Electronic and magnetic properties of the [Ni(salophen)]: An experimental and DFT study

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Introduction

Recently, there has been an increased interest in the chemistry of transition metal complexes containing N2O2 coordination sites,
such as salicylidenedes [1], due to their broad range of applications in areas such as catalysis [2], functional materials, non-linear optics [3], molecular magnetism [4], and organic electronics [5], such as light emitting diodes (OLEDs) [6] and display magnetic properties [7]. Among them, technologies based on light emission or charge transport abilities are currently receiving particular interest, leading to their use in electronic devices, such as solar cells and active components for image and data treatment storage [8]. Salicylidenedes are a type of Schiff base derived from the condensation of a primary amine with a salicylaldehyde derivative. Numerous substituents can be placed on the phenol ring, and the imine bridge allows for tuning the size and the shape of the complexes to control their self-assembly on surfaces [9]. A schematic structure can be observed in Fig. 1.

Their easiness along with the synthesis and modulation of the physical-chemical properties of salicylidenedes make them a versatile and interesting class of molecules [1,10]. Their molecular structure and capable coordinate sites make salicylidenedes flexible coordination compounds with several metal ions, such as Ni(II), Cu(II), Zn(II), Ru(II), Os(II), Pt(II), and Ir(III) [3]. A variety of metal ions (diamagnetic and paramagnetic) can be introduced in the coordination site, in most cases forming square planar array frameworks [6]. Schiff base nickel(II) salen-type complexes have been extensively used in catalysis [7,9] and for biological purposes [11].

Salophen metal complexes are planar systems composed of three aromatic rings [12]. Aromatic molecules are known for their ability to sustain diatropic currents when exposed to an external magnetic field. For instance, when applying a perpendicular magnetic field towards the plane of the aromatic system, a ring current is induced along the delocalized π electrons [13]. The strength and the pathway of the magnetically induced current flow sustained by delocalized electrons in molecular systems play an important role in nanotechnological applications, such as molecular switches or optical devices [13]. The current pathways and the flow along chemical bonds and around molecular rings reflect the electron delocalization in macrocycles, such as porphyrins and fullerenes [14]. Several methodologies are used to calculate magnetically induced current strengths [15]; however, to evaluate the effect of the modification of the central metal atom on the electronic delocalization of the salophen framework, the magnetically induced current method [16] proposed by Sulzer et al. [17] was chosen for this study.

To investigate the potential use of these compounds for optical devices such as solar cells, structural, electronic, and magnetic properties were calculated at the DFT/TD-DFT level, which has been proven useful in evaluating the electronic structure of this type of complex [7,18]. The obtained results were correlated with the experimental measurements.

**Material and methods**

All solvents and reagents were used as purchased from Sigma-Aldrich (São Paulo, São Paulo, Brazil) without further purification. The infrared spectra of KBr pellets of the complex were obtained and measured with a Varian 600-IR spectrometer (Atibaia, São Paulo, Brazil). The TG/DTA curves were obtained in a Shimadzu apparatus (Kyoto, Japan) with a heating rate of 10 °C min⁻¹ using a dynamic atmosphere of synthetic air at a flow rate of 100 mL min⁻¹ until 800 °C. The crystal structure of the salophen ligand has been reported [6]. The electronic absorption spectra of salophen and [Ni(salophen)] in DMSO solutions (1 × 10⁻⁵ mol L⁻¹) were acquired using a Hewlett-Packard 8452A diode array UV–vis spectrophotometer (Santa Clara, California, United States). The steady-state fluorescence spectrum was acquired using an ISS PC1 spectrofluorometer (Champaign, Illinois, United States) of λ exc = 378 nm in a 1.0 cm quartz cuvette (model 111-10-40, type 111-QS, Hellma Analytics, Plainview, New York, United States).

Fluorescence decay was recorded using time-correlated single photon counting and an Edinburgh Analytical Instruments FL 900 spectrofluorimeter (Livingston, Scotland) with an MCP-PMT detector (Hamamatsu R3809U-50). The excitation wavelength for [Ni(salophen)] in the DMSO solution was λ exc = 375 nm (Edinburgh model EPL-375, Livingston, Scotland, with a 10 nm bandwidth, 77.0 ps). The decay signals for this sample were collected at λ em = 420 nm. The instrument response was recorded using Ludox samples. At least 10,000 counts in the peak channel were accumulated for the lifetime measurements. The emission decays were analyzed using exponential functions.

**Synthesis**

The procedure to obtain the ligand (salophen) has been described in detail [6]. [Ni(salophen)] was obtained by dissolving the salophen ligand (158 mg; 0.5 mmol) in ethanol (20 mL) after stirring until total solubility on a round flask balloon. Then, an ethanolic solution of NaOH (40 mg; 1.0 mmol) was slowly dropped into the reaction system. After 5 min, NiSO ₄ (~ 77.5 mg; 0.5 mmol) was added to the mixture, and a [Ni(salophen)] coordination complex instantly formed. As a result, a polycrystalline deep-red powder was obtained with a 67% yield. The main infrared bands measured in the KBr pellet were ν Ni-O = 457, ν Ni-H = 545, ν C-O = 1610, ν C-N = 1197, ν C-H = 3050, and ν Br = 755 (cm⁻¹) (Fig. S1). The TGA experimental weight loss (wt %, in parenthesis calculated values) was: 47.84 (47.76) (280–468 °C) and 32.15 (32.22) (468–510 °C) ligand pyrolysis and residual 20.01 (20.02) – NiO (Fig. S2). X-ray diffraction residual characterization was performed

![Fig. 1. Molecular structures of the salophen and its Ni(II) coordination compound with principal atom labels and the position of the Cartesian axes.](image-url)
according to the X-ray Data Bank files with PDF number 01-071-1179 (Tune Cell) NiO-Bunsenite (Fig. S3).

**Computational details**

The calculations were performed within the density functional theory and its time-dependent counterpart (DFT and TD-DFT). This level of calculation has yielded reliable results in predicting the electronic spectra of chromospheres at a relatively low computational cost, and it is one of the most popular methods used for the evaluation of excitation energies [6,12]. The ground and first active singlet state structures of [Ni(salophen)] were optimized at the PBE0/6-311++G(d,p) [19–21] level of the theory using Gaussian 09 [22]. Vertical excitation energies for 30 low-lying excited states were calculated. To determine the solvent effect (DMSO, $\varepsilon = 46.826$), the polarizable continuum method - PCM approximation was used, defining the cavity unit as a universal forcefield – UFF, and the cavity type scaled the van der Wall surface ($\sigma = 1.10$ 0) [23,24].

The magnetically induced current density maps were evaluated with DIRAC [25] software using the four component relativistic Dirac-Coulomb Hamiltonian [26]. These results were obtained using perturbation-dependent basis sets that shift the gauge origin to their respective center, thereby ensuring that the calculated magnetic properties are independent of the position of the gauge origin [27]. For the large component, triple-$\zeta$ quality Dyall basis sets for the nickel(II) atom was used, while for the light atoms, an uncontracted Dunning cc-pVTZ basis set was chosen [28]. The induced current density streamline plots were visualized using the PyNGL package [29]. The integration plane was chosen to be perpendicular to the molecular plane, beginning from the center of mass and extending to 10 atomic units in all directions. This plane cuts a C–C bond and allows for obtaining the net current intensity around the molecular framework.

**Results and discussion**

**Molecular structures**

The salophen and [Ni(salophen)] ground ($S_0$) and first active excited state ($S_1$) structures were optimized at the PBE0/6-311++G(d,p) level. The research group observed the remarkable quality of [Ni(salophen)]-related molecular structures via crystal refinement obtained by Lerch et al. [32]. For [Zn(salophen)(OH$_2$)], the coordination site bond distances calculated at the PBE0/6-311G++(d,p) level are equal to 2.104 and 1.987 Å for M-N$_{12}$ and M-O$_{10}$, respectively. For [Ni(salophen)] obtained using B3LYP/6-31G(d), these values are 1.860 and 1.842 Å, respectively, which is in agreement with crystal refinement obtained by Lerch et al. [32], who also studied the electronic structure of [Ni(salophen)]-related structures, focusing on one-electron-oxidized Ni(II) salophen complex and its amino derivatives. Optimized structures obtained for [Cu(salophen)] and [Ni(salophen)] at the PBE0/def-TZVP level reported by Zarei et al. [31] showed the bond lengths to be: 1.959 and 1.910 and 1.853 Å for Cu–N$_{12}$, Cu–O$_{10}$, Ni–N$_{12}$, and Ni–O$_{10}$, respectively. Finally, Atakol et al. [33] identified the structural positions of [Ni(salophen)]-related molecular structures via crystal refinement using the DRX technique. The same value for Ni–N$_{12}$ bond length 1.867 Å was also obtained as herein reported.

**Steady-State absorption spectra and calculated electronic transitions**

Salophen and [Ni(salophen)] absorption spectra were measured in a DMSO solution (1 × 10$^{-3}$ mol L$^{-1}$), and the data obtained are presented in Table 1 and Fig. 2. Salophen electronic absorption spectra in a DMSO solution were reported in a previous work [6] to exhibit two absorption bands centered at 270 and 335 nm ($\varepsilon = 2.08$ and 1.48 × 10$^4$ L mol$^{-1}$ cm$^{-1}$, respectively) assigned to $\pi \rightarrow \pi^*$ electronic transitions, mainly involving the frontier orbitals spread over the ligand structure. Due to the increment of the electron delocalization of the macrocycle in the complex, the ligand absorption bands were redshifted to 302 and 378 nm ($\varepsilon = 1.69$ and 2.67 × 10$^4$ L mol$^{-1}$ cm$^{-1}$, respectively). In addition, a band was observed at 478 nm ($\varepsilon = 8.42 \times 10^3$ L mol$^{-1}$ cm$^{-1}$) due to the ligand had a higher symmetry (corresponding to the point group C$_2$) and was more planar than the ground state.

Similar bond lengths were observed in the literature for [M(salophen)] (M = Mn, Ni, Cu, and Zn) and related molecular structures [31,32]. For [Zn(salophen)(OH$_2$)], the coordination site bond distances calculated at the PBE0/6-311G++(d,p) level are equal to 2.104 and 1.987 Å for M-N$_{12}$ and M-O$_{10}$, respectively. For [Ni(salophen)] obtained using B3LYP/6-31G(d), these values are 1.860 and 1.842 Å, respectively, which is in agreement with crystal refinement obtained by Lerch et al. [32], who also studied the electronic structure of [Ni(salophen)]-related structures, focusing on one-electron-oxidized Ni(II) salophen complex and its amino derivatives. Optimized structures obtained for [Cu(salophen)] and [Ni(salophen)] at the PBE0/def-TZVP level reported by Zarei et al. [31] showed the bond lengths to be: 1.959 and 1.910 and 1.853 Å for Cu–N$_{12}$, Cu–O$_{10}$, Ni–N$_{12}$, and Ni–O$_{10}$, respectively. Finally, Atakol et al. [33] identified the structural positions of [Ni(salophen)]-related molecular structures via crystal refinement using the DRX technique. The same value for Ni–N$_{12}$ bond length 1.867 Å was also obtained as herein reported.

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**Table 1**

Excitation energies experimentally obtained and calculated at the PBE0/6-311++G(d,p) level for [Ni(salophen)] using DMSO as solvent.

| $\lambda_{\text{ex}}$ [nm] | $E$ [eV] | $\lambda$ [nm] | $f$ | Assignment$^a$ |
|-------------------------|--------|----------------|----|----------------|
| Salophen                | 370    | 3.28           | 378 | 0.46           | 100% H $\rightarrow$ L |
|                         | 335    | 3.71           | 334 | 0.80           | 60% H-1 $\rightarrow$ L; 40% H $\rightarrow$ L+1 |
| [Ni(salophen)]          | 478    | 2.80           | 442 | 0.28           | 100% H-1 $\rightarrow$ L |
|                         | 378    | 3.36           | 369 | 0.55           | 100% H-1 $\rightarrow$ L+1 |
|                         | 302    | 3.61           | 344 | 0.69           | 100% H-2 $\rightarrow$ L |

$^a$ H = HOMO and L = LUMO.
contribution of the atomic orbitals $d$ of the Ni(II) ion to the frontier molecular orbitals involved in the electronic transitions.

To obtain more information regarding the nature of these excitations, theoretical calculations were performed using a PBE0/6-311++G(d,p) basis set and DMSO as a solvent according to the PCM approach. As can be seen in the energy diagram given in Fig. 3, frontier molecular orbitals are degenerate; hence, all electronic transitions are mainly located in the ligand of a $\pi \rightarrow \pi^*$ type, which involves molecular orbitals mainly located over the ligand framework. There is also a contribution of the metal atom to the complex transitions, resulting in a partial metal-to-ligand charge transfer (${}^1\text{ILCT} / {}^1\text{MLCT}$), favoring the destabilization of the frontier molecular orbitals and resulting in a redshift of the absorption bands respective to the free basis ligand. Despite the larger deviation of the excitation energies of the complex from the experimental data, the results are still within the expected accuracy of TDDFT using hybrid density functionals of around 0.3 eV [37]. It has been pointed out that TDDFT also yields substantial errors for excited states of molecules with extended $\pi$-systems [38,39] as well for charge-transfer (CT) states [40,41], as observed for the complex [Ni(salophen)]. Lecarme et al. [32] also observed a similar deviation for [Ni(salophen)], where CT could be observed. According to Jacquemin et al. [37], a deviation in a TDDFT calculation can be related to a long-range charge transfer, a potential energy surface, non-Franck-Condon transitions, or a singlet-triplet transition. Although a small deviation of the excitation energies of the complex from the experimental data was observed, diffuse orbitals base set (mandatory for CT states) and a global hybrid functional PBE0 were responsible for decreasing the deviation. Despite these failures of TDDFT, it has been applied to large molecular systems [42] in which inter- or intra-molecular CT states might play important roles.

Photoluminescence spectra

Fig. 4 presents the steady-state photoluminescence (PL) spectrum of [Ni(salophen)] obtained in DMSO ($\lambda_{\text{exc}} = 378$ nm; $1 \times 10^{-5}$ mol L$^{-1}$) with only one emission band at the blue-region centered at $\lambda_{\text{PL}} = 420$ nm with a Stokes Shift respective to the excitation wavelength of $SS = 2646$ cm$^{-1}$.

As observed by Nijegorodov et al. [1], for a series of planar and non-planar molecules, due to the increment of the rigidity of the ligand after the coordination of the metal atom, the Stokes shift for [Ni(salophen)] is significantly lower (~6000 cm$^{-1}$) than the free basis ligand reported in Ref. [5]. Hence, the complex emission band appears at a lower wavelength than its free basis ligand. Its fluorescence decay ($\lambda_{\text{exc}} = 375$ nm; $\lambda_{\text{PL}} = 420$ nm) was also measured (Fig. 5). A biexponential decay profile was observed, presenting two fluorescence lifetimes: a shorter lifetime of $T_1 = 0.815 \pm 0.025$ ns (41%) and a longer lifetime of $T_2 = 1.958 \pm 0.037$ ns (59%). Thus, the two fluorescence lifetimes were attributed to the same chromophore group but with different solvent environments.

According to Atvars et al. [6], a faster decay indicates that the metal disturbs the electronic excited states of the ligand, which

![Fig. 3. Frontier molecular orbital energy diagram for salophen and [Ni(salophen)] obtained at PBE0/6-311++G(d,p).](image)

![Fig. 4. Steady-state photoluminescence spectrum of the salophen ligand (blue) and [Ni(salophen)] coordination compound (red) measured in DMSO (1 \times 10^{-5} \text{ mol L}^{-1}).](image)
is in agreement with the observations of Chavan et al. [34]. For related complexes, the emission observed for [Ni(salophen)] predominantly originated not only due to the \( \pi \rightarrow \pi^* \) intra ligand transition but also due to specific MLCT characteristics. This behavior was also observed for Zn(II) salicylidenes in solutions and solid states [35] but with different contributions of the fluorescence lifetimes. Its Ni(II) complex does not present phosphorescence emissions at room temperature in a DMSO solution.

Magnetically induced currents

To evaluate the impact of the nickel(II) atom coordination modification on the electronic delocalization of the salophen ligand, magnetically induced currents were calculated at the PBE0/cc-pVTZ (dyall.v3z for the nickel atom) level, and the results are shown in Fig. 6. According to the chosen methodology, the total probability current density can be separated into paratropic (counter-clockwise) and diatropic (clockwise) components when a magnetic field is perpendicularly directed to the plane of the aromatic system. Located at 1 Å over the molecular plane to mainly consider the contribution from the \( \pi \) orbitals of the aromatic ring, a diatropic probability current can be observed outside the carbon atoms of the ligand framework, and an opposite paratropic current inside the carbon rings is visible for both molecules. It was observed that for both salophen and the [Ni(salophen)] complex, the diatropic ring current \( \pi \) system dominates the streamline plot.

A quantitative analysis of the strength of the magnetically induced ring was performed using the numerical integration of the current density passing a C–C bond from the phenolic ring perpendicular to the XZ plane, as shown in Fig. 1. The total integrated ring current susceptibilities along with their paramagnetic and diamagnetic contributions are presented in Table 2.

According to these results, both molecules have a net ring current of the same order as benzene (\( \sim 12 \text{nA T}^{-1} \)) [36]. Also, Sundholm et al. [43] studied magnetically induced current density along Zn(II)-octaethylporphyrin. According to the authors, at the pyrrole rings, the magnetically induced current values were the same order (by \( \sim 11.9 \text{nA T}^{-1} \)). Although there was a larger value of the total integrated magnetically induced current values for the salophen and [Ni(salophen)] complex, the diamagnetic current was stronger (by \( \sim 10 \text{nA T}^{-1} \)) for the complex than for the ligand due to the planarization of the ligand framework caused by the coordination of the nickel(II) ion. These findings support the changes observed in the ligand absorption spectra that occur after metal coordination.

Conclusions

In this article, the electronic and magnetic properties of [Ni(salophen)] and the effect of the nickel(II) coordination on the ligand characteristics were theoretically and experimentally evaluated. The spectral data obtained was measured in DMSO and showed a
red shift of the ligand absorption bands, mainly composed by $\pi \rightarrow \pi^*$ electronic transitions, after the coordination of the nickel(II) ion. In addition, there was a contribution of d metal orbitals to the complex transitions, resulting in a partial metal-to-ligand charge transfer, which caused the appearance of a low-lying absorption band of around 470 nm. Furthermore, a significant increment of its band intensities was observed, favoring absorption in a broader range of the visible spectrum, a desired characteristic for applications in organic electronics, such as solar cells. This finding is related to the increment of the planarity and the consequent electron delocalization of the macrocycle in the complex, which was estimated using the calculations of the current strengths.

Conflict of interest

The authors have declared no conflict of interest.

Compliance with Ethics Requirements

This article does not contain any studies with human or animal subjects.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found, in the online version, at https://doi.org/10.1016/j.jare.2017.10.004.

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