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ARTICLE TYPE

Cation-size-controlled assembly of Ni(Ac)₂-1,4-H₂NDC system: geminal dicationic ionothermal syntheses, crystal structures and magnetic properties

Bing An,^a Yan Bai*,^{a,b} Jun-Li Wang^a and Dong-Bin Dang^{*a}

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Geminal dicationic ionic liquids with various alkyl spacer lengths regularly tune four MOF structures in two structural types of Ni(Ac)₂-1,4-H₂NDC system, which in turn bring about the magnetic properties divided into antiferromagnetic 10 and ferromagnetic couplings consistent with the structures.

Ionic liquids (ILs), which are credited with a number of unusual properties such as low vapor pressure, weakly coordinating properties and high thermal/chemical stability, have received considerable attention as "green alternatives" for synthesis and ¹⁵ application.¹ Geminal dicationic ionic liquids (DILs), which carry two positive unit charges, are a class of materials with growing portfolio of proposed applications as catalysts, solvents, lubricants and electrolytes etc.² In contrast to monocationic ionic liquids (MILs) with monovalent cations, DILs have been shown 20 to possess superior physical and chemical properties in terms of

higher densities and viscosities, greater thermal stabilities and longer liquid ranges.

The family of metal-organic frameworks (MOFs) is rapidly expanded with their rich structural aesthetics and optical, electric,

- 25 magnetic and other potential applications.³ Recently, directional design and synthesis of functional MOFs becomes a hot topic that most chemists make efforts to catch up with, such as understanding and chemically controlling the structures and the properties of these materials.⁴ The development of new synthetic
- 30 methods is undoubtedly an effective strategy to achieve the above target. Ionothermal synthesis, advantages over the traditional hydro/solvo-thermal methods, may open up new research directions: (a) ILs show a range of properties, which make them suitable for use as reaction media. (b) The variation of cations
- 35 and/or anions of ILs can often function as "structure directing agent" which may provide a means to direct the structures of MOFs. (c) ILs can act as charge-compensating groups and offer an additional and effective way of reaction control to form ionic liquid crystals (ILCs).⁵

Although a remarkable increase of ILs participating in the 40 synthesis of MOFs has been reported, all of them are MILs under investigation, especially imidazole-containing MILs. Mostly, some 1-alkyl-3-methylimidazolium MIL was used to study the influence on different metal-organic ligands systems.⁶ So far,

45 only few reports have explored how the alkyl chain length of 1-

alkyl-3-methylimidazolium cation (abbreviated $[C_nMIm]^+$) impacts on the structures.^{5c, 7} Compared to MILs, DILs contain two tunable functional terminal cation groups, which may give rise to significant changes in the MOF structures, but there is no 50 study in this aspect and it needs more research activities. In the present work, we synthesized and used four sorts of geminal 1,3bis(3-methylimidazolium-1-yl)alkyl bromide DILs with different alkyl spacer lengths from three to six carbons, as solvents. In order to isolate the effect of the cation parts, we have fixed all the 55 other variables such as the anion of DILs, the types of the reagent metal ion and the ligand. In comparison with the flexible and long organic carboxylic acid ligands, the size of rigid 1,4-H₂NDC (1,4- $H_2NDC = 1,4$ -naphthalenedicarboxylate acid) is suited to study the size of ionic liquids, and it also has the role of controlling the 60 aesthetically pleasing molecular structures.⁸ Thereby, adopting the Ni(Ac)₂-1,4-H₂NDC system, four MOFs in two structure types were obtained: $[C_3(MIm)_2][Ni(1,4-ndc)Br]_2$ 1 and $[C_4(MIm)_2][Ni(1,4-ndc)Br]_2$ (type A), $[Ni_3(1,4-ndc)_4(MIm C_5Im_2(H_2O_2)$ 3 and $[Ni_3(1,4-ndc)_4(MIm-C_6Im_2(H_2O_2))]$ 4 (type 65 B). Surprisingly, the magnetic properties of 1-4 are also divided into two types, dominant antiferromagnetic (AFM) and ferromagnetic (FM) coupling interactions, respectively, which correspond to the structures type A and type B.

The single crystal X-ray diffraction analysis reveals that the 70 structures of 1 and 2 feature a two-dimensional anionic layer sandwiched by DIL cations, $[C_3(MIm)_2]^{2+}$ and $[C_4(MIm)_2]^{2+}$, respectively (Fig. 1). Both structures consist of two [Ni(1,4ndc)Br]⁻ units and one DIL cation, in which 1,4-ndc²⁻ serves as a tetradentate ligand bridging four nickel ions through four 75 monodentate carboxylate oxygen atoms (Scheme S1). Ni(II) site lies in five-coordinated square pyramidal geometry with an O₄Br set from one Br⁻ and four 1,4-ndc²⁻ ligands. A pair of Ni centres are bridged by O-C-O rods from four cross-linked 1,4-ndc²⁻ ligands to form a paddlewheel-like cluster [Ni₂(O₂C)₄] as a 80 secondary building unit (SBU). The Ni-Ni separations are 2.85 Å for 1 and 2.84 Å for 2, respectively. Each SBU extends out to

connect the four adjacent SBUs by bridging interaction of 1,4 ndc^{2-} ligands giving rise to a 2D (4,4)-anionic layer (Fig. S3). The adjacent layers are stacked in an AB sequence, and the DIL 85 cations are located at the interlayer spaces.



Fig. 1 Schematic representations of the reactions, the SBUs and the structures of 1-4 in two types with DILs as multifunctional roles in the ionothermal synthesis. Note that the DILs tune structures to cause the magnetic properties with antiferromagnetic and ferromagnetic couplings, respectively.

The isostructural **3** and **4** are 3D network structures consisting of centro-symmetric trinuclear nickel SBUs and two crystallographically independent 1,4-ndc²⁻ ligands. The two 1,4ndc²⁻ adopt μ_2 -(η_1)-(η_1) and μ_4 -(μ_2 - η_1 : η_1)-(μ_2 - η_1 : η_1) bridging modes. What merits our attention is that the [MIm-C₅Im]⁺ and [MIm-C₆Im]⁺ segments come from the *situ* decomposition of corresponding raw material DILs, [C₅(MIm)₂]²⁺ and [C₆(MIm)₂]²⁺.Both of them lose one methyl to participate in coordination. Each of the two equivalent terminal Ni1 atoms is hexa-coordinated by four oxygen atoms of four 1,4-ndc²⁻ ligands in the equatorial plane of octahedron, one O1W of the bridging ¹⁵ water molecule and N1 of [MIm-C₅Im]⁺ or [MIm-C₆Im]⁺ locating

- in the axial positions. The coordination geometry around the central metal ion Ni2 is also a distorted octahedron involving four 1,4-ndc²⁻ ligands oxygen atoms from each of the four bridging bidentate carboxyl groups constituting the equatorial plane. The prior of the position of the two provides the two provides by the provides the prior bridging bridging and the prior of the position of the position of the two provides the two provides the prior bridging bridging bridging bridging at the position of the position o
- ²⁰ apical positions are occupied by two symmetric bridging water molecules. The linkages of Ni1 and Ni2 are two μ_2 - η_1 : η_1 carboxyl groups and μ_2 -H₂O. The distance between Ni1 and Ni2 is about 3.5 Å, indicating the absence of any bond between the two nickel centres. The Ni1–O1W–Ni2 angle in the bridge part is 113.6° for
- ²⁵ 3 and 115.9° for 4. Each Ni₃ segment formed by distorted cornersharing octahedra is double-cross-linked to eight different Ni₃ segments through two types of 1,4-ndc²⁻ backbones to generate a 3D framework. N-donor imidazole segments pointing to the voids of the framework have no contribution to dimension expansion
- ³⁰ (Fig. S10). If each Ni₃ unit acts as a node, the overall framework can be simplified as eight-connected CsCl-type topology (Fig. S11).⁹

The solid-state direct current (dc) magnetic measurements for 1-4 were measured in the temperature range of 1.8-300 K and in a

- ³⁵ magnetic field of 1 kOe (Fig. 2). The magnetic behaviors indicate both 1 and 2 (type A) are antiferromagnetic Ni(II)–Ni(II) exchange couplings inside their dimeric structures, while that of linear Ni(II) trimers in 3 and 4 (type B) exhibit the presence of weak ferromagnetic interactions. The distances between two
- ⁴⁰ nickel(II) ions within a well-isolated paddlewheel Ni(II) dimer unit connected through carboxyl groups in **1** and **2** suggest that the existence of the super-exchange coupling interactions among the metal centers.¹⁰ The magnetic data of **1** and **2** have been

analyzed using the Bleaney-Bowers (ESI eq 1) derived from the ⁴⁵ isotropic spin Heisenberg Hamiltonian $H = -2JS_1S_2$ (S = 1) and the exchange coupling constants of the fitting results are J =-29.76 cm⁻¹ for 1 and J = -3.44 cm⁻¹ for 2. Note that the magnetic coupling may also include a zero field splitting (ZFS) contribution, as is customarily observed in most Ni(II) ⁵⁰ complexes.¹¹ We also fit the magnetic properties considering ZFS factor in ESI. The result indicates that polymer 2 have a larger ZFS parameter than polymer 1, which may cause the differences ($\chi_M T vs T$ shape and J value) of 1 and 2.



⁵⁵ Fig. 2 Temperature variations of the magnetic susceptibility of 1-4 as $\chi_M T$ vs T. 1 and 2 show antiferromagnetic couplings (left); 3 and 4 (type B) exhibit the presence of weak ferromagnetic interactions (right).

For 3 and 4, we have fitted the magnetic data to the simple model derived for a centrosymmetrical S = 1 linear trimer with the ⁶⁰ Hamiltonian $H = -2J(S_1S_2 + S_2S_3)$, where S_2 is the spin state of the central Ni(II) ion. The consequence of J values is positive (J= 2.78 cm⁻¹ for 3 and J = 4.88 cm⁻¹ for 4.), which represents the exchange couplings between neighboring nickel centers are ferromagnetic. The structure of **3** and **4** shows that three Ni(II) 65 are connected by an agua and a carboxylate bridge. For the agua pathways, the Ni-O-Ni bond angles are out of the range 80-100° for ferromagnetic coupling. Hence, the overall ferromagnetic interaction should be originated from the ferromagnetic contribution of carboxylate bridges that is larger than the 70 antiferromagnetic contribution of aqua bridges.¹² In short, the structures generated from different sorts of metal centers by ionthermal synthesis have some relatives to the magnetic properties (see ESI for details).

Remarkably, the crystal structures of 1-4 reveal that multiple ⁷⁵ chemical identities of DILs participate into the architecture of the

final materials. As discussed above, type A is a typical (4,4) grid framework comprising the paddle-wheel SBUs, in which DIL cations as structure directing agents are located at the interlayer spaces as well as Br⁻ anions enter the final structures. And rich π

- s electronic imiadzole DILs have an advantage of generating $\pi \cdots \pi$ interactions to make them stable. Besides, grid size of the overall rigid framework can also be restrained by the size of ligand. Just as **1** and **2** even including the reported isostructural Cd-H₂NDC structure of [EMIm]⁺, small size ILs can perfectly and stably
- ¹⁰ accommodate inside.¹³ With increasing the length of the alkyl spacer of geminal DIL cations, owing to steric hindrance and charge repulsion, the DIL cations may be unstable to fill the voids and the structure turns to be 3D type B framework with the linear trinuclear Ni(II) SBUs. Apparently, there is a characteristic in
- ¹⁵ type B that partial decomposed DILs act as flexible ligands coordinating to Ni(II) ions. According to the previously reported about the thermal stability and degradation mechanism of ILs, imidazole MILs inclined to decompose into alkyl halogen and the residual imidazole group.¹⁴ As a consequence, this situ reaction ²⁰ of DILs provides a specific coordination access to generating
- diversity of MOF architectures.

Up to now, a large amount of chemists focus on "how anions of MILs tune the structures on different metal-carboxylic acid systems", but only few attentions on the MOFs controlled by

- 25 cation sizes with alkyl chain lengths regularly increased. Especially on that in the DILs syntheses, there is no report to the best of our knowledge. In our study, DILs as multiple functional reaction mediums are adopted at the first time to synthesize new MOFs. The systematically investigated and analyzed about the
- ³⁰ influence of the alkyl spacer lengths of DILs cations on the structures as well as the magnetic properties of the products, show that DILs can bring about a variety of magnetic properties by means of tuning the structures into different groups, which gives a possible way to synthesize functional MOF materials with ³⁵ structural controllability. Further studies on these aspects are
- called for.

In conclusion, we have explored the ionothermal synthesis of $Ni(Ac)_2-1,4-H_2NDC$ system in a host of DIL media, four geminal 1,3-bis(3-methylimidazolium-1-yl)alkyl bromide DILs with the

- ⁴⁰ alkyl spacer length varying from propyl to hexyl. Interestingly, we obtained four frameworks grouped into two structural types, and they can be master of the magnetic properties, in which type A exhibits AFM coupling interactions and type B is FM coupling interactions. It is expected to be importance in investigating the
- ⁴⁵ relationship between features which are controlled by ILs and resultant magnetic properties. Certainly, our work is just the beginning of such a paramount task, and a large volume of researches needs to be accumulated for.

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Notes and references

^aHenan Key Laboratory of Polyoxometalate Chemistry. Institute of 55 Molecular and Crystal Engineering, College of Chemistry and Chemical

Engineering, Henan University, Kaifeng 475004, P.R. China. ^bState Key Laboratory of Coordination Chemistry, Nanjing University, Nanjing 210093, P.R. China. *Fax:* (+86)-371-23881589; *Tel:* (+86)-371-23881589; *E-mail:* 60 <u>baiyan@henu.edu.cn;</u> <u>dangdb@henu.edu.cn</u>

† Electronic Supplementary Information (ESI) available: Experimental details; crystal data, additional figures, TG curves, XRPD and magnetic properties. CCDC reference numbers: 981577-981580. For ESI and crystallographicdata in CIF or other electronic format see 65 DOI: 10.1039/b000000x/

\$\$ Syntheses. A mixture of 1,4-H₂NDC (1 mmol), Ni(Ac)₂·6H₂O (1.5 mmol) and DILs (1.5 g) was transferred to a 25 mL Teflon-lined steel autoclave and kept at 180 °C for 3 days under autogenous pressure. After cooled down to room temperature, green crystals of 1-4 were obtained, 70 respectively.

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