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# Helical fluxionality: numerical frustration drives concerted low-barrier screw motions of a tricopper cluster†

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Uneven allocation of resources creates frustration, tension, and conflicts. Confronted with an apparent mismatch between the number of donor atoms and the number of metal atoms to be supported, helically twisted ligands cleverly come up with a sustainable symbiotic solution. As an example, we present a tricopper metallohelicate exhibiting screw motions for intramolecular site exchange. A combination of X-ray crystallographic and solution NMR spectroscopic studies revealed thermo-neutral site exchange of three metal centres hopping back and forth inside the helical cavity lined by a spiral staircase-like arrangement of ligand donor atoms. This hitherto unknown *helical fluxionality* is a superimposition of translational and rotational movements of molecular actuation, taking the shortest path with an extraordinarily low energy barrier without compromising the overall structural integrity of the metal-ligand assembly.

#### Introduction

The visually appealing shape of helices continues to inspire the creation of architectural objects and art forms featuring spiral structures. 1,2 In addition to macroscopic objects, helical structures abound in biological systems. Examples include doublestranded DNA<sup>3,4</sup> or protein α-helices,<sup>5,6</sup> in which shape complementarity serves as the underpinning mechanisms of information storage and spontaneous folding. To build structural mimics of such constructs, helical metal-ligand assemblies have been investigated extensively.7-17 Studies on metallohelicates have focused primarily on topological design,10,18-21 stereoselective synthesis, 22-26 chiroptical application,26-29 and DNA binding properties.30-34 Within this context, only a handful of works explored the dynamic aspects of helical metal complexes.15,35-37

As schematically shown in Fig. 1, a screw-like motion is a superimposition of translation and rotation, both of which have been realized with molecular machines and devices.<sup>38-42</sup> While conceptually appealing, spiral motion is difficult to implement with multinuclear metal complexes. Movement along the helical path (Fig. 1) requires concerted breaking and remaking of multiple metal-ligand bonds without

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compromising the overall structural integrity of the intertwined topology. To draw an analogy, a railroad track must be constantly removed and reinstalled as the train moves. This challenging synchronization is realized for the first time with a tricopper(1) cluster 1 built on a conformationally restricted  $\pi$ -conjugated ligand L (Scheme 1). The design, synthesis, and molecular mechanism of low-barrier spiral motions of the metallohelicate 1 constitute the main topics of this paper.

#### Results and discussion

#### Ligand design and cluster assembly

Our entry into the chemistry of the tricopper(i) complex 1 was aided by the ligand L (Scheme 1). Multiple aryl–aryl junctions of L not only restrict the conformational freedom of bond rotation, but also promote tight  $\pi$ - $\pi$  stacking upon metallation-induced

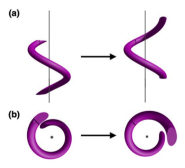
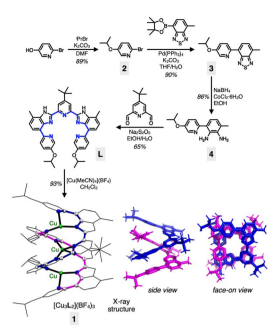


Fig. 1 Helical motion as a superimposition of translational and rotational motions viewed (a) parallel to, and (b) perpendicular to, the spiral axis.

<sup>†</sup> Electronic supplementary information (ESI) available: Experimental procedures, characterization data and supplementary figures, including <sup>1</sup>H and <sup>13</sup>C NMR spectra, crystallographic data, FT-IR spectra, HR-MS and elemental analysis. CCDC 2178526. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d3sc00851g



Scheme 1 Synthetic routes to L and  $[Cu_3L_2](BF_4)_3$  (1), and the X-ray structure of the cation. In the capped-stick representations (bottom right), the two intertwined ligand strands are color-coded (in blue and magenta) for better visualization of the double-helical topology; the metal atoms are omitted for clarity.

helical assembly. The solubility-enhancing isopropoxy and *tert*-butyl groups also served as convenient spectroscopic handles to probe the intriguing dynamic behaviour in solution (*vide infra*).

A pyridine-appended benzothiadiazole 3 was prepared by a Suzuki coupling reaction between bromopyridine 2 and the corresponding boronic ester (Scheme 1). The reduction of 3 to diamine 4, followed by an oxidative condensation reaction with the pyridine dialdehyde afforded L in a straightforward manner. The reaction between L and [Cu(MeCN)\_4](BF\_4) in CH\_2Cl\_2 produced 1 in high yield (93%). The [3 + 2]-type metal-to-ligand binding stoichiometry of this assembly was initially suggested by a Job plot analysis (Fig. S1†), and the resulting double-stranded helical structure of 1 was unambiguously established by single-crystal X-ray diffraction analysis on dark brown crystals grown by vapor diffusion of Et\_2O to a saturated CH\_3CN solution of the material.

As shown in Scheme 1 and Fig. S2 $\dagger$ , the tricopper(i) cluster 1 features a  $C_2$ -symmetric double-helical core with the tightly positioned ten N-donor atoms supporting an essentially linear alignment of the trimetallic array with  $\angle_{\text{Cu-Cu}} = 170.41(2)^{\circ}$ . The Cu···Cu intermetallic distances of 2.8000(5) Å are close to the sum of their van der Waals (vdW) radii (2.80 Å).<sup>43</sup> The metalligand bond distances range from 1.945(2) Å to 2.174(2) Å, typical for Cu<sup>I</sup>–N<sub>sp²</sub> bonds.<sup>10</sup> The helical ligand assembly is also aided by the tightly stacked pyridine···benzimidazole aromatic ring pairs with a centroid-to-centroid distance of 3.3894(14) Å.

With two pentadentate ligands supporting three metal centres, a numerical mismatch is inevitable. As shown in Scheme 1, ten N-donor atoms within 1 are allocated to three copper(1) centres to produce a combination of {3, 4, 3} as the

coordination numbers. Such inequality in the coordination environments, in conjunction with the kinetic lability of the d<sup>10</sup> electronic configuration of copper(I), promotes rapid site-exchanging motions, which is the topic of the following section.

### Helix is dynamic, not static: solvent-controlled and temperature-dependent fluxional motions

To probe whether the helical structure of **1** is also maintained in the solution phase, we proceeded to carry out NMR spectroscopic studies. In CDCl<sub>3</sub> at T=298 K, the <sup>1</sup>H NMR spectrum of **1** featured 12 different proton resonances in the aromatic region, which is consistent with the solid-state structure (Fig. 2a). In support of this notion, the 2D-ROESY NMR spectrum of **1** revealed "inter"ligand correlations between H2a/H2b and H<sub>Me</sub> protons (Fig. S4†) expected for the double-helical structure comprising two strands of **L**. The hydrodynamic radius of **1** cation determined by DOSY at different temperatures (Table S2; Fig. S5 and S6†) are also in good agreement with the crystallographically determined molecular dimensions (Fig. S7†).

Upon metallation to become part of the helicate, the symmetry-related proton pairs in the 1D <sup>1</sup>H NMR spectrum of the free ligand **L** (Fig. S8†) become no longer equivalent. For example, the two isopropyl C–H protons of **1** are now in different environments: one in the "inner groove" (H2a in Fig. 2e and f) and the other at the "outer end" (H2b in Fig. 2e

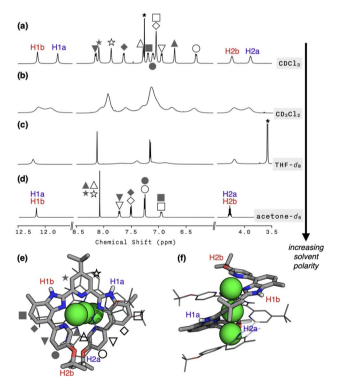
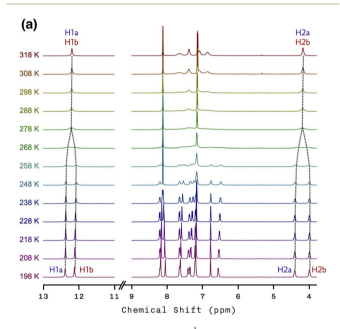


Fig. 2  $^{1}$ H NMR spectra of 1 in (a) CDCl<sub>3</sub>, (b) CD<sub>2</sub>Cl<sub>2</sub>, (c) THF- $d_8$ , and (d) acetone- $d_6$  at T=298 K. The residual solvent peaks are indicated by asterisks. (e) Top view, and (f) side view of the capped-stick (for one strand) and wireframe (for the other strand) representation of the X-ray structure of 1 cation, with the representative protons labelled. Green spheres are copper atoms.

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and f) of the helix. In stark contrast, a highly symmetric spectral pattern emerged in a polar solvent such as acetone- $d_6$ ; only six resonances were observed in the aromatic region (Fig. 2d). Solvents of intermediate polarities, such as  $CD_2Cl_2$  or THF- $d_8$ , produced unusually broadened spectra (Fig. 2b and c). Regardless of the choice of solvent, however, molecular ion peaks of m/z = 804 and 1694, corresponding to  $[Cu_3L_2](BF_4)^{2+}$  and  $[Cu_3L_2](BF_4)^{2+}$ , respectively, were consistently observed by CSI-MS (Fig. S9†).

To understand this peculiar solvent-dependent behaviour of 1, temperature-dependent <sup>1</sup>H NMR spectra were obtained in various solvents (Fig. 3a and S10–S13†). At sufficiently low temperatures, regardless of the choice of solvent, the spectral



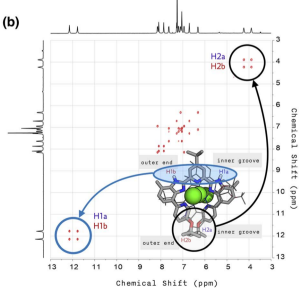


Fig. 3 (a) Temperature-dependent (from  $T=198~{\rm K}$  to 318 K)  $^1{\rm H}$  NMR spectra of 1 in THF- $d_8$ . (b) 2D  $^1{\rm H}$ - $^1{\rm H}$  EXSY NMR spectrum of 1 with  $t_{\rm m}=100~{\rm ms}$  in CDCl $_3$  at  $T=298~{\rm K}$ .

patterns of twelve well-resolved aromatic resonances are consistent with the solid-state structure of **1**. Upon increasing the temperature, however, the sharp peaks began to collapse into broad signals (Fig. 3a). In polar solvents such as THF- $d_8$  or acetone- $d_6$ , the resonances eventually coalesce into simple and symmetric patterns. Such spectral transitions could not be observed in CDCl<sub>3</sub> or CD<sub>2</sub>Cl<sub>2</sub> (Fig. S10 and S11†), presumably because the coalescence temperatures ( $T_c$ ) lie above the solvent's boiling point. Nonetheless, the overall trend of this fluxional behaviour unambiguously points toward the involvement of dynamic intramolecular ligand exchange in solution.

Further evidence for the fluxionality came from 2D <sup>1</sup>H-<sup>1</sup>H NMR exchange spectroscopy (EXSY) on 1 in CD<sub>2</sub>Cl<sub>2</sub> and CDCl<sub>3</sub> (Fig. 3b, S14, and S15†). We found that every proton of the ligand has a one-to-one correspondence to the equivalent proton of the other ligand constituting the double-helical 1. This observation implies that the protons at the inner groove rapidly exchange positions with the corresponding protons at the outer end of the double helix, which is clearly seen for the imidazolyl N-H (*i.e.*, H1a and H1b) and isopropyl C-H (*i.e.*, H2a and H2b) protons denoted in Fig. 2e, f, and 3b.

#### Mechanism and energetics of helical fluxionality

Variable-temperature (VT) and 2D NMR spectroscopic studies (Fig. 3) suggest that the thermoneutral site exchange of 1 occurs by sliding motions of one ligand along the slope of a spiral path defined by the other ligand (Fig. 4a and b). To take the path of least resistance, this intramolecular screw motion is coupled to a concerted hopping of the three copper( $\iota$ ) centres up and down inside the helical cavity. The traveling distance of the ligand along the screw-axis is estimated to be 2.8 Å, which corresponds to the Cu···Cu intermetallic separation (Scheme 1; Fig. S1†).

The energetics of this site-exchange was investigated by NMR spectroscopy. Using the  $\Delta \nu$  value obtained at the slow-exchange limit in VT 1D NMR and the coalescence temperature ( $T_c$ ), the activation barrier  $\Delta G^{\ddagger}$  of the screw motion was estimated in THF- $d_8$  and acetone- $d_6$  (Table S3†).<sup>44</sup> For low boiling point solvents CDCl<sub>3</sub> and CD<sub>2</sub>Cl<sub>2</sub>, the  $\Delta G^{\ddagger}$  values were determined by VT 2D EXSY NMR (see ESI for details†). With the  $\Delta G^{\ddagger}$  value of 14.2 kcal mol<sup>-1</sup> obtained by VT 2D EXSY NMR (Fig. S18†) and the  $\Delta \nu$  value of 100 Hz determined by VT 1D NMR (Fig. S11†), the coalescence temperature of 1 in CD<sub>2</sub>Cl<sub>2</sub> is estimated to be 298 K. This prediction is consistent with the data shown in Fig. S11†, with  $T_c$  lying just above this temperature point.

As shown in Fig. 4c, the energy barrier of the fluxional motion correlates inversely with the solvent dipole moment, with polar solvent facilitating the site exchange. In the most polar solvent acetone- $d_6$ , the  $\Delta G^{\ddagger}$  value is as low as 10.5 kcal mol<sup>-1</sup>. For comparison, the energy barrier of cyclohexane ring flip is 10.8 kcal mol<sup>-1</sup>. Considering that the fluxional motion of 1 requires concomitant dissociation and reassociation of at least four Cu-N bonds (Fig. 4a and b), this low-barrier ligand substitution cascade is quite remarkable. The spiral staircase-like arrangement of the ligand donor atoms inside the helical cavity plays a critical function in this low-barrier fluxional motion. Buried inside the tightly  $\pi$ - $\pi$ 

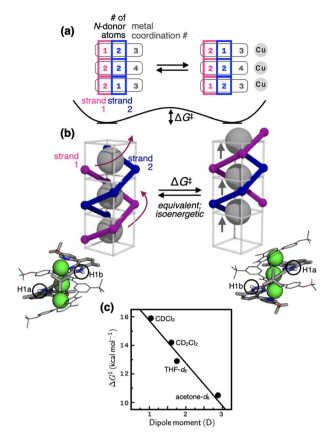


Fig. 4 (a) Changes in the number of N-donor atoms contributed by each ligand strand (blue and magenta) to the individual copper centres (gray) during the fluxional process schematically shown in (b). On each side of the schematic model in (b) is shown the corresponding view of the X-ray structure; the N-H protons are labelled as H1a (inner groove) and H1b (outer end), which exchange positions (see the EXSY spectrum in Fig. 3b) by the screw motion. (c) The relationship between the solvent dipole moments<sup>46</sup> and the activation barrier ( $\Delta G^{\ddagger}$ ) defined in (b).

stacked double helix, they enable rapid dissociation, migration, and reassociation of the Cu–N bond as the metal centres hop through the shortest path along the axis of the helix (Fig. 4). Apparently, polar solvent environments facilitate this process by assisting metal–ligand bond polarization (Fig. 4c).

Aside from the mechanistic aspects of the helical fluxionality, what is the origin of the stereochemical instability of this particular metallohelicate? Within 1, the two intertwined pentadentate ligands provide a total of ten donor atoms that need to be shared by three metal centres. This apparent numerical mismatch creates inequalities in the coordination numbers as  $\{3,4,3\}$  (Scheme 1). A constant movement of each ligand tries to resolve this imbalance by distributing its five N-donor atoms as a set of either 2–2–1 or 1–2–2 (Fig. 4a). The palindromic nature of these number combinations helps maintain the overall coordination number set of  $\{3,4,3\}$  for the three metal centres, each exchanging ligand partners as the screw oscillates. Equally important for the molecular motion here is an ideal balance between the non-stretchability (of the  $\pi$ -conjugated backbone) and the conformational adaptability (through aryl–aryl

rotations) of the ligand **L**. This structural feature ensures that as the head advances, the tail always follows by the same distance. Such a mechanical coupling is difficult to realize with typical metallohelicates, which are built with segmented bi/tridentate ligands connected by flexible alkyl/ether linkers.<sup>7-12,18,37</sup>

#### Conclusions

In sum, a double-stranded trimetallic helicate undergoes lowbarrier screw-like motions to alleviate the numerical imbalance of metal-ligand pairing. Through this hitherto unknown helical fluxionality, the metal atoms take the shortest path that traverses a spiral staircase-like arrangement of ligand donor atoms constituting the helical inner cavity.

By definition, fluxionality involves multiple conformers with no particular structural preference. An externally triggered directionality is thus needed to turn the current system into a new type of molecular actuator with programmable motions. Efforts are currently underway in our laboratory in such directions.

#### Data availability

The X-ray crystallographic coordinates for 1 have been deposited at the Cambridge Crystallographic Data Centre (CCDC), under deposition number 2178526. Experimental procedures, NMR spectra, and supplementary figures are available in the ESI.†

#### **Author contributions**

H. Kim: conceptualization, formal analysis, investigation, methodology, visualization, writing-original draft, writing-review & editing. J. Shin and S. Kim: conceptualization, formal analysis, investigation, methodology. D. Lee: conceptualization, funding acquisition, project administration, supervision, validation, writing-original draft, writing-review & editing.

#### Conflicts of interest

There are no conflicts to declare.

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