

Molecular structure and DFT investigations on new cobalt(II) chloride complex with superbase guanidine type ligand

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Abstract. The new [Co(btmgn)Cl₂] complex and the 1,8-bis(tetramethylguanidino)naphthalene (btmgn) ligand were synthesized and characterized. The X-ray single crystal investigation showed distorted tetrahedral geometry around the Co(II) ion. The geometry of the btmgn and [Co(btmgn)Cl₂] complex was optimized using the B3LYP/6–311G(d,p) method. The calculated geometric parameters at the optimized structure of the [Co(btmgn)Cl₂] complex showed good agreement with our reported X-ray structure. The two tetramethylguanidino groups are in a cis-type position to the naphthalene ring plane both in the free and coordinated btmgn. The large red shift of the $\nu_{C=N}$ mode upon coordination indicates the strong ligand–metal interactions. The calculated natural charges using natural bond orbital (NBO) analysis at the two coordinated Cl-atoms are not equivalent. Also the two LP(4)Cl \rightarrow LP*(3)Co intramolecular charge transfer interaction energies (E⁽²⁾) are 29.00 and 39.17 kcal/mol, respectively. The two Co-Cl bonds are not equivalent where the longer Co-Cl bond has more electronegative chlorine atom than the shorter one. Molecular electrostatic potential (MEP) study of the btmgn ligand showed that the N4 and N7 atoms are the most reactive nucleophilic centers for the coordination with the Co²⁺ ion. The [Co(btmgn)Cl₂] complex has higher polarizability (α_0), first hyperpolarizability (α_0) and lower energy gap (α_0 E) than the free ligand. The TD-DFT calculations predicted the transition bands at 337.2 nm (f=0.2299, H α_0 E) and 342.6 nm (f=0.1465, H-2/H α_0 E) for the btmgn and [Co(btmgn)Cl₂], respectively.

Keywords. Guanidine; superbases; Cobalt(II); DFT; NBO; MEP.

1. Introduction

In the past decades, a number of organic proton sponges and superbases were synthesized and applied for various purposes. The classical proton sponge 1,8-bis(dimethylamino)naphthalene was described already in 1968 by Alder *et al.* More recently, Sundermeyer *et al.* reported on the new superbasic proton sponge 1,8-bis(tetramethylguanidino)naphthalene (btmgn) as an example for an aromatically bridged bisguanidine. Due to resonance stabilization of the conjugated acid, guanidines are much stronger bases than amines. In coordination chemistry, the neutral guanidine ligands represent not only excellent σ -donor, but also π -donor ligands. In the last decade, the design of organic—inorganic hybrid mesoporous materials with strong basic functionalities

has attracted great attention because of their potential applications in heterogeneous catalysis for the synthesis of fine chemicals.⁶ The peralkylated bifunctional nitrogen ligands of Cu and Fe were designed and used in the field of biomimetic coordination chemistry.^{7,8} Also, the Mo and Ru complexes of these ligands were prepared and their catalytic performance in the hydrogenation process was demonstrated. Late-transition metals, especially Ni, Pd, and Pt coordinated to diimine ligands were used to catalyze polymerization of ethylene and other α -olefin molecules. ¹⁰ Moreover, guanidine groups (arginine, creatine) play an important role in biological systems, in particular for proton transfer processes in proton channels. 11,12 Studies of the structural, electronic and spectroscopic properties of guanidine ligands containing aromatic group as potentially strong electron donors and their complexes with Co(II) and Ni(II) were still rare in literature. 13-15

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In the present work the new CoCl₂ complex of the guanidine-type ligand, bis(tetramethylguanidino) naphthalene (btmgn) has been synthesized. The ligand and the new [Co(btmgn)Cl₂] complex were characterized using different spectroscopic techniques. The X-ray structure of the [Co(btmgn)Cl₂] complex is determined. The electronic and spectroscopic aspects of the ligands as well as the [Co(btmgn)Cl₂] were discussed with the aid of DFT quantum chemical calculations.

2. Experimental and Computational

2.1 General

The solvents used were of HPLC reagent grade. Melting points were determined with a Mel-Temp apparatus and are uncorrected. Magnetic resonance spectra (1 H NMR and 13 C NMR spectra) were recorded on a JOEL 400 MHz spectrometer with chemical shift values reported in δ units (ppm) relative to an internal standard. Follow-up of the reactions and checks of the purity of the compounds was done by TLC on silica gel-protected aluminum sheets (Type 60 GF254, Merck) and the spots were detected by exposure to UV-lamp at λ =254 nm for a few seconds.

2.2 Synthesis of 1,8-bis(tetramethylguanidino) naphthalene (btmgn)³

Chloroformamidinium salt 1 (TCFH, 5.7 g, 20 mmol) was added slowly to a solution of 1,8-diaminonaphthalene 2 (1.6 g, 10 mmol) and triethylamine (2.0 g, 20 mmol) in 10 mL DMF. The reaction was carried out at 90°C in a water bath for 6 h. The mixture was left to cool to room temperature, and then poured into 50 mL of ice cold water. The solid product formed was filtered directly and then washed with water. The crude product btmgn was recrystallized from CH₂Cl₂/n-hexane.

The product btmgn was obtained as a gray solid in 76% yield, M.p. 214-216°C (dec). R_f : 0.59 (CH₂Cl₂/MeOH 9:1). IR (KBr, cm⁻¹): 3050-2855 (C-H), 1640 (C=N), 1570-1540 (C=C aromatic), 1281 (C-N). ¹H-NMR (DMSO-d₆) ppm: δ 2.86 (s, 24 H, 8 CH₃), 6.49 (d, 2 H, aromatic), 7.32-7.42 (m, 4 H, aromatic). ¹³C-NMR ((CD₃)₂CO) ppm: δ 40.79, 115.12, 119.50, 123.12, 127.47, 137.93, 143.69, 160.62.

2.3 Synthesis of $[Co(btmgn)Cl_2]$

To an aqueous solution, 10 mL, of cobalt(II) chloride (1.0 g, 0.77 mmol) a 15 mL ethanolic solution of the organic ligand (btmgn), (3 g, \sim 0.85 mmol) was added dropwise with constant stirring over about 15 min. The clear solution was allowed to stand for 2 weeks (some

preparations needed longer time). Uniform, deep blue crystals, crystals suitable for X-ray measurements were collected and air-dried. There was a yield of 0.85 g, about 76% with respect to the metal. FTIR: 3059-2860 (C-H), 1558 (C=N), 1520 (C=C aromatic), 1278 (C-N) cm⁻¹. UV-Vis. 237.0, 348.0, 611.1, 648.0 nm (Nujal mull).

2.4 XRD measurement

Single crystal X-ray diffraction data were collected on an Enraf-Nonius 590 Kappa CCD single crystal diffractometer with graphite monochromated MoK α (λ = 0.71073 Å) radiation. Intensity data were collected at room temperature using φ and ω scans; the crystal to detector distance was 40 mm. Data processing was performed using the maXus program package, 16 the cell refinement and data reduction were carried with Denzo and Scalepak's programs;¹⁷ the crystal structures were solved by direct methods using the SHELXS program¹⁸ which revealed the positions of all nonhydrogen atoms. The crystal structure was refined by full matrix least square refinement based on F² using the SHELX program package. 18 The temperature factors of all non-hydrogen atoms were refined anisotropically. The hydrogen atoms were calculated using a riding model with C-H = 0.96 Å. Multiscan absorption corrections were applied to all data sets using the program SORTAV.¹⁹ Crystallographic data and data collection details are summarized in Table 1.

Supplementary data are available from the CCDC, 12 Union Road, Cambridge CB2 1EZ, UK on request, quoting the deposition number CCDC-1037138 or the DOI for this paper. These data can be obtained free of charge at www.ccdc.cam.ac.uk/conts/retrieving.html or fax: (internat.) +44-1223/336-033; E-mail: deposit@ccdc.cam.ac.uk.

2.5 Computational Details

All calculations for the [Co(btmgn)Cl₂] complex as well as the free ligand (btmgn) were carried out using Gaussian 03W software.²⁰ The quantum chemical calculations have been performed using the B3LYP level of theory supplemented with the 6–311G(d,p) basis set. GaussView4.1²¹ and Chemcraft²² have been used to draw the structures of the optimized geometries and to draw the HOMO and LUMO pictures. The input file was taken from the CIF obtained from our reported X-ray single crystal measurement. The geometries were optimized by minimizing energies with respect to all geometrical parameters without imposing any

Table 1. Crystal data and structure refinement of the [Co(btmgn)Cl₂] complex.

Empirical formula Formula weight Temperature Wavelength Crystal system Space group Unit cell dimensions	$C_{20}H_{30}Cl_{2}CoN_{6}$ 484.33 $298(2) K$ $0.71073 Å$ monoclinic $P2_{1}/c$ $a = 11.385(2) Å$ $b = 11.318(2) Å$ $c = 18.339(4) Å$	$\alpha = 90^{\circ}$ $\beta = 95.59(3)^{\circ}$ $\gamma = 90^{\circ}$
Volume	$2351.8(8) \text{ Å}^3$,
Z	4	
Density (calculated)	1.368 Mg/m^3	
Absorption coefficient	0.975 mm^{-1}	
F(000)	1012	
Crystal size	$0.12 \times 0.04 \times 0.03 \text{ mm}^3$	
Theta range for data collection	3.00 to 24.99°	
Index ranges	-13 <= h <= 13, -13 <= k <= 12, -21 <= l <= 21	
Reflections collected	7011	
Independent reflections	4115 [R(int) = 0.0408]	
Completeness to theta = 24.99°	99.30%	
Max. and min. transmission	0.9713 and 0.8920	
Refinement method	Full-matrix least-squares on F ²	
Data / restraints / parameters	4115 / 0 / 271	
Goodness-of-fit on F ²	1.009	
Final R indices $[I > 2\sigma(I)]$	R1 = 0.0397, wR2 = 0.0704	
R indices (all data)	R1 = 0.0859, wR2 = 0.0829	
Extinction coefficient	0.0020(4)	
Largest diff. peak and hole	$0.251 \text{ and } -0.267 \text{ e.Å}^{-3}$	

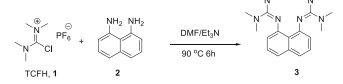
molecular symmetry constraints. The true energy minimum at the optimized geometry of the studied compounds was confirmed as indicated by the absence of any imaginary frequency modes. The electronic spectra of the studied compounds were calculated by the TD-DFT method. The natural bond orbital (NBO) calculations²³ were performed using NBO 3.1 program as implemented in the Gaussian 03W package at the DFT/B3LYP level. The second order Fock matrix was carried out to evaluate the donor–acceptor interactions in the NBO basis.²⁴

3. Results and Discussion

3.1 *Structure of the ligand*

The ligand (btmgn) was prepared by reaction of TCFH and 1,8-diaminonaphthalene using DMF as a solvent and following the reported method for preparation of guanidine using TCFH²⁵ (scheme 1). The NMR spectra are presented in figure S1 (Supplementary Information).

The spectral analysis of the btmgn ligand confirmed the expected structure. The IR spectrum of the btmgn



Scheme 1. Synthesis of Ligand.

showed an absorption peak at 1640 cm⁻¹ corresponding to the C=N groups, and the expected bands for aromatic structure. The corresponding band in the [Co(btmgn)Cl₂] is located at about 1558 cm⁻¹. The large red shift upon coordination indicates the strong ligand–metal interaction.

¹H NMR spectrum of the btmgn showed a multiplet corresponding to the four aromatic protons H-2, H-3, H-6 and H-7 in the region 7.42-7.32 ppm. A doublet (δ =6.49, J=7.1 Hz) corresponding to the two protons H-4 and H-5 was observed. A singlet at δ 2.86 ppm corresponding to the twenty four hydrogens of the eight equivalent N-methyl groups was also observed.

¹³C-NMR spectrum of btmgn showed the expected signals. A signal at 160.2 ppm corresponding to the two equivalent C=N groups was observed. Six signals

corresponding to the ten aromatic carbons showed in the region 143.69-115.12 ppm. A signal at δ 40.78 ppm assigned to the carbons of the eight equivalent methyl groups was also observed.

3.2 *X-ray Crystal Structure of* [$Co(btmgn)Cl_2$]

The complex [Co(btmgn)Cl₂], crystallizes with monoclinic symmetry in the space group P2₁/c. The asymmetric unit of the cell contains one formula unit of the monomeric complex, thus the unit cell comprises four symmetrically equivalent complex molecules. The cobalt atom is distorted tetrahedrally coordinated by two chlorine atoms and the chelating guanidine ligand. Figure 1 shows the molecular structure of this monomeric complex, the interatomic distances as well as the bond angles are listed in table S1 (Supplementary Information). The Cl-Co-Cl bond angle is 112.94(4)° and thus close to the ideal tetrahedral angle. As expected, the two Co-N distances in this complex are very similar: Co-N1: 1.994(2) Å and Co-N4: 1.996(2) Å. The N-Co-N bite angle of the chelating guanidine ligand is 90.18°. This value compares quite well with the data of 89.69° and 90.12° found for the corresponding complexes of NiCl₂ and NiBr₂, respectively.²⁶ These two compounds crystallize in the same space group and with similar unit cell dimensions and thus can be regarded to be isotypic with the presently discussed [Co(btmgn)Cl₂] complex. All compounds using 1,8-bis(tetramethylguanidino)naphthalene as chelating

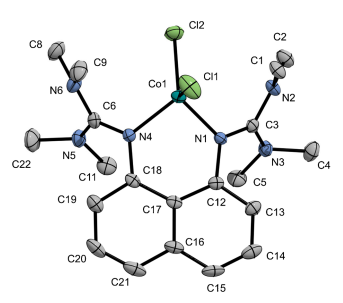


Figure 1. Structure and atom numbering scheme of the [Co(btmgn)Cl₂] complex. The hydrogen atoms have been omitted from the drawing for clarity. All atoms are drawn with ellipsoids of 30% probability level.

agent have in common, that the two tetramethyl-guanidino groups are in a cis-type position to the plane formed by the naphthalene rings. This was also found for the very similar compound bis(tetramethyl-guanidino)benzenedichlorocobalt(II).²⁷

Figure 2 depicts the packing of monomeric complex molecules in a view along the crystallographic b axis. It is evident that the crystal structure contains non-polar and polar regions. The non-polar regions are formed by the naphthalene rings located perpendicular to the bc plane at the height x=0. In the range between $x\sim 1/4$ and $x\sim 1/3$ numerous polar interactions between the complex molecules can be observed. A number of hydrogen bridge bonds between some protons of methyl residuals in the tetramethylguanidino groups and the chlorine atom with Cl-H distances well below 3.00 Å can be found (see figure 3). Additional Cl-H bridge occurs between the H15 hydrogen atom of the naphthalene ring and the Cl1 atoms of the adjacent molecular layers.

3.3 Optimized molecular structure

The optimized geometric structures and atomic numbering of the studied complex are shown in figure 4. The predicted geometric parameters (bond lengths and bond angles) of the investigated structures are collected in table S2 (Supplementary Information). Most of the bond lengths are overestimated except some of the C–N bonds which have smaller bond lengths than the experimental values. Generally, the bond lengths and bond angles are predicted very well. The maximum deviations of the calculated bond length values from the experimental data are 0.051 and 0.031 Å for the Co–Cl2 and Co–Cl3, respectively. The calculated Co–Cl2 and Co–Cl3 bond distances are 2.316 Å (exp. 2.265 Å) and 2.264 Å (exp. 2.233 Å). The percentage error in these

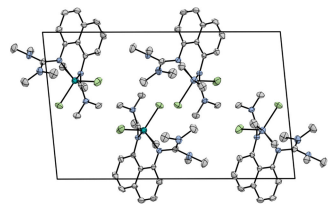


Figure 2. Unit cell content of $[Co(btmgn)Cl_2]$ complex. The structure is shown in a view along the monoclinic b-axis.

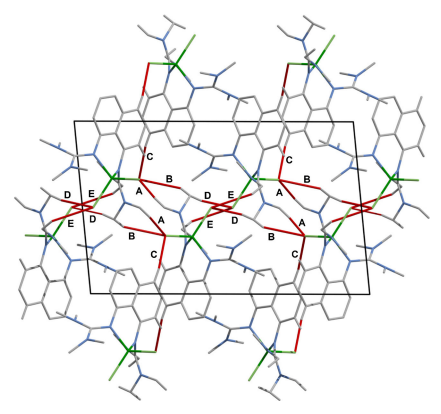


Figure 3. Hydrogen bonds in the structure of [Co(btmgn)Cl₂] complex. The Cl-H interactions shorter than 3.00 Å are shown as red bars (**A**: Cl1-H1B: 2.762 Å; **B**: Cl1-H2B: 2.841 Å; **C**: Cl1-H15: 2.731Å; **D**: Cl2-H2A: 2.930 Å; **E**: Cl2-H4B: 2.858Å). The different Cl-H interactions are marked by the capital letters**A-E** in this figure. The structure is shown in a view along the monoclinic b-axis.

bond distances does not exceed 2.3%. These deviations are mainly due to the phase difference between the calculations and the experimental data. The former is free from the crystal field effects. In agreement with the experimental X-ray structure, the calculations predicted that the two Co–Cl bond distances are not identical.

In the present section, the structural variations in the ligand geometric parameters due to coordination between the Co²⁺-ion and the ligand through N4 and N7 atoms have been discussed. Most of the ligand geometric parameters undergo very little variations due to the formation of the Co-N bonds. The C10-N4 and C11-N7 bond distances are increased by 0.04 Å while the C10-N10 and C11-N9 bond distances are decreased by the same magnitude. The rest of the bond distances are changed very little. Moreover, the most significant bond angle changes occur around the coordination sites (N4 and N7). The C10–N4–C40, N4–C10–N5, N4– C10-N6, C11-N7-C43, N7-C11-N8 and N7-C11-N9 bond angle changes are in the range of 4.36-4.74°. The C10-N4-C40-C44, C10-N4-C40-C41, C11-N7-C43-C46 and C11-N7-C43-C41 dihedral angles of the free ligand and [Co(btmgn)Cl₂] complex are calculated in the range of the values of the ranges should be 39.7– 46.2° and 32.0– 36.9° , respectively. We noted that the formation of the Co–N bonds decreases the deviation of the two –C(N(CH₃)₂)₂ from the naphthalene ring plane. Also, the two tetramethylguanidino groups are in a cistype position to the plane formed by the naphthalene rings both in the btmgn and [Co(btmgn)Cl₂].

3.4 Natural atomic charges

Distribution of positive and negative charges has vital role in the application of quantum chemical calculations to molecular system because of atomic charges affect dipole moment, molecular polarizability, electronic structure, acidity-basicity behavior and other properties of molecular system. ²⁸ In this regard, the natural atomic charges (NAC) were calculated using the DFT B3LYP/6–311G(d,p). The calculated natural charges at the different atomic sites of the [Co(btmgn)Cl₂] complex as well as the free ligand (btmgn) are given in table 2. All the H-atoms are electropositive where the natural atomic charge values at these sites are in the

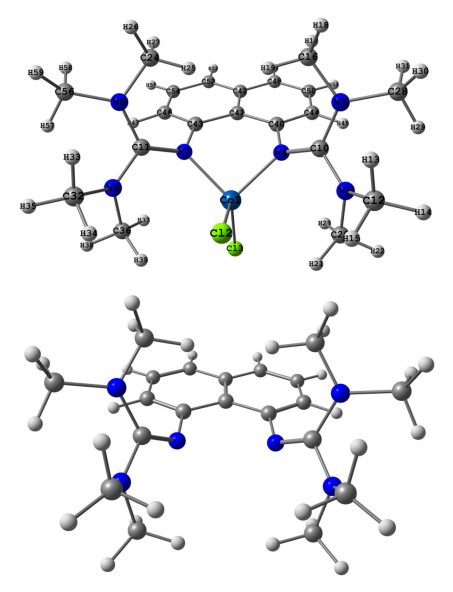


Figure 4. The optimized molecular structure of the free ligand btmgn (lower) and the complex [Co((btmgn))Cl₂] (upper).

range 0.2209-0.2641. Moreover the aromatic protons are more electropositive than the methyl H-atoms. In contrast, the aromatic and the methyl carbon atoms are electronegative. The C10, C11, C40 and C43 atoms have positive natural atomic charges as these C-sites bonded to the strong electronegative N-atoms. The charge densities at all the N-atoms of the free ligand (btmgn) are negative (-0.4859 to -0.5374). The NAC values at the N4 and N7 atomic sites are calculated to be -0.5373 and -0.5374 respectively indicating the higher basicity of these N-sites compared to the others. It is noted that the NAC values at the N4 (-0.7436) and N7 (-0.7435) of the [Co(btmgn)Cl₂] complex are more negative than the free ligand. The coordination between the Co(II) ion and the ligand (btmgn) via the N4 and N7 coordination centers increases their negative charge values by 0.21 unit. The NAC values at the other N-sites of the free and coordinated btmgn ligand are almost the same. Furthermore, the NAC values of the Co1, Cl2 and Cl3 are calculated to be +1.4314, -0.7942 and -0.7686, respectively. The two coordinated Cl-atoms are not equivalent. We noted that the longer Co-Cl2 bond has more electronegative chlorine atom than the shorter Co-Cl3.

3.5 Molecular electrostatic potential

Molecular electrostatic potential (MEP) is a very useful property for analyzing and predicting molecular reactive behavior. It is very useful three dimensional diagrams used to visualize the charge distributions and charge related properties of molecules as well as in assessing the molecules reactivity towards positively or negatively charged reactants. The MEP is typically

Table 2. The calculated natural atomic charges of the [Co(btmgn)Cl₂] complex and the free ligand (btmgn).

Atom ^a	$[Co(btmgn)Cl_2]$	Btmgn	Atom	$[Co(btmgn)Cl_2] \\$	btmgn
Co1	1.4314		H31	0.2292	0.2165
C12	-0.7942		C32	-0.4833	-0.4763
C13	-0.7686		H33	0.2428	0.2418
N4	-0.7436	-0.5373	H34	0.2600	0.2103
N5	-0.4715	-0.5041	H35	0.2209	0.2249
N6	-0.4373	-0.4860	C36	-0.4996	-0.4734
N7	-0.7435	-0.5374	H37	0.2640	0.2540
N8	-0.4715	-0.5041	H38	0.2271	0.2201
N9	-0.4376	-0.4859	H39	0.2540	0.2156
C10	0.6775	0.6395	C40	0.1764	0.1795
C11	0.6774	0.6396	C41	-0.0780	-0.0787
C12	-0.4833	-0.4763	C42	-0.0378	-0.0421
H13	0.2429	0.2418	C43	0.1763	0.1795
H14	0.2209	0.2249	C44	-0.2779	-0.2937
H15	0.2600	0.2103	H45	0.2354	0.2320
C16	-0.4726	-0.4695	C46	-0.2778	-0.2937
H17	0.2310	0.2251	H47	0.2354	0.2320
H18	0.2320	0.2226	C48	-0.2359	-0.2445
H19	0.2418	0.2379	H49	0.2374	0.2310
C20	-0.4996	-0.4734	C50	-0.2213	-0.2301
H21	0.2539	0.2156	H51	0.2406	0.2347
H22	0.2271	0.2201	C52	-0.2358	-0.2445
H23	0.2641	0.2540	H53	0.2374	0.2310
C24	-0.4725	-0.4695	C54	-0.2213	-0.2301
H25	0.2418	0.2380	H55	0.2406	0.2347
H26	0.2320	0.2226	C56	-0.4813	-0.4780
H27	0.2311	0.2251	H57	0.2485	0.2480
C28	-0.4813	-0.4780	H58	0.2293	0.2166
H29	0.2484	0.2480	H59	0.2294	0.2195
H30	0.2294	0.2195			

^a Atom numbering refers to Figure 4

visualized through mapping its values onto the surface reflecting the molecules boundaries so it allows us to visualize the size and shape of molecules. The MEP diagrams of the free ligand (btmgn) and its [Co(btmgn)Cl₂] complex are shown in figure 5. In this particular, the regions of high electron density appear in red (electrophilic reactivity) while the blue regions are related to electron deficiency (nucleophilic reactivity). The resulting overall MEP of the ligand (btmgn) leaves no doubt that the nitrogen atoms carrying methyl groups (N5, N6, N8 and N9) already are not attracted to a positive test charge such as Co²⁺ while the opposite is true for the N4 and N7 atoms. From the present investigations, we could conclude that the N4 and N7 atoms are the most reactive nucleophilic centers for the coordination with the Co²⁺ ion. Hence, the high basicity of the two guanidinic nitrogen centers (N4 and N7) is evident. Moreover, the MEP picture of the [Co(btmgn)Cl₂] complex indicated that the MEP values around the two Cl-sites

are not equivalent. These results are in good agreement with the calculated NAC values.

3.6 Nonlinear optical properties

The calculated electronic properties such as dipole moment (μ) , polarizability (α_0) , first hyperpolarizability (β_0) , HOMO and LUMO energies as well as their energy gap are given in table 3. These electronic properties have strong relations to the nonlinear optical activity of compound. The present calculations indicated that the [Co(btmgn)Cl₂] has higher dipole moment and polarizability than the free ligand. Also, the calculated first hyperpolarizability (β_0) values of the btmgn and [Co(btmgn)Cl₂] are 457.4870 a.u and 837.3271 a.u, respectively. Moreover, the lower energy gap (ΔE) indicates the more facile electronic transition which is a basic requirement for good nonlinear optical materials. The [Co(btmgn)Cl₂] has lower energy gap

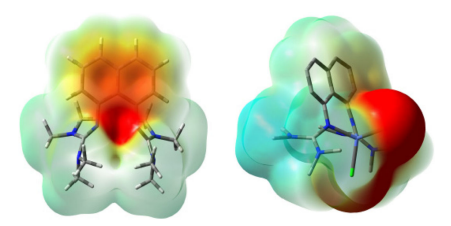


Figure 5. Molecular electrostatic potentials mapped on the electron density surface for free ligand (btmgn) and [Co(btmgn)Cl₂] complex calculated by DFT/B3LYP method. For the [Co(btmgn)Cl₂], the H-atoms are omitted for clarity.

Table 3. The dipole moments components μ (D), the average polarizability α_0 (a.u) and the first hyperpolarizability β_0 (a.u) of the studied compounds.

Parameter	$[Co(btmgn)Cl_2]$	btmgn	Parameter	$[Co(btmgn)Cl_2] \\$	btmgn
$\mu_{ ext{x}}$	-0.004	-0.006	$eta_{ m xyy}$	0.8458	-0.5237
μ_{y}	4.470	-3.950	$eta_{ m yyy}$	-232.7684	-130.1947
μ_{z}	8.683	2.759	$\beta_{ m xxz}$	-1.1379	0.8055
μ_{tot}	9.766	4.818	$eta_{ ext{xyz}}$	914.8886	591.0694
α_{xx}	358.823	327.880	$eta_{ m yyz}$	240.9203	12.5230
α_{xy}	-0.033	-8.937	$\beta_{ ext{xzz}}$	0.1853	0.2940
α_{yy}	356.760	249.173	$eta_{ m yzz}$	15.2282	236.6741
α_{xz}	0.063	26.499	β_{zzz}	-0.0390	0.0335
$\alpha_{ m yz}$	-12.255	63.602	$eta_{ ext{xyy}}$	-74.8864	-125.2531
α_{zz}	232.653	248.899	$eta_{ m yyy}$	320.3787	61.6926
α_0	316.080	275.320	β_{x}	-0.3311	0.3152
E _{HOMO}	-5.018	-4.380	$\stackrel{\sim}{eta_{ m y}}$	607.2338	335.6216
E _{LUMO}	-1.186	-0.334	$\stackrel{r}{eta_{ m z}}$	576.5272	310.8897
ΔΕ	3.832	4.046	β_0	837.3271	457.4870

 (ΔE) than the free ligand. These results indicate that the [Co(btmgn)Cl₂] has better nonlinear optical (NLO) properties than the free organic ligand.

3.7 Electronic spectra

The electronic transition from the ground to the first excited state is mainly described by one electron excitation from the highest occupied molecular orbital (HOMO) to the lowest unoccupied molecular orbital (LUMO). The excitation energy involves the difference between them ($\Delta E=E_{LUMO}-E_{HOMO}$). The HOMO and LUMO plots of the [Co(btmgn)Cl₂] complex as well as the free ligand (btmgn) drawn by DFT–B3LYP method are shown in figure 6. In both molecules the

HOMO and LUMO levels are mainly localized over the ligand π -system. There is significant orbital densities present on the Co^{2+} and the coordinated Cl-atoms in case of [Co(btmgn)Cl₂]. The electronic transition in case of the ligand is mainly π - π * excitation while in the [Co(btmgn)Cl₂] the electronic transition is mixed π - π * and d-d transitions.

The more accurate electronic transitions of the ligand (btmgn) and the complex $[Co(btmgn)Cl_2]$ have been investigated by using the time-dependent density functional theory. Electronic absorption spectra calculated by the TD-DFT are presented in figure 7. For the free ligand, the TD-DFT calculations predicted moderate intensity transition band at 337.2 nm (f=0.2299) which is mainly due to HOMO \rightarrow LUMO (83%) excitation. Of the fifty singlet-singlet electronic excitations calculated

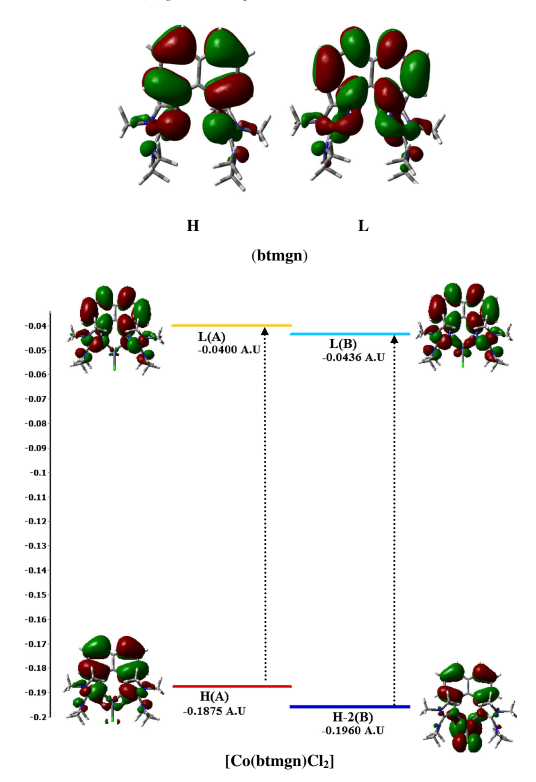


Figure 6. The ground state isodensity surface plots for the frontier molecular orbitals contributed in the electronic transition.

for the [Co(btmgn)Cl₂] complex, a very broad transition band with low intensity (f=0.1465) has been calculated at 342.6 nm (exp. 348.0 nm) due to the HOMO-2 \rightarrow LUMO (53%) and HOMO \rightarrow LUMO (28%) excitations. The rest of the transitions bands have very low intensity (table S3, Supplementary Information).

3.8 Natural bond orbital analysis

The natural bond orbital (NBO) calculations were performed in order to understand various interactions between the filled and vacant NBOs, which are measure of the intramolecular delocalization of electrons

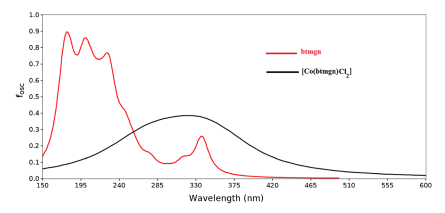


Figure 7. The calculated electronic spectra of the studied compounds using TD-DFT method.

Table 4. The second order perturbation energies $E^{(2)}$ (kcal/mol) of the most important charge transfer interactions (donor–acceptor) of the studied compounds using B3LYP method.

		E ⁽²⁾ kcal/mol	
Donor NBO (i) ^a	Acceptor NBO (j)	btmgn	[Co(btmgn)Cl ₂]
BD(2)N4-C10	BD*(2)C40-C44	7.66	5.38
BD(2)N7-C11	BD*(2)C43-C46	7.67	5.36
BD(2)C40-C44	BD*(2)C41-C42	13.20	
BD(2)C40-C44	BD*(2)C48-C50	20.90	9.35
BD(2)C41-C42	BD*(2)C40- C44	19.87	
BD(2)C41-C42	BD*(2)C43- C46	19.87	
BD(2)C41-C42	BD*(2)C48- C50	15.08	
BD(2)C41-C42	BD*(2)C52- C54	15.08	
BD(2)C43-C46	BD*(2)C41- C42	13.20	
BD(2)C43-C46	BD*(2)C52-C54	20.90	9.35
BD(2)C48-C50	BD*(2)C40-C44	15.56	8.80
BD(2)C48-C50	BD*(2)C41-C42	17.83	
BD(2)C52-C54	BD*(2)C41-C42	17.83	
BD(2)C52-C54	BD*(2)C43-C46	15.56	8.80
LP(1)N4	BD*(1)N5-C10	18.45	5.57
LP(1)N4	BD*(2)C40-C44	13.70	3.17
LP(1)N5	BD*(2)N4-C10	28.00	20.60
LP(1)N6	BD*(2)N4-C10	36.18	29.06
LP(1)N7	BD*(1)N8-C11	18.45	5.57
LP(1)N7	BD*(2)C43-C46	13.70	3.17
LP(1)N8	BD*(2)N7-C11	28.01	20.61
LP(1)N9	BD*(2)N7-C11	36.24	29.03
LP(1)C12	LP*(3)Co1		5.14
LP(4)C12	LP*(3)Co1		29.00
LP(4)C12	LP*(5)Co1		7.73
LP(1)C13	LP*(3)Co1		6.38
LP(4)C13	LP*(3)Co1		39.17
LP(4)C13	LP*(5)Co1		3.97
LP(1)N4	LP*(3)Co1		14.10
LP(1)N4	LP*(4)Co1		8.85
LP(1)N4	LP*(6)Co1		6.29
LP(1)N7	LP*(3)Co1		14.09
LP(1)N7	LP*(4)Co1		9.09
LP(1)N7	LP*(6)Co1		6.18

^{*}Bold for the significant $E^{(2)}$ variations of the $n\to\pi^*$ at N4 and N7. ^a Atom numbering refers to figure 4.

in the studied system. The stabilization energies $(E^{(2)})$ deduced from the NBO calculations for the most significant intramolecular charge transfer interactions are reported in table 4. The stabilization energy $E^{(2)}$ values associated with delocalization of electron density between the occupied and unoccupied NBOs correspond to stabilizing donor-acceptor interactions. These stabilization energies are estimated using the second-order perturbation theory. The delocalization of electrons due to the $\pi - \pi^*$, $n - \pi^*$ and $n - \sigma^*$ interactions are responsible for conjugation of the respective π -bonds leading to a maximum stabilization in the system. It is noted that most of the π - π * ICT interactions in the naphthalene ring do not occur in the [Co(btmgn)Cl₂] complex compared to the free one while the rest of the π - π * ICT interactions are destabilized. Moreover, the changes in the stabilization energies $E^{(2)}$ of the $n-\pi^*$ and $n-\sigma^*$ ICT delocalizations are in the range of 7.12-12.88 kcal/mol. The maximum changes in the E⁽²⁾ values of these interactions occur for the LP(1)N4 \rightarrow BD*(1)N5-C10/BD*(2)C40-C44 and LP(1)N7 \to BD*(2)N7-C11/BD*(2)N7-C11 ICT. It seems that the coordination between the Co(II) and the ligand (btmgn) via the N4 and N7 atoms strongly destabilize the $n-\pi^*$ interactions resulting from LP(1)N4/N7 more than the others.

Interestingly, the NBO calculations predicted the ICT LP(4)Cl2 \rightarrow LP*(3)Co and LP(4)Cl3 \rightarrow LP*(3)Co interactions having E⁽²⁾ of 29.00 and 39.17 kcal/mol, respectively. On the other hand, the LP(1)N \rightarrow LP*Co (6.29–18.85 kcal/mol) and LP(1)N4 \rightarrow LP*Co (6.18–14.09 kcal/mol) have been predicted. It is clear that the,

Table 5. The spin density at the different atomic sites of the [Co(btmgn)Cl₂] complex.

	ungn/Ci2] complex.		
Co1	2.78714	H31	0.00006
C12	0.05262	C32	-0.00015
C13	0.06195	H33	0.00001
N4	0.01957	H34	0.00021
N5	0.00516	H35	0.00044
N6	0.00591	C36	0.00063
N7	0.01958	H37	0.00013
N8	0.00515	H38	0.00084
N9	0.00590	H39	0.00006
C10	0.00354	C40	-0.00508
C11	0.00353	C41	0.01015
C12	-0.00015	C42	-0.00611
H13	0.00001	C43	-0.00509
H14	0.00044	C44	0.01202
H15	0.00021	H45	-0.00056
C16	-0.00034	C46	0.01203
H17	0.00009	H47	-0.00056
H18	0.00026	C48	0.00967
H19	0.00009	H49	-0.00042
C20	0.00063	C50	-0.00570
H21	0.00006	H51	0.00028
H22	0.00084	C52	0.00968
H23	0.00014	H53	-0.00042
C24	-0.00034	C54	-0.00570
H25	0.00009	H55	0.00028
H26	0.00026	C56	-0.00008
H27	0.00009	H57	0.00001
C28	-0.00009	H58	0.00006
H29	0.00001	H59	0.00049
H30	0.00049		

^a Atom numbering refers to figure 4

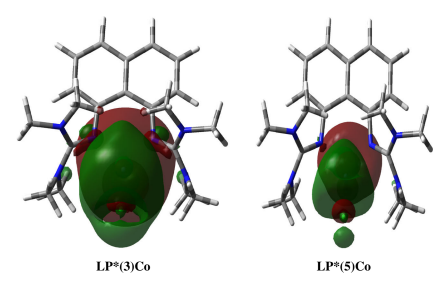


Figure 8. The second order perturbation theory computed donor–acceptor NBO plot for the HS state in complex showing the most significant donor-acceptor charge transfer Co-Ligands interactions.

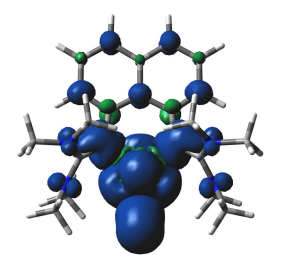


Figure 9. The computed spin density plots for the HS state of complexes; the blue and green for the positive and negative spin densities, respectively.

the two N \rightarrow Co interactions are almost equivalent while the Cl \rightarrow Co ICT interactions are not. These results are in good agreement with our reported structure investigations. The two Co-N bonds are nearly equivalent while the Co-Cl2 is weaker (longer) than the Co-Cl3. To explore the nature of the d-orbitals involved in the most significant donor acceptor interactions, the NBO plots of these interactions are shown in figure 8. The composition analysis of these NBOs showed that, the d_{x2-y2} as well as the d_{z2} are the most important metal ion d-orbitals involved in the donor-acceptor charge transfer interactions.

3.9 Spin density analysis

In order to investigate the mechanism by which the unpaired d-electrons of the studied high spin Co(II) complex can distribute some spin density at the other atoms, the spin densities at the different atoms are calculated and listed in table 5. Also, the spin density plots of the high-spin state of the studied Co(II) complex are shown in figure 9. Generally, we have two possible mechanisms: (1) the molecular orbital that hosts an unpaired electron (metal d orbital) mixes in the atomic orbitals of the ligands (mostly from the donor atoms). As a result, a distribution of positive spin density occurs through out the molecule, hence the resulting distribution of spin density is said to arise from a spin delocalization mechanism; (2) the positive spin at the paramagnetic center may induce some spin density of the opposite sign at the atoms bonded to it which is called the spin polarization mechanism. In this case, a concentration of negative spin density is favored around the ligand donor atoms. In the presented $[Co(btmgn)Cl_2]$, the spin density of the Co(II) ion is found to be 2.78714. All the atoms coordinated to Co(II) are found to have positive spin. This illustrates that the spin delocalization mechanism is operational.

4. Conclusions

The cobalt(II) chloride complex, [Co(btmgn)Cl₂] of 1,8-bis(tetramethylguanidino)naphthalene (btmgn) ligand has been synthesized and characterized. The Xray crystal structure of the [Co(btmgn)Cl₂] is reported. The geometry of the complex as well as the free btmgn ligand have been calculated using DFT B3LYP/6-311G(d,p) method. The calculated geometric parameters showed that the structure of the free ligand is almost the same as in [Co(btmgn)Cl₂] where the two tetramethylguanidino group are in a cis-type position to the plane formed by the naphthalene ring. The [Co(btmgn)Cl₂] has distorted tetrahedral geometry with two equivalent Co-N bonds while the two Co-Cl bonds are not. The calculated natural atomic charges and molecular electrostatic potential (MEP) map showed that the two guanidinic N-atoms (N4 and N7) have the highest basic character and the most favored nucleophilic centers for the Co(II) ion attack. The electronic spectra of the btmgn and [Co(btmgn)Cl₂] have been calculated using the TD-DFT method. The latter showed broad band at 342.6 nm due to the H-2/H→L excitations. The NBO calculations were used to study the different intramolecular charge transfer interactions occur in the studied systems. The FTIR spectra showed strong red shift for the $v_{C=N}$ mode due to the coordination between the Co(II) and the ligand via N4 and N7 atoms. This large red shift upon coordination indicates a strong ligand-metal interaction.

Supplementary Information

The ¹H and ¹³C NMR spectra of the ligand are given in figure S1. The experimental bond distances and angles obtained from the X-ray structure of the complex are shown in table S1. The calculated bond distances and angles are given in table S2. The TD-DFT results of the studied compounds are given in table S3. Supplementary Information is available at www.ias.ac.in/chemsci.

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