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# Azobased iminopyridine ligands and their rhenium metal complexes: syntheses, spectroscopic, trans-cis photoisomerization and theoretical studies

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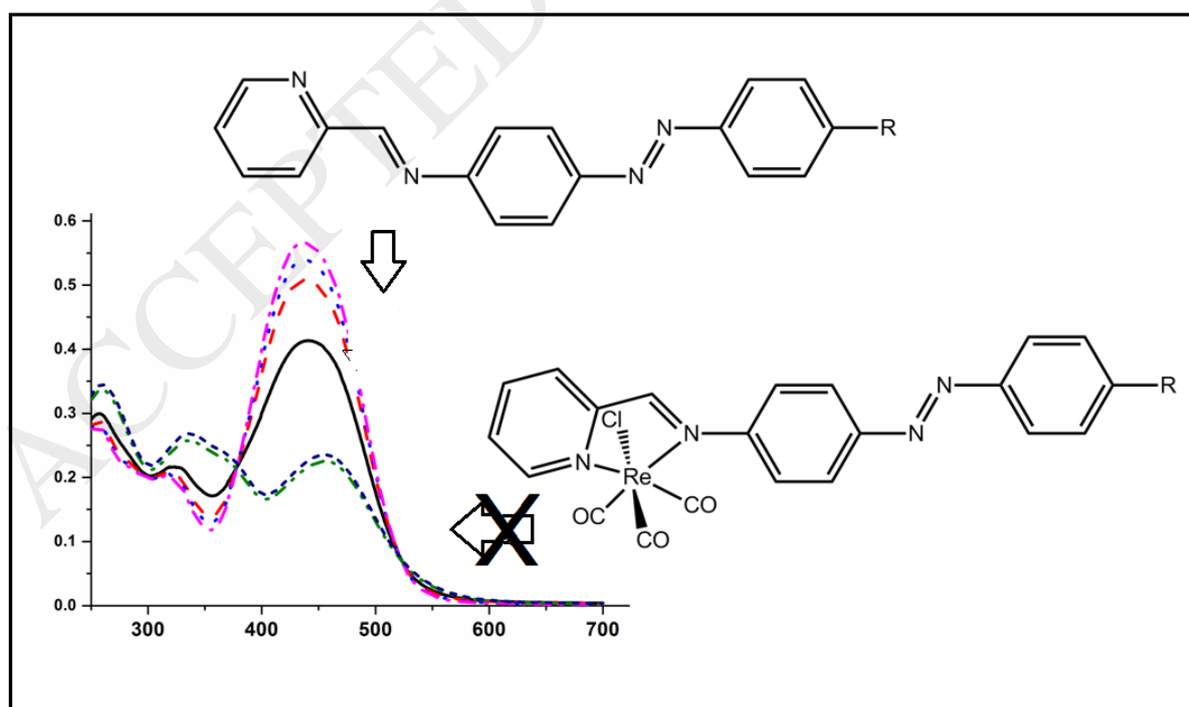
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## Graphical Abstract



## Highlights

- Two rhenium metal complexes have been prepared and structurally characterized
- Absorption spectra have been measured
- DFT calculations have been performed to support the experimental results
- Photoisomerization studies of the azo group in the ligands and in the metal complexes have been performed.

## Abstract

The reaction of rhenium(I) pentacarbonyl chloride  $\text{Re}(\text{CO})_5\text{Cl}$  with N,N-dimethyl-4-((E)-(pyridin-2-ylmethylene) amino)phenyl diazenyl) aniline **L1** and (E)-4-((E)-(4-nitrophenyl) diazenyl)-N-(pyridin-2-ylmethylene) aniline **L2** affords two rhenium metal complexes **[ReL1(CO)<sub>3</sub>Cl]** (**ReL1**) and **[ReL2(CO)<sub>3</sub>Cl]** (**ReL2**), respectively. These Re(I) complexes were characterized in detail, including their single crystal structures, absorption spectra and electronic structures using DFT calculations. *Trans* to *cis* photoisomerization of the free ligands and complexes was investigated experimentally and rationalized theoretically. Interestingly, light-induced photoisomerization of the azo fragment is observed for free ligand **L1** while inhibition of this process is produced upon coordination to Re metal cation.

**Keywords:** Iminopyridine; Azobenzene; UV-Visible spectroscopy; Rhenium complex; X-ray diffraction; DFT.

## 1. Introduction

Increasing interest is currently devoted to the photochemistry of transition metal derivatives incorporating stimuli-responsive functions for their potential applications in photonic memory, photosensing, as well as luminescence imaging [1]. In particular, photochromic metal complexes allow the specific properties of the metal center and the optical response of the photochromic group to be combined. Of course, the photoinduced isomerization of azobenzene and its derivatives is one of the most and well-studied light driven processes. The location of the azobenzene unit in the ligands has

been established as an important factor to influence the photochromic properties of azobenzene containing transition metal complexes [2-5]. The isomerization is assumed to take place from intraligand excited states localized on the azobenzene unit. Meanwhile, the choice of the metal center also participates to the photochromic properties of these azobenzene containing complexes [6]. It would be of great interest to better identify the factors which favor an efficient synergy between the intrinsic properties of the metal complex and that of the photochromic unit.

Our interest recently focused on the possible effects of coordination of iminopyridine azo-based ligands to heavy transition metals on the photochemistry of these ligands. For instance, it has been reported that organoplatinum complexes containing iminopyridine azo-based ligands can undergo *trans-cis* isomerization of the azo group [7]. In this context, we were interested in the photochemistry of Re(I) polypyridyl complexes having iminopyridine-appended azobenzene ligands. Rhenium(I) tricarbonyl polypyridine complexes generally possess attractive photophysical and photochemical properties such as large Stokes shifts, long emission lifetimes, and high photostability through the coordination of functionalized ligands. In the present paper, we aim to report on the synthesis and full characterization, including X-ray diffraction analysis, of two rhenium metal complexes. The focus of this work is on the *trans-cis* photoisomerization behavior of the free ligands, **L1** and **L2**, (scheme 1) [8] and of the corresponding complexes **ReL1** and **ReL2**. These studies are supported by DFT and TD-DFT computational studies that reveal the importance of relevant barriers on the potential energy surfaces in the *S1* excited states, which involve conical intersection, for rationalizing the observed behaviours.

## 2. Experimental

### 2.1. General remarks

All manipulations were performed under nitrogen atmosphere using commercial grade solvents. Rhenium(I) pentacarbonyl chloride was of reagent grade quality and used as received. NMR spectra were recorded on a Bruker Avance DRX 300 spectrometer operating at 300 MHz for  $^1\text{H}$  and 75 MHz for  $^{13}\text{C}$ . Chemical shifts are expressed in parts per million (ppm) downfield from external TMS. The following abbreviations are used: s, singlet; d, doublet; t, triplet; td, triplet of doublet. UV-visible spectra were recorded at room temperature in quartz cuvettes using Perkin-Elmer spectrophotometer. Mass spectra were collected with Bruker Biflex-III TM. IR spectra were recorded on a Bruker vertex 70.

### 2.2. Crystallography

X-ray single-crystal diffraction data for both complexes were collected on an Agilent SuperNova diffractometer equipped with Atlas CCD detector and mirror monochromated micro-focus Cu  $\text{K}\alpha$  radiation ( $\lambda = 1.54184 \text{ \AA}$ ). The two structures were solved by direct methods, expanded and refined on  $F^2$  by full matrix least-squares techniques using SHELX97 programs (G.M. Sheldrick, 1998).

All non-H atoms were refined anisotropically and the H atoms were included in the calculation without refinement. Multiscan empirical absorption was corrected using the CrysAlisPro program (CrysAlisPro, Agilent Technologies, V1.171.37.35g, 2014). Crystallographic data for the structural analysis have been deposited with the Cambridge Crystallographic Data Centre, CCDC 1837472 for complex **ReL1** and CCDC 1837473 complex **ReL2**. Details about data collection and solution refinement are given in Table S1.

### 2.3. Synthesis of **ReL1** and **ReL2** metal complexes

#### 2.3.1. Synthesis of Rhenium(I) complex [ReL1(CO)<sub>3</sub>Cl] (**ReL1**)

To a solution of **L1** (0.01 g, 0.03 mmol) in a solution of toluene and dichloromethane (3:1, v/v) was added [Re(CO)<sub>5</sub>Cl] (0.016 g, 0.045 mmol). The mixture was refluxed for 6 hours under nitrogen atmosphere. After cooling the resulting mixture to room temperature, the solvent was removed by a rotary evaporator. The dark residue was extracted with dichloromethane and recrystallized from acetone/hexane to yield complex **ReL1** as brown crystals with 90% yield (0.017 g, 0.027 mmol); mp 245 °C; <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ/ppm: 9.14 (d, 1H, J = 4.89 Hz), 8.89 (s, 1H), 8.16 (td, 1H, J = 7.83 Hz, J = 1.39 Hz), 8.06 (m, 1H), 8.02 (d, 2H, J = 8.7 Hz), 7.96 (d, 2H, J = 8.83 Hz), 7.66 (m, 3H), 6.82 (d, 2H, J = 9.04 Hz), 3.17 (s, 6H). <sup>13</sup>C NMR (75 MHz, CDCl<sub>3</sub>) δ/ppm: 165.6, 155.7, 155.3, 154.9, 153.5, 153.3, 152.9, 151.9, 151.8, 139.3, 132.2, 129.1, 129, 126.3, 126.2, 124.1, 123.2, 123.1, 122.8, 111.9, 111.7, 40.5, 40.4. Selected IR bands (cm<sup>-1</sup>): 2017 (CO), 1897 (CO), 1860 (CO), 1595, 1519, 1357, 1125, 819, 788, 554, 495. Anal. calcd for C<sub>23</sub>H<sub>19</sub>N<sub>5</sub>O<sub>3</sub>ClRe: C, 43.50; H, 3.02; N, 11.03, found: C, 43.31; H, 2.87; N, 10.79. MALDI-TOF MS calcd: m/z = 635.0 Da. found: m/z = 635.3; HR-MS (M): calcd for C<sub>23</sub>H<sub>19</sub>N<sub>5</sub>O<sub>3</sub>ClRe: 635.0734; found: 635.0736.

#### 2.3.2. Synthesis of Rhenium(I) complex [ReL2(CO)<sub>3</sub>Cl] (**ReL2**)

This complex was prepared by following the same procedure as for complex **1**. Yield: 88% (0.017 g, 0.027 mmol); mp 205 °C; <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ/ppm: 9.06 (d, 1H, J = 4.98 Hz), 8.83 (s, 1H), 8.34 (d, 2H, J = 8.83 Hz), 8.03 (m, 6H), 7.64 (m, 3H). <sup>13</sup>C NMR (75 MHz, CDCl<sub>3</sub>) δ/ppm: 166.6, 155.4, 155.1, 153.5, 153.0, 152.3, 149.1, 139.4, 129.5, 129.4, 124.9, 124.84, 123.7, 123.3. Selected IR bands (cm<sup>-1</sup>): 2014 (CO), 1888 (CO), 1879(CO), 1578, 1519, 1342, 1102, 859, 781, 553, 450. Anal. calcd for C<sub>21</sub>H<sub>13</sub>N<sub>5</sub>O<sub>5</sub>ClRe: C, 39.59; H, 2.06; N, 10.99, found: C, 39.28; H, 1.99; N, 10.83. MALDI-TOF MS calcd: m/z = 637.0 Da. found: m/z = 637.1; HR-MS (M): calcd for C<sub>21</sub>H<sub>13</sub>N<sub>5</sub>O<sub>5</sub>ClRe: 637.0163; found: 637.0161.

## 3. Results and discussion

### 3.1 Syntheses

The equimolar reaction between **L1** or **L2** and the [Re(CO)<sub>5</sub>Cl] precursor at refluxing toluene/dichloromethane mixture, under inert atmosphere, afforded the mononuclear neutral

rhenium(I) metal complexes **ReL1** and **ReL2** (scheme 1), described as **[ReL1(CO)<sub>3</sub>Cl]** and **[ReL2(CO)<sub>3</sub>Cl]** respectively as a brown precipitate for **ReL1** and as an orange precipitate for **ReL2**.

### 3.2. Description of crystal structures

Single crystals of **ReL1** and **ReL2** metal complexes were obtained by recrystallization from acetone/hexane solutions. Crystal structures are represented in Fig. 1. Both **ReL1** and **ReL2** complexes crystallize in the triclinic space group P-1 with one independent molecule in the unit cell.

Both ligands **L1** and **L2** acquired a *cis*-conformation of the iminopyridyl group, within the complexes. The rhenium center is surrounded by the bidentate iminopyridyl chelating fragment, three carbonyl groups, and a chlorine atom and its coordination sphere presents the expected, although slightly distorted, octahedral geometry. This minor distortion from the ideal octahedral geometry is caused by the formation of a five-membered chelate ring (the N–Re–N angle) [9]. The structures of both complexes highlight the expected facial arrangement of the three carbonyl ligands. For both complexes, the values corresponding to *cis* angles around the Re(I) ion are very close to the ideal value of 90°. Meanwhile, the N–Re–N angle shows significantly lower values, 74.33(2)° and 74.69(1)° for **ReL1** and **ReL2** respectively, as imposed by the small N–N bite distance of a  $\alpha$ -iminopyridyl type ligand (average values for **ReL1** and **ReL2** respectively). The dihedral angle between the iminopyridyl fragment and the adjacent phenyl ring of the azobenzene unit is equal to 32.80°(3) for **ReL1** and to 42.67°(3) for **ReL2**.

Within the crystal structure of **ReL1** the Re–N distances (Re–N4 = 2.206(5) Å, Re–N5 = 2.177(5) Å) are similar to those found in complexes of the {ReX(CO)<sub>3</sub>} fragment with the iminopyridyl based coordinating group (Table S2 and Figure S1 in supplementary information) [10–12]. All three Re–CO bond lengths are very close, and the Re–C–O angles present minor deviations from linear structure, values ranging from (177.50(9)° to 177.78(8)°).

In the crystal structure of **ReL2** complex, the ligands surrounding the Re(I) ion are also involved in hydrogen bonding interactions type. Thus, two of the three CO ligands establish short C–H $\cdots$ O contacts (H11 $\cdots$ O1 2.600(6) Å; H15 $\cdots$ O2 2.550(5) Å) with hydrogen atoms from pyridine linking neighbor molecules into layers parallel with the *ac* crystallographic plane (Fig. S2).

### 3.3. UV-Visible absorption spectroscopy

The UV-visible absorption spectra (Fig. 3 and Fig. 4) of the two ligands **L1** and **L2** and their corresponding metal complexes **ReL1** and **ReL2** were recorded in dichloromethane solution ( $\sim C = 5.9 \cdot 10^{-5}$  M) at room temperature. As previously reported [8], ligand **L1** exhibits two strong absorption bands at  $\lambda = 266$  nm and 438 nm that are assigned to the  $\pi \rightarrow \pi^*$  and  $n \rightarrow \pi^*$  ( $^1$ ILCT) transitions together with a slight intramolecular charge transfer (ICT) from the donating dimethylamino donor to the

accepting iminopyridine unit. The spectrum of **ReL1** complex shows the same characteristics as ligand **L1** with a bathochromic shift and a broadening of the visible absorption band. The bathochromic shift (ca. 40 nm) indicates an increase of the electron acceptor character of the iminopyridyl fragment upon complexation with rhenium (I) which acts as a strong Lewis acid while the broadening of the band can be assigned to the presence of a metal to ligand charge transfer  $\text{Re}(d\pi) \rightarrow \text{ligand}(\pi^*)$   $^1\text{MLCT}$  transition [13]. We assume that, in **ReL1**, the two absorption bands corresponding to the  $^1\text{ILCT}$  and  $^1\text{MLCT}$  are superimposed. These features are confirmed by means of the TD-DFT computations which have been carried out (*vide infra*).

In the case of ligand **L2** [8] a strong absorption band around 380 nm assigned to  $\pi\text{-}\pi^*$  and  $n\text{-}\pi^*$  transitions is observed. In the case of **ReL2**, because of the presence of a nitro electron withdrawing group in **L2**, the absorption band in the visible region is blue-shifted, by comparison with **L2**, while a broad absorption band, of lower intensity ( $\epsilon = 4500 \text{ L}\cdot\text{mol}^{-1}\cdot\text{cm}^{-1}$ ) is observed in the visible region around  $\lambda_{\text{max}} = 452 \text{ nm}$  which is characteristic of a metal to ligand charge transfer ( $\text{Re}(d\pi) \rightarrow \text{ligand}(\pi^*)$   $^1\text{MLCT}$  transition [13]. Note that no significant emission behavior has been observed for the two rhenium complexes.

### 3.4. Photoisomerization studies

It is well known that azobenzenes experience a *trans-cis* isomerization by irradiation at a light wavelength close to their absorption maxima [14]. **L1**, **L2** (that can be considered as being push-pull azobenzenes (ppAB) types [13]) and their corresponding rhenium complexes were systematically studied in order to establish if *trans-cis* isomerization occurs in these systems.

Fig. 2 exhibits the UV-visible absorption spectra of **L1** in dichloromethane, showing drastic modifications of the UV-visible absorption spectra upon irradiation at 450 nm revealing the photoisomerization of the azobenzene chromophore in **L1**. Upon irradiation the intensity of the absorbance at 440 nm, attributed to the  $\pi\text{-}\pi^*$  transition of the *trans*-azobenzene group, decreased gradually, reaching a relatively stable value at a maximum wavelength of 460 nm. At the same time, the intensity of a new signal in the 300-400 nm range, which we assign to the  $\pi\text{-}\pi^*$  transition of the *cis* isomer, increased, leading to two isobestic points at 380 and 530 nm for **L1**. These variations are consistent with the *trans-cis* isomerization of the azobenzene group in **L1**, as described for similar compounds [15], and at the photostationary state, the intensity of the main absorption band collapsed by a factor approximately 3. Indeed, the TD-DFT simulations of the UV-visible spectra of the *trans* and *cis* isomers of **L1** (Fig. S3), which compare very well with the observed ones, confirms the occurrence of the *trans-cis* isomerization under irradiation.

In the case of ligand **L2** we did not observe any photoisomerization, whatever is the solvent (dichloromethane or acetonitrile).

For complexes **ReL1** and **ReL2** we did not observe any change of the absorption spectra in acetonitrile or in dichloromethane under irradiation or in the dark. The *trans*-to-*cis* photoisomerization property is completely switched off for the complex **ReL1**, compared to the ligand **L1**. Clearly, the photoisomerization dynamics is strongly influenced by the conjugation pathway and modified upon coordination of a metal to the azobenzene containing conjugated ligand, as previously demonstrated by Fillaut and coworkers for a series of Ruthenium(II) tris(bipyridine) complexes [16]. More recently Freixa and coworkers reported Iridium(III) cyclometalated complexes, in which the use of an aliphatic spacer unit between the azobenzene and the coordinating unit of the ligand allowed the electronic communication to be disrupted and the observation of photochromic properties for these coordination complexes [2].

### 3.5. Simulated UV-Visible absorption spectra and theoretical calculations

All calculations were performed at the DFT level of theory, employing the G09 suite of programs [17]. Based on the results of several previous studies on similar systems, the B3LYP [18-20] and CAM-B3LYP functionals [21,22] have been chosen to perform the computations. The 6-31G(d) and LanL2DZ basis set [23] augmented with polarization function on all atoms, except hydrogen ones, have been used for the ligands and their corresponding complexes respectively. Solvent effects were taken into account using the polarisable continuum model (PCM) [24]. All ground state geometries were optimized at these levels of theory and checked to be true minima on the potential energy surface, the computed frequencies of the normal modes of vibration being all real. Then, time-dependent density functional theory (TD-DFT) [25] computations were performed, using the optimized ground state geometries, to obtain excitation energies and simulate the UV-visible spectra of the considered species.

The simulated UV-visible spectra of the considered ligands and complexes agree very well with the observed ones, when using the CAM-B3LYP functional, so that our discussion will be based on the CAM-B3LYP results only. This reliability of the CAM-B3LYP functional for the UV-visible spectra simulation has already been observed in the case of the ligands [8].

First, we observe that the band shifts observed for the UV-visible spectra upon complexation are well reproduced by the computations, namely the bathochromic one in the case of the complexation of ligand **L1** and the hypsochromic one in the case of ligand **L2** (Fig. 3 and Fig. 4).

The results of the TD-DFT computations which permit the assignment of the UV-visible absorption bands of the ligands are given in Table 1.

<sup>a</sup>H = HOMO; L = LUMO

The assignments of the observed UV-visible bands of the free **L1** and **L2** ligands have been described in reference [8]. To consider the photoisomerization issue, the properties of the free **L1** and **L2** ligands are compared to those of their corresponding complexes, reminding that ligand **L1** only undergoes the *trans* to *cis* isomerization. The first excited states of both **L1** and **L2** derive from transitions involving the diazo group, thus likely to trigger the *trans-cis* photoisomerization of the azobenzene group. Indeed, HOMO-2 of **L1** and HOMO-1 of **L2**, exhibit a diazo lone pair character, whereas the reached MOs upon excitation, namely LUMO and LUMO+1 for **L1**, LUMO and LUMO+2 for **L2** are  $\pi$  type MOs (Fig. 5), so that the computed excitations at 431 nm for **L1** and 468 nm for **L2** exhibit mainly a  $n-\pi^*$  character. However, the oscillator strength of the electronic transition is *ca.* 0.0050 for **L1** and *ca.* 0.0001 for **L2** which is very low, fifty times smaller than that for **L1**. This is certainly a factor contributing to the fact that ligand **L2** does not exhibit a photoisomerization, contrarily to ligand **L1**. It is worth noting that in our previous TD-DFT study of the ligands [8] we were interested in the UV-visible spectra only so that the electronic transitions of low oscillator strength were not considered. These latter transitions, *i.e.* HOMO-2 to LUMO for **L1** and HOMO-1 to LUMO for **L2** are of importance (*vide supra*) for the isomerization processes, explaining why they are considered now.

The assignment of the UV-visible absorption bands of the **ReL1** and **ReL2** complexes has also been done on the basis of the TD-DFT results given in Table 2 and Table 3 whereas the frontier MO diagrams of these species are displayed on Fig. 6.

For the **ReL1** complex, the computed excitation at 431 nm exhibits an  $n-\pi^*$  character, the involved occupied molecular orbital HOMO-4 is localized on the diazo function (Table 2). This excitation is similar to that computed for the ligand **L1**, but with a lower oscillator strength. However, it appears that this complex does not isomerize under irradiation contrarily to ligand **L1**. Thus, for **ReL2**, we observe the same situation as for the ligand **L2** alone; the computed excitation at 467 nm exhibits an  $n-\pi^*$  character with a very low oscillator strength of *ca.* 0.0007, the involved occupied HOMO-4 being localized on the azo function (Table 2). Similarly to the **L2** ligand, the relevant excited state will not be easily reached and then the photoisomerization should hardly be possible.

Another interesting difference appears between the UV-visible spectra of **ReL1** and **ReL2**, when considering the most intense absorption band computed at 419 nm for **ReL1** and at 354 nm for **ReL2**. Contrarily to **ReL1**, the absorption band of **ReL2** exhibits a significant MLCT character, since the weight of the metal in the involved occupied MOs of the corresponding transitions (Table 4) namely HOMO

of **ReL1**, is *ca.* 0.15% whereas for the HOMO-2 of **ReL2**, this weight is *ca.* 26%. In the case of **ReL1** the two excitations at 395 and 374 nm with a smaller oscillator strength exhibit also a MLCT character, the metal weights in the corresponding occupied MOs being equal to *ca.* 50% and 48% respectively. The low intensity bands computed at 402 nm and 380 nm of **ReL2** exhibit also an important MLCT character. Finally, it is interesting to note that whereas the HOMO of **ReL2** has an important metallic character (*ca.* 49%) it is not the case for **ReL1**, mainly due to the strong donating character of the NMe<sub>2</sub> group of the **L1** ligand which contributes strongly to push the ligand HOMO to higher energies than the metallic MO.

Considering in more details the photoisomerization issue, the ligand and the complexes behaviors have been investigated. The photoisomerization of azobenzene in solution has already been studied experimentally and by means of theoretical calculations [26-28]. It has been found theoretically that the isomerization from the *trans* conformation to the *cis* one after the n- $\pi^*$  vertical excitation, occurs on the S1 potential energy surface (PES), in three steps. The first two steps correspond to a relaxation to a local minimum, then to an intermediate state through an energy barrier. The third step corresponds to a S1  $\rightarrow$  S0 relaxation via a conical intersection (CI) S1 / S0 with an energy barrier of 12 kJ / mol, towards the *trans* to *cis* isomerization [26].

Note that in the singlet ground state (S0) the ligands are more stable in a *trans* conformation. The energy barrier between the two *trans* and *cis* forms is high in the ground state (S0) up to about 50 kcal/mol. Interestingly, we found that the *trans* structure undergoes a distortion in the S1 state leading to a non-planar structure of the phenyl azo moiety. Therefore, we determined for the two ligands the minima on the S1 PES, then we identified for each ligand intermediate (I-*cis* and I-*trans*) and transition states on this S1 PES using the QST2 method, in order to estimate the energy barriers between the *cis* forms and the TSs. Moreover, as expected [26-28] we observed the occurrence of a conical intersection, namely that the minimum energy structure on the S1 PES corresponds to a TS on the ground state S0 PES.

According to Fig. 7, we note that the intermediate I-*cis* (S1) state of **L2**, although lower in energy than the TS(S1) is higher in energy than the min(S1)=TS(S0) so that the relaxation of this intermediate via the S1/S0 CI leads to the *trans* conformation, whereas for **L1** (Fig. 7), the intermediate I-*cis* (S1) state is lower in energy than the min(S1)=TS(S0); then, the relaxation of this intermediate leading to a *cis* conformation is facilitated. There is also another difference between **L1** and **L2**, which is the *cis* to *trans* backward barrier on S1; as shown in Fig. 7 and Fig. 8, the barrier in the case of **L2** is *ca.* 5.8 kcal/mol which is lower than that for **L1** (23.8

kcal/mol) so that the latter ligand cannot undergo the reverse *cis-trans* conversion. Thus, either on the S1 PES or via the conical intersection, the *trans-cis* isomerization appears to be highly favoured for the **L1** ligand contrarily to the **L2** one, thus explaining the observed facts.

We have also explored the PES for the two **ReL1** and **ReL2** complexes; these complexes are more stable in the singlet ground state (S0) in a *trans* conformation, whereas in the S1 PES this conformation is distorted as already observed for the ligands. In the same way as for the ligands, we have located for each complex the transition state (TS), and then the energy barrier (*cis*-TS). We found that these complexes behave as ligand **L2**. Indeed, the backward *cis* to *trans* barrier is low, *ca.* 4 kcal/mol for **ReL1** and is *ca.* 6 kcal/mol for **ReL2**, thus of the same order of magnitude as **L2** (5.8 kcal/mol). We think that this low energy barrier can explain the non-photoisomerization for the two complexes, since it is likely to allow the return from *cis*-conformation to *trans*-conformation.

We have also investigated theoretically the isomerization process in the triplet state reminding that in our case, the experiments which have been carried out do not involve this excited state. The obtained computational results show the same trend as observed for the ligands, namely that the triplet state as well, should allow the *trans-cis* isomerization for the **L1** ligand but not for **L2** and the complexes.

## Conclusions

In this paper, we describe the synthesis, full characterization and the theoretical studies of two new rhenium complexes **ReL1** and **ReL2**, **L1** and **L2** being azo-based iminopyridine ligands. The UV-visible spectrophotometric data show a red shift for complex **ReL1** indicating an increase of the electron acceptor behavior of the iminopyridine fragment in ligand **L1** upon complexation with Re (I) and a blue shift for complex **ReL2** with appearance of new broad absorption band in the visible region indicating the occurrence of a MLCT transition. The observed bands were fully assigned using the TD-DFT results. The distinct behaviors of the ligands and complexes regarding the *trans-cis* photoisomerization process have been rationalized, using DFT and TD-DFT computations. Several parameters have been considered for that, among them the relevant barriers on the potential energy surfaces in the S1 excited states, which involve conical intersection.

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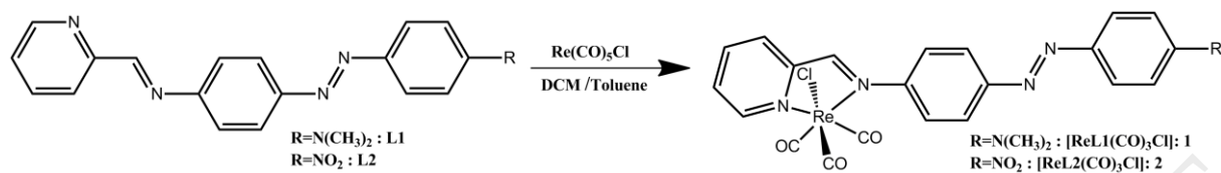
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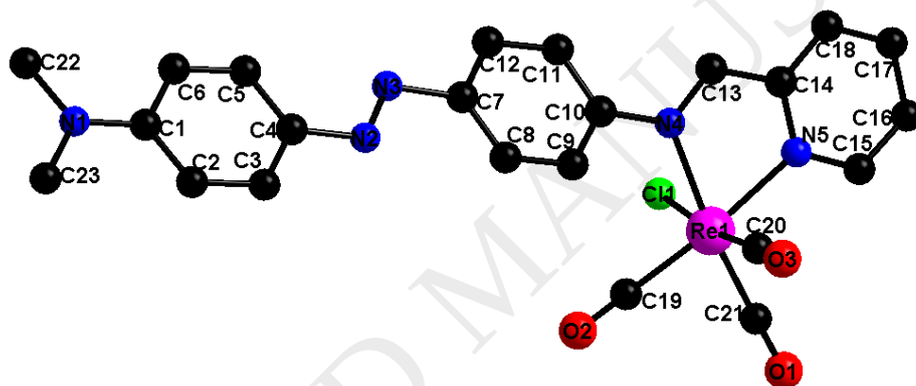
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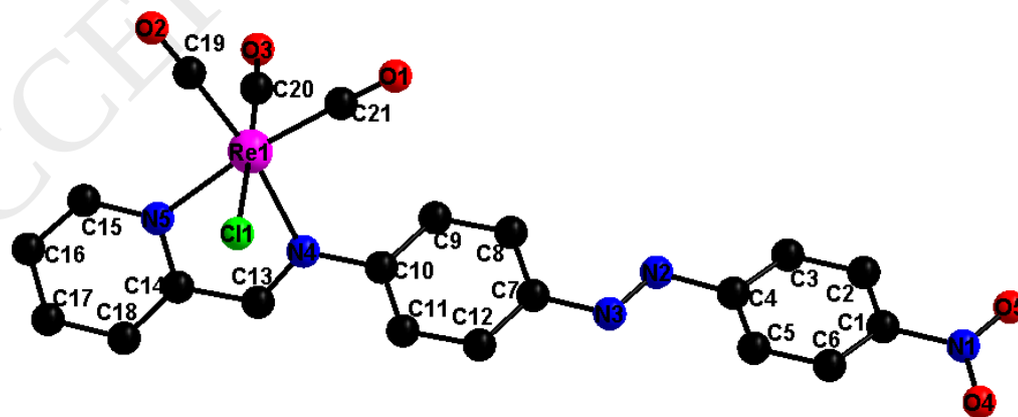
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Scheme 1. Synthetic route of rhenium complexes **ReL1** and **ReL2**.

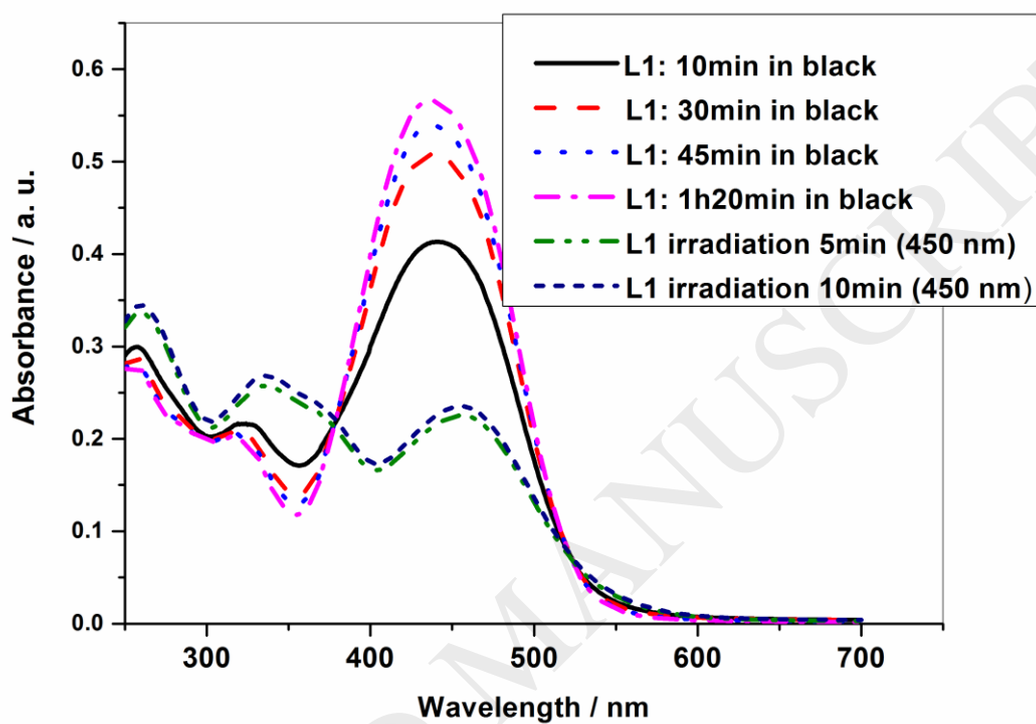


(ReL1)

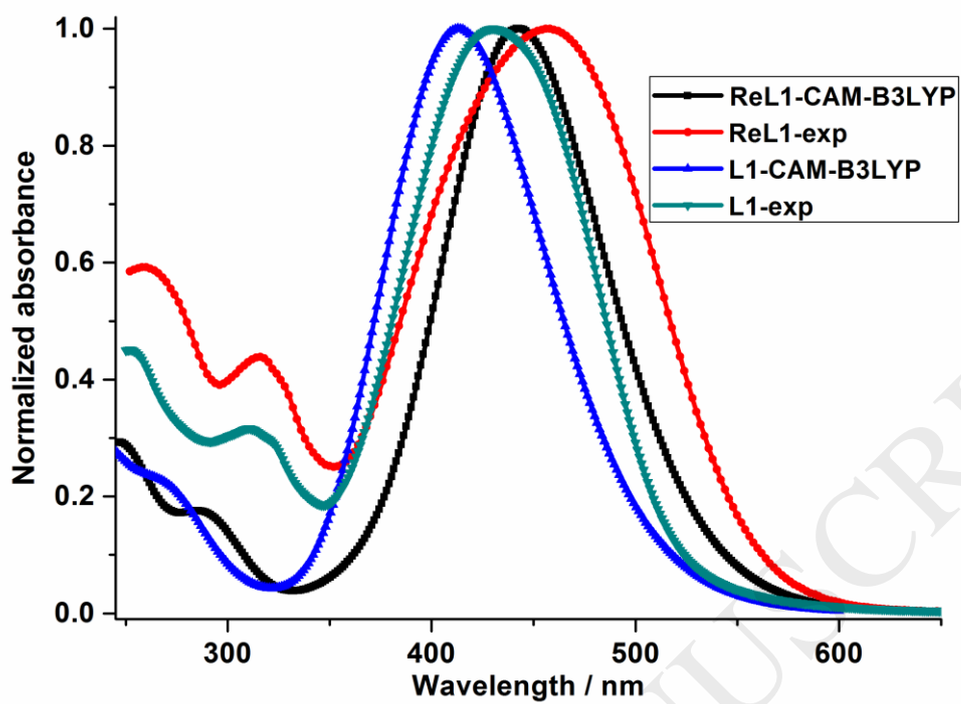


(ReL2)

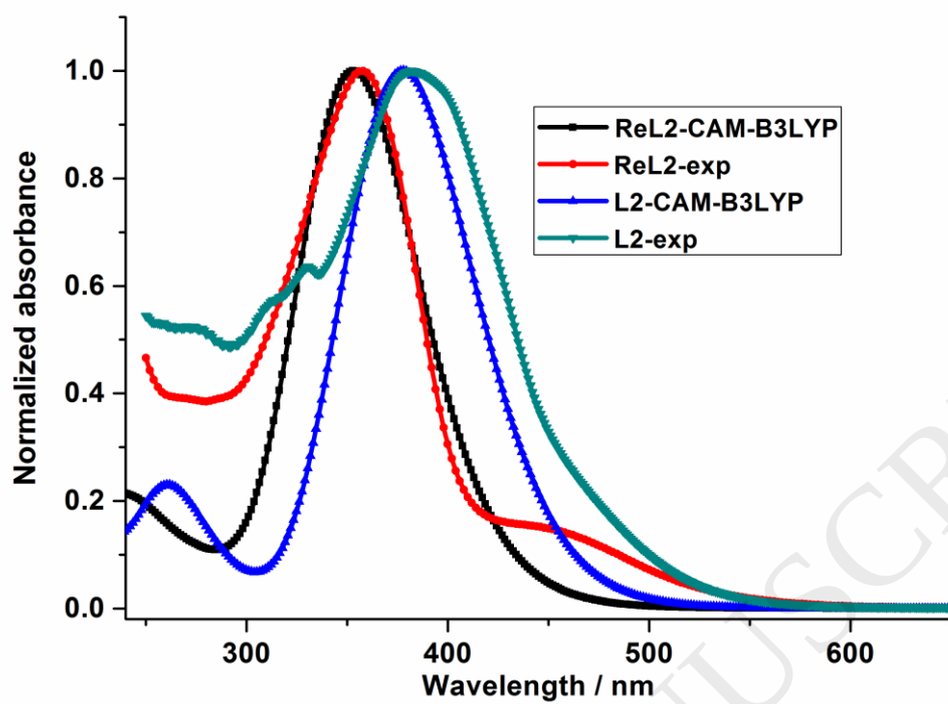
**Fig. 1.** Crystal structure of **ReL1** and **ReL2** complexes with atom numbering scheme. Hydrogen atoms were omitted for clarity.



**Fig. 2.** Changes in the UV-Visible absorption spectra upon irradiation a 450 nm of ligand **L1** in  $\text{CH}_2\text{Cl}_2$ .



**Fig. 3.** Experimental ( $C = 2.5 \cdot 10^{-5}$  M) and simulated UV-Visible absorption spectra of ligand **L1** and **ReL1** complex (in dichloromethane solution at room temperature)



**Fig. 4.** Experimental ( $C = 2.5 \cdot 10^{-5}$  M) and simulated UV-Visible absorption spectra of ligand **L2** and **ReL2** complex (in dichloromethane solution at room temperature)

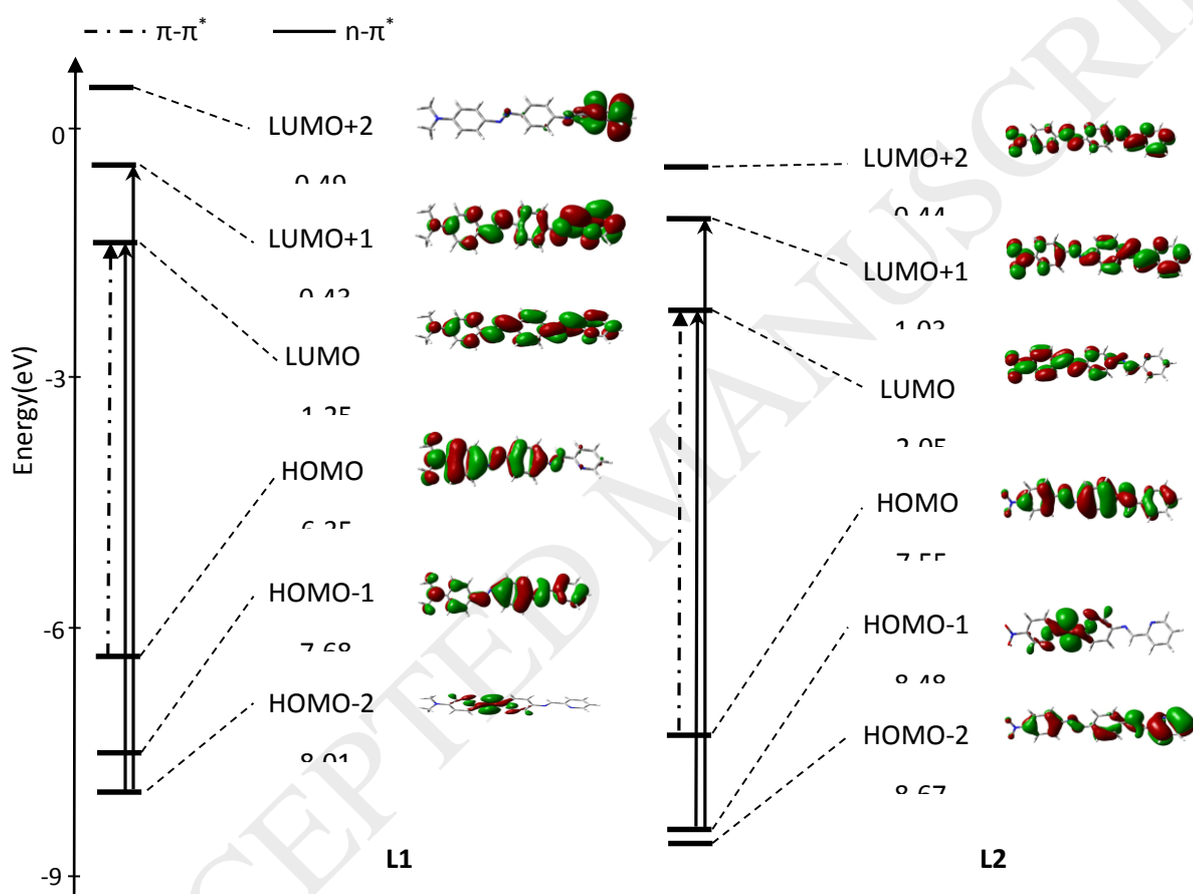
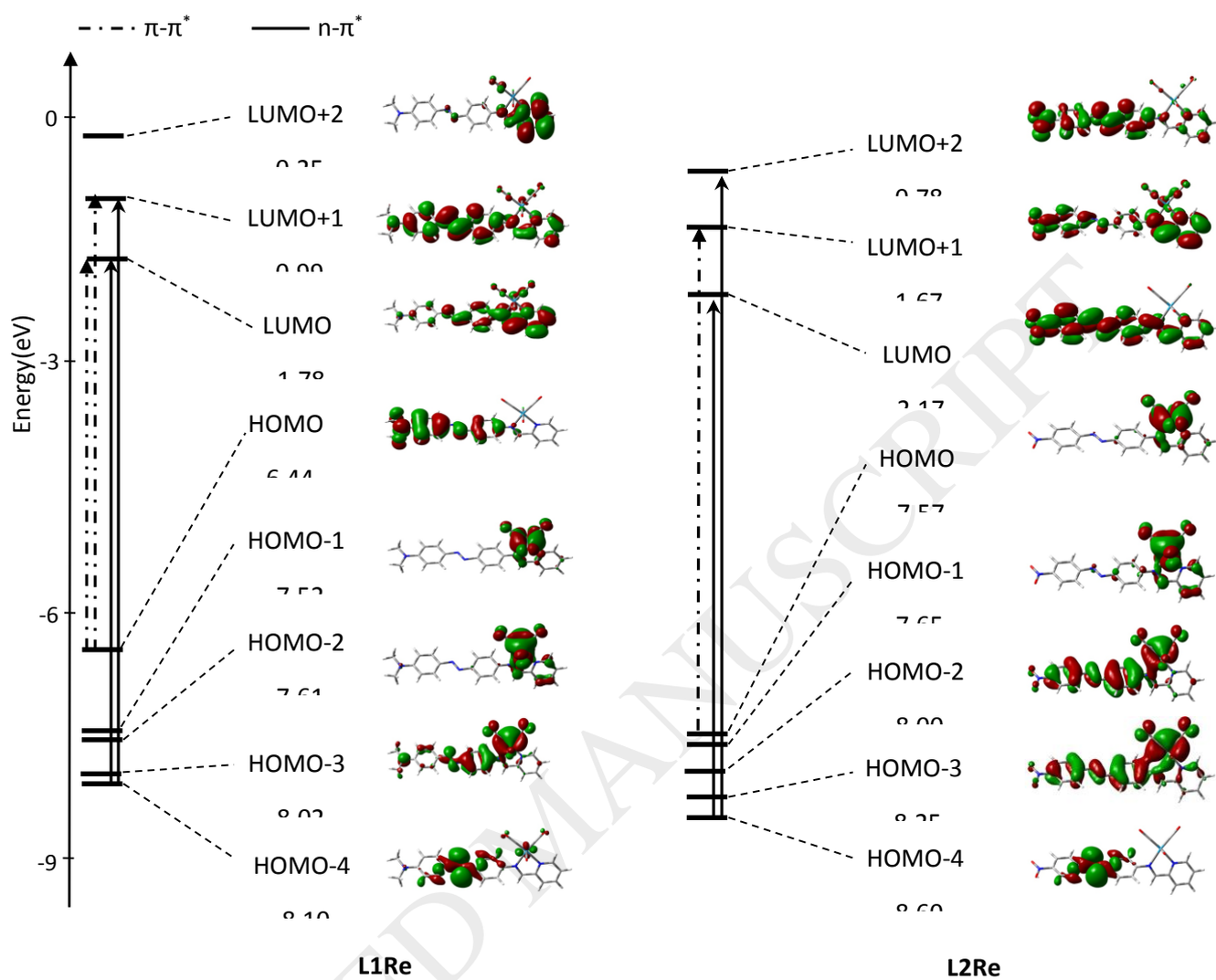


Fig. 5. Frontier MO diagrams of ligands L1 and L2 with main transitions (CAM-B3LYP level).



**Fig. 6.** Frontier MO diagrams of complexes **ReL1** and **ReL2** with main transitions (CAM-B3LYP level).

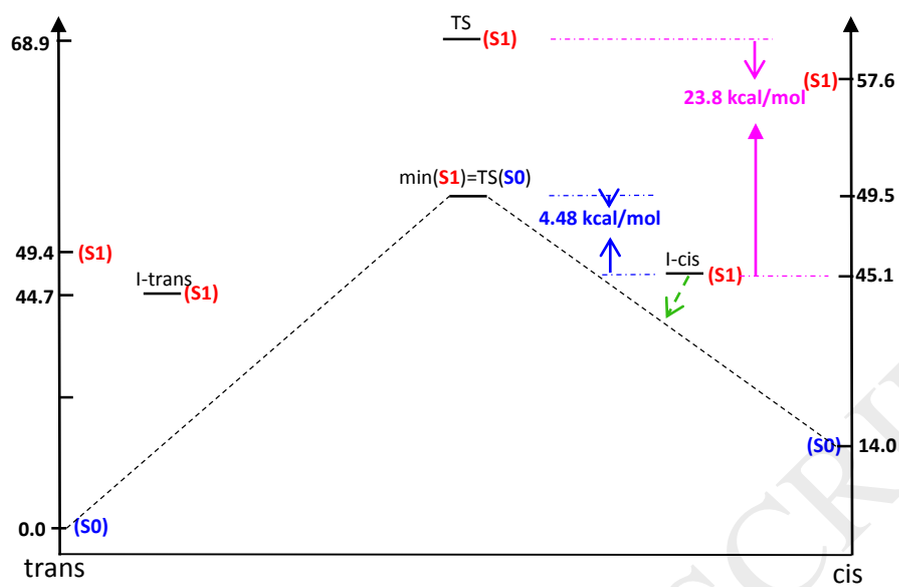


Fig. 7. Energy diagram of L1 (kcal/mol; CAM-B3LYP level)

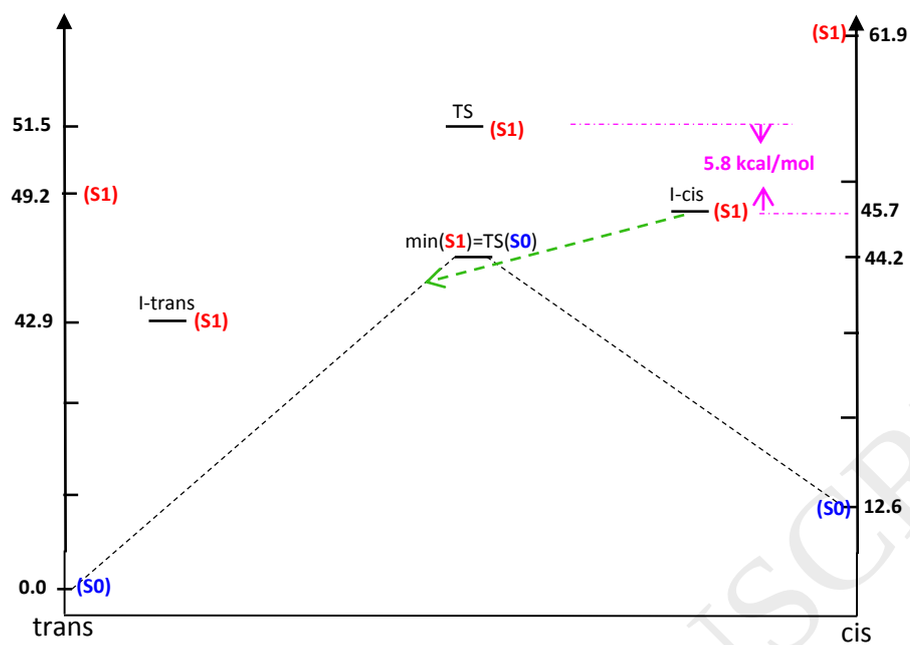


Fig. 8. Energy diagram of L2 (kcal/mol; CAM-B3LYP level)

**Table 1.** TD-DFT results (absorption wavelengths, oscillator strengths) for the ligands

Compounds	$\lambda$ in nm ( $\Delta E$ in eV)	$f$	Main transitions <sup>a</sup>
<b>L1</b>	431(2.87)	0.0050	H-2→L(72%) H-2→L+1(23%)
	418(2.97)	1.7858	H→L(85%)
	315(3.93)	0.0204	H-1→L(33%) H→L+1(32%)
<b>L2</b>	468(2.65)	0.0001	H-1→L(76%) H-1→L+2(16%)
	383(3.23)	1.6871	H→L(84%) H-2→L(10%)

<sup>a</sup>H = HOMO; L = LUMO**Table 2.** TD-DFT results for the complexes

Compounds	$\lambda$ (nm)	$f$	Main transitions
<b>ReL1</b>	431	0.0030	H-4→L(67%) H-4→L+1(21%)
	419	1.4566	H→L(52%) H→L+1(35%)
	395	0.0805	H-1→L(85%)
	374	0.0823	H-2→L(88%)
<b>ReL2</b>	467	0.0007	H-4→L(68%) H-4→L+2(21%)
	402	0.0114	H→L+1(59%) H→L(36%)
	380	0.1444	H-1→L+1(57%) H-1→L(37%)
	354	1.2107	H-2→L(69%)

Table 3. Rhenium, metal group and azo percentage weights in frontier molecular orbitals

MO	compounds	Rhenium weight %	Metal group <sup>a</sup> weight %	Azo group weight %
LUMO+2	<b>ReL1</b>	0.49	2.74	1.14
	<b>ReL2</b>	0.83	2.15	22.93
LUMO+1	<b>ReL1</b>	1.29	3.67	37.78
	<b>ReL2</b>	2.24	5.9	2.70
LUMO	<b>ReL1</b>	2.18	5.50	5.92
	<b>ReL2</b>	0.5	1.26	23.20
HOMO	<b>ReL1</b>	0.15	0.28	8.60
	<b>ReL2</b>	49.14	91.67	0.19
HOMO-1	<b>ReL1</b>	49.75	94.02	0.09
	<b>ReL2</b>	47.24	89.36	0.33
HOMO-2	<b>ReL1</b>	47.93	91.71	0.14
	<b>ReL2</b>	26.29	31.94	6.92
HOMO-3	<b>ReL1</b>	44.00	54.32	6.05
	<b>ReL2</b>	44.45	77.07	4.19
HOMO-4	<b>ReL1</b>	1.91	02.01	73.23
	<b>ReL2</b>	0.17	0.53	76.47

<sup>a</sup>Metal group = Re(CO)<sub>3</sub>Cl