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Two state "ON–OFF" NLO switch based on coordination complexes of iron and cobalt containing isomeric ligand: a DFT study†

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Coordination complexes are interesting materials for nonlinear optical (NLO) applications due to their large hyperpolarizability values. Moreover, switchable NLO response is also important in coordination complexes. Herein, we report two state ON–OFF switchable NLO contrast of coordination complexes of Fe and Co containing isomeric ligands. The optical, UV-visible, and electronic properties besides the "ON–OFF" switching effect are calculated using the CAM-B3LYP/6-31+G (d) method. The NLO responses of ligand–metal isomers are qualitatively evaluated through variation in charge transference (CT) style through TD-DFT. The higher β_o in each isomeric pair is strongly dependent on the HOMO–LUMO gap. The isomer **4b** with lowest HOMO–LUMO gap shows the highest NLO response. The charge transfer pattern in these complexes results in variation of their β_o values. The notable β_o contrast of 21.15 in isomeric pairs **3a** and **3b** makes these complexes a favorable material for genuine NLO switches. Hence, the outcome of the current investigation reveals that these ligand–metal isomeric complexes exhibit a two-state switch "ON–OFF" effect.

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Introduction

In recent years nonlinear optics (NLO) achieved notable consideration owing to their extensive usage in advanced optoelectronics,¹ optical switching,² signal processing,³ information storage,⁴ and all data processing.⁵ NLO materials generally have properties such as polarization, conjugated structure, phase, amplitude, frequency *etc.* which can be altered for their broad spectrum applications.^{6–9} Amongst NLO materials, switchable NLO molecular materials display a high potential usage in data storage, signal processing and sensing besides optoelectronic technologies. Molecules with switchable second-order nonlinear optical properties have gained a great deal of interest because of their potential use in photonic devices.^{10–12} The progress of innovative nonlinear optical materials is essential for sustainable development in nanoscience and nanotechnology.^{13–15}

The literature study revealed that various strategies including bond length alteration, metal–ligand frameworks, excess electron incorporation, extended conjugation in π -linkage, and accommodation of donor–acceptor fragments

connected *via* a π -network have been frequently adopted to improve the NLO properties of materials.^{16–18}

NLO materials, for instance organic compounds and coordination complexes, have been extensively used in the last few decades.^{15,19–24} Organic compounds are most used as NLO materials due to their ease of synthesis, low cost, and versatility in tailoring their properties. One of the key benefits of using organic materials is that through precise and distinct synthetic modifications of the molecular structures, their physical and chemical properties can be customized.^{14,16,25–28} The assimilation of donor and acceptor fragments connected by employing a π -network proves to be an effective strategy to enhance the NLO properties of organic material. Intramolecular charge transfer (ICT) affects NLO properties that originate primarily from the π -conjugated bridge due to donor to acceptor moieties. The proper acceptor and donor fragments connected using a π -network develop an electron push–pull mechanism that enhances the NLO properties of materials.^{20,29–38} However, the low thermal stability and volatile nature of organic compounds limit their potential use.

Coordination complexes have gained remarkable interest in the field of nonlinear optics due to their more flexible molecular design. The presence of a greater number of diverse charge-transfer transitions of high intensity at quite low energy is tunable by employing the nature, oxidation state, and metal center of the coordination sphere. There exist huge varieties and notably limitless opportunities for metal–ligand combinations that allow tailoring of metal complexes for various potential uses in the switching of NLO response. In these

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complexes, it is possible to switch molecular architecture, spin state, charge distribution pattern, and magnetic moment.³⁹ Mostly **4d** and **5d** metal complexes exhibit low-energy ligand-to-metal charge-transfer (LMCT), metal-to-ligand charge-transfer (MLCT), ligand-to-ligand charge-transfer (LLCT) or intra ligand charge-transfer (ILCT) excitations.⁴⁰ The metal can effectively be the donor or the acceptor moiety or the polarizable bridge of a push-pull system that can act as a dipolar NLO chromophore.⁴¹ The existence of the metal likewise is a required feature that has been exploited in efficient redox-switching of the NLO properties, which is of great attention in the field of photonics.

The coordination complexes are attractive NLO materials due to optical bands in the visible region and various oxidation states of metal ions. Coordination complexes are useful for optical applications due to the presence of an electron donor and acceptor moieties connected *via* a π -conjugate link.^{12,42–44} Using metal-ligand frameworks in these compounds, the NLO response can be enhanced. This strategy induces metal to ligand charge transfer (MLCT). In coordination complexes, charge transfer excitations occur and they differ in nature either in directions from metal towards ligand, ligand towards metal, or intra ligand.^{45–47} The coordination of the ambidentate ligand with the metal center results in structural isomerism in coordination complexes because the ligand can bind in more than one way to the metal center. The study of various ligand isomers is of potential interest for numerous applications in molecular NLO switches.^{12,15,48,49}

Literature reveals that numerous molecular switches have been reported both experimentally and theoretically to investigate their switchable NLO response. The switching NLO properties in the photochromic pair of furylfulgide Aberchrome 540 and dihydrobenzofuran derivatives studied at B3LYP/6-311++G(d,p) level of theory reveal that these systems can be used as efficient molecular NLO switches.⁵⁰ In another theoretical study, photoisomerization effect for dithiazolylarylene was studied.⁵¹ The outcome of the study demonstrates that dithiazolylarylene is not only a photochromic material but also a reversible NLO switch.⁵² The NLO switching mechanism of the scarce singlet diradical electride molecule has been reported at the CAM-B3LYP/6-311++G (2d,2p) level of DFT.⁵³ The easily driven electrons due to the electric field at two opposite ends of the molecule make the material quite suitable for use in molecular NLO-based switch design. Alkalides owing to their large NLO responses exhibit broad applications in electro-optical device fields.^{54–56} The photoactive layer materials were also used to examine the substantial role of graphene quantum dots in NLO-based molecular switches at the B3LYP/6-31G(d) level of DFT.^{57–60}

The linkage isomers have been widely investigated. These switchable organometallics have ambidentate ligands, which can have their coordination geometry altered by subjecting the crystal to outside stimuli like heat or light. Photoinduced linkage isomers exist as long-lived metastable states in the crystal (with excited states), and lifetimes that are temperature dependent.^{61–63} A variety of linkage isomers have been studied, including nitrosyl, sulfur dioxide, dinitrogen, and nitrite complexes. Recent advancements in NLO materials development include

[Ni(dppe)(NO₂)Cl], alkali metals doped 2N-atoms functionalized corannulene complexes, 5-phenylamino-isophthalic acid, 9,10-bis(phenylethynyl) anthracene, Zn complexes based on N₂O₄-type pro-ligand iridium complexes, Pt(II) dithienylethene complexes. Sulfur and nitrogen-bonded thiocyanato linkage isomers of dicarbonyl- π -cyclopentadienyl iron and tricarbonyl- π -cyclopentadienyl molybdenum have been investigated for their NLO response.^{10–12,16,64}

Therefore, we are interested to explore the NLO effects of ligand isomeric complexes of transition metals with different ambidentate ligands. Due to the nature of charge-transfer excitations, these ligand isomeric complexes are anticipated to exhibit NLO contrasts. We herein, theoretically investigated within the DFT framework the geometric, electronic, and optical properties of ligand-metal isomers of iron and cobalt with different ambidentate ligands.⁴⁵

Computational methodology

Gaussian 09 program is used for all quantum chemical simulations. Geometries of all complexes are optimized at CAM-B3LYP along with 6-31+G (d) basis set.⁶⁵ CAM-B3LYP is a range separated functional which is selected on account of capturing intra-molecular charge transfer within complexes.^{66–70} To get the most stable spin state, geometry optimization of ligand-metal complexes is performed at various spin states. The most stable structures of ligand-metal isomeric complexes are then presented for the calculations of the HOMO-LUMO gap, dipole moment (μ_0), polarizabilities (α_0) and hyperpolarizability (β_0). All properties are calculated with 6-31+G(d) basis set while using CAM-B3LYP method except hyperpolarizability which is calculated at 6-31+G(d,p) basis set. The mean polarizability (α_0) and hyperpolarizability (β_0) are documented in this manner:

$$\alpha_0 = \frac{1}{3}(\alpha_{xx} + \alpha_{yy} + \alpha_{zz}) \quad (1)$$

$$\beta_0 = \left[\beta_x^2 + \beta_y^2 + \beta_z^2 \right]^{\frac{1}{2}} \quad (2)$$

where

$$\beta_x = \beta_{xxx} + \beta_{xyy} + \beta_{xzz}$$

$$\beta_y = \beta_{yyx} + \beta_{yzz} + \beta_{yxx}$$

$$\text{and } \beta_z = \beta_{zzx} + \beta_{zxx} + \beta_{zyy}$$

For understanding the large hyperpolarizability value, a two-level model is considered.^{71,72} As stated by the two-level method

$$\beta_0 \approx \Delta\mu \times f_0 / \Delta E^3 \quad (3)$$

where, $\Delta\mu$ is variation in dipole moment, f_0 is the oscillator strength and ΔE is the transition energy.

According to the relation, hyperpolarizability is related to oscillator strength and variation in μ while it is inversely related to the transition energy.



Time-dependent density functional theory (TD-DFT) simulations are conducted by the TD-CAM-B3LYP/6-31+G (d) method to attain dominant excited states, oscillator strength (f_o), absorption maxima and energies needed for excitation of the most stable geometries.

Results and discussion

Outcomes of frequency simulations affirmed the geometries are products and not transition states.

Relative energies of ligand–metal isomers

The energy is an imperative stability parameter, complexes with lower energies are more stable than complexes having higher energies. The relative energies relating to the utmost stable complexes are displayed in Table 1.

Complex **1a** has an ONO ligand (nitrito), which exists in its isomeric NO₂ in complex **1b**. In ligand–metal isomeric complex **1a**, the nitrogen atom is bonded with cobalt metal (Fig. 1). Spin multiplicities results reveal that complex **1a** is most stable in the quintet state. The quintet spin state of **1a** is comparatively higher in stability than its singlet and triplet states by amounts of 0.5 and 6.3 kcal mol^{−1}, respectively. Furthermore, the singlet spin state of **1b**, is comparatively lower in energy compared to its triplet and quintet states by 7.9 and 2.6 kcal mol^{−1}, respectively.

Ligand–metal complexes **2a** and **2b** contain iron with nitric oxide ambidentate ligand. In complex **2a** and **2b**, oxygen and nitrogen atoms of NO are bonded with iron metal, respectively (Fig. 2). The most stable spin state of complex **2a** is quintet whereas complex **2b** is a singlet. The quintet spin state of **2a** stands more stable than its singlet and triplet states by 36.9 and 6.1 kcal mol^{−1}, correspondingly. The most stable spin state (singlet) of complex **2b**, is almost 10.1 and 13.0 kcal mol^{−1}, lower in energy than triplet and quintet states, respectively because of the strong field NO ligand.

Ligand–metal isomeric complexes **3a** and **3b**, consist of two linkage isomers of thiocyanate ambidentate ligand. In complex **3a**, the nitrogen atom of a ligand is bonded with iron whereas, in complex **3b**, the sulfur atom is bonded with iron (Fig. 3). Complex **3a** is more stable in the quintet spin state. The singlet and quintet states of **3a**, are about 90.8 and 77.8 kcal mol^{−1},

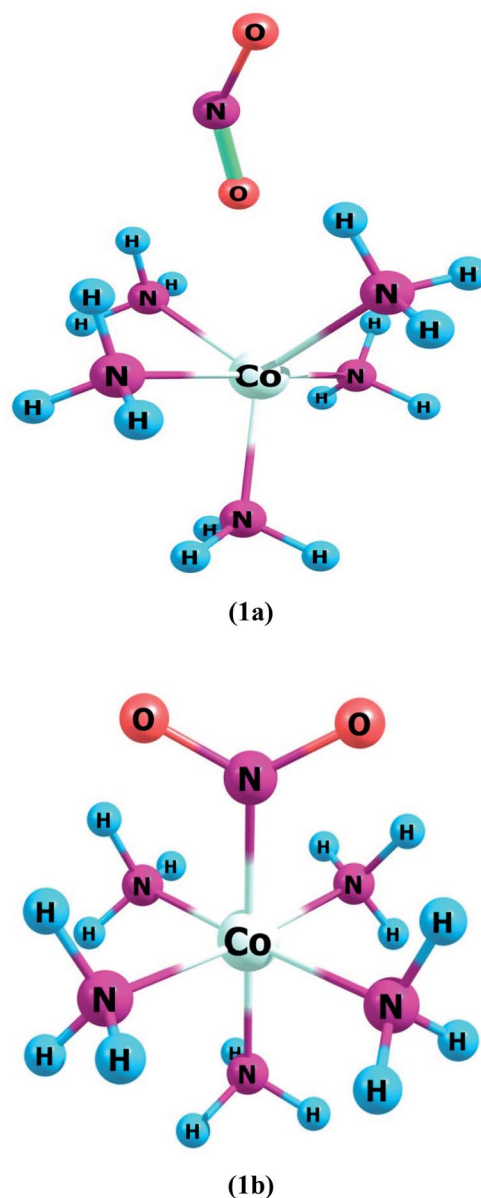


Fig. 1 Geometries of ligand–metal isomers (1a) Co–ONO and (1b) Co–NO₂.

Table 1 Relative energies (kcal mol^{−1}) of ligand–metal isomers in various spin states

Ligand–metal isomers		Relative energies		
		Singlet	Triplet	Quintet
1a	Co–ONO	0.5	6.3	0.0
1b	Co–NO ₂	0.0	7.9	2.6
2a	Fe–ON	36.9	6.1	0.0
2b	Fe–NO	0.0	10.1	13.0
3a	Fe–NSC	90.8	77.8	0.0
3b	Fe–SCN	0.0	10.8	55.2
4a	Co–SO ₃	0.0	90.5	90.4
4b	Co–OSO ₂	0.0	43.8	25.1

greater in energy than its triplet state. The S-bonded complex **3b** is more stable in the singlet spin state, which is about 10.8 and 55.2 kcal mol^{−1}, lower in energy compared to its triplet and quintet spin states, respectively.

In ligand–metal, isomeric complexes **4a** and **4b**, sulfur and oxygen atoms of ligand are bonded with cobalt metal (Fig. 4). Both **4a** and **4b** complexes remain more stable in the singlet states. The singlet state of **4a** is rather more stable as compared to its singlet and quintet states by 90.5 and 90.4 kcal mol^{−1}, respectively. The results of relative energies of most stable spin states reveal that the singlet state of **4b**, shows higher stability than triplet and quintet spin states. **4b**'s triplet and quintet states exhibit energies of 43.8 and 25.1 kcal mol^{−1}, correspondingly, which become higher than its lowest energy singlet



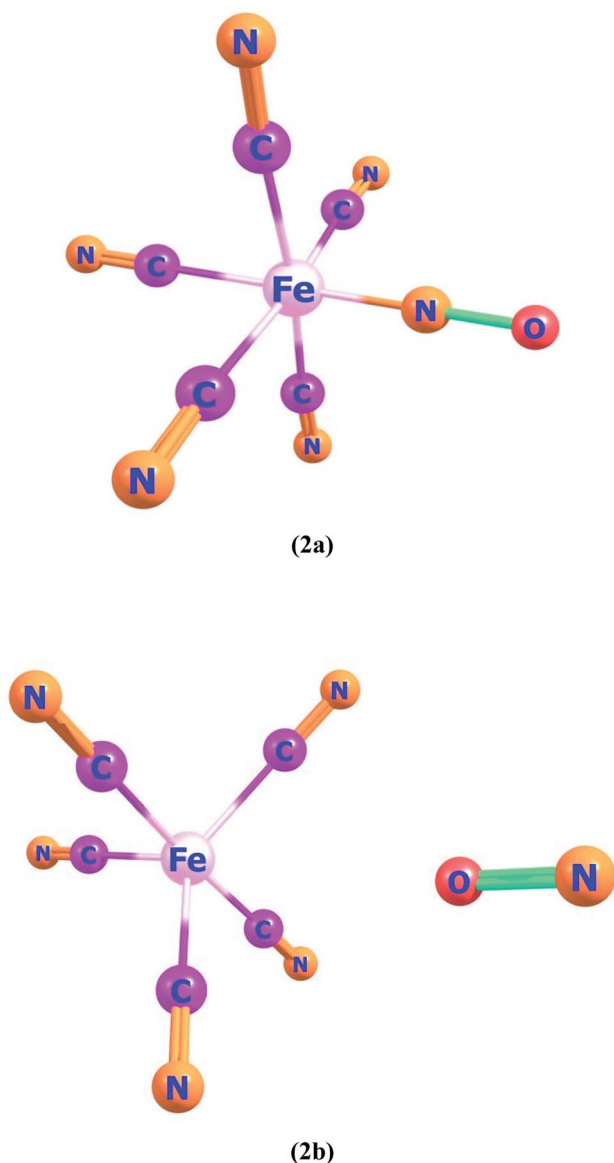


Fig. 2 Geometries of ligand–metal isomers (2a) Fe–ON and (2b) Fe–NO.

state. The OSO_2 is a strong field ligand therefore, it pairs up electrons in complex and gives stable geometry in a singlet spin state.

Electronic properties of ligand–metal isomers

FMO's investigation is carried out to get evidence concerning energies of occupied (HOMO) and virtual orbitals (LUMO) of ligand–metal isomers. Reactivity of the system is inspected by variations in energies of HOMO and LUMO. The reactivity and stability of a species are characterized by a HOMO–LUMO gap. The species with smaller H–L gap show high reactivity, and high conductivity and are generally less stable. A small ΔE value is a distinctive feature of less stable materials although easily polarizable suggesting simpler electronic transition and, therefore, improved NLO properties of the complexes.

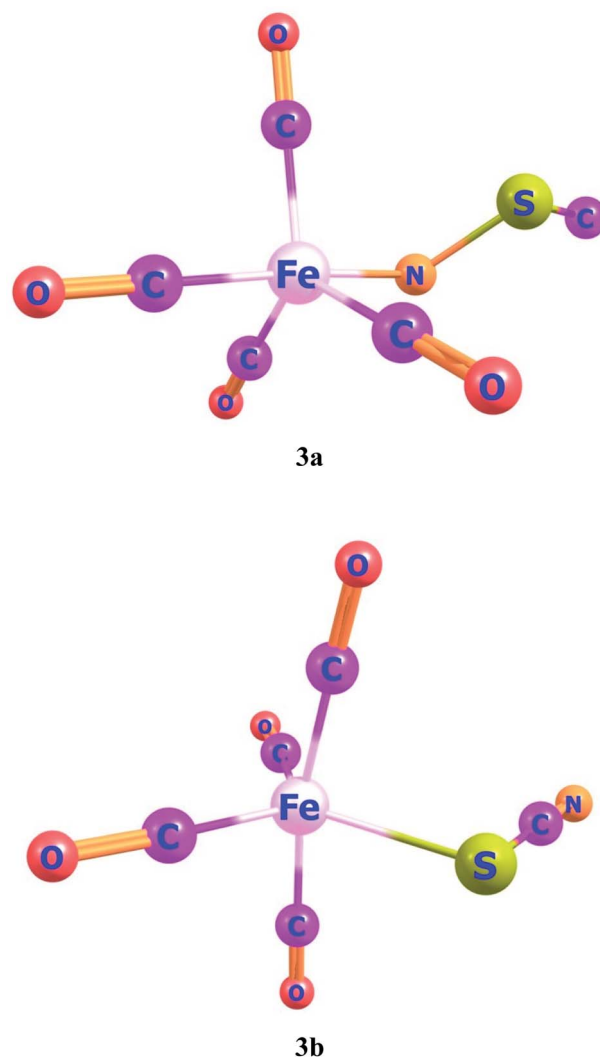


Fig. 3 Geometries of ligand–metal isomers (3a) Fe–NSC and (3b) Fe–SCN.

The results of the FMO investigation are given in Table 2. The HOMO of ligand–metal isomer **1a** appeared at -16.96 while its LUMO at -9.34 eV beside H–L gap of 7.62 eV. The HOMO and LUMO energies of ligand–metal isomer **1b** are 16.49 and 9.21 eV, respectively, with a 7.27 eV H–L energy gap. The H–L gap of **1a** (in which oxygen is bonded to Fe) is higher compared to the H–L gap of **1b** (in which nitrogen is bonded to Fe) due to efficient orbital hybridization in the latter (Fig. 5).

The HOMO of ligand–metal isomer **2a**, appeared at -11.18 whereas its LUMO was at -5.26 eV, beside the H–L energy gap of 5.92 eV. The HOMO and LUMO of ligand–metal isomer **2b**, exist at -2.71 and 4.54 eV, correspondingly beside 7.26 eV of the H–L gap (Fig. 6). The lower H–L gap in **2a**, is owing to greater orbital hybridization between the highly electronegative oxygen atom and iron centre.

HOMO and LUMO are located at 4.26 and 2.08 eV, respectively, besides the H–L gap of 6.34 eV in ligand–metal isomer **3a** (Fig. 7). The HOMO of **3b** is situated at -3.84 eV while its LUMO at 2.61 eV. The H–L gap in **3b** (sulfur bonded isomer) is 6.46 eV,

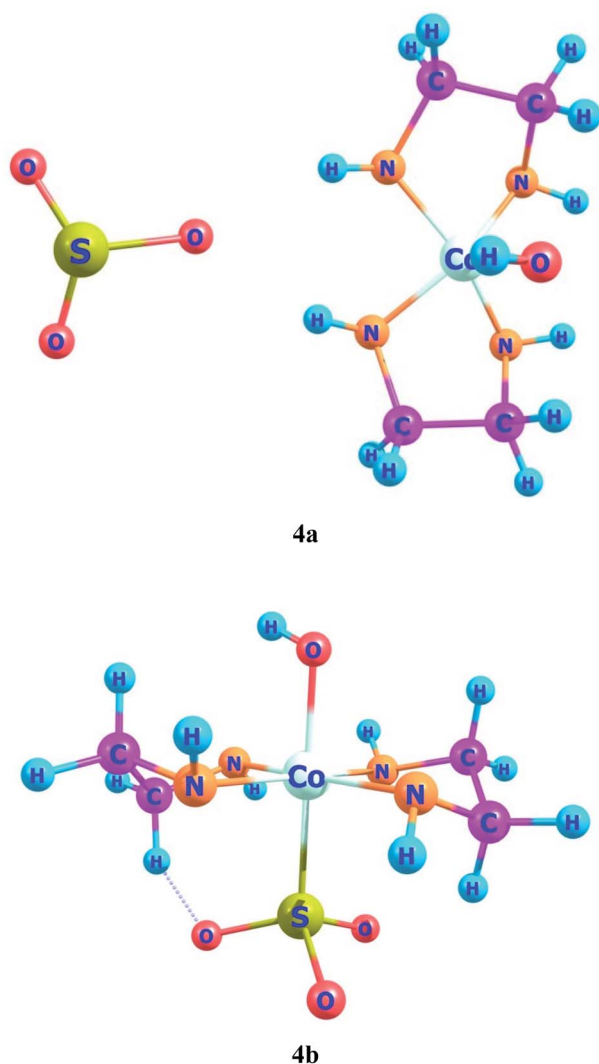


Fig. 4 Geometries of ligand–metal isomers (4a) Co–SO₃ and (4b) Co–OSO₂.

which is higher compared to the H–L gap of isomer 3a (nitrogen bonded isomer). The reduced H–L gap of 3a is because of the high electronegativity of the N atom of NSC ligand bonded with iron metal which causes efficient orbital hybridization.

The HOMO of ligand–metal isomer 4a is located at –11.99 eV whereas its LUMO is present at –5.21 eV beside H–L gap of

6.77 eV in the complex. The H–L gap in 4b is 3.84 eV which is found to be the lowest among all studied complexes. The HOMO of the 4b is located at –7.62 eV, whereas LUMO is at –3.78 eV. The substantial decrease in H–L gap in 4b compared to 4a is due to the effective orbital overlap between the highly electronegative oxygen atom of the ligand and cobalt metal center. Moreover, the results of FMO reveal that a noteworthy drop in the H–L gap is noticed in isomeric complexes where a highly electronegative atom of ligand bonded with an electropositive metal center.

Nonlinear optical properties of ligand–metal isomers

The transference of charge both from metal to ligand and from ligand to metal assures the actual NLO response of ligand–metal isomers. These complexes contain functioning centers of an additional electron that certainly be transported to an electron-scarce place. This intramolecular charge transference assures the advancement of NLO properties of ligand–metal isomers. The outcomes of the NLO investigation are listed in Table 3.

Ligand isomers complexes consist of ambidentate ligand which contains two different donor atoms of different electronegativities. As a result of the difference in electronegativities of donor atoms of ligand, the shift of electronic density is different in these isomeric complexes. Because of the different patterns of electronic densities distribution, these isomers show a contrast in NLO response.

Complex 1, comprises two isomers (1a & 1b) of NO₂ ambidentate ligand. In isomer 1a, cobalt is attached to the oxygen atom of the ligand, and in 1b, the nitrogen atom of the ligand is attached to the Cobalt center (Fig. 1).

These two isomers have different hyperpolarizability values. The β_o of 1a is 158 a.u. while that of 1b is 192 a.u. The β_o of isomer 1b, is 1.21 times higher compared to 1a, due to the charge transference pattern and high oscillator strength. The greater β_o of 1b is consistent with its small H–L gap than 1a (*vide supra*). In isomer 1a, HOMO density is mainly situated equally on metal and ligand, though LUMO density is only sited on metal. This demonstrates the transfer of electronic cloud from ligand to metal on excitation from HOMO to LUMO and hence called MLCT. In isomer 1b, HOMO and LUMO densities are equally situated on the ligand moiety. In ligand isomer 1b, the orbital overlap is inadequate to develop an effectual electronic

Table 2 FMO and UV-Vis spectroscopic outcomes of ligand–metal isomers

Ligand–metal isomers	E_{HOMO} (eV)	E_{LUMO} (eV)	H–L gap (eV)	$E_{\text{excitation}}$ (eV)	f_o (a.u.)	λ_{max} (nm)
1a	–16.96	–9.34	7.62	3.26	0.01	256
1b	–16.49	–9.21	7.27	4.82	0.23	380
2a	–11.18	–5.26	5.92	3.73	0.03	332
2b	–2.71	4.54	7.26	3.97	0.02	311
3a	–4.26	2.08	6.34	4.96	0.01	249
3b	–3.84	2.61	6.46	2.27	1.03	545
4a	–11.99	–5.21	6.77	2.47	0.01	500
4b	–7.62	–3.78	3.84	2.79	0.01	444



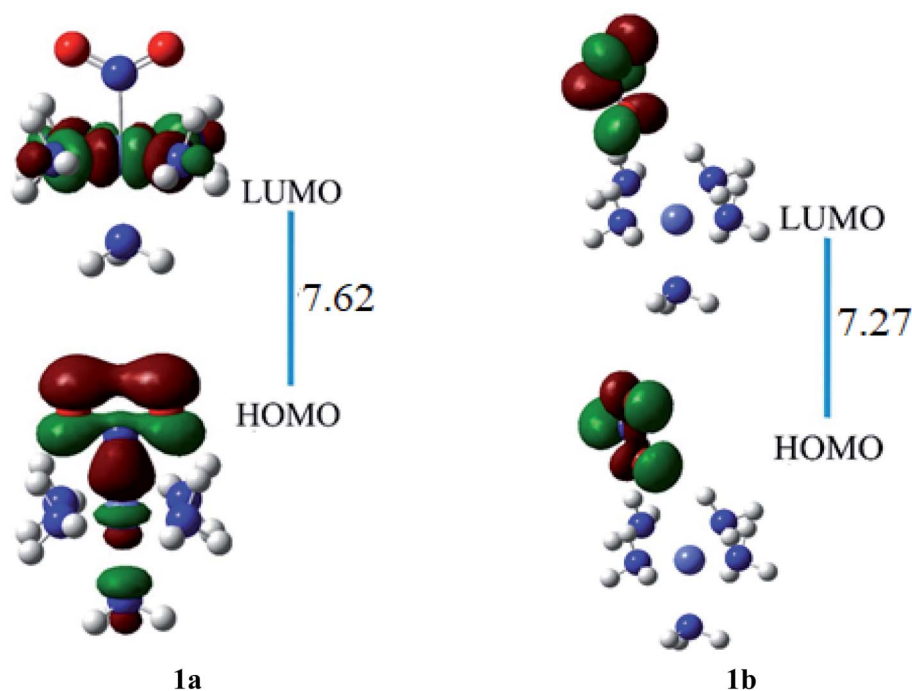


Fig. 5 The HOMO and LUMO visualization of ligand-metal isomers **1a** and **1b**.

densities distribution. The intra-ligand transference of charge transitions in **1b**, is termed ILCT. The high energy of transition beside poor distribution of charge and lower dipole moment are observed in **1a**. The donor and acceptor features between molecule may influence the NLO response and thus charge

transference cause variation in the β_o value of these ligand-metal isomers.

The β_o values of ligand-metal isomeric complexes **2a** and **2b** are 828 and 173 a.u., respectively. The β_o of **2a** stands 4.78 times higher in value than its isomer **2b**. The HOMO and LUMO

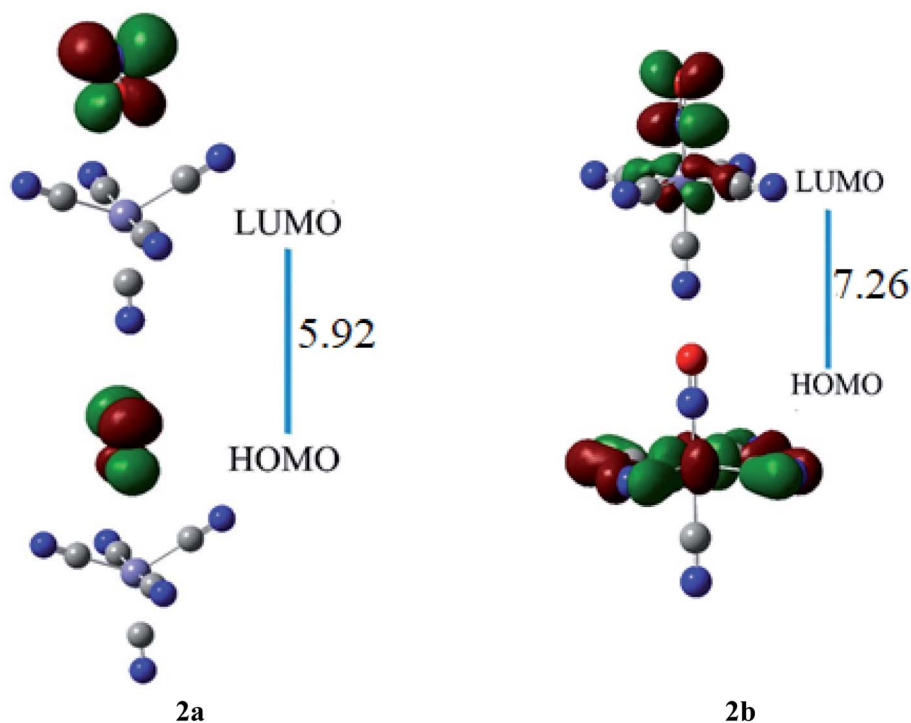


Fig. 6 The HOMO and LUMO visualization of ligand-metal isomers **2a** and **2b**.



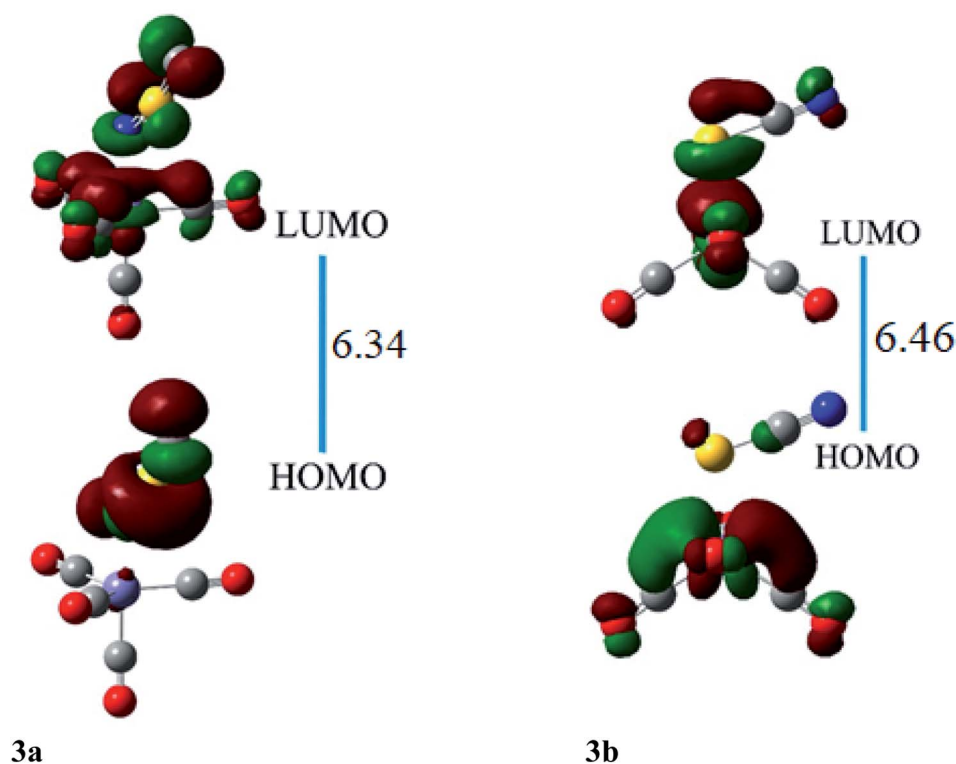


Fig. 7 The HOMO and LUMO visualization of ligand–metal isomers **3a** and **3b**.

Table 3 Dipole moment (μ_o), mean polarizabilities (α_o), mean first hyperpolarizabilities (β_o), and ratios of hyperpolarizabilities of ligand–metal isomers calculated at CAM-B3LYP/6-31G+(d,p) level

Complexes	μ_o (Debye)	α_o (a.u.)	β_o (a.u.)	State	Ratio (ON/OFF)
1a	7.82	83	158	OFF	
1b	10.61	88	192	ON	1.21
2a	5.48	138	828	ON	4.78
2b	7.23	137	173	OFF	
3a	5.35	165	3662	ON	21.15
3b	5.94	140	431	OFF	
4a	11.67	134	390	OFF	
4b	2.10	165	4184	ON	10.71

densities in **2a**, are equally present on the ligand. The intra-ligand charge transference mode (ILCT) of **2a**, is displayed in Fig. 6. In **2b**, the HOMO density is placed at the metal center, however, the LUMO density is situated at ligand as well on metal sites of the complex (MLCT). In isomer **2a**, where iron is bonded with the oxygen atom of ligand, the electronic density is shifted more towards ligand than in isomer **2b** (where Fe is bonded with the nitrogen atom of ligand). The greater shifting of electronic density towards ligand in isomer **2a** is due to the higher electronegativity of the oxygen atom. The greater β_o of isomer **2a**, compared to **2b**, is consistent with its lesser transition energies and reduced H–L gap besides bathochromic effects.

The β_o of ligand–metal isomer **3a** is 3662 a.u., however that of **3b** is 431 a.u. The β_o of isomer **3a**, is 21.15 times higher in value compared to **3b**. The greater contrast of β_o values is observed in **3a** and **3b**, among all studied isomeric complexes. The visualization of HOMO and LUMO densities of **3a** and **3b** are presented in Fig. 7. The results of charge transference excitation in isomer **3a**, show that HOMO density is entirely situated at the ligand site while LUMO densities are equally at ligand and metal moieties. The spreading of HOMO and LUMO densities in this isomeric pair suggests ligand to metal charge transfer (LMCT).

In **3b**, the density of HOMO is present at the metal part while LUMO density is situated at both metal and ligand (MLCT). The larger hyperpolarizability of **3b** than **3a** is due to efficient charge transference from Fe to the ligand in the complex.

The ligand–metal isomers, **4a** & **4b** have hyperpolarizabilities of 390 a.u. and 4184 a.u., respectively. Isomer **4b**, exhibit a high β_o compared to **4a**, beside its lesser H–L gap. The HOMO and LUMO densities are equally spread over metal in isomer **4a** (Fig. 8). The orbital overlap is insufficient to cause an efficient distribution of electronic density in **4a**, due to the low electronegativity of the S atom in the ligand. In ligand–metal isomer **4b**, HOMO density is situated at metal and ligand parts. Likewise, LUMO density is also situated equally at ligand and Co moieties in isomer **4b** where charge transfer from ligand to metal is noticed (MLCT). The higher electronegativity of the oxygen atom of ligand bonded with cobalt metal in isomer **4b**, causes effective charge distribution which results in its higher β_o . The noteworthy intra-molecular transference of charge in



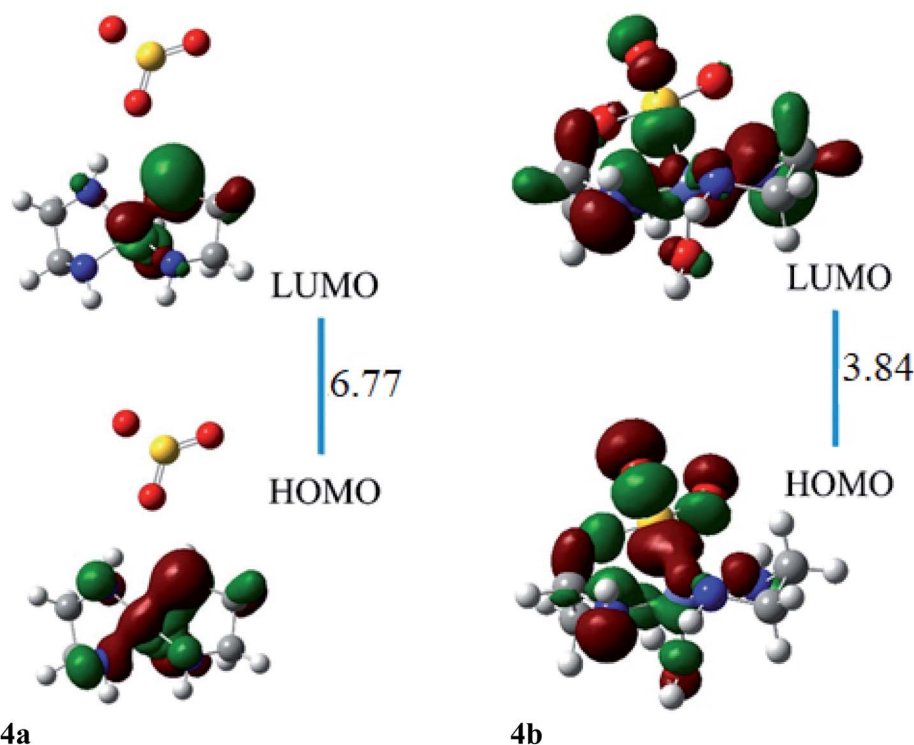


Fig. 8 The HOMO and LUMO visualization of ligand–metal isomers **4a** and **4b**.

ligand–metal isomers, declares their potential usage, as distinctive NLO materials for numerous applications. Analysis of the results in Tables 2 and 3 reveals that the H–L gap is the main deciding factor for hyperpolarizabilities. The lower the HOMO–LUMO gap, the higher is the hyperpolarizability. This statement is valid for all complexes.

Molecular absorption spectra of ligand–metal isomers

TD-DFT is a reliable method to develop ultraviolet-visible (UV-visible) absorption spectra. Ligand–metal isomers show considerable absorption in the entire ultraviolet and visible region to some extent. Results of excited state calculations of ligand–metal complexes are displayed in Table 2. The UV-visible spectra of isomers **3a** & **3b** are presented in Fig. 9, however, all remaining complexes are given in ESI.† We herein, calculated twenty electronic states to attain vital excited states. The results reveal that ligand–metal isomers **1a** and **1b**, show maximum absorption (λ_{max}) at 256 and 380 nm, respectively. The bathochromic shifting of absorption maximum is observed in **1b** compared to **1a**. The oscillator strength (f_o) or the probability of absorption for electronic transition in **1a** and **1b**, are 0.01 and 0.23 a.u. respectively. The λ_{max} of isomer **2a** is moved to a longer wavelength compared to the λ_{max} of isomer **2b**. The oscillator strength of isomers **2a** and **2b**, are 0.03 and 0.02 a.u. respectively. The red shifting of λ_{max} in **2a**, compared to its isomer **2b**, is consistent with its higher hyperpolarizability and lower excitation energy.

The isomers **3a** & **3b**, show absorption maxima at 249 and 545 nm, respectively. Isomers **3a** and **3b**, show oscillator

strength of 0.01 and 1.03 a.u. respectively. λ_{max} of **3b**, is red shifted regarding **3a**, which matches with its lower excitation energy. The electronic transitions of ligand isomers **4a** and **4b**, appeared at 500 and 444 nm, respectively. The oscillator strength in both isomers is 0.01 a.u. Isomer **4a** has lower excitation energy than corresponding isomer **4b**, which is associated with its larger bathochromic shift of λ_{max} . The predictions of UV-visible absorption spectral results we obtained reflect better NLO properties of ligand–metal isomers.

Ligand–metal isomeric NLO switches

Materials having the contrast of NLO properties are of significant attention owing to their extensive use in many devices. In the current work, we investigated the NLO switching effects of ligand–metal isomeric complexes. We here focused on the extent of hyperpolarizability values of these complexes besides their switching ratios (Fig. 10).

Ligand–metal isomer **1b** shows a β_o value which is 1.21 times more than **1a**. Therefore, isomer **1b** is switched “ON” state while **1a** is switched “OFF” state. Likewise, isomer **2a** shows 4.78 times more β_o than the corresponding **2b** isomer. Ligand–metal isomer **2a** is examined as switch “ON” whereas, **2b** as switch “OFF” state. The β_o of **3a**, is 21.15 times greater than its isomeric complex **3b**. Isomer **3a** with a greater β_o value than **3b**, is termed as switch “ON” state but **3b** as switch “OFF” state. The highest contrast in β_o values is observed in isomeric pair **3a** and **3b**. Also, ligand–metal isomer **4b**, with 10.71 times larger β_o value than **4a**, is switch “ON” state whereas **4a** is switch “OFF” state. Hence, the results reveal



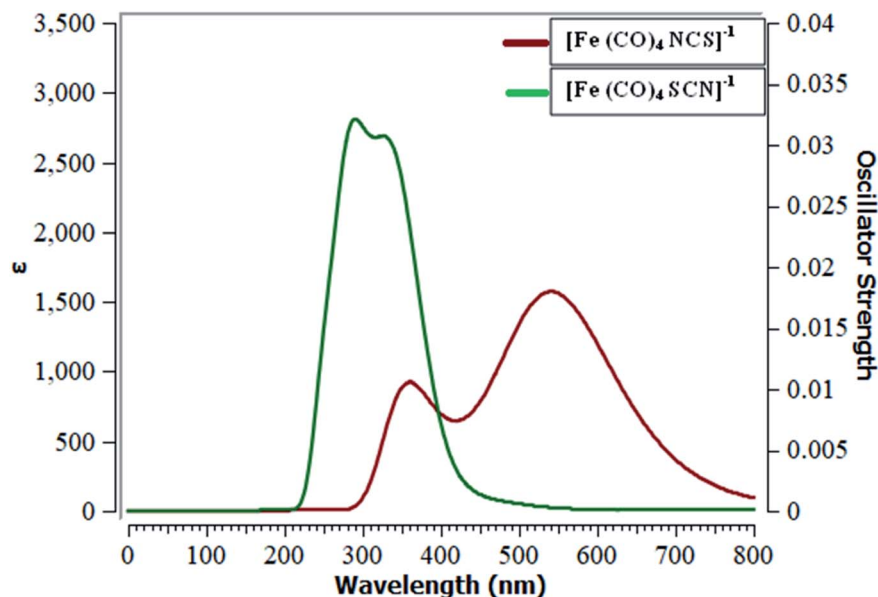


Fig. 9 UV-visible spectra of metal isomers of SCN.

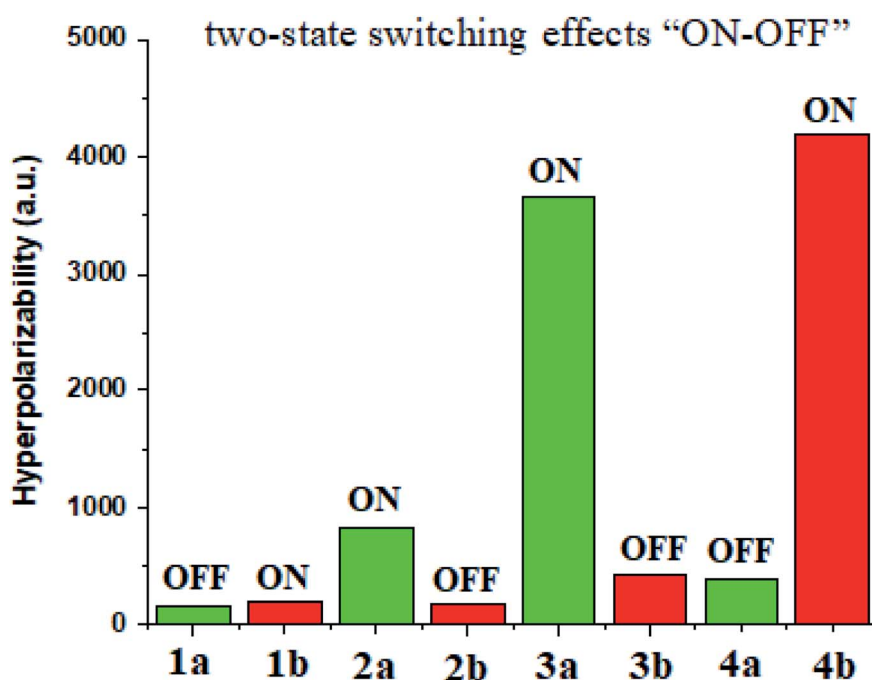


Fig. 10 Switch ON–OFF ratios of ligand–metal isomers.

that these ligand–metal isomeric pairs show two-state switching effects “ON–OFF”.

Conclusions

DFT study is performed to investigate the relative stability, molecular absorption spectra, and NLO responses besides switching effects of ligand–metal isomeric complexes. Larger contrast in β_o values of studied complexes ensure their potential

use as a switchable NLO material. The outcomes of excited state investigations show that electronic transitions and charge density distributions *via* MLCT, LMCT and ILCT, mark the difference in NLO behavior in these isomers. Transition energies and H–L gaps of these isomeric pairs mainly contribute to the comparative variations of β_o values. Larger contrast of the β_o values has been observed in these ligand–metal isomeric complexes. The highest NLO contrast in β_o values is seen in **3a** and **3b**, isomeric pairs. The highest β_o value of 4184 a.u. is seen



in **4b**, that reflects its strong NLO properties. Hopefully, coordination complexes of Fe and Co with isomeric ligands are talented to become redox-triggered switchable NLO materials for the development of NLO devices.

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Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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