## SUPPORTING INFORMATION

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

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**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, CuCl<sub>2</sub>, NaN(CN)<sub>2</sub>, NaOH and all the solvents were purchased from Alfa Aesar, respectively. The Schiff base ligands  $R-H_2L^1$ ,  $S-H_2L^1$ ,  $H_2L^2$  were synthesized according to literature procedures.

Synthesis of  $[Cu(R-HL^1)(\mu_{1,5}-dca)]_n$  (1). A mixture of  $CuCl_2 \cdot 4H_2O$  (0.207 g, 1 mmol),  $R-H_2L^1$  (0.208 g, 1 mmol),  $NaN(CN)_2$  (0.089 g, 1 mmol) and NaOH (0.040 g, 1 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  $C_{13}H_{14}CuN_4O_3$ : C, 46.22; H, 4.18; N, 16.59. Found: C, 46.53; H, 4.37; N, 16.21. IR (KBr pellet,  $cm^{-1}$ ): 3107*m*, 2318*vs*, 2242*vs*, 2188*vs*, 1641*s*, 1601*m*, 1445*m*, 1322*m*, 1301*m*, 1247*m*, 1027*m*, 975*m*, 871*w*, 742*m*, 630*w*, 567*w*.

Synthesis of  $[Cu(S-HL^1)(\mu_{1,5}-dca)]_n$  (2). This compound was prepared using the same procedure as that described above for the synthesis of using S-H<sub>2</sub>L<sup>2</sup> in place of R-H<sub>2</sub>L<sup>1</sup>. Elemental analysis (%): Calc. for C<sub>13</sub>H<sub>14</sub>CuN<sub>4</sub>O<sub>3</sub>: C, 46.22; H, 4.18; N, 16.59. Found: C, 46.46; H, 4.32; N, 16.25. IR (KBr pellet, cm<sup>-1</sup>): 3135*m*, 2321*vs*, 2259*vs*, 2200*vs*, 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  $[Cu(HL^2)(\mu_{1,5}-dca)]_n$  (3). A mixture of CuCl<sub>2</sub>·4H<sub>2</sub>O (0.207 g, 1 mmol), H<sub>2</sub>L<sup>2</sup> (0.193g, 1 mmol), NaN(CN)<sub>2</sub> (0.089 g, 1 mmol) and NaOH (0.040 mg, 1 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  $C_{12}H_{12}CuN_4O_3$  (1): C, 44.51; H, 3.74; N, 17.30. Found: C, 44.19; H, 3.81; N, 17.62. IR (KBr pellet, cm<sup>-1</sup>): 3119*m*, 2320*vs*, 2252*vs*, 2193*vs*, 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 Mo-K $\alpha$  radiation ( $\lambda = 0.71073$  Å), 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 (4000–400 cm<sup>-1</sup>). 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.

	1	2	3
Empirical Formula	$C_{13}H_{14}CuN_4O_3$	$C_{13}H_{14}CuN_4O_3$	$C_{12}H_{12}CuN_4O_3$
Formula weight	337.82	337.82	323.80
Crystal system	Orthorhombic	Orthorhombic	Orthorhombic
Space group	$P2_{1}2_{1}2_{1}$	$P2_{1}2_{1}2_{1}$	$P2_{1}2_{1}2_{1}$
<i>a</i> (Å)	9.586(2)	9.589(2)	9.596(2)
<i>b</i> (Å)	10.713(2)	10.709(2)	10.990(2)
<i>c</i> (Å)	13.527(3)	13.528(3)	12.258(3)
$\alpha, \beta, \gamma$ (°)	90, 90, 90	90, 90, 90	90, 90, 90
$V(Å^3)$	1389.1(5)	1389.2(5)	1292.7(4)
Ζ	4	4	4
<i>T</i> (K)	293(2)	293(2)	293(2)
$\rho_{\text{calcd}} (\text{g cm}^{-3})$	1.615	1.615	1.664
Abs. coefficient (mm <sup>-1</sup> )	1.588	1.588	1.702
F(000)	692	692	660
$\theta$ Range (°)	$2.43 < \theta < 28.27$	$2.43 < \theta < 28.28$	$2.49 < \theta < 27.86$
Reflections collected	10202	10187	13542
unique reflns, R <sub>int</sub>	3428, 0.0199	3422, 0.0195	3082, 0.0446
GOF on $F^2$	1.038	1.027	0.960
Final R indices $[I > 2\sigma(I)]$	$R^{a} = 0.0235$ , w $R^{b} = 0.0607$	$R^{a} = 0.0235$ , w $R^{b} = 0.0627$	$R^{a} = 0.0243$ , w $R^{b} = 0.0512$
R indices (all data)	R = 0.0272, wR = 0.0623	R = 0.0255, wR = 0.0635	R = 0.0298, wR = 0.0518
Flack parameter	0.011(10)	0.014(10)	0.009(18)

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

a)  $R = \Sigma ||F_0| - |F_c|| / \Sigma |F_0|$  b)  $wR = [\Sigma w (F_0^2 - F_c^2)^2 / \Sigma (wF_0^4)]^{1/2}$ 



Scheme S1. The structures of the bromine-substituted chelate ligands ( $R-H_2L^3$ ,  $S-H_2L^3$ ,  $H_2L^4$ ).



**Figure S1**. A perspective view of structure for  $[Cu(R-HL^3)(N(CN)_2)]_2$  (4). Hydrogen atoms are omitted for clarity. Selected bond distances [Å] and angles [°]: Cu1–O1 1.910(6), Cu1–N1 1.932(7), Cu1–N2 1.932(7), Cu1–O2 1.963(7), Cu1–O1A 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 R-H<sub>2</sub>L<sup>3</sup> in place of R-H<sub>2</sub>L<sup>1</sup>. Elemental analysis (%): Calc. for C<sub>24</sub>H<sub>22</sub>Br<sub>2</sub>Cu<sub>2</sub>N<sub>8</sub>O<sub>4</sub> (1): C, 37.27; H, 2.87; N, 14.49. Found: C, 37.61; H, 2.55; N, 14.28. IR (KBr pellet, cm<sup>-1</sup>): 2972*m*, 2316*vs*, 2255*vs*, 2179*vs*, 1637*s*, 1590*m*, 1456*m*, 1378*m*, 1293*m*, 1177*m*, 1040*m*, 830*m*, 796*w*, 672*m*, 645*w*, 550*w*.



**Figure S2**. A perspective view of structure for  $[Cu(S-HL^3)(N(CN)_2)]_2$  (5). Hydrogen atoms are omitted for clarity. Selected bond distances [Å] and angles [°]: Cu1–O1 1.915(10), Cu1–N2 1.949(14), Cu1–N1 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), O1–Cu1–O2 176.8(5), N2–Cu1–O2 90.8(5), N1–Cu1–O2 82.6(5), O3–Cu2–N5 94.1(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 S-H<sub>2</sub>L<sup>4</sup> in place of H<sub>2</sub>L<sup>1</sup>. Elemental analysis (%):Calc. for C<sub>24</sub>H<sub>22</sub>Br<sub>2</sub>Cu<sub>2</sub>N<sub>8</sub>O<sub>4</sub> (1): C, 37.27; H, 2.87; N, 14.49. Found: C, 37.59; H, 2.66; N, 14.17. IR (KBr pellet, cm<sup>-1</sup>): 3124*m*, 2333*vs*, 2262*vs*, 2183*vs*, 1653*s*, 1611*m*, 1451*m*, 1387*m*, 1289*m*, 1227*m*, 1065*m*, 948*m*, 866*w*, 730*m*, 628*w*, 563*w*.



**Figure S3**. A perspective view of structure for  $[Cu(HL^4)(N(CN)_2)]_2$  (6). Hydrogen atoms are omitted for clarity. Selected bond distances [Å] and angles [°]: Cu1–O3 1.8829(15), Cu1–N1 1.9205(18), Cu1–N2 1.943(2), Cu1–O4 2.0256(17), Cu2–O1 1.8796(16), Cu2–N8 1.9160(18), Cu2–N7 1.947(2), Cu2–O2 2.0212(18), O3–Cu1–N1 94.07(7), O3–Cu1–N2 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 H<sub>2</sub>L<sup>3</sup> in place of H<sub>2</sub>L<sup>1</sup>. Elemental analysis (%): Calc. for C<sub>22</sub>H<sub>18</sub>Br<sub>2</sub>Cu<sub>2</sub>N<sub>8</sub>O<sub>4</sub> (1): C, 35.45; H, 2.43; N, 15.03. Found: C, 35.10; H, 2.61; N, 14.89. IR (KBr pellet, cm<sup>-1</sup>): 2964*m*, 2307*vs*, 2252*vs*, 2187*vs*, 1636*s*, 1521*m*, 1458*m*, 1375*m*, 1318*m*, 1175*m*, 1087*m*, 928*m*, 883*w*, 683*m*, 648*w*, 565*w*.

	4 (CCDC 933907)	<b>5</b> (CCDC 933908)	<b>6</b> (CCDC 933909)
Empirical Formula	$C_{24}H_{22}Br_2Cu_2N_8O_4$	$C_{24}H_{22}Br_{2}Cu_{2}N_{8}O_{4} \\$	$C_{22}H_{18}Br_{2}Cu_{2}N_{8}O_{4} \\$
Formula weight	773.40	773.40	745.34
Crystal system	Triclinic	Triclinic	Monoclinic
Space group	<i>P</i> -1	<i>P</i> -1	$P2_{1}/c$
a (Å)	8.224(2)	9.518(4)	13.724(3)
<i>b</i> (Å)	9.513(2)	11.058(4)	13.478(3)
<i>c</i> (Å)	10.460(2)	14.814(6)	15.482(7)
$\alpha, \beta, \gamma$ (°)	64.03(3), 71.43(3), 83.95(3)	92.275(7), 105.925(7), 109.584(6)	90, 117.98(2), 90
$V(Å^3)$	696.8(2)	1397.6(10)	2529.0(14)
Ζ	1	2	4
<i>T</i> (K)	153(2)	296(2)	153(2)
$\rho_{\text{calcd}} (\text{g cm}^{-3})$	1.843	1.838	1.958
Abs. coefficient (mm <sup>-1</sup> )	4.440	4.427	4.889
F(000)	382	764	1464
$\theta$ Range (°)	$2.27 < \theta < 28.38$	$1.44 < \theta < 28.23$	$1.68 < \theta < 28.30$
Reflections collected	5161	10226	18375
unique reflns, R <sub>int</sub>	3445, 0.0211	6831, 0.0304	6272, 0.0204
GOF on $F^2$	1.034	0.969	1.057
Final R indices $[I > 2\sigma(I)]$	$R^{a} = 0.0759$ , w $R^{b} = 0.1943$	$R^{a} = 0.0659$ , w $R^{b} = 0.1619$	$R^{a} = 0.0271$ , w $R^{b} = 0.0656$
R indices (all data)	R = 0.0843, wR = 0.1973	R = 0.1529, wR = 0.1906	R = 0.0426, wR = 0.0707

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

a)  $R=\Sigma||F_0|-|F_c||/\Sigma|F_0|$ 

b)  $wR = [\Sigma w (F_0^2 - F_c^2)^2 / \Sigma (wF_0^4)]^{1/2}$ 



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 3



Figure S7. The TGA curves for compound 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)				

Table S3 Selected bond lengths [Å] and angles [°] for 1-3  $\,$ 

Symmetry transformations used to generate equivalent atoms: #1: 2-x, y+1/2, -z+1/2; #2: 2-x, y-1/2, -z+1/2



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