Quantum Chemical Study of Spin Transitions in the Bimetallic Fe/Co Complexes with the Bis(catecholate) Bridging Ligand

M. G. Chegerev*, A. G. Starikov*, and A. A. Starikova*

Institute of Physical and Organic Chemistry, Southern Federal University, Rostov-on-Don, Russia
*e-mail: mchegerev@sfedu.ru

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Abstract—The computational modeling of the spatial and electronic structures, energy characteristics, and magnetic properties of the bimetallic iron and cobalt complexes with 9,10-dimethyl-9,10-ethano-9,10-dihydro-2,3,6,7-tetrahydroxyanthracene and terminal tris(2-pyridylmethyl)amine bases is performed using the density functional theory method (DFT UTPSSh/6-311++G(d,p)). The chosen tetradentate redox ligand is shown to be a promising precursor for the production of magnetically active compounds. The calculations make it possible to establish a relationship between the relative energies of the electronic isomers of the complexes and the structures of the ancillary N-donor moieties. The coordination compounds prone to the manifestation of spin transitions accompanied by a change in the magnetic properties are revealed.

Keywords: iron, cobalt, tris(2-pyridylmethyl)amine, redox-active ligand, quantum chemical calculations

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INTRODUCTION

Coordination compounds of metals with redox-active ligands attract increasing interest of chemists and materials scientists due to their potential use as a basis for devices of molecular electronics and spintronics [1–10]. If the structural components are chosen rationally, these molecules can exhibit reversible mutual transformations between electronic isomers (electromers [11]) with various magnetic properties. The polystable compounds that can be switched between three and more electromeric forms [12–19] are of special interest, since they can be used in storage systems of high-density data or for performing complicated logical operations.

Spin crossover (SCO) [20, 21] caused by the rearrangement of electrons inside the valence shell of the coordinated metal ion and redox isomerism (or valence tautomerism (VT) [22–24], i.e., intramolecular electron transfer between the metal center and redox-active ligand) are among the widely studied mechanisms of switching spin states of complexes. The SCO is most frequently met in the iron complexes in the N6 or N4O2 coordination environment [25, 26] and is accompanied by transitions of the metal ion between the low-spin (LS) and high-spin (HS) states. In turn, the transformation of the LS electron containing LS\text{Co}^{III} and the dianionic catecholate (Cat) form of the redox ligand into the HS isomer HS\text{Co}^{II}SO (SO is the radical-anion semiquinone form of \text{o}-benzoquinone) occurs in the VT systems, the classical examples of which are the cobalt \text{o}-benzoquinone complexes [8, 22, 27, 28]. The binuclear metal complexes demonstrate partial, one- or two-step SCO/VT rearrangements [14].

An abundant type of coordination compounds with redox ligands is presented by the salt-like metal complexes containing the \text{o}-benzoquinone derivative and ancillary tetradentate nitrogen-containing bases [29–37]. According to the theoretical [29, 38–42] and experimental [30–37] data, thus constructed mono- and binuclear systems are capable of exhibiting spin transitions due to the VT and/or SCO. It is found that the possibility of rearrangements leading to a change in the magnetic properties and their character are determined by the nature of the metal center (Fe/Co), structure of the redox-active ligand, and type of the nitrogen-containing base.

9,10-Dimethyl-9,10-ethano-9,10-dihydro-2,3,6,7-tetrahydroxyanthracene (L) is the synthetically available but poorly studied bis(catecholate) ligand. The octauranate cage complexes [43] and manganese(III) [44] and platinum(II) [45] compounds based on ligand L are known. The possibility of the two-step VT accompanied by switching spin states has been shown previously [46, 47] by the theoretical study of the binuclear adducts of cobalt diketones with di-\text{o}-quinone L. In this work, we aimed at enlarging the range of magnetically active compounds and performed the computational modeling of the bimetallic (Fe–Fe and Fe–Co) complexes with tetrahydroxyanthracene L in which tris(2-pyridylmethyl)amine (Tpa) bases (Me_nTpa, n = 0, 2) played the role of ancillary terminal
moieties. Hexafluorophosphate anions were chosen because of their wide use for the synthesis of similar salt-like o-benzoquinone complexes with tetradentate N-donor ligands. The electronic structures were studied and the geometric, energy, and magnetic characteristics of compounds I–IV (M = Fe, Co; R = H, CH₃) presented in Scheme 1 were calculated, and the possibility of spin transitions to occur in them was studied. The metal-containing dicatonic moieties of target complexes I'–IV' (M = Fe, Co; R = H, CH₃) of the general formula [FeM(Me₉Tpa)₂(L)]²⁺ (M = Fe, Co; n = 0, 2) were also considered to reveal the role of the outer-sphere counterions.

Scheme 1.

CALCULATION PROCEDURE

The calculations were performed using the Gaussian 16 program [48] by the density functional theory (DFT) method using the UTPSSh functional [49, 50] and 6-311++G(d,p) extended basis set, a combination of which correctly reproduced the energy and magnetic characteristics of the complexes exhibiting the magnetic bistability mechanisms [14, 30, 38, 51–56]. According to the calculation results obtained in this approximation for the experimentally studied cobalt and iron compounds, the spin transitions occur at ΔE_{HS–LS} < 10 kcal/mol. The position and orientation of the hexafluorophosphate counterions in complexes I–IV were taken from the X-ray structure analysis (XSA) data of the structurally similar compounds [14]. The stationary points were localized on the potential energy surface (PES) by the full optimization of the geometry of the molecular structures checking DFT stability of the wave function and calculating the force constants. The exchange interaction parameters (J, cm⁻¹) were calculated in the framework of the broken symmetry (BS) formalism [57] using the generalized spin projection method proposed by Yamaguchi [58].

The graphical images of the molecular structures were drawn using the ChemCraft program [59].

RESULTS AND DISCUSSION

According to the calculation results, the ground state of dicatonic moiety I' (M = Fe, R = H) corresponds to a minimum on the triplet PES characterized by the localization of the spin density (qₘFe, qₘM = 0.45) on the iron(III) ions (Table 1). The inclusion of the hexafluorophosphate counterions (compound I (M = Fe, R = H)) in the calculation scheme results in an increase in qₘFe and qₘM to 0.56 indicating that one lone electron is located on each metal center, which allows one to ascribe the electronic structure lsFe₃³⁺–Cat–Cat–lsFe₃³⁺ to the discussed isomer. It should be mentioned that a significant portion of the spin density is delocalized on the oxygen atoms adjacent to the complexing agent (0.14 on each atom), which indicates (along with the determined bond lengths in the o-benzoquinone ring (Fig. 1) being intermediate between the distances characteristic of SQ and Cat [60]) the existence of the lsFe₃³⁺–SQ–SQ–lsFe₃³⁺ structure with...
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a close energy. The data of the experimental studies (XSA, Mössbauer spectroscopy) for the mononuclear iron(III) complexes of similar structure confirm that the electronic configuration $LSFe^{II}$–SQ is admixed to the more preferable state $LSFe^{III}$–Cat [61, 62], which is consistent with the obtained calculation results for complex I ($M = Fe, R = H$) and its dicationic moiety I’. The study of the exchange interactions in the $LSFe^{III}$–Cat–Cat–$LSFe^{III}$ electromer shows a strong antiferromagnetic coupling of lone electron spins ($J = –187 \text{ cm}^{-1}$).

The next in energy isomers of complex I ($M = Fe, R = H$) and its dicationic moiety I’ are presented by the minima of $LSFe^{III}$–Cat–Cat–$HSFe^{III}$ on the septet PES (Table 1, Fig. 1). The absence of spin density on the carbon atoms and an insignificant alternation of the bonds in redox ligand L indicates its Cat form. The calculated spin density on the metal in the high-spin state is somewhat lower than the value expected for five electrons ($q^M_s, q^M_s \approx 4$). A similar situation is often observed in the trivalent iron compounds with $\sigma$-benzoquinone ligands [62–65]. A moderate antiferromagnetic exchange ($J = –36 \text{ cm}^{-1}$) is expected for the $LSFe^{III}$–Cat–Cat–$HSFe^{III}$ isomer. The predicted difference in energies between the $LSFe^{III}$–Cat–Cat–$LSFe^{III}$ and $LSFe^{III}$–Cat–Cat–$HSFe^{III}$ electromers both ignoring (10.8 kcal/mol) and taking into account two anions (11.2 kcal/mol) allows one to expect that the thermally initiated SCO would occur on one of the metal centers.

The HS isomer $HSFe^{III}$–Cat–Cat–$HSFe^{III}$ is remote from the ground state $LSFe^{III}$–Cat–Cat–$LSFe^{III}$ by more than 20 kcal/mol, indicating a low probability of its population under the thermal conditions. A comparison of the data obtained by the calculation of dicationic moiety I’ and electroneutral com-

### Table 1. Spin ($S$), total energy ($E$), relative energy ($\Delta E$), spin-squared operator ($S^2$), and spin density on the metal centers ($q^M_s, q^M_s$) in the electromers of complexes I and II and their dicationic moieties I’ and II’ calculated by the DFT UTPSh/6-311++G(d,p) method

| Electrometer | $S$ | $E$, a.u. | $\Delta E$, kcal/mol | $S^2$ | $q^M_s$ | $q^M_s$ |
|--------------|-----|----------|----------------------|------|--------|--------|
| $LSFe^{III}$–Cat–Cat–$LSFe^{III}$ | 1 | -5354.937570 | 0.0 | 2.038 | 0.45 | 0.45 |
| BS | 0 | -5354.938477 | 0.972 | | | |
| $LSFe^{III}$–Cat–Cat–$HSFe^{III}$ | 3 | -5354.920405 | 10.8 | 12.048 | 0.43 | 4.05 |
| BS | 0 | -5354.921238 | 6.993 | | | |
| $HSFe^{III}$–Cat–Cat–$HSFe^{III}$ | 5 | -5354.903194 | 21.6 | 30.059 | 4.04 | 4.04 |
| BS | 0 | -5354.903804 | 5.015 | | | |
| $LSFe^{III}$–Cat–Cat–$LSFe^{III}$ | 1 | -7236.924456 | 0.0 | 2.052 | 0.56 | 0.56 |
| BS | 0 | -7236.925776 | 0.992 | | | |
| $LSFe^{III}$–Cat–Cat–$HSFe^{III}$ | 3 | -7236.906528 | 11.2 | 12.050 | 0.55 | 4.08 |
| BS | 0 | -7236.888550 | 6.993 | | | |
| $HSFe^{III}$–Cat–Cat–$HSFe^{III}$ | 5 | -7236.888550 | 22.5 | 30.048 | 4.08 | 4.08 |
| BS | 0 | -7236.888550 | 5.015 | | | |
| $LSFe^{III}$–Cat–Cat–$LSFe^{III}$ | 1 | -5512.244973 | 0.0 | 2.052 | 0.53 | 0.53 |
| BS | 0 | -5512.245776 | 0.992 | | | |
| $LSFe^{III}$–Cat–Cat–$HSFe^{III}$ | 3 | -5512.239644 | 3.3 | 12.053 | 0.52 | 4.05 |
| BS | 0 | -5512.240453 | 6.993 | | | |
| $HSFe^{III}$–Cat–Cat–$HSFe^{III}$ | 5 | -5512.234596 | 6.5 | 30.056 | 4.04 | 4.04 |
| BS | 0 | -5512.235297 | 5.007 | | | |
| $LSFe^{III}$–Cat–Cat–$LSFe^{III}$ | 1 | -7394.229462 | 0.0 | 2.075 | 0.65 | 0.65 |
| BS | 0 | -7394.222740 | 4.2 | 12.064 | 0.66 | 4.08 |
| $HSFe^{III}$–Cat–Cat–$HSFe^{III}$ | 5 | -7394.217166 | 7.7 | 30.048 | 4.09 | 4.09 |
plex I indicates that the accounting for the counterion exerts no substantial effect on the difference in energies between the electromers involved in the SCO rearrangement (\(\text{LSFe}^{III} \rightleftharpoons \text{HSFe}^{III}\)).

It has previously been shown \([14, 29, 34, 38, 42]\) that the introduction of methyl substituents in the sixth position of the pyridine ring of Tpa creates steric hindrances for the formation of the LS isomer characterized by shorter M–N coordination bonds and results in the narrowing of the energy gap between the electromers containing metal ions in different spin states. Electroneutral complexes II (M = Fe, R = CH₃) and their dicationic moieties II' (M = Fe, R = CH₃) in which Me₂Tpa act as ancillary terminal ligands were studied in order to find a compound based on L potentially capable of undergoing two-step SCO rearrangements.

As follows from the obtained calculation results, the functionalization of the tetradentate base does not change the electronic configuration of the determined
isomers: the values of spin density on the metal centers (Table 1) and calculated bond lengths in the redox moiety (Fig. 1) indicate the trivalent state of iron and catecholate form of ligand L in all structures. At the same time, the introduction of alkyl groups is expectedly [14, 29, 34, 38, 42] accompanied by a decrease in the stability of the LSFeIII–Cat–Cat–LSFeIII and LSFeIII–Cat–Cat–HSFeIII electromers containing the LS metal ions. As a consequence, the HS form HSFeIII–Cat–Cat–HSFeIII is remote from the ground state LSFeIII–Cat–Cat–LSFeIII by less than 8 kcal/mol (Table 1), which provides the occurrence of the two-step SCO transformation in compound II (M = Fe, R = CH₃).

According to the calculation of the exchange interactions, a strong antiferromagnetic coupling of the metal centers (J = –166 cm⁻¹) that can result in diamagnetism is expected in the LS LSFeIII–Cat–Cat–LSFeIII electronomer, the isomer in the intermediate spin state LSFeIII–Cat–Cat–HSFeIII exhibits a moderate antiferromagnetic exchange (J = –35 cm⁻¹), and lone electron spins in the HS structure HSFeIII–Cat–Cat–HSFeIII do not almost interact (J = –6 cm⁻¹). Thus, the total spin of the system (S) would change from S = 0 to S = 5 during the predicted spin transitions in complex II (M = Fe, R = CH₃), which allows one to consider this compound as a basis for molecular switches.

The quantum chemical study of heterometallic compounds III and IV and their dicaticonic moieties III' and IV' shows that the LS (S = 1/2) electronomers contain the catecholate form of L and three-charge metal ions (Table 2), and the single electron in the LSFeIII–Cat–Cat–LSCoIII isomers is concentrated on the iron center (q²Fe = 0.5) when a significant portion of the spin density is delocalized over the adjacent donor atoms. Two types of structures were found on the sextet PES: the LSFeIII–Cat–Cat–HSCoII isomer containing three paramagnetic centers (LSFeIII, SQ, HSCoII) and the HSFeIII–Cat–Cat–LSCoIII electronomer in which five lone electrons are localized on the iron(III) ion in the HS state. The structure with the highest spin (S = 9/2) is presented by the electronic configuration HSFeIII–Cat–Cat–HSCoII.

The calculated difference in energies between the most stable isomers of complex III and its dicaticonic moiety III' does not exceed 10 kcal/mol and allows one to expect the VT rearrangement in the cobalt moiety of the molecule: LSFeIII–Cat–Cat–LSCoIII ⇄ LSFeIII–Cat–Cat–HSCoII (Fig. 2). The discussed values of ΔE ignoring (5.3 kcal/mol) and taking into account the hexafluorophosphate anions (9.9 kcal/mol) differ nearly by a factor of 2, which is consistent with the earlier revealed [29, 30, 42, 54] overestimation of the stabilization energy of the HS electronomers by the DFT study of the cationic cobalt complexes with the redox ligands and confirms the importance of inclusion of the outer-sphere counterions in the calculation scheme. This fact is caused, most probably, by the charge redistribution in the redox moiety during the VT transformation Co³⁺–Co²⁺–SQ−, which is reflected as different strengths of the Coulomb interactions of the valence tautomers with the outer-sphere counterion. A comparison of the calculated energy characteristics of compound IV and its structural moiety IV' (Table 2) is more clear: in the case of dicaticonic system IV', the LSFeIII–Cat–SQ–HSCoII isomer turns out to be the most stable, whereas the accounting for PF₆⁻ (complex IV) results in the inversion of the relative energies and stabilization of the LSFeIII–Cat–Cat–LSCoIII electronomer.

The predicted difference in energies between the isomers of complex III (M = Co, R = H) differed in the spin state of the iron(III) ion (Table 2, Fig. 2) indicates the possibility of the SCO process LSFeIII–Cat–Cat–LSCoIII ⇄ HSFeIII–Cat–Cat–LSCoIII, which was not completed under the thermal conditions. A significant destabilization of the HS form HSFeIII–Cat–SQ–HSCoII does not allow one to expect its participation in spin transitions. Thus, heterometallic complex III can demonstrate two transformations leading to switching the magnetic properties: VT on the cobalt center and SCO on the iron(III) ion. Since the antiferromagnetic exchange interactions between LSFeIII and SQ (J = –265 cm⁻¹) in the LSFeIII–Cat–SQ–HSCoII isomer are expected to be strong, the first rearrangement would be accompanied by a change in the spin from 1/2 to 3/2 and the SCO would induce transitions between the doublet (S = 1/2) and sextet (S = 5/2) states.

The closeness of the total energies of four electronomers of complex IV (M = Co, R = CH₃) assumes the possibility of each of them (Table 2, Fig. 2) to be involved in spin transitions. The transformation of the LS form into the HS form can proceed via three equally probable mechanisms: (1) the process LSFeIII–Cat–Cat–LSCoIII ⇄ LSFeIII–Cat–Cat–SQ–HSCoII caused by the electron transfer between the redox ligand and cobalt(II) ion followed by the SCO rearrangement on the Fe(III) center LSFeIII–Cat–Cat–SQ–HSCoII, (2) sequential phenomena of SCO (LSFeIII–Cat–Cat–LSCoIII ⇄ HSFeIII–Cat–Cat–LSCoIII) and VT (HSFeIII–Cat–Cat–LSCoIII ⇄ HSFeIII–Cat–SQ–HSCoIII), and (3) synchronized mechanisms of magnetic bistability SCO + VT (LSFeIII–Cat–Cat–LSCoIII ⇄ HSFeIII–Cat–SQ–HSCoIII). In the LSFeIII–Cat–Cat–SQ–HSCoII and HSFeIII–Cat–SQ–HSCoII isomers, two metal centers do not nearly interact (J < 3 cm⁻¹) and the HS cobalt(II) ion
Table 2. Spin ($S$), total energy ($E$), relative energy ($\Delta E$), spin-squared operator ($S^2$), and spin density on the metal centers ($q_s^F$, $q_s^M$) in the electromers of complexes III and IV and their dicationic moieties III' and IV' calculated by the DFT UTPSSh/6-311++G(d,p) method

| Electomer | $S$    | $E$, a.u. | $\Delta E$, kcal/mol | $S^2$ | $q_s^F$ | $q_s^M$ |
|-----------|--------|-----------|----------------------|-------|---------|---------|
| Dicationic moiety III' (M = Co, R = H) |        |           |                      |       |         |         |
| $LSFe^{III}$–Cat–Cat–$LSCo^{III}$ | 1/2    | -5473.971117 | 0.0                 | 0.765 | 0.37    | 0.00    |
| $LSFe^{III}$–Cat–SQ–$HSCo^{II}$ | 5/2    | -5473.962600 | 5.3                 | 8.785 | 0.48    | 2.80    |
| $\beta\alpha\alpha^*$ | 3/2    | -5473.963854 | 4.702               |       |         |         |
| $\alpha\beta\alpha$ | 3/2    | -5473.961945 | 4.667               |       |         |         |
| $\alpha\alpha\beta$ | 1/2    | -5473.960567 | 2.761               |       |         |         |
| $HSFe^{III}$–Cat–Cat–$LSCo^{III}$ | 5/2    | -5473.953431 | 11.1                | 8.785 | 4.02    | 0.00    |
| $HSFe^{III}$–Cat–SQ–$HSCo^{II}$ | 9/2    | -5473.945618 | 16.0                | 24.792| 4.06    | 2.80    |
| $\beta\alpha\alpha$ | 1/2    | -5473.946583 | 4.701               |       |         |         |
| $\alpha\beta\alpha$ | 3/2    | -5473.944834 | 4.693               |       |         |         |
| $\alpha\alpha\beta$ | 3/2    | -5473.943411 | 6.766               |       |         |         |
| III (M = Co, R = H) |        |           |                      |       |         |         |
| $LSFe^{III}$–Cat–Cat–$LSCo^{III}$ | 1/2    | -7355.961933 | 0.0                 | 0.772 | 0.49    | 0.00    |
| $LSFe^{III}$–Cat–SQ–$HSCo^{II}$ | 5/2    | -7355.946107 | 9.9                 | 8.794 | 0.60    | 2.82    |
| $HSFe^{III}$–Cat–Cat–$LSCo^{III}$ | 5/2    | -7355.943379 | 11.6                | 8.780 | 4.07    | 0.00    |
| $HSFe^{III}$–Cat–SQ–$HSCo^{II}$ | 9/2    | -7355.928050 | 21.3                | 24.788| 4.10    | 2.81    |
| $\beta\alpha\alpha$ | 3/2    | -7355.928050 | 4.701               |       |         |         |
| $\alpha\beta\alpha$ | 7/2    | -5473.944834 | 6.766               |       |         |         |
| $\alpha\alpha\beta$ | 3/2    | -5473.943411 | 6.766               |       |         |         |
| Dicationic moiety IV' (M = Co, R = CH$_3$) | | | | | |
| $LSFe^{III}$–Cat–Cat–$LSCo^{III}$ | 1/2    | -5631.275676 | 0.0                 | 0.770 | 0.46    | 0.00    |
| $LSFe^{III}$–Cat–SQ–$HSCo^{II}$ | 5/2    | -5631.279327 | -2.3                | 8.792 | 0.58    | 2.81    |
| $\beta\alpha\alpha$ | 3/2    | -5631.280491 | 4.714               |       |         |         |
| $\alpha\beta\alpha$ | 3/2    | -5631.278009 | 4.695               |       |         |         |
| $\alpha\alpha\beta$ | 1/2    | -5631.276798 | 2.786               |       |         |         |
| $HSFe^{III}$–Cat–Cat–$LSCo^{III}$ | 5/2    | -5631.270361 | 3.3                 | 8.781 | 4.03    | 0.00    |
| $HSFe^{III}$–Cat–SQ–$HSCo^{II}$ | 9/2    | -5631.274188 | 0.9                 | 24.790| 4.06    | 2.81    |
| $\beta\alpha\alpha$ | 1/2    | -5631.275290 | 4.720               |       |         |         |
| $\alpha\beta\alpha$ | 7/2    | -5631.272824 | 16.702              |       |         |         |
| $\alpha\alpha\beta$ | 3/2    | -5631.271634 | 6.785               |       |         |         |
| IV (M = Co, R = CH$_3$) | | | | | |
| $LSFe^{III}$–Cat–Cat–$LSCo^{III}$ | 1/2    | -7513.263112 | 0.0                 | 0.779 | 0.59    | 0.00    |
| $LSFe^{III}$–Cat–SQ–$HSCo^{II}$ | 5/2    | -7513.260727 | 1.5                 | 8.806 | 0.61    | 2.81    |
| $HSFe^{III}$–Cat–Cat–$LSCo^{III}$ | 5/2    | -7513.258541 | 2.9                 | 8.777 | 4.06    | 0.00    |
| $HSFe^{III}$–Cat–SQ–$HSCo^{II}$ | 9/2    | -7513.255784 | 4.6                 | 24.787| 4.10    | 2.82    |

* $\alpha$ corresponds to the up orientation of spins, and $\beta$ corresponds to the down orientation: 1, Fe$^{III}$; 2, SQ; and 3, Co$^{II}$. is ferromagnetically coupled with the semiquinone radical anion ($J = 183$ cm$^{-1}$), which is consistent with the earlier performed theoretical and experimental studies of the cobalt(II) $\sigma$-benzosemiquinone complexes with the Tpa derivatives [14, 29, 34, 38, 42], whereas the exchange between the Fe(III) center and SQ is antiferromagnetic. Moderate interactions in the pair of $HSFe^{III}$ and SQ ($J = -50$ cm$^{-1}$) would not affect the total spin ($S = 9/2$) of the $HSFe^{III}$–Cat–SQ–$HSCo^{II}$ structure, but the strong antiferromagnetic
Fig. 2. Calculated data (spatial structures, bond lengths, and exchange parameters) for the electromers of complexes III and IV obtained by the DFT UTPSh/6-311++G(d,p) method.
coupling between $\text{LSFe}^{\text{III}}$ and SQ ($J = -240 \text{ cm}^{-1}$) can result in the stabilization of the $\text{LSFe}^{\text{III}}$–Cat–SQ–$\text{HSCo}^{\text{II}}$ electromer in the state $S = 3/2$. Therefore, the expected mutual transformations of the isomers of coordination compound IV ($M = \text{Co}, R = \text{CH}_3$) characterized by different values of $S$ (Table 2, Fig. 2) would be accompanied by switching of the magnetic properties in a wide range, which makes the considered heterometallic mixed-ligand complex to become a promising candidate for the design of molecular switches and spin qubits (quantum bits).

Thus, the magnetically active bimetallic coordination compounds in which 9,10-dimethyl-9,10-ethano-9,10-dihydro-2,3,6,7-tetrahydroxyanthracene acted as the bridging ligand were modeled and studied by quantum chemical calculations. The complexes that can manifest one- and two-step spin transitions and synchronized SCO and VT mechanisms were theoretically designed by the building up of the coordination sphere of the metals (Fe, Co) by tris(2-pyridyl)methylamine bases. The variation of the structural parameters in the ancillary N-donor ligands was found to exert no effect on the character and strength of exchange interactions between the paramagnetic centers but allowed one to control the relative energies of the electromers. The study of the heterospin complexes containing the cobalt ions prone to the participation in the intramolecular electron transfer with the redox ligand ascertained the importance of including the outer-sphere counterions in the calculation scheme. The compounds discussed in this article are promising building blocks for the production of magnetic materials and, hence, are being presently synthesized.

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**CONFLICT OF INTEREST**

The authors declare that they have no conflicts of interest.

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