

Theoretical Studies of Ground and Excited Electronic States in a Series of Halide Rhenium(I) Bipyridine Complexes

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Density functional theory (DFT) is applied to analyze ground- and excited-state properties of the Re(I) halide bipyridine complex $\text{ReCl}(\text{CO})_3(\text{bpy})$ (**1**) and the related complexes $\text{ReCl}(\text{CO})_3(5,5'\text{-dibromo-bpy})$ (**2**), $\text{ReCl}(\text{CO})_3(4,4'\text{-dimethyl-bpy})$ (**3**), and $\text{ReCl}(\text{CO})_3(4,4'\text{-dimethylformyl-bpy})$ (**4**) (where bpy = 2,2'-bipyridine). The electronic properties of the neutral molecules, in addition to the positive and negative ions, are studied using the B3LYP functional. Excited singlet and triplet states are examined using time-dependent DFT (TDDFT). The low-lying excited-state geometries are optimized at the ab initio configuration interaction singlets (CIS) level. As shown, the occupied orbitals involved in the transitions have a significant mixture of the metal Re and the group Cl, by the amount of metal 5d character which varies from 30 to 65%. The lowest unoccupied molecular orbital (LUMO) is a π^* orbital of the ligand bpy for the series of molecules. The TDDFT result indicates that the absorption maxima are at relatively high energy and are mainly assigned to bpy-based $\pi\pi^*$ transitions with somewhat metal-to-ligand charge transfer (MLCT) [$d(\text{Re}) \rightarrow \pi^*(\text{bpy})$] and ligand-to-ligand charge transfer (LLCT) [$p(\text{Cl}) \rightarrow \pi^*(\text{bpy})$] except for complex **3**, in which this band is mainly assigned to mixed MLCT/LLCT, and overlaps bpy $\pi\pi^*$ character. All the low-lying transitions are categorized as mixed MLCT/LLCT. The absorption bands are blue shifted when substituted by an electron-releasing group ($-\text{CH}_3$), and they are red shifted when substituted by an electron-withdrawing group ($-\text{Br}$ or $-\text{COOCH}_3$). The luminescence of all complexes is assigned as a triplet metal/chlorine to bpy charge transfer (MLCT/LLCT).

Introduction

There has been considerable recent interest in the optical properties of complexes and polymers (or oligomers),^{1–4} and particularly the luminescent properties of polypyridine complexes containing d^6 metals such as rhenium(I), ruthenium(II), and osmium(II) have been widely investigated in experiments, which shows immense applied prospect.^{5–8} Metal-based dyes have proven to perform surprisingly well, and against all odds, some dye molecules were found to be very stable under operating conditions.⁹ Their peculiar molecular properties, and the possibilities of related technological applications, are directly related to (a) the energetic spectrum of the molecules that determines the optical absorption and emission characteristics and (b) the molecules electronic structure of the ground as well as of the lowest excited states. The experimental appeal of these molecules lies in the adjustability of their electronic properties through change in the attached ligands and in their easily measured spectroscopic characteristics.¹⁰ As far as we know, very little theoretical work is available on this topic despite the potential interest of an advanced quantum chemical approach

for a better understanding of key issues, such as the nature of both the ground and the excited states involved in the absorption and/or photoemission and its tuning by different substituents.^{11–13}

The main difficulties against a reliable computational approach are related to the size of such systems and to the presence of strong electron correlation effects. Both properties are difficult to treat in the framework of the quantum mechanical methods rooted in the Hartree–Fock (HF) theory. As a matter of fact, the post-HF methods needed to obtain reliable excited-state properties have scaling properties with the number of electrons (N^6 or worse) that prevent their application to large systems. Recently, investigations have been carried out at the approximate level of theory (semiempirical models)^{14–18} or obtained by a reduction of the size of the system.¹⁹

On one hand, density functional theory (DFT) is successful at providing a means to evaluate a variety of ground-state properties with an accuracy close to that of post-HF methods.^{20,21} As a consequence, there is currently a great interest in extending DFT to excited electronic states.²² In this context, the time-dependent DFT (TDDFT) approach offers a rigorous route to the calculation of vertical electronic excitation spectra.^{23–25} Furthermore, remarkable structural predictions have been obtained, especially using the “hybrid” density functionals^{26,27} such as B3LYP and B3PW91 combining “exact exchange” with gradient-corrected density functionals. For excited states of closed-shell molecules, time-dependent DFT methods (TDDFT)

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have been developed. Applications of TDDFT approaches have recently been reported on transition-metal complexes and get a considerably good result.^{28–31}

On the other hand, Gaussian offers the configuration interaction approach, modeling excited states as combinations of single substitutions out of the HF ground state, and the method is thus named CIS.³² When paired with a basis set, it also may be used to define excited-state model chemistries whose results may be compared across the full range of practical systems. Theoretical investigations on excited states are uncommon but necessary for the molecules used in organic light-emitting diode devices (OLEDs), because the calculation of excited-state properties typically requires significantly more computational effort than is needed for the ground states.

The structure and electronic properties of the ground and lowest-excited states for the four metal polypyridyl halide complexes $\text{ReCl}(\text{CO})_3(\text{bpy})$ (**1**), $\text{ReCl}(\text{CO})_3(5,5'-dibromo-bpy) (**2**), $\text{ReCl}(\text{CO})_3(4,4'-dimethyl-bpy) (**3**), and $\text{ReCl}(\text{CO})_3(4,4'-dimethylformyl-bpy) (**4**) have been simulated. The aim of this paper is 2-fold. On one hand, we want to gain information on the physicochemical characteristics of the ground and low-lying excited states and, in particular, to understand the luminescent nature of metal Re complexes and the roles of their different substituents in tuning the dye properties. On the other hand, we want to show the potential of a quantum mechanical modeling based on DFT, in the evaluation of ground- and excited-state properties by comparison to the available experimental data.$$$

Computation Methods

Calculations on the electronic ground state of four related $\text{Re}(\text{I})$ halide complexes **1**, **2**, **3**, and **4** were carried out using B3LYP DFT. The “double- ξ ” quality basis set LANL2DZ was employed as basis set. A relative effective core potential (ECP) on Re replaced the inner-core electrons, leaving the outer-core [$(5s^2)(5p^6)$] electrons and the $(5d^6)$ valence electrons of $\text{Re}(\text{I})$. The excited-state geometries were optimized at the CIS level of theory. The transition energies will be calculated at the ground-state and excited-state geometries, and the results are compared with the available experimental data. The nature of the excited states, as well as the positive and negative ions with regard to “electron-hole” creation, is relevant to their use in OLED materials. The geometries were fully optimized with C_s symmetry constraints. All calculations are performed with Gaussian 98 using a spin-restricted formalism at the B3LYP/LANL2DZ level of theory, which has proved useful for other rhenium polypyridyl complexes.

Results and Discussion

The four $\text{Re}(\text{I})$ halide complexes **1**, **2**, **3**, and **4** are shown in Figure 1. For the calculated ground-state geometries, the electronic structure is examined in terms of the highest occupied and lowest virtual molecular orbitals. The nature of the low-lying excited states is then explored using the TDDFT approach to derive both absorption and emission spectra, which are compared to available spectroscopic data.

Ground-State Structures. The optimized geometrical parameters for all the molecules considered are reported in Table 1, together with the available X-ray data³³ for **2**. The calculated structure for complex **2** is depicted in Figure 2 as an example of the four complexes. As depicted, the rhenium(I) in each case adopts a distorted octahedral coordination geometry, and the carbonyl groups are arranged in a facial orientation. All four complexes have similar geometrical rearrangements according

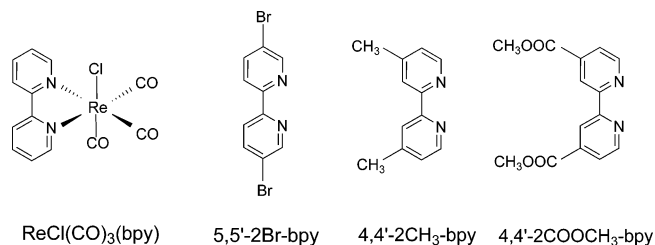


Figure 1. Schematic structures of $\text{Re}(\text{CO})_3\text{Cl}(\text{bpy})$ (**1**), $\text{Re}(\text{CO})_3\text{Cl}(5,5'$ -dibromo-bpy) (**2**), $\text{Re}(\text{CO})_3\text{Cl}(4,4'$ -dimethyl-bpy) (**3**), and $\text{Re}(\text{CO})_3\text{Cl}(4,4'$ -dimethylformate-bpy) (**4**). The full picture is depicted for parent complex **1**, and for complexes **2**, **3**, and **4** only ligands are depicted.]

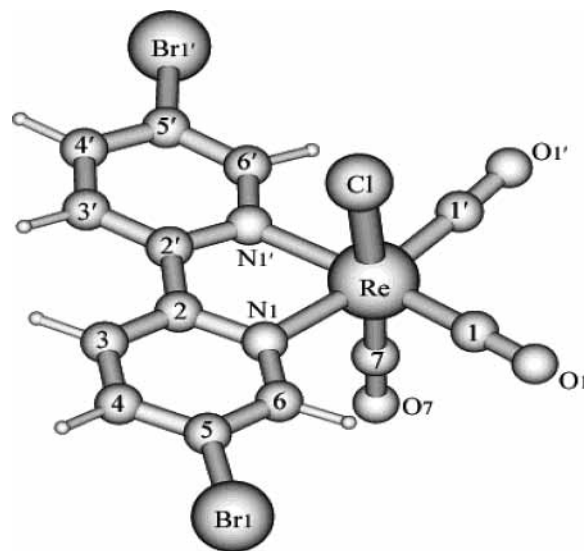


Figure 2. The calculated structure for molecule **2**.

TABLE 1: Calculated Bond Lengths for Four Re Center Complexes and the Comparison of Calculated Values with Experimental Values²⁹ from X-ray Diffraction on the Complexes 2

bonds	B3LYP/Å	bonds	B3LYP/Å	exp/Å
$\text{ReCl}(\text{CO})_3(\text{bpy})$		$\text{ReCl}(\text{CO})_3(5,5'$ -2Br-bpy)		
R[Re–N(1)]	2.172	R[Re–N(1)]	2.172	2.177
R[Re–N(1')]	2.172	R[Re–N(1')]	2.172	2.173
R[Re–C(1)]	1.925	R[Re–C(1)]	1.926	1.927
R[Re–C(1')]	1.925	R[Re–C(1')]	1.926	1.951
R[Re–C(7)]	1.910	R[Re–C(7)]	1.912	1.917
R[Re–Cl]	2.548	R[Re–Cl]	2.545	2.48
$\text{ReCl}(\text{CO})_3(4,4'$ -2CH ₃ -bpy)		$\text{ReCl}(\text{CO})_3(4,4'$ -2COOCH ₃ -bpy)		
R[Re–N(1)]	2.171	R[Re–N(1)]	2.163	
R[Re–N(1')]	2.171	R[Re–N(1')]	2.163	
R[Re–C(1)]	1.925	R[Re–C(1)]	1.928	
R[Re–C(1')]	1.925	R[Re–C(1')]	1.928	
R[Re–C(7)]	1.909	R[Re–C(7)]	1.913	
R[Re–Cl]	2.551	R[Re–Cl]	2.547	

to calculations due to the fact that they have been generated with the same approach and corresponding basis sets and the largest differences concerning bonds to the central metal atom, and these small differences can be attributed to different substituents.

The calculated Re–N bond lengths of 2.172 Å are only 0.005 and 0.001 Å shorter than the experimental values. The Re–C(1), Re–C(1'), and Re–C(7) bond lengths (1.926, 1.926, and 1.912 Å) are about 0.001, 0.025, and 0.005 Å shorter, respectively, than the measured values. The resulting Re–Cl (2.545 Å) bond length differed by 0.065 Å from the experimental values. It seems that the calculated values basically agree with experimental data, except that the axial Re–Cl bond is

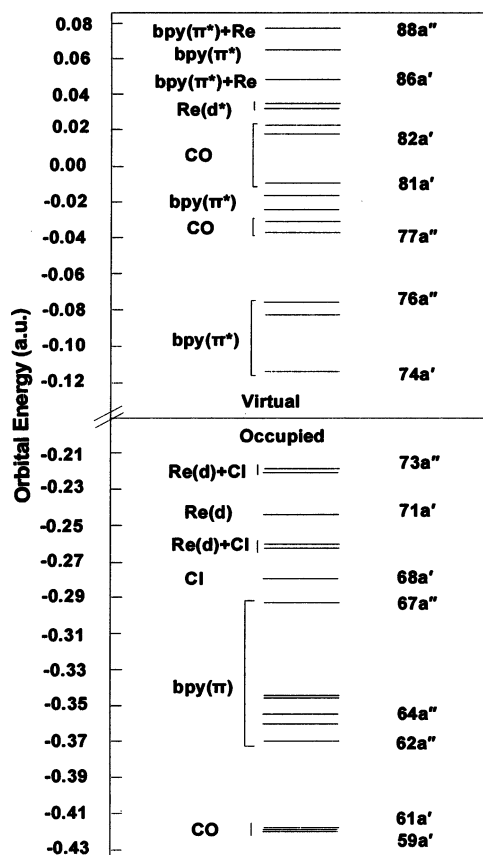


Figure 3. Energy-level diagram of $\text{Re}(\text{CO})_3\text{Cl}(\text{bpy})$ frontier molecular orbitals calculated at the B3LYP/LanL2DZ level. Labels on the left denote the dominant moiety contributing to each molecular orbital. For clarity, only a few of the molecular orbitals are numbered.

overestimated, and this point consists with the research of Turki et al.,³⁴ in which the author addressed that the drawback of DFT arises from the dynamical correlation effects, which become very important in complexes with a polar $\text{Ru}-\text{Cl}$ bond. Different substituents slightly influence the bond distances of the molecules. The $\text{Re}-\text{N}(1)$ and $\text{Re}-\text{N}(1')$ bond lengths of the complexes with substituents slightly shortened compared with parent molecule **1**. On the other hand, when substituted by $-\text{CH}_3$, the $\text{Re}-\text{C}(1)$, $\text{Re}-\text{C}(1')$, and $\text{Re}-\text{C}(7)$ bond lengths are slightly shorter, and when substituted by $-\text{Br}$ and $-\text{COOCH}_3$, they are slightly longer than that of parent molecule.

Molecular Orbitals. It will be useful to examine the highest occupied and lowest virtual orbitals for these Re complexes to provide the framework for the excited-state TDDFT and CIS calculations in the subsequent section. Moreover, frontier orbitals play a relevant role in such systems because they rule the electronic excitations and the transition character. The frontier orbitals are plotted according to their energies in Figure 3 for complex **1** as an example. The assignment of the type of each MO was made on the basis of its composition (see Table 2, in which only the most important five occupied and three virtual (unoccupied) orbitals are listed) and by visual inspection of its three-dimensional representation (e.g., Figure 4).

We find although the five highest occupied orbitals (HOMOs) have strong rhenium d orbitals (over 30%), nearly equivalent contributions come from the chlorine p orbital, with the exception that the HOMO_{-2} orbitals are mainly rhenium d orbitals, in which the Re d orbital composition exceeds 60% and displays the dominant character. In other words, these occupied MOs are mainly composed of chloride p_z orbitals that form antibonding interactions with metal d_{xz} orbitals (see Table

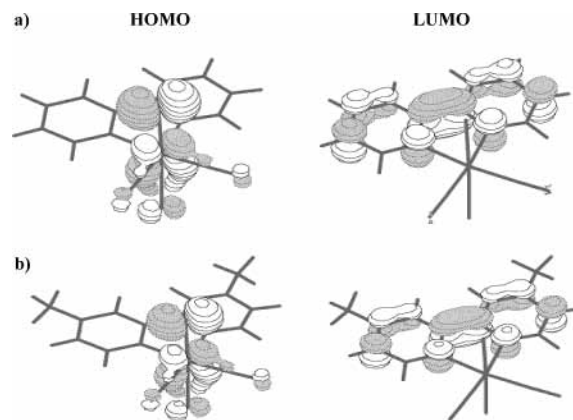


Figure 4. Contour plots of highest occupied and lowest virtual orbitals in considered molecules (a) **1** and (b) **3**.

TABLE 2: Highest Occupied and Lowest Virtual Orbitals with Character for Re Complexes

designation	character (%)	ϵ (au)
ReCl(CO) ₃ (bpy) Occupied		
69 a'	5d (37%) + p (Cl) (34%)	-0.262
70 a''	5d (34%) + p (Cl) (39%)	-0.260
71 a'	5d (64%)	-0.245
72 a'	5d (35%) + p (Cl) (37%)	-0.220
73 a'' (HOMO)	5d (36%) + p (Cl) (36%)	-0.218
	$ \Delta E(\text{HOMO}-\text{LUMO}) ^a$	0.103
ReCl(CO) ₃ (bpy) Virtual		
74 a' (LUMO)	bpy (93%)	-0.115
75 a'	bpy (97%)	-0.082
76 a''	bpy (98%)	-0.077
ReCl(CO) ₃ (5,5'-2Br-bpy) Occupied		
75 a'	5d (33%) + p (Cl) (36%)	-0.270
76 a''	5d (31%) + p (Cl) (38%)	-0.268
77 a'	5d (64%)	-0.253
78 a'	5d (36%) + p (Cl) (38%)	-0.228
79 a'' (HOMO)	5d (38%) + p (Cl) (36%)	-0.225
	$ \Delta E(\text{HOMO}-\text{LUMO}) ^a$	0.099
ReCl(CO) ₃ (5,5'-2Br-bpy) Virtual		
80 a' (LUMO)	bpy (89%)	-0.126
81 a'	bpy (97%)	-0.095
82 a''	bpy (98%)	-0.090
ReCl(CO) ₃ (4,4'-2CH ₃ -bpy) Occupied		
77 a'	5d (34%) + p (Cl) (35%)	-0.256
78 a''	5d (33%) + p (Cl) (39%)	-0.255
79 a'	5d (65%)	-0.240
80 a'	5d (36%) + p (Cl) (37%)	-0.215
81 a'' (HOMO)	5d (36%) + p (Cl) (36%)	-0.213
	$ \Delta E(\text{HOMO}-\text{LUMO}) ^a$	0.105
ReCl(CO) ₃ (4,4'-2CH ₃ -bpy) Virtual		
82 a' (LUMO)	bpy (91%)	-0.107
83 a'	bpy (95%)	-0.074
84 a''	bpy (95%)	-0.066
ReCl(CO) ₃ (4,4'-2COOCH ₃ -bpy) Occupied		
99 a'	5d (37%) + p (Cl) (33%)	-0.268
100 a''	5d (35%) + p (Cl) (38%)	-0.266
101 a'	5d (64%)	-0.251
102 a'	5d (34%) + p (Cl) (38%)	-0.226
103 a'' (HOMO)	5d (35%) + p (Cl) (37%)	-0.224
	$ \Delta E(\text{HOMO}-\text{LUMO}) ^a$	0.092
ReCl(CO) ₃ (4,4'-2COOCH ₃ -bpy) Virtual		
104 a' (LUMO)	bpy (78%) + p (COOCH ₃)	-0.132
105 a''	bpy (70%) + p (COOCH ₃)	-0.111
106 a'	bpy (92%) + p (COOCH ₃)	-0.096

^a ΔE refers to band gap.

2 and Figure 3). Some lower-energy occupied MOs are mainly centered on the bpy ligands or group CO in each complex. On one hand, Turki and co-worker³⁴ calculated the UV-vis

TABLE 3: Ionization Potentials, Electron Affinities, “Small-Polaron” Stabilization Energies, and Spin Densities for the Four Complexes (in eV)^a

	IP(v)	IP(a)	HEP	SPE(h)	Re	bpy	spin density of cation (%)					
							3CO	Cl	2Br	2CH ₃	2COOCH ₃	
1	7.59	7.36	0.23	0.23	0.53	0.02	0.10	0.35				
2	7.80	7.57	0.23	0.23	0.54	0.03	0.09	0.32	0.02			
3	7.43	7.19	0.25	0.24	0.54	0.03	0.10	0.33		0.00		
4	7.67	7.42	0.24	0.25	0.52	0.01	0.10	0.37				0.00

	EA(v)	EA(a)	EEP	SPE(e)	Re	bpy	spin density of anion (%)					
							3CO	Cl	2Br	2CH ₃	2COOCH ₃	
1	-1.63	-1.84	0.21	0.21	-0.01	0.96	0.04	0.01				
2	-2.04	-2.21	0.17	0.17	-0.01	0.95	0.04	0.02	0.00			
3	-1.49	-1.72	0.23	0.22	-0.01	0.95	0.04	0.01		0.01		
4	-2.28	-2.50	0.22	0.22	-0.00	0.76	0.03	0.01				0.20

^a See text for definitions. The suffixes v and a respectively indicate vertical and adiabatic values.

absorption spectra of the Ru complex containing Cl with CASSCF/CASPT2 and TDDFT methods; they thought DFT calculations overestimate halide contributions in metal–halide complexes, especially if a low valent metal atom is involved, although DFT approaches are remarkably good in the case of nonhalide complexes. On the other hand, Stufkens^{35,36} have analyzed that the electronic transitions to the α -diimine have strongly mixed metal-to-ligand charge transfer (MLCT)/X-to-ligand charge transfer (XLCT) character due to the interaction of $M-d_{\pi}$ and $X-p_{\pi}$ orbitals in experiment. As a result, two absorption bands are observed, the relative intensities of which depend on the metal character of the orbitals involved in the transitions. In contrast, the LUMO and two subsequent virtual (unoccupied) orbitals are essentially π^* orbitals localized on the bipyridine moieties (over 90%), but the higher LUMOs show considerable localization on the CO groups. The four other rhenium orbitals (MO 84a'', MO 85a', MO 86a', and MO 88a'') with mainly d character are still higher in energy and display nonbonding or antibonding interactions with the ligands. However, since the contribution from these orbitals is quite small for low-energy transition, we only concentrate on the orbitals listed in Table 2. In fact, due to their similar basic structure, the frontier orbitals have similar features in all the considered complexes.

For intuition, the contour plot of HOMO and LUMO orbitals for molecules **1** and **3** are depicted as example in Figure 4. In the contour plots, the mixing of 5d (Re) and Cl (p) is evident, while the LUMO is seen to be delocalized over the bpy ligand. Thus there must be a mixture between MLCT and ligand-to-ligand charge transfer (LLCT) upon transition from the ground state to the excited state.

But different substituents still influence the character of the complexes as shown in Table 2. We note that the three lowest unoccupied orbitals are essentially π^* orbitals of the bipyridine moieties, whereas the substituents are localized at bipyridine rings. So substituents have a direct effect on LUMO. From the energies listed, we observed that the electron-releasing $-CH_3$ group slightly increases the LUMO energy, resulting in the increased HOMO–LUMO band gap, which then leads to a decrease in the λ_{\max} values. On the other hand, electron-withdrawing $-Br$ and $-COOCH_3$ groups decrease the LUMO energy, resulting in the decreased band gap in the order $-Br > -COOCH_3$, which then leads to an increase in the λ_{\max} values.

Ionization Potentials and Electron Affinities. Additional information derived from our calculations provides insight into the inter-relationship of structure and electronic behavior, in particular the response of the molecule to the formation of a hole or the addition of an electron. Table 3 contains the

ionization potentials (IPs), electron affinities (EAs), both vertical (v, at the geometry of the neutral molecule) and adiabatic (a, optimized structure for both the neutral and charged molecule), and extraction potentials (HEP and EEP for the hole and the electron, respectively) that refer to the geometry of the ions. Relatively small energy changes are associated with structural relaxation. In all cases, the energy required to create a hole is ~ 8 eV, while the extraction of an electron from the anion requires ~ 2 eV. In fact, these complexes share the common features of having accessible Re-based oxidations and bipyridine-based reductions. In fact, early work has amply demonstrated that $ReX(CO)_3$ (diimine) complexes are both strong reductants and oxidants. Experimental research reveals nearly reversible oxidation and reduction waves at $\sim +1.3$ and ~ -1.3 V vs an aqueous saturated calomel electrode for the complex *fac*- $ReCl(CO)_3$ (phen).³⁷ It is found that the calculated IPs (no matter IP(a) or IP(v)) of molecules **2** and **4** are relatively larger, whereas for molecule **3**, they are relatively lower than their parent molecule, which is in agreement with the trend in HOMO energy. These all correspond to removal of an electron from the “5d” orbital. As shown in the cation spin densities in Table 3, over 50% of the spin density in all four complexes is on the Re and the remainder largely shared on group Cl and less on bpy and CO ligands, which means there will be mixed character between metal Re and group Cl for electronic transitions, and this basically consists of the analysis of HOMO composition.

Four complexes show weakly bound negative ions, corresponding to the electron affinity. In each complex, the unpaired spin density are totally on bpy ligands and less on CO and Cl ligands. This is consistent with the LUMO being primarily the bpy π^* orbitals. Although both IP(a) and EA(a) are lower than 0.23 eV on average than IP(v) and EA(v), respectively, due to structural relaxation after oxidation or reduction, the structure changes little after electron reduction or addition.

The same calculations are also used to estimate self-trapping energies of positive and negative charges in the materials. Indeed, in ref 38, the traps that characterize the electron transport in the material were identified as the states in which the injected electron is self-trapped in the individual molecules as a consequence of structure relaxation. Such a procedure, however, provides an estimate of the exciton trap energy rather than that of the injected electron. Following the same interpretation for the trap states, the correct energy in our scheme is the energy gain of the excess electron due to structural relaxation, i.e., the difference $EA(a) - EA(v)$, which we also report in Table 3, as the “small-polaron” stabilization energy (SPE) for the electron. It seems that the four complexes appear to trap the electron nearly the same efficiently. The SPE values may be compared

TABLE 4: Selected Calculated Excitation Energies (E), Wavelengths (λ), Oscillator Strengths ($\times c4$), and Dominant Excitation Character for Low-Lying Singlet (S_n) and Triplet (T_n) States of the Complex $\text{ReCl}(\text{CO})_3(\text{bpy})$ and Comparison with Experimental Data

state	excitation	E_{cal} , eV	λ_{cal} , nm	$\times c4_{\text{cal}}$	λ_{exp} , nm	character
Singlet Excited States						
1(A'')	73 \rightarrow 74(0.70)	2.04	609	0.0013		MLCT/LLCT
2(A')	72 \rightarrow 74(0.68)	2.21	561	0.0315		MLCT/LLCT
9(A')	69 \rightarrow 74(0.64)	3.35	370	0.0872	387(CH ₂ Cl ₂) 370(CH ₃ CN) ³⁷	MLCT/LLCT
14(A')	73 \rightarrow 77(0.58)	3.93	315	0.0103		MLCT/LLCT
17(A')	69 \rightarrow 75(0.65)	4.23	293	0.0134		MLCT/LLCT
18(A'')	69 \rightarrow 76(0.58)	4.28	290	0.0228		MLCT/LLCT
	67 \rightarrow 74(0.32)					LLCT
19(A')	72 \rightarrow 78(0.56)	4.30	288	0.0185		MLCT/LLCT
	73 \rightarrow 77(0.17)					MLCT/LLCT
20(A')	70 \rightarrow 76(0.65)	4.35	285	0.0614		MLCT/LLCT
	69 \rightarrow 75(0.19)					MLCT/LLCT
21(A')	73 \rightarrow 79(0.69)	4.53	274	0.0160		MLCT/LLCT
22(A'')	67 \rightarrow 74(0.43)	4.54	273	0.2318	293(CH ₂ Cl ₂)	IL
	69 \rightarrow 76(0.33)					MLCT/LLCT
	71 \rightarrow 77(0.26)					MLCT
25(A'')	72 \rightarrow 80(0.45)	4.71	263	0.1202		MLCT/LLCT
	71 \rightarrow 77(0.31)					MLCT
	72 \rightarrow 79(0.28)					MLCT/LLCT
27(A'')	68 \rightarrow 76(0.69)	4.86	255	0.0301		LLCT
28(A'')	72 \rightarrow 80(0.40)	4.90	253	0.0372		MLCT/LLCT
	71 \rightarrow 80(0.30)					MLCT
	73 \rightarrow 81(0.24)					MLCT/LLCT
29(A')	73 \rightarrow 80(0.52)	4.95	251	0.0168		MLCT/LLCT
	69 \rightarrow 78(0.26)					MLCT/LLCT
33(A'')	67 \rightarrow 75(0.53)	5.17	240	0.0827		IL
	65 \rightarrow 74(0.25)					IL
34(A'')	70 \rightarrow 78(0.41)	5.18	239	0.0250		MLCT/LLCT
	71 \rightarrow 79(0.29)					IL
35(A')	67 \rightarrow 76(0.53)	5.22	238	0.0157		IL
	72 \rightarrow 81(0.24)					MLCT/LLCT
	64 \rightarrow 74(0.21)					IL
40(A'')	72 \rightarrow 83(0.49)	5.46	227	0.0230		MLCT/LLCT
	71 \rightarrow 80(0.31)					MLCT
Triplet Excited States						
1(A'')	73 \rightarrow 74(0.71)	1.97	628	0.0000		MLCT/LLCT
2(A')	72 \rightarrow 74(0.71)	2.03	610	0.0000		MLCT/LLCT
3(A')	71 \rightarrow 74(0.71)	2.71	457	0.0000		MLCT
4(A'')	70 \rightarrow 74(0.52)	2.86	434	0.0000		MLCT/LLCT
	67 \rightarrow 74(0.40)					IL
5(A'')	73 \rightarrow 75(0.69)	2.98	416	0.0000		MLCT/LLCT

with tris-(8-hydroxyquinolate)-aluminum (Alq) another dopant for OLED applications. Similar DFT calculations³⁹ on Alq predicted 0.13 eV for SPE.

Excitation Energies. With the prerequisite ground-state DFT calculation in hand, we proceed to the time-dependent calculation of complexes **1–4** to find the characters and energies of their low-lying singlet and triplet excited states. We begin with the singlet \rightarrow singlet spin-allowed transition. Forty singlet excited states are calculated at the optimized structure of the ground state for each complex, and only the singlet excited states with the greatest oscillator strengths (above 0.01) are typically listed in Tables 4–7, together with experimental data; this information is also presented graphically for all calculated excited states in Figures 6–9, respectively. The energy of each excited is vertical excitation energy in eV from the ground state. As shown, the four complexes have relatively similar absorption character. There are significant oscillator strength throughout the ~ 2 –5-eV region, but there are also several singlet excited states with zero oscillator strength, and these states, although present in the molecule's excited-state manifold, will therefore no contribute to the compound's absorption spectra. No excited states or absorption features are found below 1 eV.

A commonly used model of an excited state corresponds to excitation of an electron from an occupied to a virtual MO (i.e.,

a one-electron picture). However, the excited states calculated herein demonstrate that excited-state electronic structures are best described in terms of multiconfigurations, wherein a linear combination of several occupied-to-virtual MO excitations comprises a given optical transition. Assignment of the character of each excited state was based on the compositions of the occupied and virtual MOs of the dominant configuration(s) for that excited state. We selected complex **1** shown in Table 4 as analysis example. For the T₃ excited state, the dominant excitation is 71 \rightarrow 74, and since the occupied orbital (71) is metal based and the virtual orbital (74) is bipyridine π^* , the transition is designated as a MLCT. Similarly, for the S₂₇ excited state, the dominant excitation is 68 \rightarrow 76, and since the occupied orbital (68) is localized on a halide chlorine and virtual orbital (76) type is bipyridine π^* , the transition is designated as a LLCT. For S₃₃, the dominant excitation is from bipyridine π (65 and 67) to bipyridine π^* (74 and 75), the transition is designated as intraligand (IL) character. After presenting the detailed theoretical results in this section, in the following section, we summarize the excited states for the four complexes and compare them to experimental absorption and emission studies.

Figure 5 illustrates the differences of MLCT, LLCT, and IL excitations for three strongly allowed transitions of complex **1**.

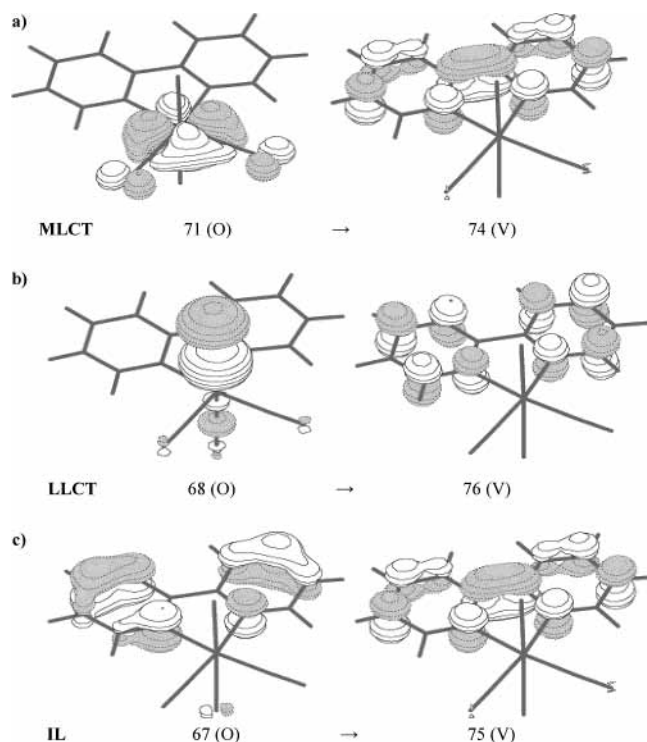


Figure 5. Example of dominant occupied and virtual orbitals for three different types of excitations of complex **1**. (a) MLCT excitation (71 → 74). (b) LLCT excitation (68 → 76). (c) IL excitation (67 → 75).

Moreover, for the S_1 excited state, the calculated spin density of positive and negative ions also has significant referenced values for transition assignment. For the majority of the excited states calculated, such an assignment can be made unambiguously. However, for the metal–halide molecules under studying

on, most of the significant excited states exhibit mixed MLCT and LLCT and, sometimes, IL characters. In fact, excited-state character as a function of the molecular structure has been reviewed from the experimental point of view, such as in the ref 34 where the author addressed that the character of the lowest excited states changes gradually from MLCT to LLCT while halide ligand changes from Cl to Br, especially, to I in $\text{Re}(\text{E})(\text{CO})_3(\alpha\text{-diimine})$ complexes, which are consistent with our calculations.

From Tables 4–7 and Figures 6–9, we see that in each complex the low-energy part of the absorption spectra originated in an electronic transition that corresponds to excitation from the rich mixed orbitals of metal Re and chlorine Cl into π^* orbital localized predominantly on bpy. And the intense bands are observed at a relative higher-energy region, which corresponds to $\pi\pi^*$ excitations. Furthermore, we compare the results of the excited states of **2**, **3**, and **4** with **1**. From the earlier discussion of the frontier orbitals (Table 2 and Figure 3), due to the influence of the electron-releasing or withdrawing substituents, the energy gaps ($|\Delta E(\text{HOMO} - \text{LUMO})|$) decrease in the order $3 > 1 > 2 > 4$, and this rule remains constant in absorption spectra. The absorption bands exhibit blue shifts in **2** and red shifts in **3** and **4** compared with **1**. It is evident for S_1 , where $2.12 (-\text{CH}_3) > 2.04 > 1.93 (-\text{Br}) > 1.75 \text{ eV} (-\text{COOCH}_3)$. But the band character basically is identical.

The 35 lowest-energy triplet excited states were also calculated, using analogous TDDFT methodology. The first five triplet excited states are listed in Table 2. MLCT, LLCT, and IL excited states are all seen, but most of them are the mixed character, as with the singlets. As expected from Hund's rule, transitions to the triplet states tend to be lower in energy than their corresponding singlets. For example, in complex **1**, the first triplet vertical transition energy is 1.97 eV lower than that of the first singlet excited state (2.04 eV), where both represent

TABLE 5: Selected Calculated Excitation Energies (E), Wavelengths (λ), Oscillator Strengths ($\times c_4$), and Dominant Excitation Character for Low-Lying Singlet (S_n) and Triplet (T_n) States of the Complex $\text{ReCl}(\text{CO})_3(5,5'\text{-2Br-bpy})$ and Comparison with Experimental Data.

state	excitation	E_{cal} , eV	λ_{cal} , nm	$\times c_4_{\text{cal}}$	λ_{exp} , nm	character
Singlet Excited States						
1(A'')	79 → 80(0.70)	1.93	643	0.0012		MLCT/LLCT
2(A')	78 → 80(0.68)	2.12	585	0.0310		MLCT/LLCT
9(A')	75 → 80(0.66)	3.26	381	0.0814	410(CH ₂ Cl ₂)	MLCT/LLCT
13(A'')	74 → 80(0.46)	3.88	320	0.1518	316(CH ₂ Cl ₂)	IL
15(A'')	76 → 81(0.29)	3.89	319	0.0582		MLCT/LLCT
	74 → 80(0.28)					IL
17(A'')	76 → 81(0.58)	4.06	306	0.1319		MLCT/LLCT
	75 → 82(0.26)					MLCT/LLCT
18(A')	75 → 81(0.64)	4.10	302	0.0105		MLCT/LLCT
19(A')	76 → 82(0.63)	4.20	295	0.0629		MLCT/LLCT
	75 → 81(0.20)					MLCT/LLCT
20(A')	78 → 84(0.53)	4.23	293	0.0232		MLCT/LLCT
	78 → 87(0.25)					MLCT/LLCT
21(A'')	75 → 82(0.57)	4.27	290	0.1933		MLCT/LLCT
	79 → 87(0.16)					MLCT/LLCT
22(A'')	79 → 84(0.48)	4.31	288	0.0551		MLCT/LLCT
	75 → 82(0.22)					MLCT/LLCT
23(A')	79 → 85(0.64)	4.37	284	0.0189		MLCT/LLCT
31(A'')	74 → 81(0.62)	4.65	267	0.1082		256(CH ₂ Cl ₂) IL
32(A'')	77 → 83(0.44)	4.65	266	0.0634		MLCT
	78 → 88(0.39)					MLCT/LLCT
33(A'')	73 → 82(0.69)	4.70	264	0.0159		IL
34(A')	74 → 82(0.61)	4.77	256	0.0297		IL
Triplet Excited States						
1(A'')	79 → 80(0.71)	1.87	664	0.0000		MLCT/LLCT
2(A')	78 → 80(0.72)	1.93	642	0.0000		MLCT/LLCT
3(A')	77 → 80(0.71)	2.61	474	0.0000		MLCT
4(A'')	76 → 80(0.62)	2.69	460	0.0000		MLCT/LLCT
5(A'')	79 → 81(0.70)	2.83	438	0.0000		MLCT/LLCT

TABLE 6: Selected Calculated Excitation Energies (E), Wavelengths (λ), Oscillator Strengths ($\times c4$), and Dominant Excitation Character for Low-Lying Singlet (S_n) and Triplet (T_n) States of the Complex $\text{ReCl}(\text{CO})_3(4,4'-2\text{CH}_3\text{-bpy})$ and Comparison with Experimental Data

state	excitation	E_{cal} , eV	λ_{cal} , nm	$\times c4_{\text{cal}}$	λ_{exp} , nm	character
Singlet Excited States						
1(A'')	81 \rightarrow 82(0.70)	2.12	586	0.0013		MLCT/LLCT
2(A')	80 \rightarrow 82(0.69)	2.28	544	0.0347		MLCT/LLCT
9(A')	77 \rightarrow 82(0.59)	3.39	365	0.0954	380(CH ₂ Cl ₂)	MLCT/LLCT
	81 \rightarrow 84(0.33)				364(CH ₃ CN) ³⁷	MLCT/LLCT
13(A')	81 \rightarrow 85(0.47)	3.95	314	0.0111		MLCT/LLCT
	76 \rightarrow 82(0.44)					LLCT
17(A')	77 \rightarrow 83(0.57)	4.28	290	0.0203		MLCT/LLCT
	81 \rightarrow 88(0.16)					MLCT/LLCT
18(A')	80 \rightarrow 86(0.46)	4.31	288	0.0257		MLCT/LLCT
	77 \rightarrow 83(0.37)					MLCT/LLCT
19(A'')	77 \rightarrow 84(0.54)	4.36	284	0.0454		MLCT/LLCT
	75 \rightarrow 82(0.36)					IL
20(A')	78 \rightarrow 84(0.66)	4.45	279	0.0521	290(CH ₂ Cl ₂)	MLCT/LLCT
	81 \rightarrow 87(0.18)					MLCT/LLCT
21(A')	81 \rightarrow 87(0.67)	4.55	272	0.0281		MLCT/LLCT
22(A'')	80 \rightarrow 87(0.51)	4.57	271	0.0186		MLCT/LLCT
	79 \rightarrow 85(0.32)					MLCT
24(A'')	80 \rightarrow 87(0.38)	4.59	270	0.1698		MLCT/LLCT
	75 \rightarrow 82(0.36)					LLCT
25(A'')	80 \rightarrow 88(0.43)	4.71	263	0.1773	260(CH ₂ Cl ₂)	MLCT/LLCT
	80 \rightarrow 87(0.30)					MLCT/LLCT
	79 \rightarrow 85(0.15)					MLCT
27(A'')	80 \rightarrow 88(0.40)	4.90	253	0.0640		MLCT/LLCT
	81 \rightarrow 89(0.21)					MLCT/LLCT
28(A')	81 \rightarrow 88(0.51)	4.95	251	0.0171		MLCT/LLCT
	80 \rightarrow 90(0.19)					MLCT/LLCT
	80 \rightarrow 86(0.18)					MLCT/LLCT
29(A'')	76 \rightarrow 84(0.68)	4.98	249	0.0110		LLCT
33(A'')	75 \rightarrow 83(0.49)	5.12	242	0.0674		IL
	74 \rightarrow 82(0.39)					IL
40(A'')	80 \rightarrow 91(0.45)	5.46	227	0.0302		LLCT
	79 \rightarrow 88(0.32)					MLCT
	78 \rightarrow 86(0.22)					MLCT/LLCT
Triplet Excited States						
1(A'')	81 \rightarrow 82(0.71)	2.06	602	0.0000		MLCT/LLCT
2(A')	80 \rightarrow 82(0.71)	2.12	586	0.0000		MLCT/LLCT
3(A')	79 \rightarrow 82(0.70)	2.79	444	0.0000		MLCT
4(A'')	78 \rightarrow 82(0.52)	2.89	429	0.0000		MLCT/LLCT
	75 \rightarrow 82(0.43)					IL
5(A')	77 \rightarrow 82(0.50)	3.04	408	0.0000		MLCT/LLCT
	80 \rightarrow 83(0.44)					MLCT/LLCT

(predominantly) a MO 73 \rightarrow MO 74 transition. The TDDFT results do not provide information on triplet–singlet absorption intensities since spin–orbit coupling effects are not included in current TDDFT approaches, which is the reason the oscillator strengths in triplet excited states are all zero. Spin–orbit coupling can mix singlet and triplet states, allowing the latter to acquire intensity in both absorption and emission. A second effect is that the triplet energies are shifted through coupling with higher singlet (or other triplet) states. For third-row transition metals, one⁴⁰ estimates the lowest triplet states to be lowered by ~ 0.2 – 0.3 eV from interactions with higher states through spin–orbital coupling. The TDDFT results should still provide a reasonable description of the overall orbital excitations that would be coupled in a subsequent spin–orbit treatment.

Comparison with Experimental Results. Combined with experimental data gathered over time, computed spectra may provide a reference for monitoring the behavior and possible degradation of the organic material in the device. The results of the TDDFT calculations for complexes **1**–**4** are compared with experimental absorption data (see Tables 4–7) and are depicted in Figures 6–9. The spectral assignment is based on a comparison of experimental band maxima with calculated energies of transitions with significant oscillator strengths,

including the simulated spectra figures. The principal experimental studies cited in the table have been carried out by Ming Zhang et al. in the Key Lab for Supramolecular Structure and Materials of Ministry of Education at Jilin University.

To independently check that these calculations produce reasonable results, the absorption spectra of the four complexes are simulated based on the TDDFT calculations (Figure 6–9). Here, each excited-state having $\times c4 \geq 0$ (as shown in Tables 4–7) was modeled as a Gaussian feature in the program OriginPro70. The numbers of maximum peaks in a certain region are decided as the number of peaks. Selection of the initial half-width estimate of each Gaussian was based upon the default values. Each graph of calculated molar absorptivity (Figures 6–9) shows a Gaussian corresponding to each of the 40 calculated singlet excited states. For example, for complex **1**, starting from the lowest transition calculated at 609 nm to 227 nm are taken into the simulation. It seems that the simulated spectra are all in reasonable agreement with the experimental spectrum. In fact, there have been numerous spectroscopic studies on Re(I) bpy complexes. Meyer and co-workers⁴¹ reported the near-UV–vis absorption spectrum of the “model” complex, (4,4'-X₂-2,2'-bipyridine)Re(CO)₃Cl. The absorption maxima are at relatively high energy and overlap with bpy-

TABLE 7: Selected Calculated Excitation Energies (E), Wavelengths (λ), Oscillator Strengths ($\times c_4$), and Dominant Excitation Character for Low-Lying Singlet (S_n) and Triplet (T_n) States of the Complex $\text{ReCl}(\text{CO})_3(4,4'\text{-2COOCH}_3\text{-bpy})$ and Comparison with Experimental Results

state	excitation	E_{cal} , eV	λ_{cal} , nm	$\times c_4_{\text{cal}}$	λ_{exp} , nm	character
Singlet Excited States						
1(A'')	103 \rightarrow 104(0.70)	1.75	707	0.0040		MLCT/LLCT
2(A')	102 \rightarrow 104(0.67)	1.95	635	0.0369		MLCT/LLCT
4(A')	103 \rightarrow 105(0.53)	2.48	500	0.0136		MLCT/LLCT
	101 \rightarrow 104(0.34)					MLCT
5(A'')	102 \rightarrow 105(0.70)	2.53	489	0.0211		MLCT/LLCT
9(A')	99 \rightarrow 104(0.66)	3.10	400	0.1237	433(CH ₂ Cl ₂)	MLCT/LLCT
13(A'')	99 \rightarrow 105(0.68)	3.65	340	0.0217		MLCT/LLCT
14(A')	100 \rightarrow 105(0.68)	3.67	338	0.1110		MLCT/LLCT
15(A'')	100 \rightarrow 106(0.57)	3.90	318	0.0379		MLCT/LLCT
19(A')	99 \rightarrow 106(0.67)	4.01	309	0.0326		MLCT/LLCT
22(A'')	97 \rightarrow 104(0.41)	4.12	301	0.1688	313(CH ₂ Cl ₂)	IL
	100 \rightarrow 106(0.35)					MLCT/LLCT
23(A'')	98 \rightarrow 105(0.60)	4.16	298	0.1244		MLCT/LLCT
	97 \rightarrow 104(0.26)					IL
25(A')	102 \rightarrow 109(0.57)	4.28	290	0.0169		MLCT/LLCT
	103 \rightarrow 108(0.23)					MLCT/LLCT
29(A')	97 \rightarrow 105(0.50)	4.57	271	0.0149		IL
	98 \rightarrow 106(0.33)					MLCT/LLCT
30(A'')	102 \rightarrow 111(0.49)	4.66	266	0.0159		MLCT/LLCT
	101 \rightarrow 108(0.39)					MLCT
35(A'')	102 \rightarrow 111(0.42)	4.90	253	0.0120		MLCT/LLCT
	103 \rightarrow 112(0.19)					MLCT/LLCT
37(A')	102 \rightarrow 110(0.45)	5.03	246	0.0190		MLCT/LLCT
	99 \rightarrow 109(0.19)					MLCT/LLCT
	100 \rightarrow 108(0.18)					MLCT/LLCT
38(A'')	97 \rightarrow 106(0.67)	5.04	246	0.1942		IL
Triplet Excited States						
1(A'')	103 \rightarrow 104(0.71)	2.06	602	0.0000		MLCT/LLCT
2(A')	102 \rightarrow 104(0.72)	2.12	586	0.0000		MLCT/LLCT
3(A')	103 \rightarrow 105(0.70)	2.79	444	0.0000		MLCT/LLCT
4(A')	101 \rightarrow 104(0.71)	2.89	429	0.0000		MLCT
5(A'')	100 \rightarrow 104(0.69)	3.04	408	0.0000		MLCT/LLCT

based, $\pi\pi^*$ bands. The lowest energy absorption band (or shoulder) is assigned to a MLCT transition ($\text{Re}^I \rightarrow \text{bpy}$). The MLCT absorption spectra are sensitive to the nature of the substituents on the polypyridyl ligand. The electron-donating substituents increase the energy of the π^* ($4, 4'\text{-X}_2\text{-bpy}$) acceptor orbitals to a greater degree than they do the $d\pi$ (Re) orbitals. At the same time, Keith et al.¹³ thought that the absorption bands for complex $\text{ReCl}(\text{CO})_3(4,4'\text{-[bis-[2,5-(dimethoxyphenyl)ethynyl]}-2,2'\text{-bipyridine]})$ are dominated by IL $\pi\pi^*$ transitions although the low-energy absorption band contains a contribution from an MLCT transition. They considered there is configuration interaction between the IL $\pi\pi^*$ and MLCT transitions, and because of such configuration interactions, the low-energy transition will have characteristics that are a composite of the two transitions.

For $\text{ReCl}(\text{CO})_3(\text{bpy})$, the calculated excitation energy (3.35 eV) with strong intensity (0.0872) for S_9 (A') arising from the mixed MLCT $d \rightarrow \pi^*$ (bpy)/LLCT $p \rightarrow \pi^*$ (bpy) excitation is close to the onset of absorption (3.20 eV, 387 nm). Experimentally, this band was assigned to MLCT $d \rightarrow \pi^*$ (bpy); TDDFT results label this mixed character. This should be qualified by the fact that 5d orbitals are highly mixed with p character on the halide Cl as discussed earlier. The other higher-energy absorption peaks at 4.23 eV (293 nm) occur in the same region of a singlet state S_{22} (A'') calculated at 4.54 eV with large oscillator strength, which corresponds to $\pi\pi^*$ excitation and considerable MLCT/LLCT. In general, the simulated spectra exhibit slight blue shifts compared with experimental ones.

For $\text{ReCl}(\text{CO})_3(4,4'\text{-2Br-bpy})$, the two relative higher-energy absorption at 256 (4.84) and 316 nm (3.92 eV) corresponds to $\pi\pi^*$ excitation, and the latter mix considerable MLCT/LLCT,

at calculated values of 293 (4.23) and 320 nm (3.88 eV) from $S_{21}(A'')$ and $S_{13}(A'')$, respectively. The absorption peak at 410 nm (3.02 eV) agrees well with the lower singlet state S_9 (A') arising from d (Re) and p (Cl) to π^* (bpy) excitation (3.26 eV).

From Figure 3b, we noted that at nearly 250–260 nm, there exists one shoulder peak at 260 nm (4.76 eV) observed in experiment for $\text{ReCl}(\text{CO})_3(4,4'\text{-2CH}_3\text{-bpy})$, but because in this region the oscillator strength is similar, it cannot present in the simulated spectra. We still assign it to mixed MLCT/LLCT character arising from S_{25} (A''). High-energy absorption peaks at 290 nm (4.28 eV) occur in the same region of S_{20} (A'), and the low-energy ones at 380 nm (3.26 eV) correspond to S_9 (A'), and they both arise from MLCT $d \rightarrow \pi^*$ (bpy)/LLCT $p \rightarrow \pi^*$ (bpy).

$\text{ReCl}(\text{CO})_3(4,4'\text{-2COOCH}_3\text{-bpy})$ have two absorption bands. The low-energy one at 432 nm (2.87 eV) corresponds to S_9 (A') at 400 nm (3.10 eV) arising from MLCT $d \rightarrow \pi^*$ (bpy)/LLCT $p \rightarrow \pi^*$ (bpy). The other calculated broad one 313 nm (3.96 eV) corresponds to S_{22} (A'') at 301 nm (4.12 eV) arising from IL excitation with considerable MLCT/LLCT. The calculated absorption bands exhibit slight blue shifts compared with experiment.

Furthermore, the spectra illustrate the considerable effect that different substituents at the chromophoric ligand have on the spectral properties of the complexes. The main bands are red shifted in the complexes substituted by the electron-withdrawing groups $-\text{Br}$ and $-\text{COOCH}_3$ and blue shifted in the electron-releasing group $-\text{CH}_3$ substituted complex. This variation is consistent with the orbital characters analyzed above.

Lowest Triplet States. The lowest triplet states of the four molecules, T_1 , have been fully optimized by carrying out ab

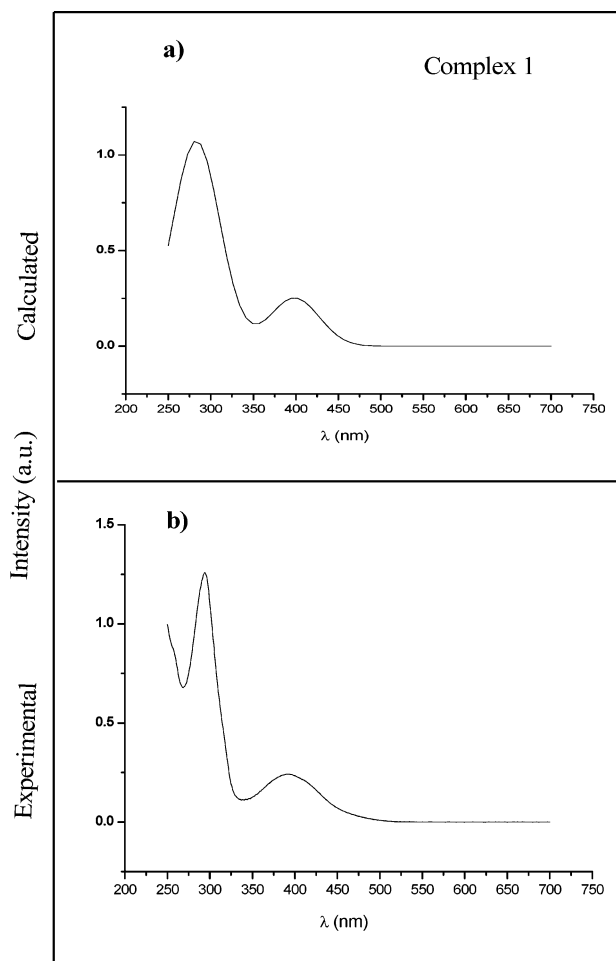


Figure 6. Comparison of calculated (a) and experimental (b) absorption spectra of the complex $\text{ReCl}(\text{CO})_3(\text{bpy})$. Experimental absorption data are measured in CH_2Cl_2 solution.

initio CIS. The optimized structure parameters are shown as Table 8. The geometrical parameters for excited states have small differences from those of the ground state, and a general elongation of all the metal–ligand bond lengths is observed. For example, the $\text{Re}-\text{N}$ bond distance in the triplet state is longer than that in the ground state by about 0.01–0.05 Å, and the distance between the metal atom and carbon atom ($\text{Re}-\text{C}$) is also longer in the triplet state than in the ground state. As far as nonmetal bonds are concerned, some decrease and some increase.

We can predict the differences of the bond lengths between the ground (S_0) and triplet excited state (T_1) from nodal patterns.⁴² Because the triplet state corresponds to an excitation from the HOMO to the LUMO in all considered complexes, we explore the bond-length variation by analyzing the HOMOs and LUMOs (see Figure 10). In the following section, we employed the complex $\text{ReCl}(\text{CO})_3(4,4'\text{-dibromo-bpy})$ as the example to analyze the bond lengths changes (see Table 6).

The LUMO has nodes across the $\text{N}(1, 1')-\text{C}(2, 2')$, $\text{C}(4, 4')-\text{C}(5, 5')$, $\text{C}(2, 2')-\text{C}(3, 3')$, and $\text{C}(5, 5')-\text{C}(6, 6')$ bonds, but the HOMO is bonding in these regions. Therefore one would expect elongation of these bonds; Table 9 shows that these bonds are in fact considerably longer in the excited state. The HOMO has a node across the $\text{C}(3, 3')-\text{C}(4, 4')$ and $\text{C}(2)-\text{C}(2')$ bonds, while the LUMO is bonding. The data in Table 9 confirms the anticipated contraction of these bonds.

The spin density map for the lowest triplet states of complex 2, reported in Figure 11, shows well the sharing of the two

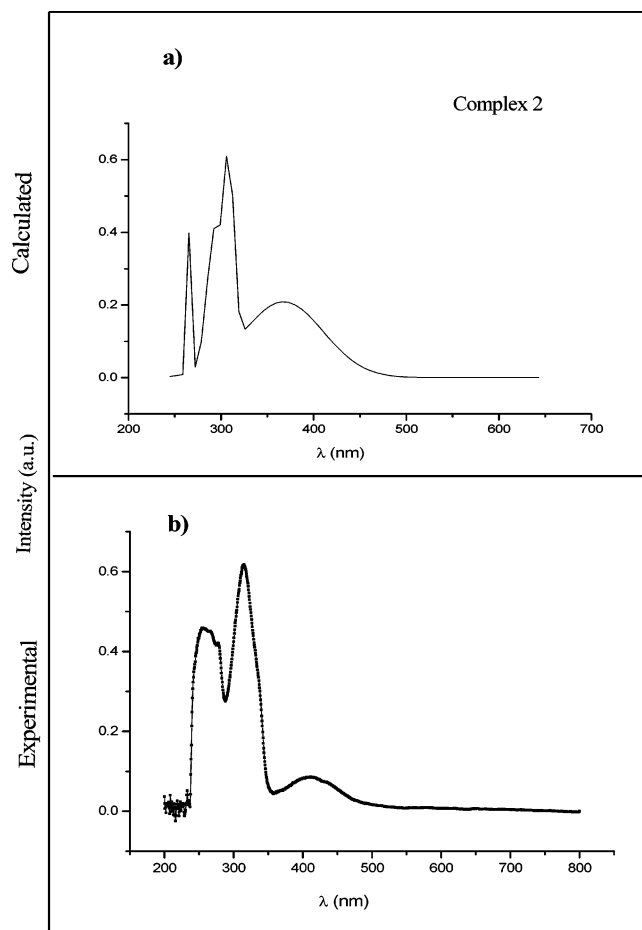


Figure 7. Comparison of calculated (a) and experimental (b) absorption spectra of the complex $\text{ReCl}(\text{CO})_3(5,5'\text{-2Br-bpy})$. Experimental absorption data are measured in CH_2Cl_2 solution.

unpaired electrons among the metal (d orbital) and group Cl and the bipyridine ligand (π^* orbital), with smaller contributions from the group CO. The four complexes have the similar behaviors. In general, all the geometry variations are consistent with the occupation of the π^* orbitals of the bpy ligands depicted in Figure 10, and the variation path follows the bonding/antibonding scheme in such plots.

Emissive Spectrum. We employed the ab initio CIS method to optimize the lowest triplet state geometry, and at the excited triplet state geometry, TDDFT is used to calculate the emission spectra. The long-lived decay lifetime from experiment revealed that the photoluminescence for the four complexes is assigned as triplet-state charge transfer, and thus we only optimize the lowest triplet state (T_1) and calculate the triplet emission spectra. The results of the TDDFT calculations for complexes 1–4 are summarized in Table 10 and are compared with experimental emission data as depicted in Figure 12.

We list the lowest three triplet excited states and the photoluminescence for each complex corresponding to the lowest triplet T_1 , which consists of the transition from HOMO to LUMO, and thus assigned as the mixed character between MLCT [$d(\text{Re}) \rightarrow \pi^*(\text{bpy})$] and LLCT [$p(\text{Cl}) \rightarrow \pi^*(\text{bpy})$]. Shih-Sheng Sun et al.⁴³ account the participation of these excited states largely for the decreasing emission quantum yields and shorter lifetimes of the emissive³ MLCT excited states. Recent studies on rhenium(I) carbonyl systems have indicated that

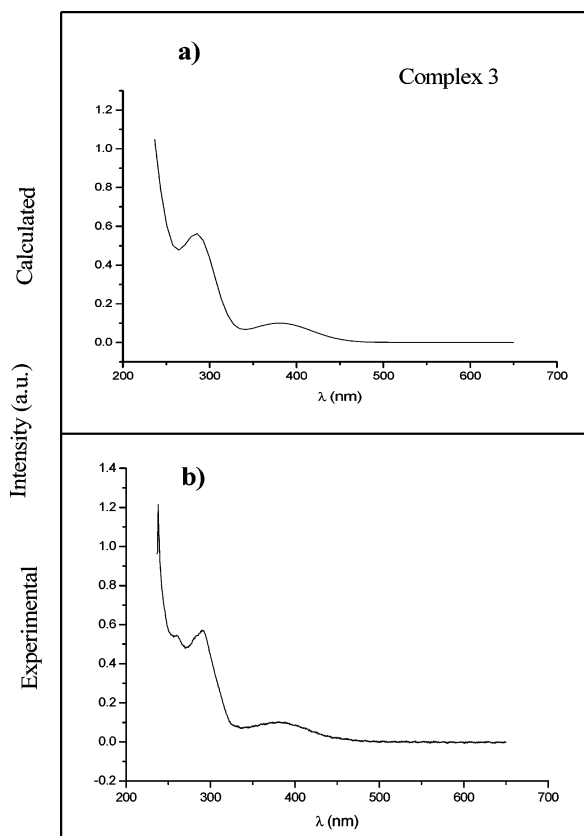


Figure 8. Comparison of calculated (a) and experimental (b) absorption spectra of the complex $\text{ReCl}(\text{CO})_3(4,4'\text{-2CH}_3\text{-bpy})$. Experimental absorption data are measured in CH_2Cl_2 solution.

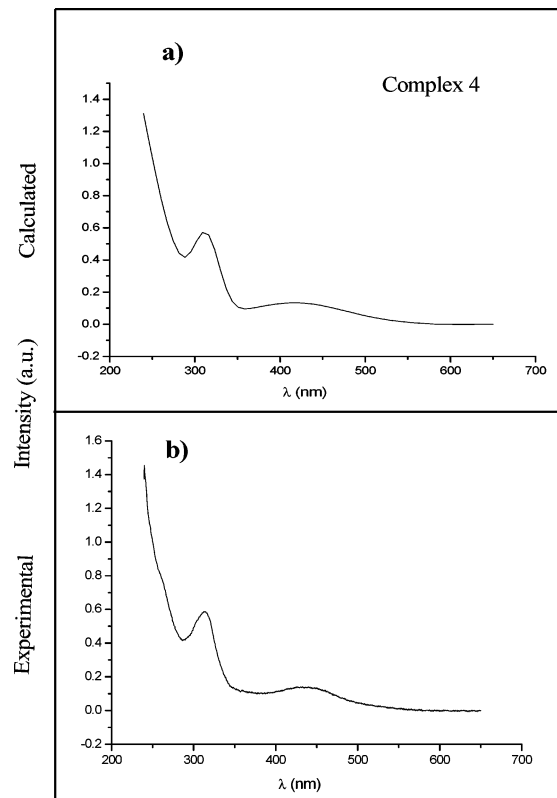


Figure 9. Comparison of calculated (a) and experimental (b) absorption spectra of the complex $\text{ReCl}(\text{CO})_3(4,4'\text{-2COOCH}_3\text{-bpy})$. Experimental absorption data are measured in CH_2Cl_2 solution.

emissive excited states of Re diimine complexes often have mixed MLCT and LLCT and, sometimes, IL characters. The

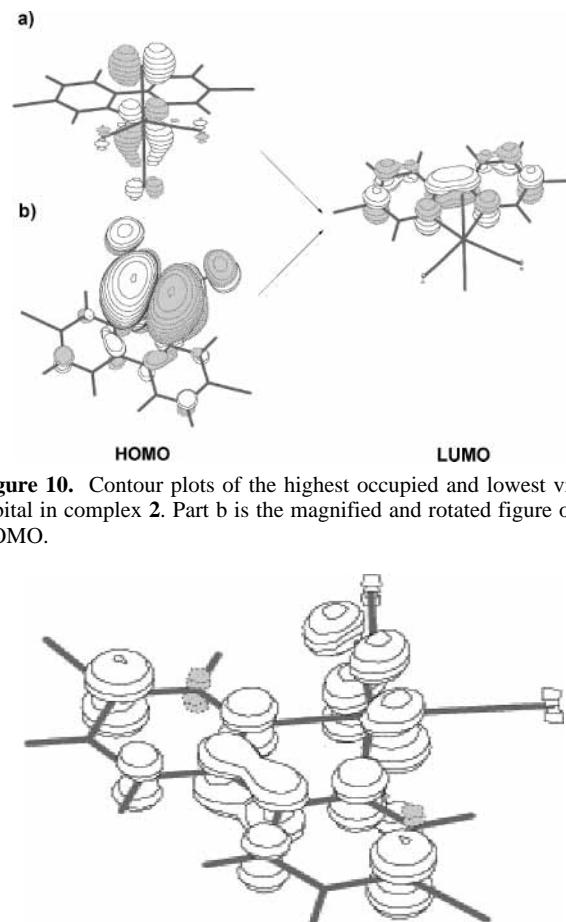


Figure 10. Contour plots of the highest occupied and lowest virtual orbital in complex 2. Part b is the magnified and rotated figure of the HOMO.

Figure 11. Spin-density map for the lowest triplet state of $\text{ReCl}(\text{CO})_3(5,5'\text{-2Br-bpy})$.

TABLE 8: Main Geometrical Parameters (\AA) of the Lowest Triplet State of the Four Complexes

bonds	1	2	3	4
$\text{R}[\text{Re}-\text{N}(1,1')]$	2.1789	2.1852	2.1852	2.1863
$\text{R}[\text{Re}-\text{C}(1,1')]$	1.9596	1.9594	1.957	1.9598
$\text{R}[\text{Re}-\text{C}(7)]$	1.9303	1.9298	1.9266	1.9301
$\text{R}[\text{Re}-\text{Cl}]$	2.5612	2.5658	2.5728	2.5674

TABLE 9: Calculated Bond Lengths of the Lowest Triplet State (T_1) and the Comparison to Corresponding Ground State Geometries (S_0) for Complex (2)

bond	S_0	T_1
$\text{R}[\text{N}(1,1')-\text{C}(2,2')]$	1.3733	1.4155
$\text{R}[\text{C}(4,4')-\text{C}(5,5')]$	1.4009	1.4332
$\text{R}[\text{C}(3,3')-\text{C}(4,4')]$	1.4005	1.367
$\text{R}[\text{C}(2,2')-\text{C}(3,3')]$	1.4075	1.4545
$\text{R}[\text{C}(5,5')-\text{C}(6,6')]$	1.4049	1.4058
$\text{R}[\text{C}(2)-\text{C}(2')]$	1.473	1.4082

detailed excited-state character influences relaxation pathways and rates.⁴⁴

Apart for transition character discussed above, our calculations also reproduce (even if not quantitatively) the red or blue shift, which affected by substituents changes observed in going from the four complexes having identical chemical environments. Consistent with experimental data, analysis for molecular orbitals and absorption spectra, an electron-releasing group leads to a blue shift and an electron-withdrawing group leads to red shift. The calculated emission peak at 757 nm (1.64 eV) for complex 3 is blue shifted, and the peaks at 797 (1.56) and at

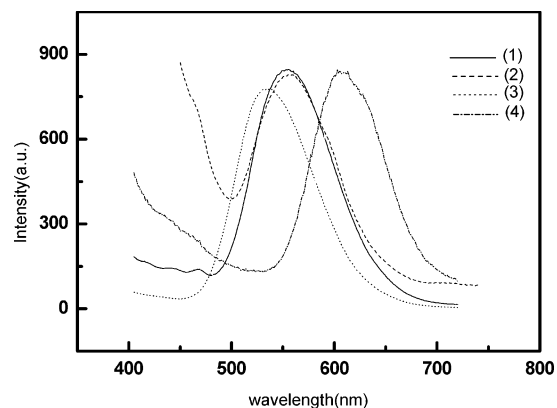


Figure 12. Experimental photoluminescence spectra of complexes 1–4 measured in polycarbonate film.

TABLE 10: Computed TDDFT Excitation Energies (nm) for the Lowest Triplet States of All the Considered Complexes with Experimental Data

	excitation	E_{cal} , eV	λ_{cal} , nm	E_{exp} , eV	λ_{exp} , nm	character
ReCl(CO) ₃ (bpy)						
T ₁ (A'')	73 → 74	1.66	743	2.24	555	MLCT/LLCT
T ₂ (A')	72 → 74	1.71	726		540 ^[36]	
T ₃ (A')	71 → 74	2.23	557			
ReCl(CO) ₃ (5,5'-dibromo-bpy)						
T ₁ (A'')	79 → 80	1.56	797	2.23	556	MLCT/LLCT
T ₂ (A')	78 → 80	1.69	734			
T ₃ (A')	77 → 80	2.25	552			
ReCl(CO) ₃ (4,4'-dimethyl-bpy)						
T ₁ (A'')	81 → 82	1.64	757	2.33	533	MLCT/LLCT
T ₂ (A')	80 → 82	1.70	729		530 ^[36]	
T ₃ (A')	79 → 82	2.33	531			
ReCl(CO) ₃ (4,4'-dimethylformyl-bpy)						
T ₁ (A'')	103 → 104	1.51	832	2.05	605	MLCT/LLCT
T ₂ (A')	102 → 104	1.61	768			
T ₃ (A')	101 → 104	2.15	576			

832 nm (1.51 eV) for complexes **2** and **4**, respectively, are red shifted compared with complex **1**, the peak at 743 nm (1.66 eV).

From Table 10, there exist some deviations between calculated and experimental data for emission spectra. Concerning the reasons of producing such deviations, we want to stress several points. First is the solvent effect. Usually the studied system is put in a gas phase environment for a quantum method, whereas it is laid in a solution environment in experiment, and our limited studies did not take the solvent effects into account and thus not add effects rectification. Second is the limit of the quantum chemical approach. CIS was employed to optimize the lowest triplet excited state, and despite the name, CIS represents for excited states a general zeroth-order method and does not include double and third excitations, whereas it is necessary for discussing electron correlation. Other DFT calculations overestimate halide contributions in metal–halide complexes, especially if a low-valent metal atom is involved.³⁴ In addition, the more detailed interpretation of phosphorescence properties will await the inclusion of spin–orbit coupling effects that are not included in these TDDFT results. We hope to investigate these effects in future studies.

Conclusions

In this paper, we have applied DFT methods to analyze ground- and excited-state properties of the complex **1** and the

related complexes **2**, **3**, and **4**. The ground-state structures are calculated by B3LYP density functional and are in good agreement with available crystallographic studies. Most of the highest-occupied orbitals are Re (5d) in character, with an almost equal admixture of Cl group with exception for HOMO₋₂, which is dominated by Re (5d). The spin densities in the cations show over 0.50 electrons on the metal center and over 0.30 electrons on the group Cl for each molecule, which is basically consistent with this analysis. The lowest virtual orbitals are totally π^* (bpy) in character, as also reflected in the spin densities in the anion.

Forty singlet and thirty-five triplet excited states are examined using TDDFT. The low-energy absorption, which the light-emitting is concerned about, is all assigned as the mixture of MLCT [d(Re) → π^* (bpy)] and LLCT [p(Cl) → π^* (bpy)]. Looking at Table 3, a good agreement between the computed absorption bands and the reported experimental bands in the ref 35 can be found. The lowest triplet excited-state structures of the four molecules are obtained using CIS method. The emission band has been supposed to be an electronic transition between the ground and the triplet excited state T₁ with the character of the mixture of MLCT and LLCT for all considered complexes.

The results of absorption and emission spectral fitting give insight into the effect that changes in X have on the structure of the bpy acceptor ligand. As might have been expected, the effect is less profound at the d orbitals than at the π^* orbitals since the substituent changes are made at the ligand. The electron-releasing group –CH₃ leads to both absorption and emission spectra blue shifts due to the increased energy of LUMO which results in the increased HOMO–LUMO band gap, and electron-withdrawing groups –Br and –COOCH₃ lead to red shift due to the decreased energies of LUMOs which decrease the energy band gaps. Furthermore, we find large Stoke shifts for the four complexes. The results indicate that application of TDDFT calculations is reliable for studying the system containing transition metals, but at the same time, it has some drawbacks, such as overestimating the compositions of halides in low-valent metal complexes and without considers spin–orbit coupling effects.

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