μ -Phenoxo- μ -pseudohalide and μ -pseudohalide dinuclear, tetranuclear and one-dimensional complexes: magneto-structural correlation and interesting type of solid state isomerism

SUJIT SASMAL and SASANKASEKHAR MOHANTA*

Department of Chemistry, University of Calcutta, 92, A P C Road, Kolkata 700 009, India e-mail: sm_cu_chem@yahoo.co.in

Abstract. Five Schiff base ligands have been utilized to explore metallo-pseudohalide (azide or cyanate) systems. These ligands are the 1:1 condensation products of 3-methoxysalicylaldehyde with ethanolamine $(H_2L^1)/1$ -(2-aminoethyl)-piperidine $(HL^2)/4$ -(2-aminoethyl)-morpholine (HL^3) or salicylaldehyde with 1-(2-aminoethyl)-piperidine $(HL^4)/4$ -(2-aminoethyl)-morpholine (HL^5) . The derived complexes are as follows: Four heterobridged μ -phenoxo- $\mu_{1,1}$ -azide/cyanate dinickel(II) compounds of composition $[Ni_2^{II}(HL^1)_3(\mu_{1,1}-N_3)]\cdot 3H_2O$ (1), $[Ni_2^{II}(L^2)_2(\mu_{1,1}-N_3)(N_3)(H_2O)]\cdot CH_3CH_2OH$ (2), $[Ni_2^{II}(L^3)_2-(\mu_{1,1}-N_3)(CH_3CN)(H_2O)](CIO_4)\cdot H_2O\cdot CH_3CN$ (3) and $[Ni_2^{II}(HL^1)_3(\mu_{1,1}-NCO)]\cdot 2H_2O$ (4); Two $\mu_{1,3}$ -azide bridged tetranickel(II) compounds $[\{Ni_1^{II}(L^4)(\mu_{1,3}-N_3)(H_2O)\}_4]$ (5) and $[\{Ni_1^{II}(L^5)(\mu_{1,3}-N_3)(H_2O)\}_4]$ (6); Two $\mu_{1,3}$ -azide/cyanate one-dimensional compounds $[Cu_1^{II}L^5(\mu_{1,3}-NCO)]_n\cdot 2nH_2O$ (7) and $[Cu_1^{II}L^5(\mu_{1,3}-N_3)]_n\cdot 2nH_2O$ (8). Except compound 5 which shows overall antiferromagnetic coupling, other compounds exhibit overall ferromagnetic interaction. Syntheses, crystal structures, magnetic properties, density functional theoretical (DFT) calculations and experimental/theoretical magneto-structural correlations have been carried out which have revealed some interesting observations on composition/topology, magneto-structural correlations and solid state isomerism. The results have been already published. The present report deals with a review of the salient and interesting features of these works.

Keywords. Metallo-pseudohalides; magneto-structural correlations; DFT; heterobridged.

1. Introduction

Molecular magnetism is a frontier research area. 1-5 One major focus in this area is to determine magnetostructural correlations, both experimental and theoretical. 1-4 In an experimental correlation, J versus one parameter (for example, bridge angle) relationship is determined for a few similar complexes. However, it is not possible to keep other governing parameters constant for these complexes, which may be considered as a limitation of experimental magneto-structural correlations. Therefore, it deserves importance if at least a pair of similar exchange-coupled systems are obtained for which all but one governing parameters are identical. On the other hand, at least for symmetrically bridged compounds, it is easily possible to change one governing parameter keeping others constant theoretically in the model systems and thus the effect of the change of other parameters is nullified in the theoretical correlations in symmetrically bridged systems. However, it is difficult to avoid the above mentioned limitation even in the theoretical correlations of

The research in structural chemistry focusing supramolecular chemistry/crystal engineering is another area which is being attracted vast fascination in recent years because of two major objectives of the self-assemblies: aesthetic beauty and possibility of being functional materials. Another perspective of solid state structural chemistry is to explore the little investigated types of isomerism as well as to find out new types of isomerism. Regarding non-classical types of isomerism that have been developed in recent years, the following may be mentioned as examples: Jahn–Teller isomerism, supramolecular isomerism, molecular isomerism, molecular isomerism due to difference in number of interstitial solvent molecules, catenation isomerism.

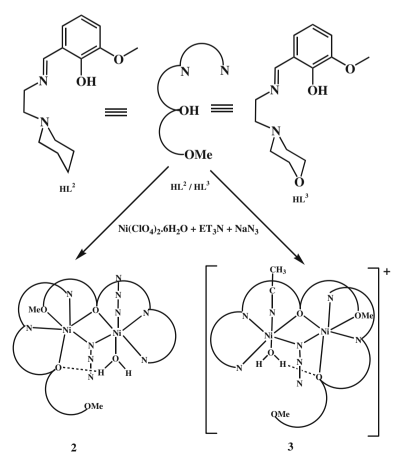
heterobridged systems (for example, μ -phenoxo- $\mu_{1,1}$ -azide dinuclear complexes) because it is not possible to change one parameter keeping all others constant (for example, the change of metal-phenoxo-metal angle is associated with change of metal-phenoxo distance in μ -phenoxo- $\mu_{1,1}$ -azide dinuclear complexes). Thus, determination of even the theoretical magneto-structural correlations in heterobridged systems deserves importance in molecular magnetism.

^{*}For correspondence

Scheme 1. Syntheses of 1 and 4.

Metallo-pseudohalide, azide or cyanate, species have been always of great interest in molecular magnetism and supramolecular chemistry/crystal engineering due to a number of coordination modes of these potentially bridging ligands as well as due to their ability to mediate ferro or antiferromagnetic interactions among metal centres. The most usual bridging modes are the end-on $(\mu_{1,1}$ -, commonly expressed as EO) $^{1a,1d,11-17}$ and end-to-end $(\mu_{1,3}$ -, EE), $^{1\hat{a},1d,18,19}$ but some others like, $\mu_{1,1,1}$, 13,16,20,21 $\mu_{1,1,3}$, 14,21 $\mu_{1,1,1,1}$, 22 and $\mu_{1,1,3,3}$ have been also described. This way, a number of azide/cyanate-bridged systems of different nuclearity (dinuclear, oligonuclear and polynuclear clusters) and also of 1D, 2D, and 3D topologies have been achieved and there is a considerable amount of information available on experimental and theoretical magneto-structural correlations. 1a,1d,11-23

Despite all the encouraging results, it is difficult to control the composition/topology of the metallopseudohalide species, which depends on factors like organic ligand, metal ion, solvent, stoichiometric ratio of reactants and other reaction conditions. To explore the metallo-pseudohalide species, we have



Scheme 2. Syntheses of **2** and **3**.

used five Schiff base ligands H₂L¹, HL², HL³, HL⁴ and HL5, which are the [1+1] condensation products of 3-methoxysalicylaldehyde with ethanolamine (for H_2L^1) / 1-(2-aminoethyl)-piperidine (for HL^2) / 4-(2-aminoethyl)-morpholine (for HL³) or of salicylaldehyde with 1-(2-aminoethyl)-piperidine (for HL⁴) / 4-(2-aminoethyl)-morpholine (for HL⁵). We have synthesized the following compounds using these ligands: Four heterobridged μ -phenoxo- $\mu_{1,1}$ azide/cyanate dinickel(II) compounds of composition $[Ni_2^{II}(HL^1)_3(\mu_{1,1}-N_3)]\cdot 3H_2O$ (1), $[Ni_2^{II}(L^2)_2(\mu_{1,1}-N_3)]\cdot 3H_2O$ $[Ni_2^{II}(L^3)_2(\mu_{1,1}-N_3)]$ $(N_3)(H_2O)$]·CH₃CH₂OH (2), $(CH_3CN)(H_2O)](ClO_4)\cdot H_2O\cdot CH_3CN$ (3) and $[Ni_2^{II}]$ $(HL^{1})_{3}(\mu_{1,1}-NCO)]\cdot 2H_{2}O$ (4); Two single end-toend azide bridged cyclic tetranickel(II) clusters of composition $[\{Ni^{II}(L^4)(\mu_{1,3}-N_3)(H_2O)\}_4]$ (5) and $[\{Ni^{II}(L^5)(\mu_{1,3}-N_3)(H_2O)\}_4]$ (6); Two single end-toend azide/cyanate bridged one-dimensional copper(II) systems of composition $[Cu^{II}L^5(\mu_{1,3}-NCO)]_n \cdot 2nH_2O$ (7), $[Cu^{II}L^{5}(\mu_{1,3}-N_{3})]_{n}\cdot 2nH_{2}O$ (8). Single crystal X-ray structure determination and variable-temperature/field magnetic studies have been done for all these eight

Scheme 3. Syntheses of **5** and **6**.

Scheme 4. Syntheses of **7** and **8**.

compounds. Density functional theoretical calculations on magnetic properties have been carried out for 1–3, 5 and 6. Interesting experimental and theoretical magneto-structural correlations have been determined and new types of isomerism have been observed. The results have been already published. ^{24–28} A review of the salient features of these works is being presented here.

2. Outline of syntheses

The dinickel(II) or tetranickel(II) complexes 1–6 are readily obtained in high yield from the reaction of the corresponding Schiff base ligand H₂L¹/HL²/HL³/HL⁴/HL⁵, nickel(II) perchlorate hexahydrate, triethylamine and sodium azide/cyanate in 3:2:6:2 ratio (for 1 and 4) or in 1:1:1:4 ratio (for 2, 3, 5 and 6). For the copper(II) systems 7 and 8, the Schiff base: copper(II) perchlorate hexahydrate: sodium azide/cyanate ratio of 1:1:4 was used; the use of the base triethylamine was not required in these cases. The syntheses of the eight compounds are demonstrated in schemes 1, 2, 3 and 4.

3. Crystal structures, and magneto-structural correlations of heterobridged $\mu\text{-phenoxo-}\mu_{1,1}\text{-}$ azide/cyanate dinickel(II) compounds: Combined experimental and theoretical exploration $^{24\text{--}26}$

Crystal structures of $Ni_2^{II}(HL^1)_3(\mu_{1,1}-N_3)]\cdot 3H_2O$ (1) and $[Ni_2^{II}(L^2)_2(\mu_{1,1}-N_3)(N_3)(H_2O)]\cdot CH_3CH_2OH$ (2) are shown in figures 1 and 2, while the crystal structures of $[Ni_2^{II}(L^3)_2(\mu_{1,1}-N_3)(CH_3CN)(H_2O)](ClO_4)\cdot H_2O\cdot CH_3CN$ (3) and $[Ni_2^{II}(HL^1)_3(\mu_{1,1}-NCO)]\cdot 2H_2O$ (4) are shown in figures S1 and S2 (supplementary informations). The structures reveal that 1–3 are μ -phenoxo- $\mu_{1,1}$ -azide dinickel(II) compounds, while 4 is a μ -phenoxo- $\mu_{1,1}$ -cyanate dinickel(II) compound.

The azide analogue 1 and the cyanate analogue 4 are derived from the same ligand and have similar type of structures containing three monodeprotonated ligands, [HL¹]⁻, in which the phenoxo moiety is deprotonated. Among the three phenolate oxygen atoms of three [HL¹]⁻, one bridges the two metal ions, while each of the remaining two is coordinated to each of the two metal ions in monodentate fashion. Of the three alcohol oxygen atoms, one is uncoordinated, while each of the other two coordinates to each of the two metal ions in monodentate fashion. Among the three imine

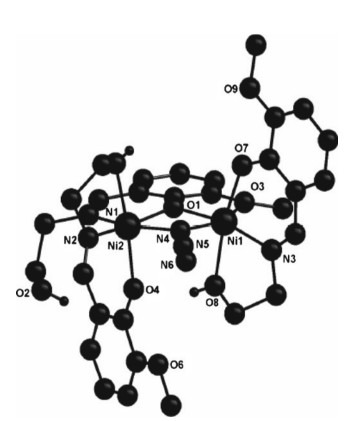


Figure 1. Crystal structure of $[Ni_2^{II}(HL^1)_3(\mu_{1,1}-N_3)]\cdot 3H_2O$ (1). Hydrogen atoms, except of three alcohol moieties, and three water molecules are deleted for clarity. ^{24,29}

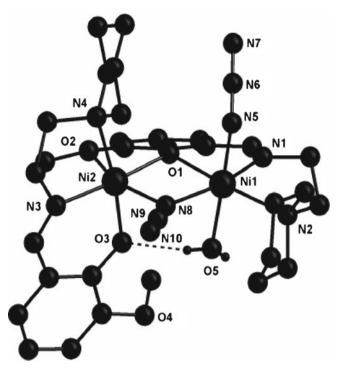


Figure 2. Crystal structure of $[Ni_2^{II}(L^2)_2(\mu_{1,1}-N_3)(N_3)(H_2O)]\cdot CH_3CH_2OH$ (2). Hydrogen atoms, except those of water molecule, and solvated ethanol molecule are omitted for clarity. 26,30

nitrogen atoms, two are coordinated to one metal ion, while the third coordinates to the second metal centre. Of the three ethereal oxygen atoms, one is coordinated to one metal ion, while the other two remain uncoordinated. In addition to the phenoxo bridge, the two metal centres are also bridged by the nitrogen atom of an end-on azide (in 1) or cyanate (in 4) ligand.

Compounds **2** and **3** contain two deprotonated ligands, $[L^2]^-/[L^3]^-$. While one of the two phenoxo oxygen atoms of two $[L^2]^-/[L^3]^-$ bridges the two metal ions, the second phenoxo oxygen atom coordinates one metal ion. The two metal ions are additionally bridged by the nitrogen atom of an end-on azide ligand. Among the two methoxy oxygen atoms of two $[L^2]^-/[L^3]^-$, one is non-coordinated, while the second is coordinated to one metal centre. Each of the two metal ions are coordinated also to the imine and piperidine (for **2**)/morpholine (for **3**) nitrogen atoms of each of the two $[L^2]^-/[L^3]^-$.

Magnetic properties of the heterobridged μ -phenoxo- $\mu_{1,1}$ -azide/cyanate dinickel(II) compounds **1–4** are shown in figures **3**, **4**, supplementary figures **S3** and **S4**, respectively, as $\chi_{\rm M}T$ versus T plots. The profiles indicate that the metal centres in all the four compounds are ferromagnetically coupled. Using the Hamiltonian $\mathbf{H} = -2J(\mathbf{S}_1 \cdot \mathbf{S}_2) + D_1[S_{Z,1}^2] + D_2[S_{Z,2}^2]$, good

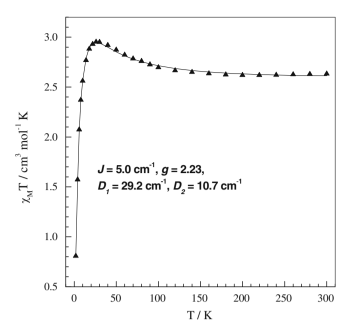


Figure 3. $\chi_{\rm M}T$ vs. T plots for $[{\rm Ni}_2^{\rm II}({\rm HL}^1)_3(\mu_{1,1}-{\rm N}_3)]$ $3{\rm H}_2{\rm O}$ (1). Symbols and solid lines represent the observed and calculated data, respectively. 24,29

quality simulations are obtained (figures 3, 4, supplementary figures S3 and S4) with following parameters: $J = 5.0 \,\mathrm{cm}^{-1}$, g = 2.23, $D_1 = 29.2 \,\mathrm{cm}^{-1}$ and $D_2 = 10.7 \,\mathrm{cm}^{-1}$ for 1; $J = 16.6 \,\mathrm{cm}^{-1}$, g = 2.2 and $D_1 = D_2 = -7.3 \,\mathrm{cm}^{-1}$ for 2; $J = 16.92 \,\mathrm{cm}^{-1}$, g = 2.2 and $D_1 = D_2 = -6.41 \,\mathrm{cm}^{-1}$ for 3; $J = 3.33 \,\mathrm{cm}^{-1}$, g = 2.24 and $D_1 = D_2 = |6.68| \,\mathrm{cm}^{-1}$ for 4.

Both the phenoxo and azide routes are responsible for the superexchange between the metal centres in a

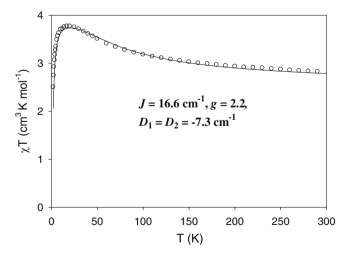


Figure 4. $\chi_M T$ vs. T plots for $[Ni_2^{II}(L^2)_2(\mu_{1,1}-N_3)(N_3)(H_2O)]\cdot CH_3CH_2OH$ (2). Symbols and solid lines represent the observed and calculated data, respectively. 26,30

heterobridged μ -phenoxo- $\mu_{1,1}$ -azide compound, such as **1–3**. So, the nickel–azide–nickel and nickel–phenoxo–nickel angles and two nickel–azide and two nickel–phenoxo distances should be the governing factors for the J values in these compounds. Magnetic and structural parameters of the **1–3** and also of the only two previously reported μ -phenoxo- $\mu_{1,1}$ -azide dinickel(II) compounds are summarized in table **1**.

For the two compounds 1 and IA (table 1), 31a J values are drastically different; 5.0 cm⁻¹ for 1 and 25.6 cm⁻¹ for **IA**. Of the four governing parameters, the nickel-azide-nickel and nickel-phenoxo-nickel angles and both the nickel-phenoxo distances are practically identical for these two compounds. Only the nickelazide bond distances are different; 2.09 and 2.21 Å for 1 and 2.13 and 2.15 Å for IA. Clearly, the significant difference in the J values arises due to difference in nickel-azide bridge distances. Thus, interestingly, these two compounds represent a unique pair for which all but one governing parameters are identical and also represent a unique pair demonstrating the bridge distance dependency of exchange integral. The asymmetry (0.12 Å in 1, 0.02 Å in IA) in the two nickel–azide bond distances are greater in 1 than in IA. Thus, it seems that more the asymmetry in the two nickel-azide bond distances less the strength of ferromagnetic interaction. Such J versus asymmetry correlation is also valid for compound 2 (table 1; $J = 16.6 \,\mathrm{cm}^{-1}$, asymmetry = $0.06 \,\text{Å}$), although other governing parameters of 2 are not identical with those of 1 or IA and thus the J versus asymmetry correlation can be considered as more strengthened. On the other hand, the J and asymmetry of compounds 3 and IB^{31b} are not in line with this correlation. So, it seems that although asymmetry in the two nickel-azide bond distances has some profound roles, the overall interaction is a composite effect of several parameters. Therefore, we have performed density functional theoretical calculations on the magnetic properties of 1-3 and IA. We have also determined theoretical magneto-structural correlations to get more insight regarding the role of the possible governing parameters, including asymmetry in the two nickel-azide bond distances.

Broken symmetry density functional calculations 32 of exchange interaction provide good numerical estimate of J values compared to experiments (table 1). Taking compound 2 as the model system, we have developed three different correlations. As mentioned in introduction section, the presence of dissimilar bridges prevents variation of only one structural parameter at a time to develop correlations. Therefore, we have defined three parameters (α , β and γ) to vary two parameters (angles as well as the Ni–O or Ni–N

Table 1. Magnetic and structural parameters of the five μ -phenoxo- μ_1 -azide dinickel(II) compounds. 26,30

	$J_{ m Expt.}$	$J_{ m DFT}$	Ni-O-Ni	N_{i-N-N}		Average	Asymmetry in		Average	Asymmetry in	
Compounds	(cm^{-1})	(cm^{-1})	(°)	(0)	Ni-0 (Å)	Ni-0 (Å)	Ni-O (Å)	Ni–N (Å)	Ni–N (Å)	$N_{i-N}(A)$	Ref.
1	5.0	14.1	106.9		1.99, 1.99	1.99	0.00	2.09, 2.21	2.15	0.12	24
2	16.6	15.8	103.0	0.86	2.01, 2.06	2.035	0.05	2.08, 2.14	2.11	90.0	56
3	16.9	15.35	104.7		1.99, 2.02	2.005	0.03	2.12, 2.11	2.115	0.01	56
IA	25.6	18.5	106.7		1.98, 1.99	1.985	0.01	2.13, 2.15	2.14	0.02	31a
IB	2.85	12.65/10.05	102.3		2.00, 2.04	2.02	0.04	2.11, 2.14	2.125	0.03	31b

distances) at a time in our correlations. To underpin the role of asymmetry on the bond lengths, the asymmetry present in the bond distances were maintained same throughout.

Ni - O - Ni	$\rho = Ni - N - Ni$	Ni - N - Ni
$\alpha = \frac{1}{d(avg \cdot Ni - 0)}$	$\frac{d}{d(avg.Ni - N)}$	$Y - \frac{d(Ni_B - N)}{d(Ni_B - N)}$

According to the J versus α correlation (figure S5), the magnitude of ferromagnetic J is decreased with the increase of the Ni-O-Ni angle or the decrease of the Ni–O distance. As shown in figure S6, the J versus β correlation indicates that the ferromagnetic interaction is increased as the nickel-azide-nickel angle increases or nickel-azide distance decreases, with a maximum at $\beta = 106^{\circ}$. The J versus ν correlation (figure 5) reveals that ferromagnetic J is increased with the increase of γ to reach saturation at very large γ values. To understand the dependency of J on the asymmetry (Δ) in two nickel-azide distances, which is our major goal, the change in the Ni–N–Ni angle in the J versus γ correlation is ignored to get a J versus asymmetry profile, as plotted in figure 6. The asymmetry values in this graph are both positive and negative with reference to the fixed Ni_A-N distance of 2.082 Å. The asymmetry is negative when $d_{NiB-N} > 2.082$ Å, while a positive asymmetry means $d_{NiB-N} < 2.082 \,\text{Å}$. Although the shape of the curve in both the positive and negative asymmetry regions is closely linear, the role of asymmetry differs; while increase in negative asymmetry decreases the ferromagnetic interaction, increase in positive asymmetry increases the ferromagnetic interaction, albeit this saturates at larger value. As evident from this plot, if the Ni_B-N distance is larger than 2.082 Å (i.e., if the asymmetry is negative), an increase in asymmetry will lead

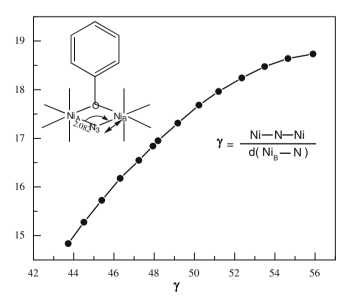


Figure 5. DFT-computed J versus γ correlation. ^{26,30}

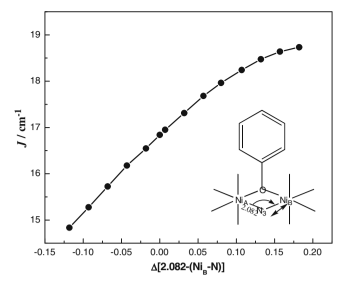


Figure 6. DFT-computed J versus asymmetry in the two Ni–N distances correlation (the variation in the Ni–N–Ni angle is ignored). 26,30

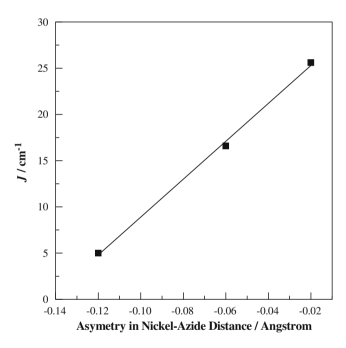


Figure 7. Linear magneto-structural correlation between experimental J and asymmetry in two nickel-azide distances. ^{26,30}

to a smaller ferromagnetic J and vice versa. In fact, such a trend (where $\Delta =$ shorter Ni–N bond distance – longer Ni–N bond distance) has been observed experimentally in $\mathbf{1}$ (Δ is -0.12 Å and J = 5.0 cm⁻¹), $\mathbf{2}$ (Δ is -0.06 Å and J = 16.6 cm⁻¹) and \mathbf{IA} (Δ is -0.02 Å and J = 25.6 cm⁻¹); the linear magneto-structural correlation, on the basis of the experimental data of $\mathbf{1}$, $\mathbf{2}$ and \mathbf{IA} , between exchange integral and asymmetry is

shown in figure 7. Thus, asymmetry in the two nickel–azide bond distances has been established both experimentally and theoretically as a major factor to govern the extent of ferromagnetic interaction in heterobridged μ -phenoxo- $\mu_{1,1}$ -azide dinickel(II) compounds.

4. Crystal structures and magneto-structural correlation studies and theoretical calculations of a unique family of single end-to-end azide bridged Ni₄¹¹ cyclic clusters²⁷

The crystal structures of $[\{Ni^{II}(L^4)(\mu_{1,3}-N_3)(H_2O)\}_4]$ (5) and $[\{Ni^{II}(L^5)(\mu_{1,3}-N_3)(H_2O)\}_4]$ (6) are shown in figures 8 and 9. Overall, the structures of both species 5 and 6 can be described in a similar way; where all Ni^{II} centres within each molecule are hexacoordinated and bounded to $[L^4]^-$ or $[L^5]^-$ through the phenoxo oxygen, imine and piperidine/morpholine nitrogen atoms of the corresponding ligand. The remaining coordination sites are satisfied by one molecule of H2O and two nitrogen atoms from N₃⁻ anions. The latest act as bridges between Ni^{II} ions and eventually, only four azido groups are linked to the same number of Ni^{II} centres resulting in the formation of cyclic [Ni^{II}₄] systems. Interestingly, compounds 5 and 6 are the sole examples of tetranuclear clusters generated exclusively by end-to-end azide bridging ligands to date.

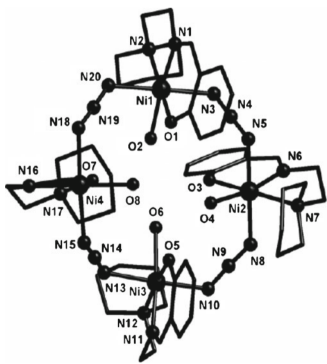


Figure 8. Crystal structure of $[\{Ni^{II}(L^4)(\mu_{1,3}-N_3)(H_2O)\}_4]$ (5). All the hydrogen atoms are omitted for clarity. ^{27,33}

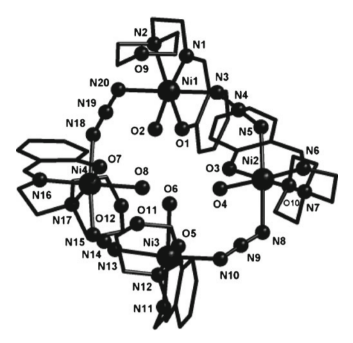


Figure 9. Crystal structure of $[\{Ni^{II}(L^5)(\mu_{1,3}-N_3)(H_2O)\}_4]$ **(6)**. All the hydrogen atoms are omitted for clarity. ^{27,33}

The magnetic data of **5** and **6** are shown in figures 10 and 11 as $\chi_M T$ versus T plots. It is interesting that the magnetic behaviour of the two complexes is drastically different; Interaction is antiferromagnetic in one, 5, but ferromagnetic in the second, 6. For the end-toend azide bridged systems, two key parameters which govern the magnetic properties are Ni-N-N angles and Ni–N···N–Ni torsion angles, τ . 1,18,19 Between these two, the Ni-N-N angles range almost similarly in the two complexes and so the difference in magnetic properties is expected to originate from the difference in the τ values: three types in 5 (87.5/90.6°, 50.4° and 3.3°) but two types in **6** (85.4/86.1° and 54.7/57.8°). It is known that the interaction should be antiferromagnetic if τ is 0° and the antiferromagnetism decreases as τ increases from 0° to 90° . Hence, compound 5 has an overall moderate antiferromagnetic behaviour due to the presence of an exchange pathway with an unprecedented Ni–N··· N–Ni torsion angle (3.3°) close to 0° , while complex 6 exhibits a predominant ferromagnetic behaviour because the torsion angles in this case lie in between $ca. 50^{\circ}$ and 90° .

Depending on the types of the τ values, the magnetic data of 5 were modelled/fitted with three-J Hamiltonian $\mathbf{H} = -2J_a(\mathbf{S_1} \cdot \mathbf{S_2} + \mathbf{S_3} \cdot \mathbf{S_4}) - 2J_b(\mathbf{S_2} \cdot \mathbf{S_3}) -2J_c(\mathbf{S_4} \cdot \mathbf{S_1})$, while the two-J Hamiltonian $\mathbf{H} = -2J_a(\mathbf{S_1} \cdot \mathbf{S_2} + \mathbf{S_3} \cdot \mathbf{S_4}) -2J_b(\mathbf{S_2} \cdot \mathbf{S_3} + \mathbf{S_4} \cdot \mathbf{S_1})$ are used for **6**. The converging fitting parameters are listed in table 2. Importantly, J = -35.25 cm⁻¹ corresponding to $\tau = 3.3^\circ$ and is responsible for antiferromagnetism in **5**.

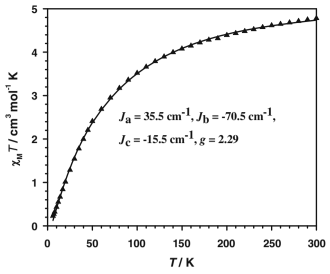


Figure 10. Observed (symbols) and calculated (solid lines) $\chi_M T$ versus T plots for $[\{Ni^{II}(L^4)(\mu_{1,3}-N_3)(H_2O)\}_4]$ (5). ^{27,33}

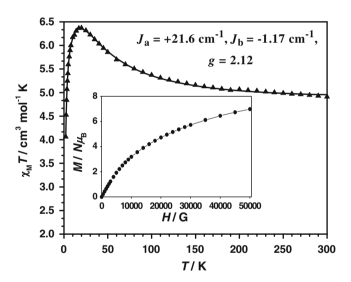


Figure 11. Observed (symbols) and calculated (solid lines) $\chi_M T$ versus T plots for $[\{Ni^{II}(L^5)(\mu_{1,3}-N_3)(H_2O)\}_4]$ (6). The inset is an isothermal $M/N\mu_B$ vs. H/G plot at 2.0 K. ^{27,33}

To model the magnetic properties theoretically, density functional theoretical calculations ³⁶ have been performed on **5** and **6** with the four-J Hamiltonian $\mathbf{H} = -2J_{12}(\mathbf{S}_1 \cdot \mathbf{S}_2) - 2J_{23}(\mathbf{S}_2 \cdot \mathbf{S}_3) - 2J_{34}(\mathbf{S}_3 \cdot \mathbf{S}_4) - 2J_{41}(\mathbf{S}_4 \cdot \mathbf{S}_1)$. The theoretical J values are compared with the experimental values in table 2; the trend is nicely matched. Interestingly, DFT-computed J values reproduce the trend of the $\chi_{\rm M}T$ versus T profile nicely (figure S7), indicating the ability of DFT methods in reproducing exchange Integrals.

Table 2.	Torsion	angles	and	experimental	and	density	functional	theoretical	J
values for	single en	d-to-end	d azio	de bridged tetra	anick	el(II) co	mpounds 5	and 6 .	

Ni–N··· N–Ni torsion angle (°)	J (DFT)/	cm ⁻¹	J (Expt)/ c	cm^{-1}
Antiferromagnetic compound [{I	$Ni^{II}(L^4)(\mu_{1,3}-N_3)$	$(H_2O)_{4}$ (5	5)	
90.6	+7.45	J_{12}	+17.75	J_a
3.3	-42.65	J_{23}	-35.25	J_b
87.5	+8.55	J_{34}	+17.75	J_a
50.4	-9.3	J_{41}	-7.75	J_c
Ferromagnetic compound [{Ni ^{II} ($(L^5)(\mu_{1.3}-N_3)(H_2)$	O)} ₄] (6)		
85.4	+7.05	J_{12}	+10.8	J_a
54.7	-2.25	J_{23}	-0.58	J_b
86.1	+7.75	J_{34}	+10.8	J_a
57.8	-0.6	J_{41}	-0.58	J_b

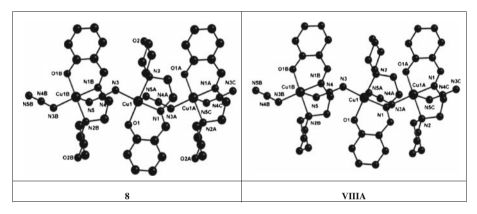


Figure 12. Crystal structures of $[Cu^{II}L^5(\mu_{1,3}-N_3)]_n \cdot 2nH_2O$ (**8**) and $[Cu^{II}L^5(\mu_{1,3}-N_3)]_n \cdot 2nH_2O$ (**VIIIA**). Hydrogen atoms and water molecules are not shown for clarity. Symmetry codes for **8**: A, -x, -0.5+y, 0.5-z; B, -x, 0.5+y, 0.5-z; C, x, -1+y, z. Symmetry codes for **VIIIA**: A, 2-x, 0.5+y, 0.5-z; B, 2-x, -0.5+y, 0.5-z; C, x, y-1, z. $2^{8,34,35}$

5. Crystal structures and magnetic properties of two one-dimensional end-to-end azide/cyanate bridged copper(II) compounds: new type of solid state isomerism²⁸

The perspective views of the end-to-end azide/cyanate bridged one-dimensional copper(II) systems $[Cu^{II}L^5(\mu_{1,3}\text{-NCO})]_n\cdot 2nH_2O$ (7) and $[Cu^{II}L^5(\mu_{1,3}\text{-NCO})]_n\cdot 2nH_2O$ (8) are shown in figures S8 and S12, respectively. While the phenoxo oxygen atom and imine and morpholine nitrogen atoms of one deprotonated ligand, $[L^5]^-$, satisfy the three coordination positions, the remaining two coordination positions of the pentacoordinated and distorted square pyramidal metal centre are satisfied by two nitrogen atoms of two end-to-end bridging azide ligands for 8 and by one nitrogen atom and one oxygen atom of two end-to-end bridging cyanate ligands for 7.

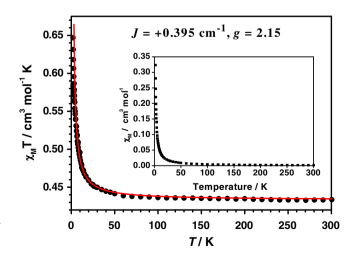


Figure 13. Fittings of the $\chi_M T$ vs. T of $[Cu^{II}L^5(\mu_{1,3}-N_3)]_n \cdot 2nH_2O$ (8) between 2.0 and 300.0 K. The experimental data is shown as squares and the solid lines correspond to the theoretical values. Inset shows χ_M vs. T data. ^{28,35}

Table 3. Comparison of the crystallographic data of $[Cu^{II}L^5(\mu_{1,3}-N_3)]_n \cdot 2nH_2O$ (**8**) and $[Cu^{II}L^5(\mu_{1,3}-N_3)]_n \cdot 2nH_2O$ (**VIIIA**).

	8	VIIIA
Formula	C ₁₃ H ₁₇ N ₅ O ₄ Cu ^a	C ₁₃ H ₂₁ N ₅ O ₄ Cu ^a
Formula weight	370.86 ^a	374.90 ^a
Crystal colour	Blue	Blue
Crystal system	Monoclinic	Monoclinic
Space group	$P2_1/c$	$P2_1/c$
a, Å	9.4934(15)	9.497(6)
b, Å	9.2962(14)	9.410(2)
c, Å	18.377(3)	18.232(8)
$V, Å^3$	1606.4(4)	1604.5(13)
α , $^{\circ}$	90.00	90.00
β , \circ	97.902(2)	100.01(3)
γ , °	90.00	90.00
Z	4	4

^aFour water hydrogen atoms in **8** were not located. Actual formula and formula weight are same for both.

The magnetic data of 7 and 8 are shown in figure S9 and figure $13,^{28,35}$ respectively, as $\chi_{\rm M}T$ versus T plots. The profiles indicate the existence of ferromagnetic interaction in both the compounds. The magnetic data were fitted/modelled with the Hamiltonian $\mathbf{H} = -2J\Sigma\mathbf{S}\mathbf{A_i}\cdot\mathbf{S}\mathbf{A_{i+1}}.^{1a,37}$ As shown by the solid lines in figure S9 and figure 13, good quality fitting has been obtained with J = +0.095 cm⁻¹ and g = 2.13 for the cyanate analogue 7 and J = +0.395 cm⁻¹ and g = 2.15 for the azide analogue 8. It may be mentioned that compound 7 is the sole example of ferromagnetically coupled EE cyanate bridged 1-D copper(II) system.

Synthesis, crystal structure and magnetic properties of a compound $[Cu^{II}L^5(\mu_{1,3}-N_3)]_n \cdot 2nH_2O$ (VIIIA) have been reported previously.³⁴ Compound $[Cu^{II}L^{5}(\mu_{1,3}-N_{3})]_{n}\cdot 2nH_{2}O$ (8) and $[Cu^{II}L^{5}(\mu_{1,3}-N_{3})]_{n}\cdot 2nH_{2}O$ N_3]_n·2nH₂O (VIIIA) are derived from the same ligand (HL⁵) and both have same composition. The azidebridged one-dimensional topology of both 8 and VIIIA are also same (figure 12). So, it may seem that 8 and **VIIIA** is the same compound. However, interestingly, the magnetic behaviour of 8 and VIIIA are drastically different; 8 exhibits ferromagnetic interaction with $J = +0.395 \,\mathrm{cm}^{-1}$, while **VIIIA** exhibits antiferromagnetic interaction with J = -2.15 cm⁻¹. Thus, on the basis of the drastic difference in magnetic properties, it is clear that 8 and VIIIA is not the same compound but they are magnetic isomers. Interestingly, such type of magnetic isomerism is unprecedented. Now, comparison of the crystallographic data in table 3 reveals another interesting and unprecedented aspect: except β , other parameters are practically identical, revealing that this pair represents another new type of solid state isomerism which may be called crystallographic isomerism. Again, comparison of the structural parameters (table S1) indicates that some parameters are identical, while some others are different. On the basis of these similarities-dissimilarities of the structural parameters, the pair can also be called crystallographic isomerism. Now, let us look into the position of the two solvated water molecule in figure 14, revealing that positions of the water molecule are different. Moreover, the number/types of hydrogen bonds and overall supramolecular structures are also different; 8 is threedimensional (figures \$10 and \$11), while VIIIA is

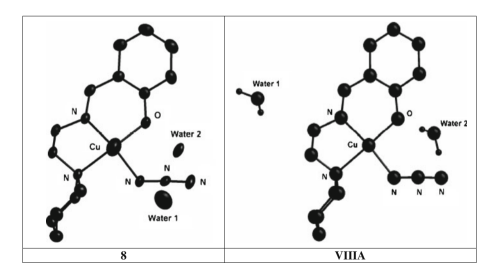


Figure 14. Perspective views in almost same orientation of $[Cu^{II}L^5(\mu_{1,3}-N_3)]_n \cdot 2nH_2O$ (**8**) and $[Cu^{II}L^5(\mu_{1,3}-N_3)]_n \cdot 2nH_2O$ (**VIIIA**) demonstrating the difference in location of the two solvated water molecule in the two structures. ^{28,34,35}

two-dimensional (figure \$12). Thus, **8** and **VIIIA** also represent rare example of supramolecular isomerism.

6. Conclusions

The outcome of the research work reviewed here can be summarized as follows: (i) unique observation of a pair of compounds for which all but one parameters, governing magnetic exchange interaction, are identical; (ii) unique example of bridge distance dependency of exchange integral; (iii) rare determination of several density functional theoretical magneto-structural correlations in heterobridged systems; (iv) profound role of asymmetry in the two nickel-azide bond distances in governing the exchange integral in heterobridged μ -phenoxo- $\mu_{1,1}$ -azide dinickel(II) systems, as obtained both experimentally and theoretically; (v) unique examples of only single end-to-end azide bridged tetranuclear complexes; (vi) observation of unprecedented small value, 3.3°, of Ni–N··· N–Ni torsion angle; (vii) demonstration of overall antiferromagnetic interaction in one single end-to-end azide bridged Ni^{II} system and overall ferromagnetic interaction in the second similar system as the function of the Ni-N···N-Ni torsion angles; (viii) reasonable/excellent matching of the DFT-computed and experimental J values; (ix) nice matching of the trends of the experimental $\chi_{\rm M}T$ data with the $\chi_M T$ data obtained from the DFT-computed J values; (x) first example of a ferromagnetically coupled end-to-end cyanate bridged one-dimensional copper(II) system; (xi) a unique pair representing new type of magnetic isomerism, new type of crystallographic isomerism and rare type of supramolecular isomerism.

Supplementary material

Figures S1–S12 and table S1 are available at http://www.ias.ac.in/chemsci.

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