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Unexpected Disruption of the Dimensionality-driven Two-Photon Absorption Enhancement within a Multipolar Polypyridyl Ruthenium Complexes Series.

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The dimensionality-driven two-photon absorption (2PA) enhancement effect is investigated in a series of functionalized bipyridyl Ru-complexes. Our design strategy leads to very high 2PA responses up to ~ 1500 GM. However, we highlight that the 2PA performance *vs.* dimensionality correlation reaches an unexpected limit stemming from 'anti-cooperative' interchromophoric couplings.

Molecules with optimized 2PA efficiency are highly demanded in the fields of photonics and biology¹. Indeed, 2PA materials are extensively widespread across many applications including microfabrication², high density data storage^{3, 4}, bio-imaging⁵⁻⁷ or photodynamic therapy⁸. Several critical factors determine the 2PA performances of full organic systems : the symmetry of the structure, the extent of the π -conjugation length, the occurrence of an efficient intramolecular charge transfer (ICT) both modulated by the nature and the position of electron donor and acceptor groups within the designed molecule9. Alternatively, coordination chemistry offers a versatile strategy for designing two-photon absorbers whose NLO properties are associated to charge-transfer processes such as intraligand charge-transfer (ILCT) and metal-ligand charge-transfer (MLCT)¹⁰⁻¹³. The metal center acts as a powerful template for self-organizing 2PA chromophores in a three-dimensional arrangement^{9, 14}. Very high 2PA amplifications can then be observed due to the synergistic interplay and extent of electronic couplings between the organic branches¹⁵. In line with this strategy, we designed a series of homoleptic and heteroleptic $[RuL_n(bpy)_{3-n}]^{2+}$ (n = 1, 2, 3) complexes whose geometrical structuration accounts for two main objectives : (i) Quantifying the 2PA enhancement upon a methodical

Electronic Supplementary Information (ESI) available: Additional figures and tables. Experimental procedures. Synthesis and characterization of new compounds

change of the ligand conjugation length and the complex coordination number, n. (ii) Identifying the cooperative and/or 'anti-cooperative' electronic interactions emerging from this incremental metal-template building strategy. With this respect, we have designed several 2,2'-bipyridinebased ligands which are 4,4'-substituted by distinctive electron-rich arylamine groups such as triphenylamine (T), carbazolyl (C) and diphenylaminofluorenyl (F) subunits through a vinylene linker (Scheme 1). These peripheral π conjugated subunits have been selected since their electron-donor ability as well as their π -conjugation length can be regulated through a planarization effect. Such a structural constraint together with the progressive increase of 3D-dimensionality will give rise to competing interactions between 'through-bond' and 'through-space' couplings^{16, 17} within the complexes series leading to a severe disruption in the dimensionality-induced 2PA enhancement effect.



Scheme 1. Molecular structures of the Ru-complexes.

The absorption spectra of the ligands along with their corresponding Ru-complexes in tetrahydrofuran (THF) are shown in **Figure 1**. The absorption of Ru(bpy)₃²⁺ used as a reference complex is also displayed. The spectroscopic data are summarized in **Table 1**. All the ligands present similar spectral features particularly with the presence of a longest wavelength absorption band located in the 330-450 nm range. This band is mainly dominated by the $S_0 \rightarrow S_1$ electronic transition which implies a long range electronic

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delocalization within the π -conjugated structure with a significant ICT character from the peripheral donor amino groups to the central acceptor bipyridine moiety^{12, 18}. This CT band is both sensitive to the nature of the donor subunit and the conjugation length. For instance, the better donor ability of the triphenylamine moiety as compared to the carbazole one^{19, 20} leads to a clear bathochromic effect on going from **C** to **T**. Moreover, such a CT process is all the more amplified when considering the dipole moment change between the ground and the excited emitting state.



Fig.1 Absorption spectra (full lines) of the ligands and their corresponding Ru-complexes in THF. Dashed lines: Absorption spectrum of $Ru(bpy)_{3}^{2+}$ in acetonitrile.

As shown in Figures S1-S3, all the ligands are fluorescent and undergo very large amplitude Stokes shifts up to one order of magnitude on going from apolar to highly polar solvents. Based on these solvatochromic measurements, the dipole moment change between ground and relaxed excited state ($\Delta \mu_{01}$) has been estimated using Lippert-Mataga formalism^{21, 22} (see Figure S4 and Table S1). Therefore, we show that all the ligands display very high $\Delta\mu_{01}$ with values of 13.6 D for **C**, 18.5 D for **T**, and 27.9 D for **F**. Note that $\Delta \mu_{01}$ is multiplied by a factor 2 between **C** and **F** which confirms the very strong ICT enhancement promoted by our proposed structural changes. Analogously, the coordination to Ru²⁺ strongly influences the ligands absorption spectra (Figure 1). All the metal-complexes display a well distinctive band in the 270-320 nm range similarly to that observed for $Ru(bpy)_3^{2+}$. This band presumably corresponds to the ligand-centred (LC) $\pi \rightarrow \pi^*$ transitions²³ (bipyridine-centred transitions). Note that this LC band shifts to the red region and increases in intensity with the coordination number of functionalized-bipyridyl ligands (n). Similar effects can be observed when considering the lowest energy band which progressively developed in the 350-630 nm range. This very broad and structured band encompasses several electronic transitions^{10, 12}: the intraligand charge transfer (ILCT)

transitions and the metal-to-ligand charge transfer (MLCT) ones. The ILCT band is located at the blue side of the lowest energy band and increases in intensity with n.

	$\lambda_{IPA}(\epsilon_{abs})$	λ_{2PA}	δ_{MAX}	$\delta_{MAX} / N_{eff.}^2$
	$/ \text{ nm} (10^3 \text{ M}^{-1} \text{ cm}^{-1})$	/ nm	/ GM	/GM
С	362 (51)	705	200	0.25
RuC ₁	415 (34.5), 469 (29.9)	795	395	0.27
RuC ₂	415 (80.2), 485 (55.2)	840	655	0.37
RuC ₃	420 (141.4), 503 (86.1)	855	1381	0.58
Т	392 (46.1)	780	187	0.35
RuT ₁	455 (39.5), 489 (37.3)	855	311	0.38
RuT ₂	445 (65.7), 505 (58.9)	855	822	0.67
RuT ₃	437 (94.5), 504 (82.0)	855	1465	0.91
F	395 (76.2)	800	208	0.22
RuF ₁	450 (66), 493 (61.7)	855	596	0.48
RuF ₂	426 (100.1), 507 (88.0)	870	889	0.44
RuF ₃	429 (138.9), 502 (122.1)	870	1315	0.46

Table. 1 1PA and 2PA properties of the ligands and Ru-complexes (Solvent : THF). The experimental uncertainty for δ is ± 15 %.

When comparing the free ligand to its first heteroleptic complex, the ICT band clearly redshifts in concomitance with a substantial decrease in intensity. Whereas this bathochromic effect can be confidently attributed to the electron withdrawing exaltation of the branched bipyridine coordinated to Ru²⁺, the observed band decrease should be ascribed to a severe reduction of the conjugation length within the entire structure of the ligand probably due to a loss in planarity of the 2,2'-bipyridine which undergoes a trans-to-cis conformation change upon complexation. The MLCT bands which strongly overlap with the ILCT ones are typically positioned at the low energy side of the complexes absorption band. For all the heteroleptic complexes, two distinctive MLCT transitions can be observed due to the presence of two types of bipyridine ligands: i) The $d\pi(Ru) \rightarrow$ $\pi^*(bpy)$ transitions are presumably located in the 430-460 nm range as suggested by the MLCT band of $Ru(bpy)_{3}^{2+}$ (see **Figure 1**). This band has a moderate intensity ($\varepsilon_{MAX} \approx 14000$) M⁻¹ cm⁻¹) and can be distinguished on the structured absorption band of the heteroleptic complexes. ii) The dπ(Ru) \rightarrow π^* (bpy-substituted) transitions should correspond to the broad and intensive band at the red edge of the absorption spectrum. This red-shifted MLCT band grows incrementally with n in the same manner as for the ILCT band. The luminescence properties of the metalcomplexes in THF are presented in Figure S5 and in Table S2. All the complexes are weakly emissive with a luminescence quantum yield (Φ_L) lower than 10⁻² in nondegassed THF. The luminescence is very sensitive to oxygen with values of Φ_l enhanced by factor ~ 10 when operating in N_2 -saturated THF. This is an indication that the Published on 15 September 2020. Downloaded by Cornell University Library on 9/15/2020 12:46:18 PM

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luminescence of the complexes arises from the triplet ³MLCT state. It is noteworthy that the luminescence bands are strongly redshifted with respect to that of $Ru(bpy)_3^{2+}$. This clearly underlines the electronic effect of the bipyridine functionalization. Indeed, the 4, 4'-substitution of bipyridine with electron donating groups stabilizes the ³MLCT state^{24, 25}. This electronic configuration implies a drift of electron density from the metal to the ligands with a dominant electronic localization on the substituted bipyridines. This assumption is both consistent with the observed luminescence spectral changes and the strong stabilization of the ³MLCT state energy ($\Delta E = 0.18-0.26 \text{ eV}$) as compared to Ru(bpy)₃²⁺. Increasing the dimensionality of the complex with functionalized bipyridines also induces a 3-fold enhancement of the complexes emissivity (see Table S2). This luminescence 'switch-on' of the 3D complexes presumably stems from a shielding effect of the peripheral donor branches from the dynamic oxygen quenching.

Increasing the dimensionality of metal-complexes also impacts the 2PA properties. Figure 2 shows 2PA spectra of the metal-complexes in THF, alongside the corresponding one-photon absorption (1PA) spectra. Figure S6 also presents the 1PA and 2PA spectra of the ligands in THF. According to our spectral resolution, the ligands display a 2PA band in the 700-900 nm range which reasonably matches their respective lowest energy 1PA band. However, the 2PA cross-sections are moderate for all ligands with δ_{MAX} hardly reaching ~ 200 GM whatever the conjugation length or the donor group nature (see Table 1). Such modest values presumably originate from the very close centrosymmetric geometry of the ligands (i.e. $D-\pi$ -A- π -D) leading to a theoretically two-photon forbidden character of the S_0 \rightarrow S_1 transition. By contrast, the complexation with Ru²⁺ induces a very strong amplification of the 2PA cross-sections by more than a factor 6 between the free ligands and their respective homoleptic complexes. Moreover, increasing the number of π -functionalized bipyridines gives rise to very high δ_{MAX} up to 1465 \pm 220 GM for RuT₃. Such a strong 2PA enhancement proceeds through a synergy effect which combines an effective increase of the intramolecular charge transfer per branch and the cooperative interbranch coupling within the entire multipolar structure^{13, 26}. As illustrated in Figure 2, the 2PA spectra of all the complexes feature a very intensive band which nicely coincides with the ILCT transitions and a second 2PA resonance in the lowest energy region which matches the MLCT transitions. However, the evolution of the relative intensities of these two bands all along our complexes series suggests that the MLCT resonance increases to the detriment of the ILCT one especially for the F series. Of particular interest, the ILCT-to-MLCT band ratio undergoes very strong changes which both depend on the coordination number and the nature of the ligand. For

instance, this ratio is about ~ 8 for RuC_3 but progressively decreases to 3.6 and 1.9 for RuT_3 and RuF_3 respectively.



Fig.2 1PA (full lines) and 2PA (symbols) spectra of Ru-complexes in THF. The one-photon molar absorption is shown at the 1PA wavelength (lower horizontal scale). The 2PA cross section is shown as a function of the excitation laser wavelength (upper horizontal scale).

Such a monotone spectral effect from **C** to **F** series illustrates the increase of dipole-dipole interchromophoric couplings¹⁶. However, in the present case, the magnitude and the orientation of the interbranch cross-talk within the complexes give rise to an unexpected disruptive effect of the 2PA enhancement course with dimensionality. In order to quantify such a disruption, one should first consider that δ_{MAX} increases quadratically with the effective number of π -electrons (N_{eff}) contributing to the nonlinear response^{27, 28}. Therefore, δ_{MAX} has been normalized by N_{eff}² in order to access to the intrinsic 2PA cross-section which indeed accounts for the real 2PA performance of a dye. For determining N_{eff}, we used the relevant counting method

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proposed by Kuzyk *et al*^{27, 28} which consists in calculating N_{eff} by geometrically weighting the number of electrons in each conjugated path of the molecular system. According to this methodology, **Figure 3** shows a histogram showing the 2PA performance of all the chromophores as a function of the ligands conjugation length and the coordination number. Note that these two coordinates respectively depict the 'through-bond' (*via* conjugation) and 'through-space' (*via* dipole-dipole couplings) interactions^{15, 16}.



Fig.3 Intrinsic 2PA performance of all the dyes as function of ligands π -conjugation length and complexes coordination number.

According to this methodology, it can be inferred that the triphenylamine series exhibits the optimal behaviour in term of 2PA cooperative effect. Indeed, δ/N_{eff} ² increases gradually with the coordination number (n) and RuT₃ exhibits the highest 2PA performance amongst its homologues. The 2PA performance of the carbazole series presents a similar behaviour but with a much weaker 2PA amplification. Finally, one of the most striking results concerns the diphenylaminofluorenyl series whose 2PA performance rapidly saturates for n = 2 and 3. Even though RuF₁ constitutes the most promising 2PA active candidate as compared to RuC1 and RuT1 with the highest 2PA enhancement (up to 118 %), it is clearly demonstrated that the further addition of F ligands has no effect on the 2PA performance. By contrast to the other ligands, the increase of interchromophoric interactions emerging from the close proximity of F within RuF2 or RuF3 paradoxically generates 'anti-cooperative' effects¹⁶ which collapse the 2PA performance. These detrimental interactions should originate from the crowded and rigid character of F which drastically limits the fine orientation and the relative positioning of the polar branches within the complex. With this respect, there is no doubt that T appears as an optimal prototype which displays the good compromise between large dipole moment change and reasonable flexibility within a sterically constrained architecture. Such structural conditions undoubtedly establish a guideline for the future

design of new multipolar metal-complexes allowing a real gain in term of 2PA performance.

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