 |
| V / Å$^3$ | 803.74(4) | 1556.53(7) |
| Z | 2 | 4 |
| $D_{calc}$ / g cm$^{-3}$ | 1.627 | 1.680 |
| $\mu$ / mm$^{-1}$ | 1.562 | 1.604 |
| $\theta$ range for the data collection / ° | 4.04–27.00 | 3.96–30.00 |
| h, k, l range | –8:8, –11:11, –17:17 | –17:16, –13:14, –16:16 |
| **Scan type** | $\omega$ | $\omega$ |
| **F(000)** | 400 | 800 |
| **No. reflections collected** | 9791 | 11628 |
| **No. independent reflections** | 3484 | 4356 |
| **No. observed reflections,** | | |
| $l \geq 2\sigma(I)$ | 3123 | 3527 |
| **No. refined parameters** | 242 | 243 |
| $R^1, wR^2$ [I $\geq 2\sigma(I)$] | 0.0251, 0.0667 | 0.0312, 0.0736 |
| $R$, $wR$ (all data) | 0.0297, 0.0692 | 0.0429, 0.0829 |
| $g_\ell$, $g_{\ell'}$ in w$^{\ell}$ | 0.0384, 0.2809 | 0.0536, 0.1086 |
| **Goodness of fit on $F^2$,$S^d$** | 1.035 | 1.015 |
| **Max., min. electron density / e Å$^{-3}$** | 0.433 / –0.222 | 1.742 / –0.288 |
| **Maximum $\Delta/\sigma$** | <0.001 | <0.001 |
| **Range of transmission factors min, max** | 0.545, 0.666 | 0.412, 0.645 |
| **Extinction coefficient** | none | none |

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*R = \sum |F_o| - |F_c| / \sum |F_o|

$wR = \left[ \sum (\frac{F_o^2 - F_c^2}{w} )^2 / \sum w(F_o^2)^2 \right]^{1/2}$

$w = \frac{1}{\sigma^2(F_o^2) + \left[ g_\ell(F_o^2) + g_{\ell'}F_o^2 \right]}$ where $P = F_o^2 + 2F_c^2$

$S = \sum w(F_o^2 - F_c^2)^2 / \left( N_{obs} - N_{param} \right)$

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coil. The spectra were acquired in dms-0d₆ solution. Solvent signals were used as secondary references assuming δ(H) = 2.49 ppm and δ(C) = 39.5 ppm with respect to TMS signal (0 ppm). Typically, ¹H NMR, ¹³C NMR, ¹H,¹H COSY and ¹³C,¹H HSQC spectra were acquired in order to identify all signals. Routine parameter sets were applied for the measurements. In addition, the GIAO NMR chemical shift calculations were performed for Zn(NCO)₂(pia)₂; calculated ¹H and ¹³C chemical shifts, although slightly different from experimental values, allowed to identify signal orders. The calculations were performed at the B3LYP/6-31G(2d), LANL2DZ (geometry optimization) and B3PW91/6-311G++(2d,p), LANL2DZ (GIAO NMR calculations) assuming isolated molecule. The shielding scales were converted to chemical shift scales applying reference shielding of 31.8821 and 182.4656 ppm for ¹H and ¹³C in TMS, respectively. All calculations were performed with the Gaussian 09 package. Compound numbering scheme used in NMR account, as well as computed chemical shifts are presented in Scheme 1.

Table 2. Selected bond distances (Å) and angles (°) for complexes 1 and 3.

| Complex | Bond/Angle | Distance (Å) | Angle (°) |
|---------|------------|--------------|-----------|
| 1       | Zn1–O3     | 2.362(1)     | O3–Zn1–O4 85.69(5) |
| 1       | Zn1–O4     | 2.263(1)     | O3–Zn1–N1 168.63(6) |
| 1       | Zn1–N1     | 2.040(2)     | O3–Zn1–N2 89.68(7)   |
| 1       | Zn1–N3     | 2.032(2)     | O3–Zn1–N3 72.64(5)   |
| 1       | Zn1–N5     | 2.124(1)     | O3–Zn1–N5 88.34(5)   |
| 1       | Zn1–N7     | 2.116(2)     | O4–Zn1–N1 87.83(6)   |
| 1       | Zn1–O1     | 2.193(1)     | O1–Zn1–O2 88.2(1)    |
| 1       | Zn1–O2     | 2.194(3)     | O1–Zn1–N1 74.7(1)    |
| 1       | Zn1–N1     | 2.165(4)     | O1–Zn1–N3 89.6(1)    |
| 1       | Zn1–N3     | 2.164(4)     | O1–Zn1–N5 163.8(2)   |
| 1       | Zn1–N5     | 2.096(3)     | O1–Zn1–N8 90.3(1)    |
| 1       | Zn1–N8     | 2.070(3)     | O2–Zn1–N3 74.3(1)    |
| 1       | O2–Zn1–N5  | 94.4(1)      | O2–Zn1–N8 163.4(2)   |
| 1       | O2–Zn1–N8  | 156.2(2)     | N1–Zn1–N3 155.6(2)   |
| 1       | O2–Zn1–N1  | 89.5(2)      | N1–Zn1–N5 89.5(2)    |
| 1       | O2–Zn1–N8  | 109.6(2)     | N1–Zn1–N8 109.6(2)   |
| 1       | N3–Zn1–N5  | 106.5(2)     | N3–Zn1–N8 89.2(2)    |
| 1       | N3–Zn1–N8  | 91.7(1)      | N5–Zn1–N8 91.7(1)    |

3. Results and Discussion

The complexes 1–4 were obtained in situ from an aqueous solutions of all reagents in a molar ratio 1 : 2 : 2 at room temperature. Only the single crystals of zinc compounds 1 and 3 were of suitable quality for the X-ray experiment, allowing structure determination and ensuring a quality refinement. Unfortunately, the crystals of compounds 2 and 4 were of poor crystallinity preventing a successful refinement. Yet, the structure of 2 is determined in the triclinic space group P1 revealing a neutral polymeric coordination compound in which cadmium atoms are doubly bridged with two cyanate-N ions in an end-on mode thus forming 1D endless chains along the a axis.

3.1. Structure Descriptions

The crystals of both 1 and 3 were grown by slow evaporation of the reaction mixtures under ambient conditions in a period from two days to one week. Selected bond distances, angles and details on hydrogen bond geometry are presented in Tables 2 and 3. The molecular structures are shown as ORTEP-3 drawings in Figs 1a and 2a.
The cyanato complex of zinc with picolinamide (1) crystallizes in the triclinic space group \( P \bar{1} \) with the zinc ion in a general position. The crystal structure of 1 comprises neutral mononuclear complex units where zinc ion is octahedrally coordinated by two \( cis \)-oriented picolinamide and cyanate ligands. The picolinamide, having the amide group in the \( o \)-position to the endocyclic nitrogen atom, binds to the metal core in its usually observed \( N,O \)-chelating mode, while cyanate coordinates via its nitrogen atom (Fig. 1a), thus a distorted octahedral coordination sphere around the zinc ion is achieved through four nitrogen and two oxygen atoms. All the coordination bond distances and angles (Table 2) are in accordance with the reported ones for the related structures.9–16;23

Although no significant conformational difference between the two picolinamide ligands were observed their engagement in supramolecular interactions is considerably different (Fig. 1b). The carboxamide moiety of the pia ligand defined by the atoms N3/N4/O3/C3-C8 participates in the formation of the amide head-to-head motifs (Table 3), typically observed in the amide structures (Scheme 2a). The amide moiety belonging to the other, symmetrically independent pia ligand (N5/N6/O4/C9-C14) does not participate in the formation of the common supramolecular amide ring structures, but instead the nitrogen atom donates both of its hydrogens in hydrogen bonds to two cyanate oxygen atoms from two neighbouring molecules thus extending the supramolecular network into 3D (Table 3). Additionally, the oxygen atom is involved in an N–H···O interaction, accepting the anti-amide hydrogen atom that remains after the formation of the amide-amide dimers giving rise to the overall stabilization of the crystal structure. Interestingly, one of the cyanate oxygen atoms acts as a bifurcated acceptor and accepts two amide hydrogens forming centrosymmetrical tetrameric \( R_{2}^{2}(4) \) motifs (Scheme 2d).24 Furthermore, the crystal structure is additionally stabilized by the off-set \( \pi \)--\( \pi \) interactions involving two symmetrically related pyridine rings (N3/C3-C8) from two adjacent molecules (Table 4).

Interestingly, the only pseudohalide metal complex with picolinamide reported so far with the observed amide head-to-head motifs was one of the two thiocyanate zinc complexes.15 As the data on the pseudohalide complexes are scarce so far, it is early to draw any conclusion.

The azido complex of zinc with picolinamide (3) crystallizes in the orthorhombic space group \( Pn\bar{a}_{2}1 \), with zinc ion located in a general position. Here again, the crystal structure is composed of neutral complex units in which the zinc ion is coordinated by two \( cis \)-related picolinamide and azide ligands. The picolinamide ligands are, here as well, bound to the metal centre through its

### Table 3. Hydrogen bonds (Å and °) for compounds 1 and 3.

| D–H···A            | D–H  | H···A  | D···A  | \( \angle \)DHA | Symmetry code |
|--------------------|------|-------|-------|----------------|---------------|
| **Compound 1**     |      |       |       |                |               |
| N4–H14N···O3       | 0.90(3) | 2.08(3) | 2.969(2) | 176(2)  | 2 – \( x \), 1 – \( y \), 2 – \( z \) |
| N6–H16N···O2       | 0.85(3) | 2.05(3) | 2.889(3) | 170(2)  | –1 + \( x \), 1 + \( y \), \( z \) |
| N4–H24N···O4       | 0.79(3) | 2.45(3) | 3.182(2) | 156(3)  | 1 – \( x \), 1 – \( y \), 2 – \( z \) |
| N6–H26N···O1       | 0.82(3) | 2.14(3) | 2.920(3) | 157(3)  | \( x \), 1 + \( y \), \( z \) |
| C4–H4···O4         | 0.93  | 2.42  | 3.192(3) | 140     | 1 – \( x \), 1 – \( y \), 2 – \( z \) |
| **Compound 3**     |      |       |       |                |               |
| N2–H12N···N7       | 0.85(8) | 2.33(4) | 3.167(6) | 167(4)  | \( 1/2 – x \), \( 1/2 + y \), 1/2 + \( z \) |
| N4–H14N···N7       | 0.84(5) | 2.24(5) | 3.018(6) | 154(5)  | \( x – 1/2, 1/2 – y, z \) |
| N2–H22N···N5       | 0.85(4) | 2.50(4) | 3.329(5) | 166(4)  | \( x – 1/2, 3/2 – y, z \) |
| N4–H24N···N10      | 0.87(4) | 2.13(4) | 2.993(7) | 174(5)  | \( 1/2 – x, y – 1/2, z – 1/2 \) |
| C2–H2···N7         | 0.93  | 2.41  | 3.333(6) | 174     | \( 1/2 – x, 1/2 + y, 1/2 + z \) |
| C3–H3···O1         | 0.93  | 2.58  | 3.226(5) | 127     | \( 1/2 – x, y – 1/2, z + 1/2 \) |
| C5–H5···N5         | 0.93  | 2.51  | 3.075(6) | 120     | \( x, y, z \) |
| C8–H8···N10        | 0.93  | 2.39  | 3.305(6) | 168     | \( 1/2 – x, y – 1/2, z – 1/2 \) |
| C10–H10···N10      | 0.93  | 2.49  | 3.314(6) | 148     | \( 1 – x, 2 – y, z – 1/2 \) |
| C11–H11···N8       | 0.93  | 2.52  | 3.065(6) | 117     | \( x, y, z \) |
Table 4. \( \pi \text{--} \pi \) interactions (Å and °)

| R(\(i\)) \(\rightarrow\) R(\(j\)) | Cg(\(i\))···Cg(\(j\)) | \(\alpha^a\) | \(\beta^b\) | Cg(\(i\))···plane[Cg(\(j\))] |
|---------------------|-----------------|------|------|-----------------|
| Compound 1          | Cg(N3,C3→C7)···Cg(N3,C3→C7)\(^i\) | 3.852(2) | 7.8  | 23.0            | 3.313 |
| Compound 3          | Cg(N3,C7→C11)···Cg(N1,C1→C5)\(^ii\) | 3.732(1) | 0  | 22.4            | 3.450 |

\(^i\) = 1 – \(x\), \(−y\), \(2−z\), \(^ii\) \(\frac{1}{2}−x\), \(\frac{1}{2}+y\), \(−\frac{1}{2}+z\).

\(^a\alpha = \) angle between planes of two aromatic rings \(^b\beta = \) angle between Cg···Cg line and normal to the plane of the first aromatic plane

Scheme 2. Supramolecular synthons present in the picolinamide structures.\(^{23}\)

\(N,O\)-donor set of atoms thus forming a distorted octahedron around the zinc ion via four nitrogen and two oxygen atoms (Fig. 2a).

The survey of the Cambridge Structural Database (CSD, version 1.16, 2013) reveals only seven octahedral bispicolinamide complexes with pseudohalide ions bound to metal centres; of these, there are five thiocyanato and two azido complexes. For the reported pseudohalide complexes the cis-orientation of the ligands is more pronounced; only for the nickle(II) complex\(^{12,14}\) both isomers are found. Therefore, the molecular structure of the complexes 1 and 3 are in accordance with literature data.
On analogy with other crystal structures of azide complexes with picolinamide reported so far\textsuperscript{12,13} none of the typically observed hydrogen bonding motifs are found (Scheme 2a–c), but instead, R\textsubscript{2}(16) dimers formed (Scheme 2e) between the amide group and the azide ligands are present in the complex \textsuperscript{3} (Fig. 2b). The same motifs are observed in the cis-[Ni(N\textsubscript{3})\textsubscript{2}(pia)\textsubscript{2}] complex.\textsuperscript{12} Additionally, one more interesting hydrogen bonding motif is observed in \textsuperscript{3}. It is the R\textsubscript{3}(10) hydrogen bonding motif (Scheme 2f) that resembles the tetrameric motif R\textsubscript{4}(8) which is most frequently present in the picolinamide structures (Scheme 2c), just in this particular motif the three azide atoms replace one pseudohalide acceptor atom (Table 3).

Moreover, there are also π–π stacking interactions between two pyridine rings (N1/C1–C5 and N3/C7–C11) belonging to two neighbouring molecules (Table 4) giving rise to the overall stability of the crystal structure.

3. 2. Spectroscopic Properties

3. 2. 1. IR Spectra

The absorptions assigned as ν(CN) in the IR spectra of the cyanate complexes \textsuperscript{1} and \textsuperscript{2} are shifted towards higher wavenumbers when compared to those of the free cyanate (2170 cm\textsuperscript{-1}). The strong ν(CN) in the spectrum of \textsuperscript{1} is observed at 2216 cm\textsuperscript{-1}. The splitted CN stretching frequency bands observed in \textsuperscript{2} (2203 and 2166 cm\textsuperscript{-1}) suggest bridging coordination of the cyanate groups.\textsuperscript{25} In spectrum of \textsuperscript{1}, the amide C–O stretching frequency is observed at 1683 cm\textsuperscript{-1}, indicating involvement of the amide oxygen atom into, probably more than one, hydrogen bonding. In the spectrum of \textsuperscript{2}, the splitting of the C–O band occurred, indicating different environment of the two amide oxygen atoms. The IR spectra of the complexes \textsuperscript{3} and \textsuperscript{4} display the strong characteristic asymmetric stretching vibrations ν\textsubscript{s}(N\textsubscript{3}) of the azido ligands over the range 2166–2216 cm\textsuperscript{-1}. In the spectrum of \textsuperscript{3}, splitting of the N≡N band (2166 and 2095 cm\textsuperscript{-1}) is observed as a result of asymmetric azido groups while in \textsuperscript{4} splitting is not present suggesting symmetry and μ\textsubscript{1,3}–N\textsubscript{3} bridging coordination of the azide ligands.\textsuperscript{26}

3. 2. 2. NMR Measurements

Signal assignments were straightforward. Aromatic \textsuperscript{1}H spin systems were identified by COSY experiments assuming doublet-like signals for H2 and H6, and triplet-like signals for H3 and H4. Signal order was established with the use of calculated chemical shifts. The \textsuperscript{13}C signals were unambiguously identified by \textsuperscript{1}H, \textsuperscript{13}C-HSQC and \textsuperscript{1}H, \textsuperscript{13}C HMBC 2D experiments. The presence of two NH signals in each \textsuperscript{1}H NMR spectrum is worth to note. Still, all data from NMR spectra corroborate with the monomeric species in DMSO solution.

3. 3. Thermal Analysis

At ambient conditions the crystals \textsuperscript{1}, \textsuperscript{2} and \textsuperscript{4} possess good stability and do not show any hygroscopic effects (Figs. S5–S8). The compound \textsuperscript{1} decomposes continuously starting form 150 °C without any pronounced mass loss step and the decomposition is followed by an endothermic process in a wide temperature range, approximately 25–500 °C, with only one emphasised peak at 152 °C. The analysis of the TG and DTA curves of the compound \textsuperscript{2} shows that it is stable up to approximately 180 °C. After that temperature, it decomposes and in the temperature range 180–600 °C two endothermic peaks in the DTA curve were observed, the first small at 201 °C, and second larger one at 260 °C. The analysis of TG curve of compound \textsuperscript{3} showed that its thermal decomposition starts readily by heating indicating possible hygroscopic effect and consists of three poorly separated steps. The first step is followed by small endothermic DT peak at 211 °C, it cor-
roborates with the loss of the azide ligands (observed loss 21.50%; calc. 21.35%), and is the only step that can be resolved from the overall endothermic process occurring through the whole temperature range. The thermogravimetric data reveals that complex 4 is stable up to 160 °C. Afterwards it decomposes, and the first step accompanied with the sharp endothermic DT peak at 259 °C corresponds to the loss of one water molecule and one azido ligand (found: 17.57%, calculated 17.84%). Furthermore, less resolved decomposition steps in the temperature range between 270 and 410 °C are characterized by one endothermic (small peak at 280 °C) and two exothermic processes (sharp peaks at 363 and 387 °C). In the temperature range 410 to 600 °C, a further exothermic process occurs with broad maximum around 530 °C. The process is followed by mass loss recorded in the TG curve, with the final residue after 600 °C of 13.66%. Due to unfinished decomposition processes and sublimation of Cd compounds in the temperature range up to 600 °C we were not able to analyse masses of the residues for all the compounds.

4. Conclusion

Here we reported on four picolinamide complexes of zinc and cadmium, [Zn(NCO)₂(pia)]₂ (1), [Cd(NCO)₂(pia)]₂ (2), [Zn(N₃)₂(pia)]₂ (3) and [Cd(N₃)₂(pia)]₂·H₂O (4). Complexes are characterized by IR and NMR spectra and TG/DTA thermal methods. The X-ray analysis revealed that the crystal structures of 1 and 3 are composed of mononuclear complex units with cis-oriented picolinamide- and pseudohalide ligands, while for 2 and 4 it was concluded that they are polymeric in the solid state. NMR data are not supportive of existence of 2 and 4 in the polymeric form. In the solid state, for none of the reported zinc complexes, 1 and 3, the tetrameric R₂(8) motifs typically present in the complexes with the picolinamide-thiocyanate set of ligands are found. Instead, in the crystal structure of the compound 1 typical supramolecular amideamide dimers, described by the R₂(8) graph-set designator, are observed. Interestingly, the same supramolecular synthons was present in one of the two zinc polymorphs with the thiocyanate as a pseudohalide ligand. Unfortunately, due to the lack of data on picolinamide complexes of cadmium with azide and cyanate ligands we are not able to draw any conclusion yet. Therefore, it is worthy to emphasize the need of further research, particularly a theoretical one, in order to put some additional light to experimental data and possible inter-ligand interactions.

5. Supplementary Data

Crystallographic data for the structures reported in the paper have been deposited with the Cambridge Crystallographic Data Centre as supplementary publication no, CCDC 1028428–1028429. Copies of the data can be obtained free of charge at www.ccdc.cam.ac.uk/conts/retrieving.html or from the Cambridge Crystallographic Data Centre (CCDC), 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44(0)1223-336-033; or e-mail: deposit@ccdc.cam.ac.uk.

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Povzetek

Pripravili in okarakterizirali z IR in NMR spektroskopijo ter termičnimi (TG/DTA) metodami smo štiri nove kompleksne Zn in Cd s pikolinamidom (pia), $[\text{Zn(NCO)}_2(pia)_2]$ (1), $[\text{Cd(NCO)}_2(pia)_2]$ (2), $[\text{Zn(N3)}_2(pia)_2]$ (3) in $[\text{Cd(N3)}_2(pia)_n] \cdot n\text{H}_2\text{O}$ (4). Rentgenska struktura analiza na monokristalih je za spojine 1 in 3 razkrila enojedrno zgradbo z oktaedrično $\text{N}_4\text{O}_2$ cis-razporeditvijo donorskih atomov okrog cinkovega(II) iona. Le amid v spojini 1 je mogoče razvrstiti v skupino »head-to-head« supramolekularnih motivov (oznaka je $\text{R}_2\text{2}(8)$). V strukturi spojine 3 pa nismo prepoznali nobenega od značilnih strukturnih motivov za amid. Čeprav je mogoče iz IR spektrov spojine 2 in 4 sklepati na polimerno zgradbo, pa NMR spektri kažejo na enojedno zgradbo kompleksnih zvrsti v raztopinah.