## SUPPORTING INFORMATION

# Homochiral crystallization of single-stranded helical coordination polymers: generated by the structure of auxiliary ligands or spontaneous symmetry breaking 

Shuai Ding, Yanfei Gao, Yufei Ji, Yanqin Wang and Zhiliang Liu*

College of Chemistry and Chemical Engineering; Key Laboratory of Nanomagnetic and Functional Materials, Inner Mongolia University, Hohhot, 010021, P. R. China.

Synthesis. All reagents and solvents were received from commercial supplies without further purification. (R)-2-amino-1-propanol, (S)-2-amino-1-propanol, 2-aminoethanol, o-vanillin, $\mathrm{CuCl}_{2}$, $\mathrm{NaN}(\mathrm{CN})_{2}, \mathrm{NaOH}$ and all the solvents were purchased from Alfa Aesar, respectively. The Schiff base ligands R- $\mathrm{H}_{2} \mathrm{~L}^{1}, \mathrm{~S}-\mathrm{H}_{2} \mathrm{~L}^{1}, \mathrm{H}_{2} \mathrm{~L}^{2}$ were synthesized according to literature procedures.

Synthesis of $\left[\mathbf{C u}\left(\mathbf{R}-\mathbf{H L}^{\mathbf{1}}\right)\left(\mu_{1,5} \text {-dca) }\right]_{\mathbf{n}}\right.$ (1). A mixture of $\mathrm{CuCl}_{2} \cdot 4 \mathrm{H}_{2} \mathrm{O}(0.207 \mathrm{~g}, 1 \mathrm{mmol}), \mathrm{R}-\mathrm{H}_{2} \mathrm{~L}^{1}$ $(0.208 \mathrm{~g}, 1 \mathrm{mmol}), \mathrm{NaN}(\mathrm{CN})_{2}(0.089 \mathrm{~g}, 1 \mathrm{mmol})$ and $\mathrm{NaOH}(0.040 \mathrm{~g}, 1 \mathrm{mmol})$ in 20 mL of methanol/ethanol/water (1:2:1) was stirred for 0.5 h . The resulting dark blue solution was left unperturbed to a slow evaporation of the solvent. After three days, dark green block-shaped crystals, suitable for X-ray diffraction analysis, were obtained. Elemental analysis (\%): Calc. for $\mathrm{C}_{13} \mathrm{H}_{14} \mathrm{CuN}_{4} \mathrm{O}_{3}$ : C, 46.22; H, 4.18; N, 16.59. Found: C, 46.53; H, 4.37; N, 16.21. IR (KBr pellet, $\left.\mathrm{cm}^{-1}\right): 3107 m, 2318 v s, 2242 v s, 2188 v s, 1641 s, 1601 m, 1445 m, 1322 m, 1301 m, 1247 m, 1027 m$, 975m, $871 w, 742 m, 630 w, 567 w$.

Synthesis of $\left[\mathbf{C u}\left(\mathbf{S}-\mathbf{H L}^{\mathbf{1}}\right)\left(\mu_{\mathbf{1 , 5}} \mathbf{- d c a}\right)\right]_{\mathbf{n}}$ (2). This compound was prepared using the same procedure as that described above for the synthesis of using $\mathrm{S}-\mathrm{H}_{2} \mathrm{~L}^{2}$ in place of $\mathrm{R}-\mathrm{H}_{2} \mathrm{~L}^{1}$. Elemental analysis (\%): Calc. for $\mathrm{C}_{13} \mathrm{H}_{14} \mathrm{CuN}_{4} \mathrm{O}_{3}$ : C, 46.22; H, 4.18; N, 16.59. Found: C, 46.46; H, 4.32; N, 16.25. IR (KBr pellet, $\mathrm{cm}^{-1}$ ): $3135 m, 2321 v s, 2259 v s, 2200 v s, 1620 s, 1587 m, 1429 m, 1380 m$, $1289 m, 1199 m, 1067 m, 954 m, 857 w, 739 m, 637 w, 580 w$.

Synthesis of $\left[\mathbf{C u}\left(\mathbf{H L}^{\mathbf{2}}\right)\left(\mu_{1,5} \text {-dca) }\right]_{\mathbf{n}}\right.$ (3). A mixture of $\mathrm{CuCl}_{2} \cdot 4 \mathrm{H}_{2} \mathrm{O}(0.207 \mathrm{~g}, 1 \mathrm{mmol}), \mathrm{H}_{2} \mathrm{~L}^{2}$ $(0.193 \mathrm{~g}, 1 \mathrm{mmol}), \mathrm{NaN}(\mathrm{CN})_{2}(0.089 \mathrm{~g}, 1 \mathrm{mmol})$ and $\mathrm{NaOH}(0.040 \mathrm{mg}, 1 \mathrm{mmol})$ in 20 mL of methanol/ethanol/water (1:2:1) was stirred for 3 h . The resulting dark blue solution was left unperturbed to a slow evaporation of the solvent. After 24 hours, dark green block-shaped crystals,
suitable for X-ray diffraction analysis, were obtained. Elemental analysis (\%): Calc. for $\mathrm{C}_{12} \mathrm{H}_{12} \mathrm{CuN}_{4} \mathrm{O}_{3}$ (1): C, 44.51; H, 3.74; N, 17.30. Found: C, 44.19; H, 3.81; N, 17.62. IR (KBr pellet, $\mathrm{cm}^{-1}$ ): $3119 m, 2320 v s, 2252 v s, 2193 v s, 1643 s, 1601 m, 1441 m, 1397 m, 1299 m, 1217 m$, $1075 m, 968 m, 856 w, 740 m, 638 w, 573 w$.

Crystallography. Single crystals of the complexes were selected and mounted on a Bruker ApexII CCD diffractometer with graphite-monochromated $\mathrm{Mo}-\mathrm{K} \alpha$ radiation $(\lambda=0.71073 \AA$ ), operating in $\omega-2 \theta$ scanning mode using suitable crystals for data collection. Lorentz-polarization correction was applied to the data. The structure was solved by direct methods (SHELX-97) and refined by full-matrix least-squares procedures on $F^{2}$ using SHELX-97. Hydrogen atoms were added theoretically and refined with riding model position parameters and fixed isotropic thermal parameters. Experimental details for the structural determinations are summarized in Table 1S.

Physical Measurements. Fourier transform infrared (FTIR) spectra (KBr disk) were measured with a Vertex 70 FTIR on a spectrophotometer $\left(4000-400 \mathrm{~cm}^{-1}\right)$. Elemental analyses for C, H and N were obtained from a Perkin-Elmer 2400 elemental analyzer. Circular dichroism (CD) spectra were measured as KBr pellet with a J-715 spectropolarimeter in the range of 200-900 nm at 298 K.

Table S1. Crystallographic Data and Structure Refinement for Complexes 1-3.

|  | $\mathbf{1}$ | $\mathbf{2}$ | $\mathbf{3}$ |
| :--- | :--- | :--- | :--- |
| Empirical Formula | $\mathrm{C}_{13} \mathrm{H}_{14} \mathrm{CuN}_{4} \mathrm{O}_{3}$ | $\mathrm{C}_{13} \mathrm{H}_{14} \mathrm{CuN}_{4} \mathrm{O}_{3}$ | $\mathrm{C}_{12} \mathrm{H}_{12} \mathrm{CuN}_{4} \mathrm{O}_{3}$ |
| Formula weight | 337.82 | 337.82 | 323.80 |
| Crystal system | Orthorhombic | Orthorhombic | Orthorhombic |
| Space group | $P 2_{1} 2_{1} 2_{1}$ | $P 2_{1} 2_{1} 2_{1}$ | $P 2_{1} 2_{1} 2_{1}$ |
| $a(\AA)$ | $9.586(2)$ | $9.589(2)$ | $9.596(2)$ |
| $b(\AA)$ | $10.713(2)$ | $10.709(2)$ | $10.990(2)$ |
| $c(\AA)$ | $13.527(3)$ | $13.528(3)$ | $12.258(3)$ |
| $\alpha, \beta, \gamma\left({ }^{\circ}\right)$ | $90,90,90$ | $90,90,90$ | $90,90,90$ |
| $V\left(\AA^{3}\right)$ | $1389.1(5)$ | $1389.2(5)$ | $1292.7(4)$ |
| $Z$ | 4 | 4 | 4 |
| $T(\mathrm{~K})$ | $293(2)$ | $293(2)$ | $293(2)$ |
| $\rho_{\text {calcd }}\left(\mathrm{g} \mathrm{g} \mathrm{cm}^{-3}\right)$ | 1.615 | 1.615 | 1.664 |
| Abs. coefficient $\left(\mathrm{mm}^{-1}\right)$ | 1.588 | 692 | 1.702 |
| $F(000)$ | 692 | $2.43<\theta<28.28$ | 660 |
| $\theta$ Range $\left({ }^{\circ}\right)$ | $2.43<\theta<28.27$ | 10187 | $2.49<\theta<27.86$ |
| Reflections collected | 10202 | $3422,0.0195$ | 13542 |
| unique reflns, $R_{\text {int }}$ | $3428,0.0199$ | 1.027 | $3082,0.0446$ |
| GOF on $F^{2}$ | 1.038 | $\mathrm{R}^{a)}=0.0235, \mathrm{wR}{ }^{b)}=0.0627$ | $\mathrm{R}^{a)}=0.0243, \mathrm{wR}{ }^{b)}=0.0512$ |
| Final R indices $[\mathrm{I}>2 \sigma(\mathrm{I})]$ | $\mathrm{R}^{a)}=0.0235, \mathrm{wR}{ }^{b)}=0.0607$ |  |  |
| $R$ indices $($ all data $)$ | $\mathrm{R}=0.0272, \mathrm{wR}=0.0623$ | $\mathrm{R}=0.0255, \mathrm{wR}=0.0635$ | $\mathrm{R}=0.0298, \mathrm{wR}=0.0518$ |
| Flack parameter | $0.011(10)$ | $0.014(10)$ | $0.009(18)$ |

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Scheme S1. The structures of the bromine-substituted chelate ligands $\left(\mathrm{R}-\mathrm{H}_{2} \mathrm{~L}^{3}, \mathrm{~S}-\mathrm{H}_{2} \mathrm{~L}^{3}\right.$, $\mathrm{H}_{2} \mathrm{~L}^{4}$ ).


Figure S1. A perspective view of structure for $\left[\mathbf{C u}\left(\mathbf{R}-\mathbf{H L}^{3}\right)\left(\mathbf{N}(\mathbf{C N})_{2}\right)\right]_{2}(4)$. Hydrogen atoms are omitted for clarity. Selected bond distances $[\AA]$ and angles [ ${ }^{\circ}$ ]: $\mathrm{Cu} 1-\mathrm{O} 11.910(6), \mathrm{Cu} 1-\mathrm{N} 1$ 1.932(7), $\mathrm{Cu} 1-\mathrm{N} 2$ 1.932(7), $\mathrm{Cu} 1-\mathrm{O} 2$ 1.963(7), $\mathrm{Cu} 1-\mathrm{O} 1 \mathrm{~A}$ 2.580(7), O1-Cu1-N1 94.3(3), N2-Cu1-O2 91.4(3), N1-Cu1-O2 81.9(3), O1-Cu1-O1A 88.7(3), N2-Cu1-O1A 91.0(3), N1-Cu1-O1A 95.3(3), O2-Cu1-O1 91.8(3) (symmetry code A: $-x+2,-y+1,-z$ ). This compound was prepared using the same procedure as that described above for the synthesis of using $\mathrm{R}-\mathrm{H}_{2} \mathrm{~L}^{3}$ in place of $\mathrm{R}-\mathrm{H}_{2} \mathrm{~L}^{1}$. Elemental analysis (\%): Calc. for $\mathrm{C}_{24} \mathrm{H}_{22} \mathrm{Br}_{2} \mathrm{Cu}_{2} \mathrm{~N}_{8} \mathrm{O}_{4}$ (1): C, 37.27; H, 2.87; N, 14.49. Found: C, $37.61 ; \mathrm{H}, 2.55$; N, 14.28. IR ( KBr pellet, $\mathrm{cm}^{-1}$ ): 2972m, 2316vs, 2255vs, 2179vs, 1637s, 1590m, 1456m, 1378m, 1293m, 1177m, 1040m, 830m, 796w, $672 m, 645 w, 550 w$.


Figure $\mathbf{S 2}$. A perspective view of structure for $\left[\mathbf{C u}\left(\mathbf{S}-\mathbf{H L}^{\mathbf{3}}\right)\left(\mathbf{N}(\mathbf{C N})_{2}\right)\right]_{2}(5)$. Hydrogen atoms are omitted for clarity. Selected bond distances $\left[\AA\right.$ ] and angles [ ${ }^{\circ}$ ]: $\mathrm{Cu} 1-\mathrm{O} 11.915(10), \mathrm{Cu} 1-\mathrm{N} 2$ $1.949(14), \mathrm{Cu}-\mathrm{N} 1$ 1.949(13), Cu1-O2 1.971(11), Cu2-O3 1.914(10), Cu2-N5 1.938(13), Cu2-N6 1.941(14), Cu2-O4 1.960(11), O1-Cu1-N2 92.4(5), O1-Cu1-N1 94.2(5), N2-Cu1-N1 171.2(6), $\mathrm{O} 1-\mathrm{Cu} 1-\mathrm{O} 2176.8(5), \mathrm{N} 2-\mathrm{Cu} 1-\mathrm{O} 290.8(5), \mathrm{N} 1-\mathrm{Cu} 1-\mathrm{O} 282.6(5), \mathrm{O} 3-\mathrm{Cu} 2-\mathrm{N} 594.1(5)$, O3-Cu2-N6 91.9(5), N5-Cu2-N6 171.0(6), O3-Cu2-O4 175.4(5), N5-Cu2-O4 81.3(5), N6-Cu2-O4 92.7(5). This compound was prepared using the same procedure as that described above for the synthesis of using $\mathrm{S}-\mathrm{H}_{2} \mathrm{~L}^{4}$ in place of $\mathrm{H}_{2} \mathrm{~L}^{1}$. Elemental analysis (\%):Calc. for $\mathrm{C}_{24} \mathrm{H}_{22} \mathrm{Br}_{2} \mathrm{Cu}_{2} \mathrm{~N}_{8} \mathrm{O}_{4}$ (1): C, 37.27; H, 2.87; N, 14.49. Found: C, 37.59; H, 2.66; N, 14.17. IR ( KBr pellet, $\mathrm{cm}^{-1}$ ): $3124 m, 2333 \mathrm{vs}, 2262 v s, 2183 v s, 1653 \mathrm{~s}, 1611 \mathrm{~m}, 1451 \mathrm{~m}, 1387 \mathrm{~m}, 1289 \mathrm{~m}, 1227 \mathrm{~m}$, $1065 m, 948 m, 866 w, 730 m, 628 w, 563 w$.


Figure S3. A perspective view of structure for $\left[\mathbf{C u}\left(\mathbf{H L}^{4}\right)\left(\mathbf{N}(\mathbf{C N})_{2}\right)\right]_{2}$ (6). Hydrogen atoms are omitted for clarity. Selected bond distances $\left[\AA\right.$ ] and angles [ ${ }^{\circ}$ ]: $\mathrm{Cu} 1-\mathrm{O} 31.8829(15)$, $\mathrm{Cu} 1-\mathrm{N} 1$ $1.9205(18), \mathrm{Cu} 1-\mathrm{N} 2$ 1.943(2), Cu1-O4 2.0256(17), Cu2-O1 1.8796(16), Cu2-N8 1.9160(18), $\mathrm{Cu} 2-\mathrm{N} 7 \quad 1.947(2), \quad \mathrm{Cu} 2-\mathrm{O} 2$ 2.0212(18), $\quad \mathrm{O} 3-\mathrm{Cu} 1-\mathrm{N} 1 \quad 94.07(7), \quad \mathrm{O} 3-\mathrm{Cu} 1-\mathrm{N} 2 \quad 94.56(8)$, N1-Cu1-O4 82.72(7), N2-Cu1-O4 90.07(8), O1-Cu2-N8 94.72(8), O1-Cu2-N7 91.45(9), N8-Cu2-O2 82.41(8), N7-Cu2-O2 92.66(9). This compound was prepared using the same procedure as that described above for the synthesis of using $\mathrm{H}_{2} \mathrm{~L}^{3}$ in place of $\mathrm{H}_{2} \mathrm{~L}^{1}$. Elemental analysis (\%): Calc. for $\mathrm{C}_{22} \mathrm{H}_{18} \mathrm{Br}_{2} \mathrm{Cu}_{2} \mathrm{~N}_{8} \mathrm{O}_{4}$ (1): C, 35.45 ; $\mathrm{H}, 2.43$; $\mathrm{N}, 15.03$. Found: $\mathrm{C}, 35.10 ; \mathrm{H}$, 2.61; $\mathrm{N}, 14.89$. IR ( KBr pellet, $\mathrm{cm}^{-1}$ ): 2964m, 2307vs, 2252vs, 2187vs, $1636 s, 1521 m, 1458 m$, $1375 m, 1318 m, 1175 m, 1087 m, 928 m, 883 w, 683 m, 648 w, 565 w$.

Table S2. Crystallographic Data and Structure Refinement for Complexes 4-6.

|  | 4 (CCDC 933907) | 5 (CCDC 933908) | 6 (CCDC 933909) |
| :---: | :---: | :---: | :---: |
| Empirical Formula | $\mathrm{C}_{24} \mathrm{H}_{22} \mathrm{Br}_{2} \mathrm{Cu}_{2} \mathrm{~N}_{8} \mathrm{O}_{4}$ | $\mathrm{C}_{24} \mathrm{H}_{22} \mathrm{Br}_{2} \mathrm{Cu}_{2} \mathrm{~N}_{8} \mathrm{O}_{4}$ | $\mathrm{C}_{22} \mathrm{H}_{18} \mathrm{Br}_{2} \mathrm{Cu}_{2} \mathrm{~N}_{8} \mathrm{O}_{4}$ |
| Formula weight | 773.40 | 773.40 | 745.34 |
| Crystal system | Triclinic | Triclinic | Monoclinic |
| Space group | $P-1$ | $P-1$ | $P 2_{1} / c$ |
| $a(\AA)$ | 8.224(2) | 9.518(4) | 13.724(3) |
| $b$ ( ( ) | 9.513(2) | 11.058(4) | 13.478(3) |
| $c(\AA)$ | 10.460(2) | 14.814(6) | 15.482(7) |
| $\alpha, \beta, \gamma \quad\left({ }^{\circ}\right)$ | 64.03(3), 71.43(3), 83.95(3) | 92.275(7), 105.925(7), 109.584(6) | 90,117.98(2), 90 |
| $V\left(\AA^{3}\right)$ | 696.8(2) | 1397.6(10) | 2529.0(14) |
| Z | 1 | 2 | 4 |
| $T$ (K) | 153(2) | 296(2) | 153(2) |
| $\rho_{\text {calcd }}\left(\mathrm{g} \mathrm{~cm}^{-3}\right)$ | 1.843 | 1.838 | 1.958 |
| Abs. coefficient ( $\mathrm{mm}^{-1}$ ) | 4.440 | 4.427 | 4.889 |
| $F(000)$ | 382 | 764 | 1464 |
| $\theta$ Range ( ${ }^{\circ}$ ) | $2.27<\theta<28.38$ | $1.44<\theta<28.23$ | $1.68<\theta<28.30$ |
| Reflections collected | 5161 | 10226 | 18375 |
| unique reflns, $R_{\text {int }}$ | 3445, 0.0211 | 6831, 0.0304 | 6272, 0.0204 |
| $\text { GOF on } F^{2}$ | 1.034 | 0.969 | 1.057 |
| Final R indices [ $\mathrm{I}>2 \sigma(\mathrm{I})$ ] | $\mathrm{R}^{a)}=0.0759, \mathrm{wR}^{\text {b) }}=0.1943$ | $\mathrm{R}^{\text {a) }}=0.0659, \mathrm{wR}^{\text {b) }}=0.1619$ | $\mathrm{R}^{\text {a) }}=0.0271, \mathrm{wR}^{\text {b }}=0.0656$ |
| $R$ indices (all data) | $\mathrm{R}=0.0843, \mathrm{wR}=0.1973$ | $\mathrm{R}=0.1529, w \mathrm{R}=0.1906$ | $\mathrm{R}=0.0426, \mathrm{wR}=0.0707$ |

a) $R=\Sigma| | F_{0}\left|-\left|F_{c}\right|\right| \Sigma\left|F_{0}\right| \quad$ b) $w R=\left[\Sigma w\left(F_{0}{ }^{2}-F_{\mathrm{c}}{ }^{2}\right)^{2} / \Sigma\left(w F_{0}{ }^{4}\right)\right]^{1 / 2}$


Figure S4. PXRD patterns and simulated data from Crystallographic Information File for compound 1


Figure S5. PXRD patterns and simulated data from Crystallographic Information File for compound 2


Figure S6. PXRD patterns and simulated data from Crystallographic Information File for compound $\mathbf{3}$


Figure $\mathbf{S 7}$. The TGA curves for compound 1-3.

Table S3 Selected bond lengths $[\AA]$ and angles $\left[{ }^{\circ}\right]$ for 1-3

| Compound 1 |  |  |  |
| :---: | :---: | :---: | :---: |
| Cu1-N1 | 1.9302(16) | N1-Cu1-O3 | 82.31(6) |
| Cu1-N2 | 1.9426(18) | O2-Cu1-O3 | 159.62(6) |
| Cu1-N4 \#1 | 2.475(2) | N2-Cu1-O3 | 91.41(7) |
| Cu1-O2 | $1.9332(13)$ | N1-Cu1-N4 \#2 | 90.49(8) |
| Cu1-O3 | 2.0457(14) | O2-Cu1-N4 \#2 | 105.21(8) |
| N1-Cu1-O2 | 93.20(6) | N2-Cu1-N4 \#2 | 88.88(9) |
| N1-Cu1-N2 | 173.61(7) | O3-Cu1-N4 \#2 | 94.74(8) |
| O2-Cu1-N2 | 93.10(6) |  |  |
| Compound 2 |  |  |  |
| Cu1-N1 | 1.9318(15) | N1-Cu1-O3 | 82.26(6) |
| Cu1-N2 | 1.9421(17) | O2-Cu1-O3 | 159.60(6) |
| Cu1-N4 \#1 | 2.473(2) | N2-Cu1-O3 | 91.38(6) |
| Cu1-O2 | 1.9352(13) | N1-Cu1-N4 \#2 | 90.57(8) |
| Cu1-O3 | 2.0471(14) | O2-Cu1-N4 \#2 | 105.31(8) |
| N1-Cu1-O2 | 93.19(6) | N2-Cu1-N4 \#2 | 88.91(8) |
| N1-Cu1-N2 | 173.55(7) | O3-Cu1-N4 \#2 | 94.65(8) |
| O2-Cu1-N2 | 93.15(6) |  |  |
| Compound 3 |  |  |  |
| Cu1-N1 | 1.9196(19) | N1-Cu1-O3 | 82.05(8) |
| Cu1-N2 | 1.928(2) | N2-Cu1-O3 | 91.52(7) |
| Cu1-N4 \#1 | 2.359(2) \#1 | O2-Cu1-O3 | 161.94(6) |
| Cu1-O2 | 1.9356(15) | N1-Cu1-N4 \#2 | 89.28(8) |
| Cu1-O3 | 2.0412(16) | N2-Cu1-N4 \#2 | 92.07(8) |
| N1-Cu1-N2 | 173.57(9) | O2-Cu1-N4 \#2 | 95.49(7) |
| N1-Cu1-O2 | 93.41(8) | O3-Cu1-N4 \#2 | 101.89(7) |
| N2-Cu1-O2 | 92.72(7) |  |  |

Symmetry transformations used to generate equivalent atoms: \#1: $2-\mathrm{x}, \mathrm{y}+1 / 2,-\mathrm{z}+1 / 2 ; \# 2: 2-\mathrm{x}, \mathrm{y}-1 / 2,-\mathrm{z}+1 / 2$


Figure S8 The complex $\mathbf{3}$ was prepared under static (unstirred) condition, solid-state CD measurements for the result bulk materials


[^0]:    a) $R=\Sigma| | F_{0}\left|-\left|F_{c}\right| / \Sigma\right| F_{0} \mid \quad$ b) $w R=\left[\Sigma w\left(F_{0}{ }^{2}-F_{\mathrm{c}}{ }^{2}\right)^{2} / \Sigma\left(w F_{0}{ }^{4}\right)\right]^{1 / 2}$

