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Hydrazone-based cobalt complexes toward multielectron redox and spin crossover[†]

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Hydrazone-based derivatives modified by substitution at different positions were utilized to prepare a series of bis-homoleptic cobalt complexes. One species, $[\text{Co}^{\text{III}}(\text{L1})_2]^+$ (1), which incorporated deprotonated ligands, adopted a Co^{III} diamagnetic ground state. However, the substituent of a hydrogen atom with a methyl group precluded the possibility of deprotonation upon metal coordination, which led to two species, $[\text{Co}^{\text{II}}(\text{L2}^{\text{Me}})_2]^{2+}$ (2) and $[\text{Co}^{\text{II}}(\text{L3}^{\text{NO}_2})_2]^{2+}$, (3) which underwent a gradual spin crossover with an adjustable substituent effect and a mixed character of low-spin (doublet) and high-spin (quartet) populations in wide temperature ranges. Depending on the electronic effects of the substituents on the ligand, the multielectron redox behavior of the cobalt center was systematically modulated as well. This result demonstrates redox-switchable spin crossover in a new hydrazone-based Co^{II} system, in which the deprotonation of the coordination pocket and substituent groups in aromatic ligands can have a profound effect on the redox potential and spin state of the metal center.

Introduction

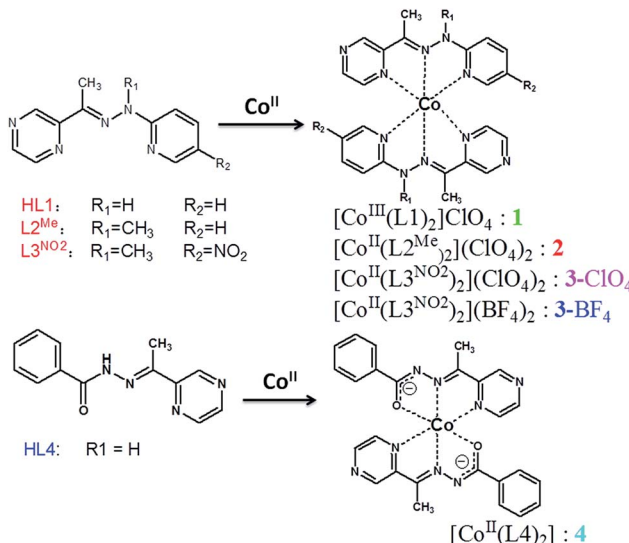
The study of molecular complexes based on a redox-controllable metal center has been a hot topic in research because of the potential application in the field of redox shuttles for dye-sensitized solar cells (DSSC),¹ anolytes for redox flow batteries, and as a redox mediator for water splitting and CO_2 reduction.^{2,3} The function of such complexes has been proved to be strongly dependent on the electronic property of the N-donor ligands.⁴ Among these subjects, polypyridyl complexes containing iron and ruthenium metal centers have been widely investigated as photosensitizers during the catalytic cycle.⁵ Although great achievements have been made concerning terpyridine ligands, the main shortcoming lie in the lack of efficient tools to investigate the new structure–function relationship, in particular for those with electronically tunable substituents, which still deserve to be currently studied.⁶ Another interesting point for cobalt complexes is that their spin state, electronic properties, and redox chemistry can be tailored by optimizing the ligand environment.^{7–10} For the purpose of potential application, it is of paramount importance to tune the donor type and ligand field strength of the ligands available to form the cobalt coordination sphere.^{11–13} Just recently, Yang *et al.* reported the

redox tuning and electronic effects of proximal nonredox active cations in cobalt Schiff-base complexes. The tunability in the reversible redox potential was ascribed to the control of the absolute energy of the molecular orbital participating in electron transfer.¹⁴ In the previous work, we investigated how the electronic effect of the ligand influences the spin crossover of Co^{II} ion within a metallogrid complex.¹⁵ Based on the electrochemical results, modification of the ligand field strength would give rise to a variation in the electronic structure (or molecular orbitals) of the redox active cation. However, spin crossover in a multinuclear system is not easy to control due to the synthetic difficulty of modifying the ligand field of the bis-multidendate ligand. In contrast, the mononuclear system could provide a convenient tool to tune the electrostatic effect, which would uniformly shift the molecular orbitals on the redox active metal.^{16–19} To further elucidate the source of the change in redox potential and to optimize the spin crossover of the Co^{II} ion starting from Schiff-base pyrazine ligands (Scheme 1, $\text{R}_1 = \text{R}_2 = \text{H}$: HL1 , $\text{R}_1 = \text{Me}$, $\text{R}_2 = \text{H}$: L2^{Me} , $\text{R}_1 = \text{Me}$, $\text{R}_2 = \text{NO}_2$: L3^{NO_2}), we synthesized a series of mononuclear cobalt complexes, aiming at achieving control of the local spin state as well as the redox chemistry. The ligand environments with the N_4O_2 donor permitted facile comparisons of the electronic structure. In compound 1, ligand **HL1** was deprotonated upon coordination and Co^{II} was oxidized to Co^{III} . In other cases, L2^{Me} and L3^{NO_2} were not deprotonated, whereas **HL4** in compound 4 was deprotonated, and divalent cobalt ion was stable. The ligand provided a coordination pocket of a transition-metal center, and thus could have a minimal effect on the coordination geometry of the cobalt center. Through this framework, a multielectron redox couple, such as $\text{Co}^{\text{III/II}}$ or $\text{Co}^{\text{II/I}}$, and ligand-based

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Scheme 1 General structures of the ligands and complexes in this work.

reduction were reversible, conveniently allowing systematic tuning of the electronic effect on the redox potential.

Although a handful of mononuclear 2,2':6',2''-terpyridine-derivative Co(II)-center spin-crossover compounds have been previously investigated,²⁰⁻²⁵ other systems have been rarely reported and should be further developed.²⁶⁻²⁸ In the previous work, we investigated the deprotonation effects at the coordination pocket, such as how L₁^H forms diamagnetic [Co(L₁^H)₂]X complexes irrespective of the counterion X and how the introduction of substituents at the imine position has an effect on the variation of the spin state.¹⁵ As a result, the cobalt(II) complex of 2,5-bis[1-[2-methyl-2-(pyridin-2-yl)-hydrazono]ethyl]pyrazine (L^{Me}), in which the methyl substituent was adopted to prevent deprotonation on the imine group, showed an incomplete spin crossover at higher temperature. In this paper we continue to expand the new ligand system and describe the synthesis of a number of ligands with modified substituents, aiming at achieving an enhanced spin crossover property of their Co(II) complexes, and also we discuss the effect of the substituents on the spin state of the cobalt(II) center.²⁹

Experimental section

Materials and general procedures

All the reagents employed were commercially available and used without further purification. Methanol and acetonitrile were dried using standard procedures. Predeuterated dimethyl sulfoxide (DMSO-*d*₆) was purchased from Alfa Aesar Co. Ltd., while 2-hydrazinylpyridine, benzohydrazide, 2-Br-5-nitropyridyl, and 2-acetylpyrazine were obtained from a commercial source and used directly without any purification. 1-Methyl-1-pyridylhydrazine was prepared according to reported procedures.³⁰

UV-visible studies were performed on a Perkin-Elmer Lambda 950 UV-vis instrument. Elemental analyses (C, H, and

N) were conducted with a Perkin-Elmer 2400 analyzer. Micro-IR spectroscopy studies were performed on a Nicolet Magna-IR 750 spectrophotometer in the 4000–400 cm⁻¹ region (w, weak; b, broad; m, medium; s, strong) with a KBr disc. ¹H NMR spectra were obtained from a solution in deuterated dimethyl sulfoxide using a Bruker-400 spectrometer. (s, singlet; d, doublet; t, triplet; m, multiplet; dd, double doublet). Magnetic susceptibility measurements were carried out using a superconducting quantum interference device (SQUID) magnetometer (Quantum Design MPMS-XL) under applied magnetic fields of 2500 Oe. Corrections for diamagnetism were applied using Pascal's constants.³¹ Cyclic voltammetry (CV) and differential pulse voltammetry (DPV) experiments were undertaken in acetonitrile (0.05 M [Bu₄N][PF₆]) at 293 ± 2 K using a CHI620E computer-controlled electrochemical workstation and a standard three-electrode cell employing a glassy carbon microelectrode as the working electrode, a Pt wire counter electrode, and a Ag wire in a tip separated from the bulk solution, which was used as the quasi-reference electrode. All the potentials given in this paper are referred to the ferrocene/ferrocenium ([FeCp₂]^{0/+}) reference couple under the same conditions as an internal reference.

Data collection and refinement

Single-crystal X-ray data for complexes 1–4 were collected on a Bruker APEX-2 CCD using Mo-K α radiation ($\lambda = 0.71073$ Å). Data collection, data reduction, and cell refinement were performed by using the Bruker Instrument Service v4.2.2 and SAINT V8.34A software, respectively.^{32,33} The structure was solved by direct methods using the SHELXS program, and refinement was performed using SHELXL based on F^2 through a full-matrix least squares routine.³⁴ Empirical multi-scan absorption corrections using equivalent reflections were performed with the SADABS program.³⁵ All the non-hydrogen atoms were refined with anisotropic displacement parameters. Hydrogen atoms were set in the calculated positions and refined as a riding model.^{36,37}

Synthetic procedures

Synthesis of 1-methyl-1-(4-nitropyridyl)hydrazine. The compound was synthesized according to the revised literature method.³⁰ To a round flask containing 2-Br-5-nitropyridyl (32 mmol, 6.5 g) was added excessive 40% methylhydrazine (480 mmol, 55 g) under a nitrogen atmosphere, which was stirred for 3 h at boiling temperature. After cooling to room temperature, the mixture was evaporated under reduced pressure to dry. The residue was dissolved in ethyl acetate (250 mL), and washed with sodium carbonate (10% aqueous solution, 50 mL × 3) and saturated brine (50 mL × 3). The organic phase was dried with anhydrous Na₂CO₃ for 2 h and was then evaporated under vacuum to produce a yellow solid. Yield: 3.39 g, 63%. The solid was directly used for the synthesis of L₃^{NO2} without NMR characterization.

Synthesis of [1-[2-(pyridin-2-yl)hydrazono]ethyl]pyrazine (HL1). A methanol solution mixture of 2-acetylpyrazine (1.2212 g, 0.01 mol) and 1-pyridylhydrazine (1.0913 g, 0.01 mol) was refluxed for 4 h, and after cooling to room temperature,



a pale-yellow crystal was obtained by filtration. The crude product was washed with cold methanol and dried *in vacuo*. Yield: 1.4721 g, 69%. ^1H NMR (400 MHz, DMSO- d_6): δ (ppm) = 2.37 (s, 3H), 6.88 (m, 1H), 7.44 (s, 0.5H), 7.47 (s, 0.5H), 7.71 (m, 1H), 8.20 (m, 1H), 8.52 (d, J = 2.64, 1H), 8.58 (m, 1H), 9.36 (d, J = 1.56, 1H), 10.22 (s, 1H). Elemental analysis (%) calculated for $\text{C}_{11}\text{H}_{11}\text{N}_5$: C, 61.96; H, 5.20; N, 32.84. Found: C, 62.12; H, 5.12; N, 32.78. MS (m/z) [M + H]⁺ calcd for $\text{C}_{11}\text{H}_{11}\text{N}_5$ 214.25, found 214.63.

Synthesis of [1-[2-methyl-2-(pyridin-2-yl)-hydrazono]ethyl] pyrazine (L2^{Me}). The pale-yellow ligand L2^{Me} was obtained by following the same procedure as that described for **HL1** except that 1-methyl-1-pyridylhydrazine was used instead of 1-pyridylhydrazine. Yield: 1.6798 g, 74%. ^1H NMR (400 MHz, DMSO- d_6): δ (ppm) = 2.49 (s, 3H), 3.49 (s, 3H), 6.89 (t, 1H), 7.18 (d, J = 8.40, 1H), 7.66 (t, 1H), 8.26 (d, J = 4.88, 1H), 8.68 (d, J = 7.04, 1H), 9.34 (s, 1H). Elemental analysis (%) calculated for $\text{C}_{12}\text{H}_{13}\text{N}_5$: C, 63.42; H, 5.77; N, 30.82. Found: C, 63.34; H, 5.65; N, 30.93. MS (m/z) [M + H]⁺ calcd for $\text{C}_{12}\text{H}_{14}\text{N}_5$ 228.27, found 228.84.

Synthesis of [1-[2-(5-nitropyridin-2-yl)-2-methylhydrazono]ethyl]pyrazine (L3^{NO2}). A toluene solution containing 2-acetylpyrazine (1.2212 g, 0.01 mol) and 1-methyl-1-(4-nitropyridyl)hydrazine (1.9217 g, 0.01 mol) was refluxed for 24 h, and after cooling to room temperature, a yellow solid was obtained by filtration. The crude product was washed with cold methanol and dried *in vacuo*. Yield: 2.2032 g, 81%. ^1H NMR (400 MHz, CDCl_3): δ (ppm) = 2.57 (s, 3H), 3.65 (s, 3H), 7.03 (s, 0.5H), 7.06 (s, 0.5H), 8.30 (dd, 1H), 8.64 (m, 2H), 9.18 (d, J = 2.60, 1H), 9.46 (d, J = 1.40, 1H). Elemental analysis (%) calculated for $\text{C}_{12}\text{H}_{12}\text{N}_6\text{O}_2$: C, 52.94; H, 4.44; N, 30.87. Found: C, 53.06; H, 4.52; N, 30.78. MS (m/z) [M + H]⁺ calcd for $\text{C}_{12}\text{H}_{13}\text{N}_6\text{O}_2$ 273.27, found 273.37.

Synthesis of [1-(benzohydrazone)ethyl] pyrazine (HL4). A methanol solution mixture of 2-acetylpyrazine (1.2212 g, 0.01 mol) and benzohydrazide (1.3615 g, 0.01 mol) was refluxed for 4 h, and after cooling to room temperature, a white crystal was obtained by filtration. The crude product was washed with cold methanol and dried *in vacuo*. Yield: 1.8718 g, 78%. ^1H NMR (400 MHz, DMSO- d_6): δ (ppm) = 2.43 (s, 3H), 7.52 (m, 2H), 7.59 (m, 1H), 7.76–8.01 (m, 2H), 8.63 (d, J = 9.88, 2H), 9.22 (s, 1H), 11.04 (s, 1H). Elemental analysis (%) calculated for $\text{C}_{13}\text{H}_{12}\text{N}_4\text{O}$: C, 64.99; H, 5.03; N, 23.32. Found: C, 64.58; H, 4.95; N, 23.56. MS (m/z) [M + H]⁺ calcd for $\text{C}_{13}\text{H}_{13}\text{N}_4\text{O}$ 241.27, found 241.22.

Caution! Perchlorate salts are potentially explosive and should be treated with great caution. Only small amounts were used in the present work.

Synthesis of complex $[\text{Co}^{\text{III}}(\text{L1})_2]\text{ClO}_4$ (1). $\text{Co}(\text{ClO}_4)_2 \cdot 6\text{H}_2\text{O}$ (1.6 mg, 4.5 μmol) was dissolved in 2 mL of distilled water and added to a test tube. Another 2 mL of water : MeOH (v/v = 1 : 1) solvent was carefully layered on the top of this solution as a buffer layer. Finally, a 2 mL solution of L1^H (1.9 mg, 9 μmol in MeOH) was layered on top of the buffer layer. The tiny tube was kept in the dark and left undisturbed without protection from air. Over the course of two weeks, dark-red block-shaped crystals suitable for X-ray diffraction studies were obtained at the buffer layer. Yield: 1.1 mg, 37%. IR (KBr pallet, cm^{-1}): 628(m),

780(m), 1092(s), 1316(m), 1472(m), 1621(m). Elemental analysis (%) calculated for $\text{C}_{22}\text{H}_{20}\text{ClCoN}_{10}\text{O}_4$: C, 45.33; H, 3.46; N, 24.03. Found: C, 45.44; H, 3.35; N, 24.23.

Synthesis of complex $[\text{Co}^{\text{II}}(\text{L2}^{\text{Me}})_2]\text{ClO}_4$ (2). L2^{Me} (0.7 mg, 3 μmol) was dissolved in a 2 mL of ethyl acetate and added to a test tube. Another 2 mL of ethyl acetate : MeOH (v/v = 1 : 1) solvent was carefully added to the top of this solution as a buffer layer. Finally, a 2 mL solution of $\text{Co}(\text{ClO}_4)_2 \cdot 6\text{H}_2\text{O}$ (0.5 mg, 1.5 μmol in MeOH) was layered on top of the buffer layer. The tiny tube was kept in the dark and left undisturbed without protection from air. Over the course of two weeks, dark-red block-shaped crystals unsuitable for X-ray diffraction studies were obtained at the interface. Yield: 0.17 mg, 16%. IR (KBr pallet, cm^{-1}): 625(m), 773(m), 850(w), 943(w), 1091(s), 1316(m), 1469(m), 1607(m). Elemental analysis (%) calculated for $\text{C}_{24}\text{H}_{26}\text{Cl}_2\text{CoN}_{10}\text{O}_8$: C, 40.46; H, 3.68; N, 19.66. Found: C, 40.72; H, 3.82; N, 19.44.

Synthesis of complex $[\text{Co}^{\text{II}}(\text{L3}^{\text{NO2}})_2]\text{ClO}_4 \cdot \text{CH}_3\text{OH}$ (3_{ClO4}). To a solution of L3^{NO2} (27.2 mg, 0.1 mmol) in acetonitrile (5 mL) was added a methanol solution (5 mL) of $\text{Co}(\text{ClO}_4)_2 \cdot 6\text{H}_2\text{O}$ (18.3 mg, 0.05 mmol). The resulting clear dark-red solution was stirred for 30 min at 55 °C and filtered. Dark-red flake-shaped crystals suitable for X-ray diffraction studies were obtained by the slow diffusion of ether vapor into the dark-red solution after 5 days without protection from air. The crystals were filtered off, washed with acetonitrile, and dried in air. Yield: 28.07 mg, 70%. IR (KBr pallet, cm^{-1}): 565(w), 624(m), 823(w), 1085(s), 1349(s), 1466(w), 1577(m), 1619(m). Elemental analysis (%) calculated for $\text{C}_{25}\text{H}_{28}\text{Cl}_2\text{CoN}_{12}\text{O}_{13}$: C, 35.99; H, 3.38; N, 20.14. Found: C, 35.92; H, 2.98; N, 20.88.

Complex $[\text{Co}^{\text{II}}(\text{L3}^{\text{NO2}})_2](\text{BF}_4)_2 \cdot 0.5\text{CH}_3\text{OH}$ (3_{BF4}) was prepared in a similar manner as for 3_{ClO4} except that $\text{Co}(\text{BF}_4)_2 \cdot 6\text{H}_2\text{O}$ was used instead of $\text{Co}(\text{ClO}_4)_2 \cdot 6\text{H}_2\text{O}$. Yield: 27.12 mg, 68%. Elemental analysis (%) calculated for $\text{C}_{24.5}\text{H}_{26}\text{B}_2\text{CoF}_8\text{N}_{12}\text{O}_{4.5}$: C, 37.10; H, 3.30; N, 21.19. Found: C, 36.92; H, 3.12; N, 21.76.

Synthesis of complex $\text{Co}^{\text{II}}(\text{L4})_2$ (4). $\text{Co}(\text{ClO}_4)_2 \cdot 6\text{H}_2\text{O}$ (3.7 mg, 10 μmol) was dissolved in 2 mL of distilled water and added to a test tube. Another 2 mL of water : MeOH (v/v = 1 : 1) solvent was carefully added to the top of this solution as a buffer layer. Finally, a 2 mL solution of **HL4** (4.8 mg, 20 μmol in MeOH) was layered on top of the buffer layer. The tiny tube was kept in the dark and left undisturbed without protection from air. Over the course of two weeks, dark-red block-shaped crystals suitable for X-ray diffraction studies were obtained at the interface. Yield: 2.0 mg, 38%. IR (KBr pallet, cm^{-1}): 626(w), 709(m), 1068(m), 1405(s), 1621(m). Elemental analysis (%) calculated for $\text{C}_{26}\text{H}_{22}\text{CoN}_8\text{O}_2$: C, 58.10; H, 4.13; N, 20.85. Found: C, 58.11; H, 4.21; N, 21.04.

Results and discussion

The non-symmetric ligands used in this work were prepared through reacting 1-pyridylhydrazine and commercially available 2-acetylpyrazine. It should be noted that the reaction to produce L3^{NO2} was difficult to occur, possibly due to the relatively low electron density on the carbonylhydrazide nitrogen, which was unfavorable to nucleophilic attack, and so forcing conditions



were required, namely 120 °C in toluene for 1 day, to complete the reaction. This method, previously used to generate the symmetric pyrazine-bridged bis-tridentate ligand (L^R),¹⁵ should open up access to a wide variety of non-symmetric pyrazine-based ligands. **HL1** and **HL4** were readily deprotonated so the complexes $[\text{Co}^{\text{III}}(\text{L1})_2]\text{ClO}_4$ (**1**) and $[\text{Co}^{\text{II}}(\text{L4})_2](\text{ClO}_4)_2$ (**4**) were initially prepared by simply combining a 2 : 1 ratio of the ligand and $[\text{Co}^{\text{II}}(\text{H}_2\text{O})_6](\text{ClO}_4)_2$ salt in methanol. However the oxidation of cobalt(II) in complex **1** cannot be avoided even by using a Schlenk flask in an inert atmosphere of nitrogen. So all the synthetic operations were undertaken in air. Single crystals were grown by a layering method of a CH_3OH solution of ligand **HL1** and **HL4** and, on one occasion, of an ethyl acetate–MeOH solution for **L2^{Me}**. However, the reaction between electron-deficient ligand **L3^{NO2}** and cobalt salt was not easy to occur at room temperature due to the dynamic inertia. Thus the syntheses of compounds **3_{ClO4}** and **3_{BF4}** were further optimized by holding the reaction suspension containing **L3^{NO2}** and cobalt salt in acetonitrile at 55 °C for 30 min, which was then diffused with ether vapor to produce higher-quality, more uniform single crystals suitable for X-ray diffraction analysis. It is necessary to note that the reaction to produce **1**, **2**, and **4** should be operated in a low concentration of reactants. The higher concentration was also attempted, but the diffusion reaction was so rapid that the precipitate with high impurity rather than crystals was obtained. Hence, to obtain spectroscopic, magnetic, and electrochemical data, we had to set up many reactions in parallel to accumulate enough sample.

The UV-vis absorption spectra of all the ligands and their cobalt complexes are shown in Fig. 1, where the features closely match those of the previously investigated $[\text{Co}_4]$ complexes.¹⁵ For the complexes **1** and **4**, two transitions appear at ~ 280 nm and ~ 320 nm that can also be observed in the free ligand spectra (see Fig. 1(a)). These could be expected to be ligand-

based $\pi \rightarrow \pi^*$ transition in nature. A similar transition was also observed in complexes **2** and **3**. The transitions can be observed at ~ 390 nm for **1** and 415 nm for **4**, and are red-shifted relative to the free ligand, and could possibly be assigned as a deprotonated ligand-based transition. The most striking point that can be observed in $[\text{Co}^{\text{III}}(\text{L1})_2]^+$ (**1**) is a strong absorption that spans from 450 to 650 nm, reminiscent of a similar feature in the $[\text{Co}^{\text{III}}_4]^{4+}$ spectrum.¹⁵ This observation in solution at room temperature distinguishes these species into three categories: **1**, **2** and **3**, and **4** in view of the ground state electronic absorption spectra, which is consistent with the later analysis of the valence state.

X-ray diffraction data were collected for single crystals of the cobalt complexes at 298 K. In addition, the data at 150 K were collected for complex **3_{ClO4}** considering the potential spin-state transition-induced structure changes. The structure refinement parameters are collected in Table 1. The cobalt(III) complex **1** crystallized in the monoclinic space group $P2_1/c$. Here, the central cobalt atom is coordinated by two deprotonated **L1** ligands binding *via* chelating in a *mer* fashion the pyridyl nitrogens (N5 and N10), imide N8 and N3, and pyrazine nitrogens (N1 and N6) (Fig. 2(A)). Furthermore, the bond lengths of C7–N4 (1.312 Å) and N4–N3 (1.332 Å), as well as another pair of bond lengths (1.330 Å for C18–N9 and 1.310 Å for N9–N8) on the ligand, are intermediate between a normal single bond and a double one, suggesting that the protons on the N4 and N9 sites are lost upon metal binding. The lengths of the Co–N bonds are quite characteristic of the respective spin and oxidation states of the coordinated metal ion and, therefore, can be used as diagnostic tools for determining the electronic state of the metal ion.³⁸ The Co–N bond distances range from 1.817(2) to 1.877(2) Å, typical of LS Co^{III} ,³⁹ and the angular distortion parameter (Σ , defined as the sum of deviation from 90° of 12 *cis*-N–Co–N angles about the Co atom) is 76.68°. The resultant structural models of Co(II) complexes display geometries that are consistent with a singlet ground state for complex **1** at room temperature, while a quartet at the same temperature for complexes **3** and **4**, highlighting the chemical modification of the electronic configuration of the metal-center orbitals.

The cobalt(II) complexes **3_{ClO4}** and **3_{BF4}** crystallized in the monoclinic space group $P2_1/c$ and had similar structures except for the anions (Fig. 2(B) and Fig. S9, ESI†). Here, the binding mode of ligand **L3^{NO2}** to Co^{II} is identical between complexes **3_{ClO4}** and **3_{BF4}** and so only the structure of **3_{ClO4}** will be described herein. Single-crystal X-ray structures were determined at 296 K (**3_{ClO4}^{RT}**) and 150 K (**3_{ClO4}^{LT}**), respectively (Table 1). As in the case of **1**, the coordination sphere of the Co^{II} ion adopts a distorted octahedron with a CoN_6 chromophore. Four equatorial N atoms, including N2, N3, N5, N9 with the average Co–N distance of 2.034(6) Å and two axial N atoms of N7 and N11 with Co–N distance of 2.114(4) and 2.122(3) Å that come from the pyrazine and pyridine group of the individual ligand consist of an elongated octahedral environment, which is characteristic of $\text{Co}^{\text{II}}\text{-HS}$ systems. In the crystal packing mode, some significant intermolecular weak interactions were found, such as $\pi \cdots \pi$ stacking between two neighboring pyrazine rings with the shortest atom–atom distance of 3.317 Å (Fig. 2(D)). In

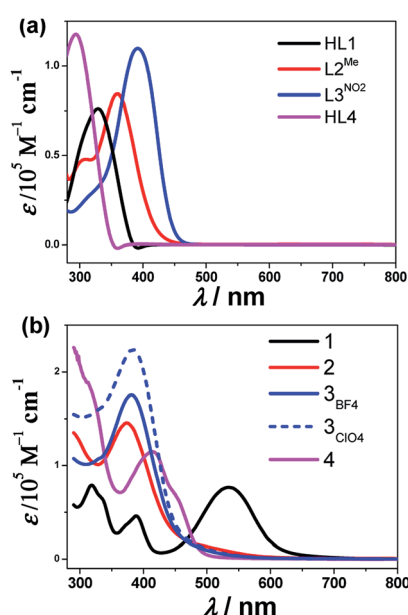


Fig. 1 UV-vis absorption spectra for all the ligands (a) and the corresponding cobalt complexes (b) in acetonitrile at room temperature.

Table 1 Data collection and structure refinement parameters for complexes 1–4

	Complex 1	Complex 3 _{BF4}	Complex 3 _{ClO4}	Complex 4	
	150 K	296 K	296 K	150 K	296 K
Formula	C ₂₂ H ₂₀ ClCoN ₁₀ O ₄	C _{24.5} H ₂₆ B ₂ CoF ₈ N ₁₂ O _{4.5}	C ₂₅ H ₂₄ Cl ₂ CoN ₁₂ O ₁₃	C ₂₅ H ₂₈ Cl ₂ CoN ₁₂ O ₁₃	C ₂₆ H ₂₂ CoN ₈ O ₂
F.W.	582.86	793.12	830.39	834.42	537.45
Crystal system	Monoclinic	Monoclinic	Monoclinic	Monoclinic	Orthorhombic
Space group	P ₂ 1/n	P ₂ 1/n	P ₂ 1/n	P ₂ 1/n	Aba2
<i>a</i> [Å]	8.6063(5)	14.713(8)	14.826(4)	14.286(8)	12.408(2)
<i>b</i> [Å]	9.7092(8)	13.235(7)	13.366(4)	13.218(7)	19.368(3)
<i>c</i> [Å]	28.092(2)	18.309(9)	18.433(5)	18.043(10)	9.8080(16)
α [°]	90	90	90	90	90
β [°]	98.083(2)	102.214(12)	102.714(7)	102.175(11)	90
γ [°]	90	90	90	90	90
Volume [Å ³]	2324.1(3)	3485 (3)	3563.1(18)	3331(3)	2357.0(7)
<i>Z</i>	4	4	4	4	4
Cal. Density (g cm ⁻³)	1.666	1.512	1.548	1.664	1.515
μ/mm^{-1}	0.909	0.587	0.710	0.760	0.771
<i>F</i> (000)	1192	1608	1636	1724	1108
θ range [°]	2.93–25.36	0.25–30.87	0.23–30.32	1.04–30.54	0.33–30.12
Goodness of fit on <i>F</i> ²	1.132	1.035	1.055	1.032	1.028
Final <i>R</i> indices	<i>R</i> ₁ ^a = 0.0466	<i>R</i> ₁ = 0.0903	<i>R</i> ₁ = 0.1001	<i>R</i> ₁ = 0.1061	<i>R</i> ₁ = 0.0427
[<i>I</i> > 2σ(<i>I</i>)]	w <i>R</i> ₂ = 0.0828	w <i>R</i> ₂ = 0.2357	w <i>R</i> ₂ = 0.2484	0.2512	w <i>R</i> ₂ = 0.1522
<i>R</i> Indices (all data)	<i>R</i> ₁ = 0.0801	<i>R</i> ₁ = 0.1508	<i>R</i> ₁ = 0.2189	0.2142	<i>R</i> ₁ = 0.0435
	w <i>R</i> ₂ ^b = 0.0905	w <i>R</i> ₂ = 0.2621	w <i>R</i> ₂ = 0.3089	0.2945	w <i>R</i> ₂ = 0.1530

^a $R_1 = \sum(|F_o| - |F_c|)/\sum|F_o|$. ^b $wR_2 = \{\sum[w(F_o^2 - F_c^2)^2]/\sum[w(F_o^2)^2]\}^{1/2}$

addition, the intermolecular contacts through the nitro-group also support the framework, such as O···O contact of 2.952 Å and O···N contact of 3.001 Å.

In view of the spin-crossover behavior, we continued to determine the structure of 3_{ClO4} at low temperature (150 K). The coordination environment of the metal center was analogous to that observed in 3_{ClO4}^{RT}. However, the average Co···N bond lengths (Table 2) within the equatorial plane now shrink to 1.942(5) Å, and here the striking point is that the axial coordination bonds of 2.086(8) and 2.119(7) Å were not significantly changed, giving rise to more elongated octahedron as expected for a Jahn-Teller distorted low-spin octahedral d⁷ system. The observed distortion for the low-spin cobalt(II) complex was reminiscent of other low-spin mononuclear cobalt(II) complexes of 2,2'-6',2''-terpyridine (terpy) and other N-donor complexes.^{20–24} The distortion situation is also consistent with our recently reported SCO Co₄^{II} metallogrid [Co₄^{II}(L^{Me})₄]⁻ClO₄₈·4MeCN·H₂O,¹⁵ but contrasts with that recently observed in the LS Co₄^{II} grid [Co₄^{II}(HL³)₄][ClO₄]₄·8H₂O, where a compressed octahedron was observed.⁴⁰ Alternatively, the distortion parameter, Σ , defined as the sum of the deviation from 12 *cis*-N-Co-N angles, is sensitive to the changes of the spin state of the center cobalt atom.⁴¹ In general, HS Co(II) compounds have a larger value of Σ than that in LS ones because HS metal compounds allow for more distorted coordination spheres. In the reported Co-terpy SCO compound,^{20–24} the values of Σ ranged from 92° corresponding to the LS state to 119° for the HS cobalt(II) center, respectively. For compound 3_{ClO4}, the angular distortion parameters, Σ (deg), about the cobalt atom shrank from 113.2(1)° at room temperature corresponding to HS Co(II) to the value of 102.8(4)° at 150 K in a LS

structure, indicating the heavier deviations from the ideal octahedral for the LS cobalt(II) center in this system. The low-spin Σ value was unusually large for this complex and, to the best of our knowledge, is at the high end among all cobalt(II) spin-crossover active complexes.^{20–24}

Cobalt(II) complex 4 crystallized in the orthorhombic system Aba2 with structural data collected at room temperature. The central Co^{II}, which lies on a center of inversion, is surrounded by two deprotonated L4 ligands binding *via* the chelating N₂O pocket (Fig. 2(D)). Compound 4 featured a distorted octahedral sphere due to the chelating character of the L4 ligand with *cis*-N-Co-N and *cis*-N-Co-O angles ranging between 75.70(4)–110.15(2)° and 75.12(4)–100.10(2)°. The Co-N (pyrazine) bond distance was 2.160(2) Å, while the Co-N (imine) and Co-O (carboxylate) bond distances were 2.050(2) Å and 2.091(2) Å. The bond valence sum (BVS) value of 2.66 for the cobalt center in 4 supported the 2+ valence assignment on the basis of bond lengths.^{42,43} There were no solvent molecules in the outer coordination sphere or the anions in the lattice, indicating that deprotonation occurs on the N4 atom upon metal coordination.

On the basis of X-ray diffraction data and electronic absorption spectra, magnetic susceptibility data of all the complexes were further collected in the studied temperature ranges in the solid state (Fig. 3(a)). As expected, complex 1 was diamagnetic over the measured temperature region, corresponding to the LS Co^{III} ions. The room temperature $\chi_M T$ value of compound 4 was equal to 2.34 cm³ K mol⁻¹, which was obviously higher than the expected values of 1.875 cm³ K mol⁻¹ corresponding to HS d⁶ (⁵T_{2g}) ground state, reflecting the substantial orbital contribution to the magnetic moment. As



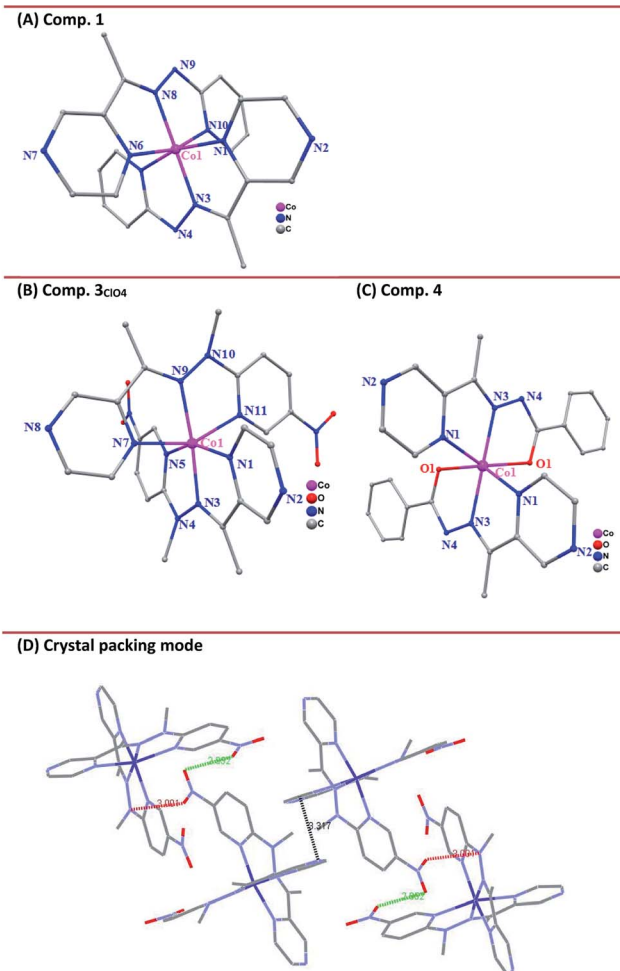


Fig. 2 Crystal structure of the Co-containing complexes in 1 (A), 3_{ClO_4} (B), 4 (C) and crystal packing mode in 3_{ClO_4} (D) with the selected atom labels with ball-and-stick representation. Co = pink ball, C = gray, N = blue, O = red. Hydrogen atoms have been removed for clarity.

the temperature decreased, $\chi_M T$ did not significantly decrease until 40 K, then decreased more rapidly to reach a value of $1.8 \text{ cm}^3 \text{ K mol}^{-1}$ at 5 K, due to zero field splitting effects on the resultant $^5\text{A}_{2g}$ state. However, the magnetic moment of

Table 2 Selected interatomic distances (Å) and distortion parameters Σ (deg) for the selected compounds at the different temperature

Comp.	1	3_{ClO_4}	3_{BF_4}	4
Temp.	150 K	150 K	298 K	298 K
Co–N _{imine}	1.866(3)	1.884(9)	2.020(7)	2.007(6)
	1.871(3)	1.962(9)	2.053(6)	2.038(6)
Co–N _{pyrazine}	1.901(3)	1.934(1)	2.028(7)	2.012(7)
	1.902(3)	2.086(8)	2.114(6)	2.127(5)
Co–N _{pyridine}	1.932(3)	1.988(1)	2.037(7)	2.034(6)
	1.934(3)	2.119(7)	2.122(5)	2.143(5)
Co–N/O _{av} ^b	1.901(3)	1.996(1)	2.061(3)	2.067(5)
Σ (deg) ^c	66.4(5)	102.8(4)	113.2(1)	111.6(5)
				131.4(8)

^a The value corresponds to the bond distance of Co–O. ^b av means the average bond distance. ^c Σ (deg) is defined as the sum of deviation from 90° of twelve *cis*-N–Fe–N angles about the cobalt atom.

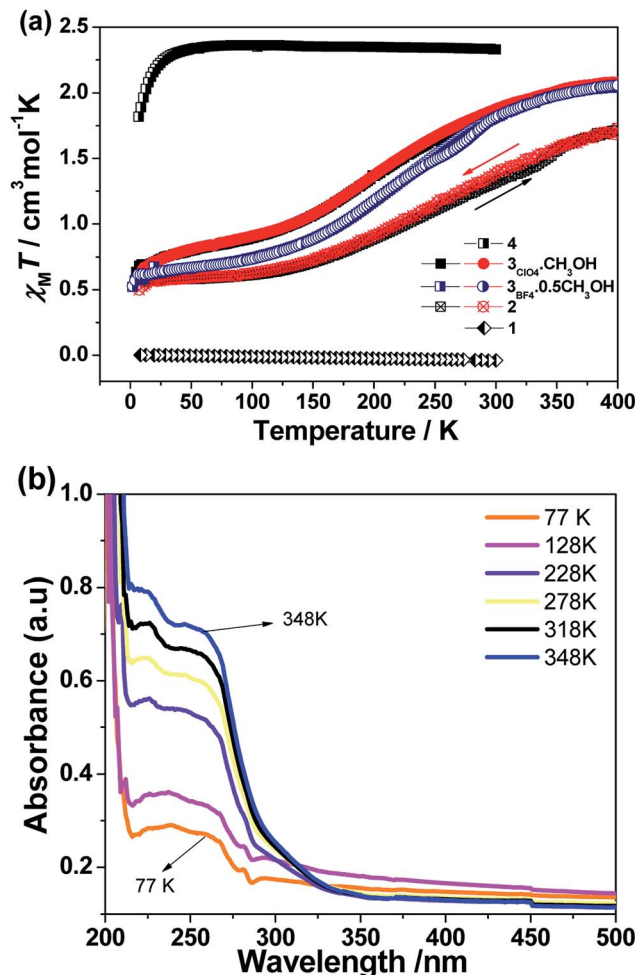


Fig. 3 (a) Temperature dependence of the product $\chi_M T$ for solids 1–4 under an applied magnetic field of 2500 Oe. (b) Solid UV-visible absorption spectra of 3_{ClO_4} powder at temperatures between 77 K and 348 K.

complexes 2 and 3 with the different anions showed different results. The sample was first measured from room temperature to 5 K, followed by the sequential heating/cooling cycles. The temperature-dependent magnetic data showed the gradual spin-crossover phenomenon for all three compounds. Compared to compound 2, compound 3 evidenced the more complete transition from the spin state ($s = 1/2$) at low temperature to high spin ($s = 3/2$) at high temperature. As a representative, complex 3_{ClO_4} showed an initial $\chi_M T$ value of $2.08 \text{ cm}^3 \text{ mol}^{-1} \text{ K}$ at 400 K, corresponding to the main population of the HS Co^{II} ground state, then a decrease in $\chi_M T$ occurred as the sample was cooled to 150 K where a $\chi_M T$ value of $0.88 \text{ cm}^3 \text{ mol}^{-1} \text{ K}$ was observed. Further cooling beyond this point gave rise to a sloping plateau in $\chi_M T$, reaching $0.58 \text{ cm}^3 \text{ mol}^{-1} \text{ K}$ at 5 K. The plot is typical of a gradual $S = 1/2 \leftrightarrow S = 3/2$ spin crossover. No obvious hysteresis occurred upon the sequential heating–cooling cycles in the temperature range of 5–400 K. The investigation of magnetic data for compound 3_{BF_4} gave a similar result. However, at low temperature, the compound showed a more complete spin transition. For

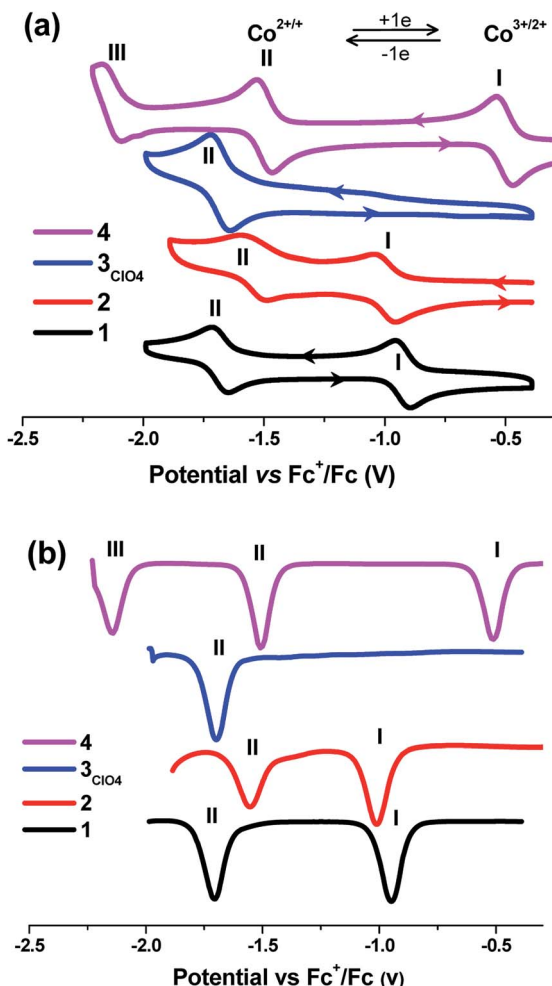


Fig. 4 Cyclic voltammogram (a) and DPV (b) of complexes 1–4 (0.1 mM) in MeCN/0.05 mM NBu_4PF_6 referenced against Fc^+/Fc at a scan rate of 100 mV s^{-1} .

comparison, the magnetic data for the grid compound were aligned with the data of 3_{BF_4} (Fig. S10, ESI[†]). In the reported tetranuclear cobalt grid complexes,¹⁵ the typical $\chi_{\text{M}}\text{T}$ value at low temperature is *ca.* $2.2 \text{ cm}^3 \text{ mol}^{-1} \text{ K}$, corresponding to four low-spin states ($s = 1/2$). With increasing temperature, but the $\chi_{\text{M}}\text{T}$ value could not reach the saturation value even at the highest temperature of 400 K, indicating the rather incomplete spin transition in the Co_4 grid complex. However, for 3_{BF_4} , the high temperature $\chi_{\text{M}}\text{T}$ value above 350 K almost plateaued at *ca.* $2.1 \text{ cm}^3 \text{ mol}^{-1} \text{ K}$, indicating the complete spin transition for the mononuclear complex.

Given the potential of temperature-dependent metal–ligand interactions occurring on the SCO complex 3_{ClO_4} , there should be a partial population of the ligand-center ground state and concomitant $\pi \rightarrow \pi^*$ transition within a wide temperature range.⁴⁴ To further check this point, we collected UV-vis absorption spectra in the solid state as a function of temperature over the range 77–348 K. As expected, the ultraviolet absorption feature increased in intensity as the sample was heated. On subsequent cooling to room temperature, the

spectra could reversibly recover to the original one (Fig. 3(b)). This temperature-dependence behavior reflected an equilibrium between the high- and low-spin populations, where the low-spin state is favored enthalpically and the high-spin state dominate at increasing temperature due to the contributions of entropy to the total Gibbs free energy.

This electrochemical switching provides an alternative stimuli for spin crossover from *ca.* $4 \mu_{\text{B}}$ for $\text{Co}^{\text{II}}\text{-HS}$ by oxidation to $0 \mu_{\text{B}}$ for $\text{Co}^{\text{III}}\text{-LS}$.^{24,45–47} Cyclic voltammetric studies of a 0.1 mM solution of all the cobalt complexes under transient conditions were performed by using cyclic voltammetry (CV) and differential pulse voltammetry (DPV) at $25 \pm 2 \text{ }^{\circ}\text{C}$ in acetonitrile containing 0.05 M of $[\text{Bu}_4\text{N}][\text{PF}_6]$ as the supporting electrolyte, and showed one to three well-separated processes labeled as I, II, and III, respectively, in Fig. 4 (Table 3). The formal reversible potentials ($E(\text{II}) = (E_{\text{p}}^{\text{ox}} + E_{\text{p}}^{\text{red}})/2$) were -946 mV and -1010 mV *versus* $[\text{Fc}]^{0/+}$ for complexes 1 and 2, which could be attributed to the one-electron reduction of $\text{Co}(\text{III})$ to $\text{Co}(\text{II})$ for process I. The peak current is proportional to the square root of the scan rates in the range of 0.05 V s^{-1} to 0.5 V s^{-1} , indicating this process is diffusion controlled at the peak potential (Fig. S11–14, ESI[†]). A second potentially chemically reversible reduction process (II) with an approximate $E(\text{I}) = -1706$ and -1554 mV for 1 and 2, respectively, was observed when the potential was scanned to more negative values, which was associated with the reduction of the $\text{Co}(\text{II})$ to $\text{Co}(\text{I})$. The $I_{\text{p}}^{\text{ox}}/I_{\text{p}}^{\text{red}}$ ratio at all scan rates studied was almost unity and ΔE_{p} ($\Delta E_{\text{p}} = E_{\text{p}}^{\text{ox}} - E_{\text{p}}^{\text{red}}$, where the superscripts ox and red denote the oxidation and reduction peaks respectively) was close to the theoretical value of 57 mV at $25 \text{ }^{\circ}\text{C}$ for a simple one-electron process, indicating all the processes were one-electron chemically and electrochemically reversible. However, complex 3 evidenced only one redox process, II, with the formal potential of -1698 mV . Interestingly, an additional process, III, was observed for complex 4 with the formal potentials ($E(n)$, $n = \text{III}$, II, I) *versus* $[\text{Fc}]^{0/+}$ of -2140 , -1510 , and -510 mV . The current magnitude for processes I, II, and III were similar. Process III, ($E_{1/2}(\text{III}) = -2140 \text{ mV}$), exhibited a negative or reduction current, and hence was ascribed to a ligand-centered redox process on the basis of previous assignments of pyrazine-substituted cobalt complexes.^{48,49} From the electrochemical data, there existed a slight difference in the electronic behavior between complex 1 and 2 and the highest oxidation state corresponding to Co^{III} ion required a quite negative oxidation potential. In this situation, complex 2 was still isolated as Co^{II} species and did not induce any partial oxidation, as evidenced by magnetic characterization and X-ray diffraction analysis. However, the oxidation of the $\text{Co}(\text{II})$ complex required the most negative potential in complex 2 compared to those in the cases of 1 and 4, suggesting that the ligand in 2 provided the most weak ligand field in the order of $\text{L2}^{\text{Me}} < \text{L1} < \text{L4}$. This is normal because deprotonation should render the delocalized aromatic system of the ligand as a weaker π acceptor, thus weakening the ligand strength in 1 and 4. This result is consistent with the observation of cobalt metallogrid complexes in the previous report.¹⁵

Table 3 The potentials ($E_{1/2}$, V vs. Fc^+/Fc) of all the cobalt complexes in this work and the selected cobalt complexes

Solvent	Comp.	$E_{1/2}$ (V) ^a	Co ^{III/II}	Co ^{II/I}	Ligand-base	
CH ₃ CN	[Co ^{III} (L1) ₂]ClO ₄ (1)	-0.93	-1.66	-	- ^c	This work ^b
	Co ^{II} (L2 ^{Me}) ₂ (2)	-0.98	-1.53	-	-	
	[Co ^{II} (L3 ^{NO₂}) ₂]·CH ₃ OH (3 _{ClO₄})	-	-1.67	-	-	
	Co ^{II} (L ₄) ₂ (4)	-0.49	-1.52	-	-2.14	
CH ₃ CN	C1-C8	-0.21 to +0.84	-0.82 to -1.92	-	-1.61 to -2.49	Ref. 10
	[Co(BTSC) (L) ₂] ⁺	-0.69 to -1.10	-1.00 to -1.19	-	-	Ref. 12
DMF	Co(salen-OMe)	-	-1.71	-	-	Ref. 14
	2M ^d	-	-1.58 to -1.41	-	-	
CH ₃ CN	[Co ^{II} (dpzca) ₂] ^e	-0.22	-	-	-1.46, -1.84	Ref. 24
	Co ^{III} Co ₃ ^{II} /Co ₄ ^{II}	Co ₂ ^{III} Co ₂ ^{II} /Co ^{III} Co ₃ ^{II}	Co ₃ ^{III} Co ^{II} /Co ₂ ^{III} Co ₂ ^{II}	Co ^{III} ₄ /Co ^{III} ₃ Co ^{II}	Co ^{III} ₄ /Co ^{III} ₃ Co ^{II}	Ref. 15
CH ₃ CN	[Co ^{III} ₄ (L ^H) ₄][ClO ₄) ₄	0.63	0.52	-	-0.77	-0.83
	[Co ^{II} ₄ (L ^{Me}) ₄][ClO ₄)	0.30	0.02	-	-0.60	-0.74
	[Co ^{II} ₄ (L ^{B+}) ₄][ClO ₄) ₈	-0.40	-0.49	-	-0.62	-0.87

^a Scan rate = 100 mV s⁻¹. ^b Concentration in all cases = 0.1 mM. ^c Not observed. ^d M = Na⁺, K⁺, Ca²⁺, Sr²⁺, and Ba²⁺. ^e The potential values were referenced against Ag/AgNO₃.

Conclusion

In conclusion, we prepared a series of new tridentate Schiff-base ligands bearing a variety of substituting moieties through a condensation reaction between 2-acetylpyrazine and different hydrazone derivatives. The corresponding cobalt complexes with bis-substitute were isolated and structurally characterized. The analysis of the crystal structures provided insights into understanding the differences in the Co–N bond lengths observed in the crystal structures, assigning them to differences in terms of the spin state and valence state. In particular, the Co–N bond length can be diagnostic for differentiating the low-spin doublet from the high-spin quartet state. In addition, we showed that the redox potential of the Co(III)/Co(II), Co(II)/Co(I), and pyrazine-base couples could be easily modulated over a wide range by tuning the substituents on the ligands, allowing a broad range of potential applications. Study of the catalytic proton reduction by applying this class of complexes as a catalyst is currently ongoing in our laboratory.

Conflicts of interest

There are no conflicts to declare.

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