Iron and Ruthenium Alkynyl Complexes with 2-Fluorenyl Groups: Some Linear and Nonlinear Optical Absorption Properties

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Abstract: Four new mononuclear alkynyl complexes featuring a terminal 2-fluorenyl group, namely $Fe(n^5-C_5Me_5)(CO)_2[C=C(2 C_{24}H_{25}$)] (2) $Ru(\kappa^2-dppe)_2(CI)[C\equiv C(2-C_{21}H_{25})]$ (3). $Ru(\kappa^2$ dppe)₂[C=C(4-C₆H₄NO₂)][C=C(2-C₂₁H₂₅)] (4), and [Fe(η^{5} -C₅Me₅)(κ^{2} -(**5**[PF₆]), dppe){C=C(C₅H₄N)-CH₂(2-C₁₃H₉)}][PF₆] have been synthesized and characterized and their redox, absorption and emission properties have been studied. For the two ruthenium derivatives (3-4), these studies are complemented bv spectroelectrochemical investigations, Z-scan measurements and DFT calculations. Fluorimetric studies reveal that these compounds are poorly or not luminescent, and, when luminescent, that the weak luminescence detected most likely originates from a higher lying (LC) excited state presumably located on fluorene. Finally, the thirdorder NLO properties of 3 and 4 are reported. It is shown that the bis-alkynyl complex 4 is significantly more active than 3 and that both compounds exhibit two-photon absorption (TPA) around 860-1050 nm, with TPA cross-sections above 350 GM. In addition, it is shown that both species should give rise to a marked switching of their cubic nonlinear optical (NLO) properties in this spectral range upon oxidation.

electroluminescent displays^[1] for more than forty years.^[2] Among them, redox-active iron(II) or ruthenium(II) alkynyl complexes were identified very early as outstanding electrophores.^[3] These species, which were initially demonstrated to constitute key building blocks for molecular electronics,^[3f,4] have revealed more recently a very promising potential in molecular photonics.^[5] largely attributable to their remarkable electrochromism,^[6] which extends to their nonlinear optical (NLO) properties.^[7] So far, however, only a handful of investigations have been concerned with their luminescence.^[8] The latter appears usually to be weak or non-existent, even when a good luminophore is incorporated into the alkynyl ligand.^[8b,8e,f,9] This is regrettable, since luminescence, in association with the large cubic NLO properties exhibited by some of these compounds (notably two-photon absorption), would constitute a highly attractive feature that would significantly enlarge the applied scope of these molecules.[10]

interest in diverse fields ranging from photovoltaic conversion to

Introduction

Inorganic or organometallic photo-active molecules presenting facile electron-transfer capabilities have attracted sustained

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Scheme 1. Selected Alkynyl Complexes Containing 9H-Fluorenyl Units.

In this context, we have recently reported a series of $[Fe(\eta^5-C_5Me_5)(\kappa^2-dppe)]$ -based organoiron complexes containing 2ethynylfluorenyl ligands such as **1** (Scheme 1), which present strong (and redox-switchable) third-order NLO activities around 800 nm.^[11] Remarkably, when excited around 300 nm, these Fe(II) chromophores are weakly emissive, but not from their first excited state, with luminescence quantum yields around a few percent (0.4 % for **1**). In line with previous proposals in the

literature,^[8c,8e,f] a redox trapping mechanism of the photogenerated fluorene-centred (LC) excited state can be tentatively proposed to rationalize their weak and unusual fluorescence (Scheme 2). This proposal, which rests on the statement that Fe(II) endgroups can be readily oxidized (and reduced with considerable difficulty), has however not been experimentally challenged so far.



Scheme 2. Proposed Redox-Trapping Mechanism of the Fluorene-Based Luminescence in 2-Ethynylfluorenyl Complexes such as 1.

According to Marcus theory,^[12] the rate (k_{eT}) of any photoinduced electron transfer process should be strongly influenced by the redox potentials of the redox-active sites experiencing this electron transfer. Based on eq. 1, the detection of some luminescence indicates that the non-radiative decay (k_{eT} + $k_{\rm NR}$) does not completely overcome the radiative decay ($k_{\rm lum}$) in these alkynyl complexes. If the rate of the oxidative quenching process (k_{eT}) dominates the rates of the other non-radiative deactivation processes ($k_{eT} >> k_{NR}$), the non-radiative decay should be controlled by the oxidation potential of the d⁶ transition metal center. Changing this potential could improve the luminescence of these species. We therefore decided to measure the luminescence of selected 2-ethynyl-9,9dibutylfluorenyl complexes with higher oxidation potentials than 1, such as 2-4 (Scheme 1), hoping that such species would be more luminescent than 1. In addition, Z-scan investigations should reveal how the nature and redox potential of the d⁶ metal center influence their third-order NLO properties. Finally, to probe the influence of the spacer between the metal center and the fluorene luminophore on the quenching process, this series was complemented by 5[PF₆], a cationic Fe(II) compound incorporating the 9H-fluorenyl group as a side-group, i.e. not π conjugated with the metal center.

$$\Phi_{\text{lum}} = k_{\text{lum}} / (k_{\text{lum}} + k_{\text{eT}} + k_{\text{NR}})$$
(1)

We thus report hereafter (i) the synthesis and characterization of the new complexes **2**, **3**, **4** and **5**[PF₆], (ii) selected linear and nonlinear optical properties for some of these species, and (iii) the results of spectroelectrochemical investigations specifically conducted on the Ru(II) derivatives **2** and **3**. These data will be compared to those of **1** and discussed with the help of DFT calculations, a particular emphasis being placed on (i) establishing (or not) the occurrence of a reductive-trapping mechanism of the fluorene-based luminescence (Scheme 2) and on (ii) determining the influence of the metal center and spacer structure on the third-order NLO properties of such 2-ethynylfluorenyl complexes.

Results and Discussion

Synthesis and Characterization of the Fe(II) Carbonyl Complex 2. The synthesis of the Fe(II) alkynyl complex 2 was effected from the known Fe(II) bis-carbonyl chloride precursor 6 and the organic 2-alkynyl-9,9-dibutyl-9H-fluorene (7) previously reported,^[11] following a workup recently developed in our group (Scheme 3). Due to the very high solubility of this complex in common organic solvents most of the product was lost during the purification procedure. This complicated seriously its isolation in pure form and resulted in a rather low yield (31 %).



Scheme 3. Synthesis of the Fe(II) Complex 5.

Complex **2** was characterized by mass spectrometry (ESI). Thus, a molecular peak corresponding to a cationic adduct with sodium, which forms during the ionization process, was detected, along with fragments corresponding to the successive loss of the two carbonyl ligands. The presence of the C₅Me₅ and of the 9,9-dibutyl-*9H*-fluorenyl fragments in the correct stoichiometry is evidenced by ¹H NMR and by ¹³C NMR, while the two carbonyl ligands and the alkynyl bridge are revealed by diagnostic stretching modes in the IR spectrum (v_{CO} at 2017 and 1967 cm⁻¹; v_{C=C} at 2093 cm⁻¹ in dichloromethane) and also by characteristic peaks in the ¹³C NMR spectrum of the complex (Experimental Part).^[3g]

Synthesis and Characterization of the Ru(II) Complexes 3-4. The synthesis of the alkynyl complex **3** was effected from the Ru(II) chloride precursor **8** and the organic 2-alkynyl-9,9-dibutyl-*9H*-fluorene (**7**) previously reported,^[11] following a classic workup (Scheme 4).^[13] The vinylidene complex (**3**-*v*) isolated by precipitation as an intermediate was not characterized, but readily deprotonated with triethylamine to give the yellow complex **3** in fair yields (61 %) after chromatographic separation. The two butyl groups at the 9-position of the fluorene group ensure a good solubility of this complex.



Scheme 4. Synthesis of the Ru(II) Complex 3.

The red bis-acetylide complex **4** was obtained in fair yields from the known monoalkynyl complex **9** by substitution of the chloride with **7**, after deprotonation, in the presence of NaPF₆ and Et₃N in dichloromethane (Scheme 5).^[14] In this case, the intermediate vinylidene complex was not isolated but rather was deprotonated *in situ* before the chromatographic purification. **4** could not be isolated pure by reacting **3** with ethynyl(4nitrobenzene) following a similar reaction pathway (Scheme 4). In contrast, it has been reported that the synthesis of such dissymmetrical complexes are often more facile from the monoacetylide precursor with the more electron-releasing arylalkynyl ligand.^[3a,13b]



Scheme 5. Synthesis of the Ru(II) Complex 4.

The two Ru(II) complexes 3 and 4 were characterized by mass spectrometry (ESI-MS), elemental analysis and by the usual spectroscopies.^[13b,14b,15] The ³¹P NMR singlets observed for 3 and 4 at 50.2 ppm and 53.2 ppm, respectively, are characteristic of these mono- and bis-alkynyl complexes.^[13b] The presence of an additional fluorenyl group is readily established by ¹H NMR and ¹³C NMR spectroscopies, when the spectra of 3 and 4 are compared to those of the starting complexes 8 and 9, while the presence of the triple bond(s) is confirmed by observation of a strong $v_{C=C}$ absorption in the infrared (IR) spectrum. This band is observed at 2066 cm⁻¹ for 3 and at 2046 cm⁻¹ for 4 in dichloromethane. Interestingly, only one band is observed for the latter complex, in spite of the presence of two alkynyl bonds. A closer examination of the infrared spectrum of 4 in KBr, however, reveals a broader absorption band with a shoulder at ca. 2070 cm⁻¹, suggesting that both vibrational modes certainly overlap, as previously observed in related bis-alkynyl Ru(II) complexes.^[3a,14a,15a,16] The presence of the nitro group is confirmed by the appearance of diagnostic symmetric and asymmetric NO₂ stretches near 1580 cm⁻¹ and 1324 cm⁻¹, respectively. Small red crystals of 4 were grown by slow evaporation of methanol-dichloromethane mixtures and the resulting X-ray structure definitively established the identity of this compound (Figure 1).



Figure 1. ORTEP representation of the dissymmetric complex **4** at the 50 % probability level. Selected distances (Å) and angles (deg): Ru1-P1 2.3717(17), Ru1-P2 2.3603(16), Ru1-P3 2.3740(16), Ru1-P4 2.3584(16), Ru-C53 2.039(6), Ru1-C61 2.062(6), C53-C54 1.229(8), C54-C55 1.435(8), C61-C62 1.219(8), C62-C63 1.422(8), C58-N1 1.459(7), N1-O1 1.234(7), N1-O2 1.216(7), P1-Ru1-P2 82.24(6), P3-Ru1-P4 81.63(6), P1-Ru1-C61 98.44(16), C53-Ru1-C61 179.3(2), P2-Ru1-C53 89.76(16), Ru1-C53-C54 176.0(5), C53-C54-C55 176.9(7), Ru1-C61-C62 175.8(5), C61-C62-C63 177.3(7), O1-N1-O2 123.3(6).

Solid-state Structure of 4. The complex 4 (Figure 1) crystallizes in the triclinic P-1 symmetry group, with one molecule of complex in the asymmetric unit and two dichloromethane molecules as solvates (see Experimental Section for details). The bond lengths and angles of this complex are not unusual in comparison to available X-ray data for related mononuclear bis-dppe Ru(II) bis-acetylide complexes and warrant no further comments.[3a,15a] As often stated for related complexes,^[15a-c,17,18] the nitro group is coplanar with the nearby phenyl ring (torsion angle less than 0.04°), in line with a possible interaction between these fragments.^[19] Also, the mean planes for the fluorenyl and phenyl ring are nearly coplanar exhibiting an angle of less than 2°, in a conformation allowing for optimal interaction of the π -manifold through the central ruthenium atom. These planes roughly bisect the two P-Ru-P' quadrants defined by the phosphorus atoms belonging to distinct dppe ligands, in line with a conformation minimizing any intramolecular steric congestion.[3a]



Scheme 6. Synthesis of the Fe(II) Complex 5[PF6].

Synthesis of the Fe(II) Pyridinium Complex 5[PF₆]. The complex 5[PF₆], with an appended fluorenyl group, was obtained

from reaction of the known pyridine complex 10[20] with 2bromomethyl-9H-fluorene (11), following a reaction previously used to access the corresponding methylpyridinium complexes (Scheme 6).^[21] This purple compound was isolated by selective precipitation in good yield (77 %) from the reaction medium and was characterized by elemental analysis, mass spectrometry ³¹P, ¹H, and ¹³C NMR and IR spectroscopies (Experimental Part). In the ¹H NMR spectrum, the bridging methylene-pyridinium protons appear as a singlet at 5.30 ppm, to low field of the two methylene protons at the 9 position of the fluorenyl group. Both the singlet observed at 98.1 ppm in the ³¹P NMR spectrum and the $v_{C=C}$ band observed at rather low-energy in the IR spectrum for a Fe(II) acetylide complex (1987 cm⁻¹ in dichloromethane)^[22] are typical signatures of such pyridinium alkynyl complexes.^[21] In order to have an organic model of the alkynyl ligand of complex 5[PF₆], we have also isolated the corresponding pyridinium salt 12[PF₆] (Figure 6), which was fully characterized by the usual means as well as by X-ray crystallography (Figure 2 and ESI). This salt crystallizes in the monoclinic $P 2_1/n$ symmetry group.



Figure 2. ORTEP representation of the pyridinium salt $12[PF_6]$ (conformation A) at the 50 % probability level. Selected distances (Å) and angles (deg): C20-N15 1.340(3), C20-C19 1.370(3), C19-C18 1.387(3), C18-C17 1.385(3), C17-C16 1.374(3), C16-N15 1.332(3), N15-C14 1.502(3), C14-C1 1.511(8), N15-C14-C1 111.8(3).

Cyclic Voltammetry. Cyclic voltammograms were recorded for 2-5[PF₆]. For the carbonyl complex 2, a fully irreversible event can be observed near 0.9 V which certainly corresponds to metal-centred oxidation, while a fully irreversible reduction is observed at -1.40 V. Similar to 1,[11] 3 and 4 exhibit pseudoreversible metal-centered oxidations (Table 1). As expected from the literature, $^{[3a,14a]}$ relative to $\mathbf{1}$, $^{[23]}$ this oxidation potential is significantly shifted to more positive values, the shift being more pronounced for the complex 4 due to the presence of the nitrophenylalkynyl ligand. The other redox event observed for this compound (at ca. -1.22 V) corresponds to the reduction of the nitro group.^[14a] Finally, and unsurprisingly,^[21] the potential of the metal-centered oxidation of the cationic pyridinium complex 5[PF₆] is shifted toward positive values relative to 1. The pyridinium-based reduction could not be observed, but was seen as an irreversible event at -1.37 V for the organic pyridinium derivative **12**[PF₆].

$$\Delta G_{CT} = \mathsf{E}^{\circ}(\mathsf{D}^{+}/\mathsf{D}) - \mathsf{E}^{\circ}(\mathsf{A}/\mathsf{A}^{-}) + (Z_{\mathsf{A}} - Z_{\mathsf{D}} - 1)\mathsf{e}^{2}/4\pi\varepsilon_{\theta}\varepsilon\mathsf{d} \tag{1}$$

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Based on the Rehm-Weller) equation (1),^[24,25] these data were used to derive an estimate of the free enthalpy of formation of the intramolecular CS state (ΔG_{CS}) resulting from the oxidative quenching of the LC state (Scheme 2).[26] The latter is given by the difference between the metal-based oxidation potential and the first reduction potential of the fluorenyl group in CH₂Cl₂ (Table 1), corrected by an electrostatic term. Since the fluorene reduction peak cannot be observed in CH₂Cl₂, the reduction potential considered was that reported for fluorene in DMF (-2.9 V vs. SCE),^[27] corrected for the change in the dielectric constant of the solvent (ca. -3.0 V in CH₂Cl₂). For the organic salt 12[PF₆] which has no metal center, a ΔG_{CS} value resulting from electron transfer from the fluorene ligand to the pyridinium group was determined instead, based on the oxidation potential reported for fluorene in CH₂Cl₂ (1.48 V vs. SCE) and the reduction potential experimentally found for that salt.^[27] In line with our expectations, we can see that significant changes in ΔG_{CS} take place between 1 and 3, 4 and $5[PF_6]$. Note also that for both 4 and $5[PF_6]$, two other CS states can be found at lower energy (denoted CS' and CS", respectively, in the following). Thus, for 4, there is a state in which the fluorene ligand is oxidized and the 4-nitrophenyl moiety is reduced (CS') and a state in which the Ru(II) center is oxidized and the 4-nitrophenyl moiety is reduced (CS"), while for 5[PF₆], there is a state in which the fluorene ligand is oxidized and the pyridinium moiety is reduced (CS') and a state in which the Fe(II) center is oxidized and the pyridinium moiety is reduced (CS"). Based on a reductive-trapping instead of an oxidativetrapping mechanism of the fluorenyl-based LC state, the CS' states offer a competitive deactivation pathway. Such a reductive quenching process is certainly operative in the compound 12[PF₆] (see later). In contrast, the CS" states are not relevant for the quenching process under consideration, since they cannot be generated by a single electron transfer from the photogenerated CS state.

Table 1. Electrochemical Data for the 2-Fluorenyl Complexes $1\text{--}5[\mathsf{PF}_6]$ and for $12[\mathsf{PF}_6].$

1					
Cmpd	E ^{0 [a]}	$\Delta E_{p}^{[b]}$	i _c /i _a	$\Delta G_{CS}^{[c]}$	Ref.
7	-0.17	0.08	1	2.63	[11]
2	-1.41	/	NR	>3.70	this work
	0.90	/	NR		
3 ^[d]	0.41	0.07	1	3.21	this work
4 ^[d]	-1.22	0.11	1	3.30	this work
	0.50	0.07	1	2.61 ^[e]	
				1.54 ^[f]	
5 [PF ₆]	0.25	0.08	1	3.26	this work
				2.87 ^[g]	
				1.66 ^[h]	
12[PF6]	-1.39	/	NR	2.87 ^[g]	this work

[a] All E° values are in V vs. SCE. Conditions (unless specified): CH₂Cl₂ solvent, 0.1 M [N*n*Bu₄][PF₆] supporting electrolyte, 25 °C, Pt electrode, sweep rate 0.1 V s⁻¹. Unless specified, the ferrocene complex was used as an internal reference for potential measurements. [b] Difference between cathodic and anodic peak potentials. [c] Computed (in eV) according to eq. 1 (see text). [d] The Fe(κ^2 -dppe)(η^5 -C₅Me₅)(C≡CPh) complex was used as an internal reference for potential measurements (E° = -0.15 vs. SCE).^[28] [e] Computed considering the reduction of the nitrophenyl group by the fluorenyl group by the Ru(II) center (see text). [g] Computed considering the reduction of the pyridinium group by the Fluorenyl group by the Fluorenyl group by the reduction of the pyridinium group by the Fluorenyl group by the Fluorenyl group by the reduction of the pyridinium group by the Fluorenyl group by the Fluorenyl group by the Fluorenyl group by the Fluorenyl group of the pyridinium group by the Fluorenyl grou

Table 2. UV-vis Absorption and Emission Data for Selected M(II) and M(III) Complexes in CH₂Cl₂.

Cmpd	Absorption:		Emis			
	λ_{max} / nm (10 ⁻³ ϵ in M ⁻¹ cm ⁻¹)	$\lambda_{max}{}^{[a]}/eV$	$\lambda_{ex}{}^{[b]} / nm / eV$	$\lambda_{\text{em}}{}^{[c]} / \text{nm} / \text{eV}$	${\it P}_{\lambda lum}$ ^[d]	Ref.
1	278 (sh, 33.4), 296 (sh, 30.0), 404 (20.0)	3.07	291 / 4.26	334 / 3.71	0.4 %	[11]
2 ^[e]	338 (30.6) ^[I]	3.67	/	/	/	This work
3	251(37.6), 286 (sh, 15.9), 352 (sh, 22.2), 373 (24.9)	3.32	290 / 4.28	331 / 3.74	0.8 %	This work
4	275 (sh, 59.8), 367(29.8), 489 (18.3)	2.53	290 / 4.28	319 / 3.89	1 %	This work
5 [PF ₆]	266 (49.8), 304(27.6), 322 (sh, 18.7), 540 (sh, 18.1), 586 (21.1)	2.12	285 / 4.35	NL ^[g]	1	This work
3[PF ₆] ^[h]	276 (38.2), 309 (sh, 18.3), 439 (16.8), 454 (sh, 12.7), 640 (3.5), 902 (14.5)	1.37	/	/	/	This work
4 [PF ₆] ^[h,i]	269 (47.1), 317 (26.2), 440 (27.0), 451(sh, 23.9), 680 (0.5), 992 (sh, 11.0),1155 (16.9)	1.07	/		1	This work
12 [PF ₆]	266 (31.3), 304 (14.1)	4.08	265 / 4.68	320 / 3.87 520 / 2.38	2 % 1 %	This work
13	282 (13.7), 376 (sh, 1.0)		/	1	/	This work
14 [PF ₆]	266 (sh, 18.4), 316 (13.1), 546 (sh, 17.9), 578 (19.6)		/	1	/	[21, 36]

[a] Lowest absorption maximum. [b] Excitation wavelength. [c] Emission wavelength. [d] Luminescence yield. [e] Photosensitive product which decomposes during the measurement. [f] Vibronically structured peak. [g] No luminescence detected. [h] Values obtained by spectroelectrochemistry in the presence of 0.2 M [Bu₄N][PF₆]. [i] Spectroelectrochemistry run at 243 K (see text).

Linear Optical Properties of the M(II) Complexes. The UV-Vis/near-IR absorption spectra of **2-4** and **5**[PF₆] were next recorded in dichloromethane (Table 2 and Figure 3).

Absorption Spectroscopy. The spectrum of the Fe(II) carbonyl complex 2 is quite simple, consisting mostly of an absorption band at 338 nm (3.67 eV). This absorption likely corresponds to the $(\pi^*)_{Flu} \leftarrow d_{Fe}$ MLCT band, since its energy matches fairly well with the energy (ΔG_{CS}) of the corresponding CS state at 3.70 eV (Table 1).^[29] The shoulders on the high energy side of this absorption band could result from an unresolved vibronic structure or from overlap with other absorptions. The intensity of this band is higher than that of the corresponding MLCT band of 13 (Scheme 7), the known analogue of 2,[29-30] suggesting that this band certainly overlaps with another weaker band which might correspond to a fluorene-based LC band. However, further studies with this complex were not attempted, since this compound was photosensitive and decomposed in the UV cell under irradiation or even in daylight. This unexpected instability is possibly related to the well-known photolability of the CO ligand in these piano-stool compounds.[31]

For the Ru(II) complexes, the lowest energy absorptions at 373 nm for **3** and at 489 and 367 nm for **4** correspond to transitions with a strong metal-to-ligand charge transfer (MLCT) character ($\pi^* \leftarrow d_{Ru}$), as indicated by DFT calculations (see later). The MLCT transition at lowest energy for **4** is at the origin of the deep red color of this compound. It corresponds to a MLCT transition toward the nitrophenyl group which is the most electron deficient alkynyl ligand of this complex and usually gives rise to transitions in this energy range.^[15a-c,32,3] The second MLCT absorption of this complex resembles that observed for **3** and corresponds to a charge transfer directed toward the 2-ethynylfluorenyl ligand. We note again a fairly good

match with the energy of the corresponding CS states (ΔG_{CS}) derived from electrochemical data (compare Table 1 and Table 2). Finally, the next-higher-energy transitions of **3** and **4** are attributed to a set of overlapped transitions including the fluorenyl-based ($\pi^* \leftarrow \pi$) LC charge transfer transition. The latter corresponds possibly to the shoulder on the high energy side of this huge absorption band, near 300 nm, as has already been seen for the iron derivative **1**.^[11]

Scheme 7. Organoiron Alkynyl Complexes Related to 2 and 5[PF₆].



Finally, the cationic pyridinium Fe(II) complex **5**[PF₆] presents electronic transitions at significantly lower energies. While no calculations have been conducted on that complex, this transition at lowest energy must also correspond to an MLCT transition from the metal center toward the pyridinium moiety ($\pi^*_{Py} \leftarrow d_{Fe}$), as indicated by the good match with the energy (ΔG_{CS}) of the corresponding CS state (compare Table 1 and Table 2).^[34] The next-higher-energy (and sizeable) absorption band, which appears as a huge and composite band in the far UV range, peaks at 266 nm. It certainly overlaps with the fluorene LC band, which could correspond to the shoulder on this band observed on its low energy side. In accordance with such an hypothesis, a shoulder of similar intensity was not observed with the methylpyridinium analogue **14**[PF₆] (Scheme 7) of **5**[PF₆].^[35]



Figure 3. UV-Vis Spectra for Complexes 1 and 2-4, 5[PF₆] in CH₂Cl₂ at 25 °C.

Emission Spectroscopy. No attempts were made to detect fluorescence for complex **2** due to its aforementioned photochemical instability. However, upon excitation in the 285-295 nm range, the Ru(II) complexes **3** and **4** were luminescent. The yields found for **3-4** in CH₂Cl₂ solutions are roughly twice that previously found for the Fe(II) complex **1** (Figure 4),^[11] with an emission maximum located slightly to the blue of that of **1**. Their fluorescence yields are significantly lower than that of 9,9-dibutyl-4-ethynylfluorene (**7**) ($\Phi_{lum} = 58\%$)^[11] and also significantly lower than that reported for fluorene in cyclohexane ($\Phi_{lum} = 72\%$).^[37]



Figure 4. UV-Vis emission spectra for complexes 1 and 3-4 in CH₂Cl₂ at 25 °C.

Under similar conditions, no luminescence could be detected for $5[PF_6]$, while that of the organic pyridinium salt $12[PF_6]$ was much lower than that of 7.^[38] Based on the electrochemical data (Table 1), intramolecular oxidative quenching of the LC state in $12[PF_6]$ by the pyridinium is possibly taking place and contributing to the weak luminescence yield, the remote fluorene-group this time playing the role of reductant. Pyridinium derivatives are often poor fluorophores^[39] because their relatively low reduction potential makes them efficient luminescence quenchers.^[40]

Linear Optical Properties of the Ru(III) Complexes. The Ru(III) complexes 3-4[PF₆] were generated from their Ru(II) parents by spectroelectrochemistry in an OTTLE cell and their electronic spectra were recorded (Table 2, Figure 5 and ESI). In contrast to 3[PF₆] for which the original spectrum could be restored upon back-reduction to the starting complex 3, the oxidation of 4 was only chemically partially reversible at 25 °C and had to be performed at lower temperature (-30 °C) to become fully reversible in the chemical sense. The spectrum of these complexes is typical of Ru(III) bis-alkynyl complexes,[41] and reveals that oxidation switches on an intense absorption with a strong LMCT character in the near-IR range for both compounds. According to calculations made on related complexes,^[14a,41] the next (higher) transitions are often multiconfigurational and should be described as $\pi^* \leftarrow \pi$ transitions involving aryl alkynyl carbon and metal MOs.



Figure 5. Spectroelectrochemistry in the near UV-Vis-near-IR range for Complex 4 in CH_2Cl_2 (0.2 M TBAH) at -30 °C.

Nonlinear Optical Properties. We next determined the cubic hyperpolarizabilities of 3 and 4 by Z-scan, using femtosecond laser pulses at various incident wavelengths between 550/600 and 1600 nm in CH₂Cl₂ solutions (Figure 6, Table 3). These compounds have TPA maxima coinciding approximately with twice the UV-Visible λ_{max} values (Table 2), indicating that the allowed excited states at lowest energy give rise to two-photon absorption. Thus, the first MLCT state of 4 has an apparent TPA cross-section twice stronger than that of compound 3. This is because these transitions correspond to different charge transfers; that in 4 is directed from the metal toward the nitroaryl group, whereas that in 3 is directed toward the fluorenyl group $(\pi^*_{PhNO2} \leftarrow \pi_{C=CPh}/d_{Ru})$, as demonstrated by DFT calculations (see later). The second MLCT band of 4 at 367 nm corresponds to the first MLCT band of **3** ($\pi^*_{Flu} \leftarrow \pi_{C=CFluo}/d_{Ru}$). When these are compared, we see that upon progressing from 3 to 4, the replacement of the chloride by the nitrophenylalkynyl ligand causes a four-fold increase in the apparent TPA cross-section near 700 nm. The strong linear absorptivity between 500 and 600 nm gives a saturable absorption (SA) effect at these wavelengths which complicates the determination of apparent TPA for 4 in this spectral range, resulting in larger experimental errors for this second TPA maximum.

Table 3. Nonlinear Optical Data at Wavelengths of Apparent 2PA Extrema for 1, 2-4 and 15 in CH₂Cl₂ (unless otherwise indicated).

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	λ_{TPA} (nm)	$\gamma_{re} (10^{-34} \text{ esu})$	γ _{im} (10 ⁻³⁴ esu)	γ (10 ⁻³⁴ esu)	σ ₂ (GM)	Ref
1	740	-160 ± 17	84 ± 16	180 ± 25	2400 ± 470	[11]
1 ^[a,b]	740	-25 ± 15	14 ± 2	28 ± 15	330 ± 50	[11]
3	720	-15 ± 5	12 ± 2	19 ± 5	360 ± 60	This work
4	700	-490 ± 55	51 ± 7	490 ± 55	1600 ± 230	This work
	1000	-35 ± 70	55 ± 19	65 ± 73	860 ± 290	This work
15	700	-85 ± 10	31 ± 5	90 ± 11	1700 ± 30	[11]
	760	-87 ± 9	14 ± 2	88 ± 9	370 ± 60	[11]

[a] Determined in THF. [b] Data obtained at Wroclaw University of Technology. All other data were obtained at the Australian National University.



Figure 6. Cubic hyperpolarizabilities determined for 3 (a) and 4 (b) in CH_2CI_2 at 25 °C by open and closed-aperture Z-scan measurements.

DFT Calculations. Calculations on *Cs*-symmetric models **3-Me** and **4-Me** have been performed. These differ from **3-4** by replacement of the dppe ligands by dmpm (bis(dimethylphosphino)methane) and the *n*-butyl groups on the fluorenyl unit by methyl groups. After structural optimization (ESI), the calculations reveal that the frontier molecular orbitals (FMO) of the mono-alkynyl derivative **3-Me** consist of a HOMO mostly localized on the ruthenium alkynyl fragment and a LUMO localized on the fluorenylalkynyl fragment. For the bis-alkynyl

derivative **4-Me**, the HOMO is now localized on the ruthenium bis-alkynyl fragment and the LUMO is localized on the nitrophenyl fragment, resulting in a significant reduction in the HOMO-LUMO (*ca.* 1.7 eV) in **4-Me** compared to **3-Me** (Figure 8). The assignment of the first oxidation step as metal-localized for these complexes is therefore mostly formal^[42] and perhaps less accurate than in the case of the Fe(II) complex **1** previously studied.^[11]

TD-DFT calculations were then performed on the Ru(II) complexes in order to gain some insight into the nature of the allowed low-energy transitions in the visible range (Table 4). As previously observed for 1, the energies of the various allowed transitions are underestimated by ca. 2000-3000 cm⁻¹. This is attributable to some extent to the neglect of solvent and structural simplifications brought on the compounds. Nevertheless, some useful qualitative information can be obtained on the nature and energy ordering of the main allowed singlet-singlet transitions. Thus, the first strongly allowed transition found for 3-Me corresponds to the LUMO \leftarrow HOMO transition with a dominant MLCT character ($\pi^*_{Fluorene} \leftarrow d_{Ru} + \pi_{C=C}$). For 4-Me, the same picture holds except that the MLCT transition at lowest energy corresponds now to charge transfer transitions towards the nitrophenyl ring rather than toward the fluorenyl fragment ($\pi^*_{C6H4NO2} \leftarrow d_{Ru}$), while the next-higherenergy allowed transitions correspond to a set of MLCT transitions toward the fluorenyl fragment (Figure 9). While the former transition corresponds formally to population of the CS" state at lowest energy (Table 1), the second one corresponds to that at higher energy for that particular compound (CS). Note however that a long range ligand-ligand charge transfer (LLCT) transition from the fluorene to the nitrophenyl ($\pi^*_{C6H4NO2}$ \leftarrow $\pi_{\text{Fluorene}})$ that would formally give rise to the intermediate CS' state was not found among the allowed transitions. Finally, for both compounds, transitions with more pronounced LC character on the fluorene are found at higher energy. In contrast to the corresponding transition in 1, which was nearly free of any metal character, our calculations indicate that some metal character is still present in these, as might have been expected given the more pronounced d/p mixing usually observed with Ru(II) alkynyl complexes compared to their isoelectronic Fe(II) counterparts.

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Cmpd	Vmax ^[a] [<i>E</i>] ^[b]	Vmax ^[a] [<i>f</i>] ^[c]	Composition	Major Assignment
	(Exp)	(Calc)	(Weight %)	
3-Me	26810 [0.25]	23832 [0.13]	57a" → 58a" (87%)	Ru_{dyz} (24%) + π_{C1} + $\pi_{C2Fluo} \rightarrow \pi^*_{C2Fluo}$
	28409 [sh, 0.18]	27948 [0.04]	56a" → 58a" (85%)	$Ru_{dyz} (12\%) + p_{pz} + \pi_{Cl} \rightarrow \pi^*_{C2Fluo}$
	39841 [0.38]	33410 [0.09]	55a'' → 58a'' (58%)	Ru_{dyz} (6%) + p _{Pz} + $\pi_{C2Fluo} \rightarrow \pi^*_{C2Fluo}$
4-Me	20450 [0.18]	15968 [0.05]	125a' → 126a' (70%)	Ru _{dxy} (30%) + π_{C2C6H4} + $\pi C2 \rightarrow \pi^*_{C2C6H4NO2}$
		20056 [0.11]	124a' → 126a' (65%)	$\pi_{C2Ph} + \pi_{C2} + p_y \rightarrow \pi^*_{C2C6H4NO2}$
	27248 [0.30]	23269 [0.10]	66a'' → 67a'' (85%)	Ru_{dxz} (30%) + π_{C2} + $\pi_{C2PhFluo} \rightarrow \pi^*_{C2Fluo}$
		28856 [0.11]	65a" → 67a" (78%)	$Ru_{dxz} (14\%) + p_y + \pi_{C2} + \pi_{C2C6H4Fluo} \rightarrow \pi^*_{C2Fluo}$
	36364 [0.27]	30502 [0.03]	121a' → 126a' (79%)	Ru _{dxz} (35%) + π_{C2} + $\pi_{C6H4} \rightarrow \pi^*_{C2C6H4NO2}$
		33813 [0.07]	66a'' → 73a'' (48%)	Ru _{dxz} (30%) + π c2 + π c2C6H4Fluo $\rightarrow \pi^*$ C2C6H4Fluo
			64a'' → 67a'' (40%)	$p_y + \pi_{C2} + \pi_{C2C6H4Fluo} \rightarrow \pi^*_{C2C6H4Fluo}$

[a] Calculated (for 3-Me and 4-Me) and observed (for 3-4) v_{max} in cm⁻¹. [b] Extinction coefficient in 10⁵ M⁻¹ cm⁻¹. [c] Calculated oscillator strength [f].



Figure 8. Frontier molecular orbitals for 3-Me and 4-Me illustrating the first allowed calculated singlet-singlet transitions (red arrows). The metal-centred MOs are also shown.



Figure 9. Plots of the frontier molecular orbitals for 3-Me (a) and 4-Me (b) predominantly involved in the first allowed transition. Contour values are \pm 0.03 (e/bohr³)^{1/2}.

Discussion

The targeted compounds 2-4 and 5[PF₆] could be isolated and characterized. While 2 turned out to be photosensitive in solution, fluorimetric studies on 3 and 4 revealed a marginally higher luminescence than for 1, whereas (5[PF₆]) was not emissive at all in solution. The luminescence of 3 and 4 is too weak to measure their TPA cross-sections by two-photon excited fluorescence (TPEF). In line with previous investigations conducted on 1 and 15 (Scheme 8),[11] we find that the luminescence of 3 and 4 does not originate from their lowest MLCT excited state, which turns out to be essentially nonemissive in solution at 25 °C, but rather from an excited state higher in energy. Although such an emissive behavior is rare,^[43] we note that related observations have been independently reported for other alkynyl complexes such as 16a-c or 17 (Scheme 8).^[8e, 44] Such unusual behaviour could be induced by the electronic decoupling of the fluorenyl fragment from the rest of the molecule,^[44] a situation perhaps achieved for selected rotamers of 1, 3 and 4 in solution.



Scheme 8. Related Organoiron Complexes Showing Dual Fluorescence.

Energy

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Hypothetical Marcus Analysis of а **Redox-Based** Luminescence-Quenching Process. Considering that, as proposed in Scheme 2, the weak quantum yields of 1, 3 and 4 originate from an intramolecular redox-quenching reaction, we have tried to learn more about this particular process. To this aim, the effect of a change in metal-centred oxidation potential on the luminescence quantum yields was closely examined for these compounds. As already mentioned above, when the latter are compared to that of 2-ethynylfluorene (7), the presence of the organometallic Ru(II) center in 3 or 4 appears to result in a luminescence quenching nearly as efficient as that induced by the presence of the Fe(II) center in 1. Based on eq. 1 and assuming that the radiative rate (k_{lum}) remains constant between 1, 3 and 4 (a likely assumption considering that the LC state is fluorene-based),^[46] this means that the global non-radiative decay rate ($k_{eT} + k_{NR}$) has roughly been halved for **3** and **4**. In comparison, the absence of fluorescence for 5[PF6] is less informative since the weak luminescence found for 12[PF₆] indicates that the non-radiative decay is already quite efficient in the pyridinium moiety. Unsurprisingly, the presence of the metal center in 5[PF₆], by opening additional non-radiative deactivation pathways, renders this compound totally non-emissive.

Focusing on 1, 3 and 4, we see that estimates of the ΔG_{CS} values (Table 1) match fairly well with the energies of one of the MLCT $[(\pi^*)_{Flu} \leftarrow d_M]$ states of these compounds (Table 2), considering that the latter correspond to "vertical" energies and therefore represent higher bounds of these values (Figure 10).^[47] This indicates that these particular MLCT states will mediate the electron-transfer process underlying the non-radiative decay. After estimating the energy of the LC state (λ_{00}) from the absorption and emission curves obtained for 3 and 4 (see ESI), the thermodynamic driving forces ($\Delta G_{\text{eT}})$ for the reductive electron-transfer trapping process can be obtained. These are always negative (-0.86 eV and -0.83 eV vs. -1.29 eV for 1), in line with an exergonic process. For 4, because of the presence of the easily reducible 4-nitrophenyl group in this complex, an oxidative quenching process of the LC state is also possible via the CS' state (Figure 10). The specific thermodynamic driving force $(\Delta G_{eT'})$ of this second process has been determined. It is even more exergonic (-1.52 eV) than the former ΔG_{eT} .



Figure 10. Comparative energetic ordering of the ground (GS) and fluorene-centred excited states (LC) for **1**, **3** and **4**. The two additional CS states CS' and CS" of **4** (in grey) are drawn in the inset ([Ru] = Ru(κ^2 -dppe)₂). Vertical energies of the MLCT state(s) are given in blue (and purple). These energies were derived from the absorption maxima and should be considered as higher bound values. Energy differences (in eV) between the LC and the vertical energy of the closest MLCT state are given (in red) as well as ΔG_{eT} values (in black or grey).

According to Marcus,^[12b] the electron transfer rate (k_{eT}) of the quenching process is determined mostly by its activation energy (ΔG^{\neq}) , given that the electronic coupling (H_{ab}) between the metal center and the fluorene chromophore should not vary dramatically within this series of compounds (eq. 3).^[3g,48,49] Thus, ΔG^{\neq} values are essentially determined by ΔG_{eT} values and by the mean reorganization energies (λ) of the redox process (eq. 4). When sensible values are considered for λ , such as 0.6 ± 0.1 eV for $\mathbf{1}^{[50]}$ and 0.9 ± 0.1 eV for **3** and $\mathbf{4}^{[32]} \Delta G^{\neq}$ values of 0.0730 eV, 0.0005 eV and 0.0010 eV, respectively, can be derived. Also, a ΔG^{\neq} value of 0.0787 eV can be derived for the second and trapping mechanism (via the CS' state) in 4.[51] This analysis indicates that, contrary to our initial belief, the most exergonic electron-transfer processes, corresponding also to complexes with the largest M^{II/III} oxidation potentials, should be the slowest ones.

$k_{eT} = C(H_{ab})^2 \exp(\Delta G^{\neq}/k_bT)$	(3)
$\Delta G^{\neq} = (\Delta G_{eT} + \lambda)^2 / (4\lambda)$	(4)

Based on these data, several statements can be made: (i) For 4, and in spite of its larger exergonicity, the oxidative quenching mechanism of the LC state (via the CS' state) will not be competitive with the reductive quenching (via the CS state) initially proposed (Scheme 2). (ii) The redox quenching process proposed should be significantly faster for the Ru(II) complexes 3 and 4 than for the Fe(II) derivative 1, at odds with what is suggested by the quantum yields measured for these compounds. (iii) Given that the ΔG^{\neq} values span nearly two orders of magnitude, the differences between the corresponding $k_{\rm eT}$ rates should be much larger. As a result, based on eq. 1, much more pronounced differences should have been observed between the fluorescence quantum yields (eq. 1) of 1, 3 and 4 if k_{eT} was controlling the non-radiative relaxation and k_{lum} was remaining fairly constant. In fine, this analysis suggests that reductive electron-transfer does not constitute the dominant nonradiative relaxation process of the presumed luminescent LC state - another mechanism different from the photoinduced electron-transfer hypothesized in Scheme 2, such as internal conversion for instance,[52] must also play a determining role in the luminescence quenching process. Improvement of the luminescence yield of such complexes, if possible, will therefore require more than just fine-tuning their metal-based oxidation potential.

Third-Order NLO Activity of 3 and 4. In Table 3, the results are compared to those of 1 and 15 (Scheme 8) previously obtained under similar conditions. As often observed in the literature,^[7e] the bis-acetylide derivative (4) which possesses an extended and more polarized π -manifold, appears to be significantly more active than 1 and 3 at 700 nm. This is apparently due to the larger refractive properties (γ_{re}) of these compounds which dominate their third-order NLO response over the wavelength range probed (ESI).

$$\gamma \propto -\mu_{\rm ge}^4/{\rm E_{ge}}^3 + \mu_{\rm ge}^2\mu_{\rm ee'}^2/{\rm E_{ge}}^2{\rm E_{ge'}} + \mu_{\rm ge}^2(\mu_{\rm ee} - \mu_{\rm gg})^2/{\rm E_{ge}}^3$$
 (5)

Based on the perturbation expression often used to express γ (eq 5),^[7a, 54] the ordering found (4 > 1 > 3) can be inversely related to the HOMO-LUMO gaps of these complexes (Figure 8),

which also determines the energy of the first allowed transition(s). Thus, **3** appears less NLO-active than **1** (and **15**), possibly consistent with its larger gap mostly induced by the larger electronegativity of Ru(II) over Fe(II).^[11] In contrast, due to the presence of the low-lying LUMO centered on the nitro group which narrows the HOMO-LUMO gap, **4** exhibits the strongest NLO activity at 700 nm, and is even more active than **15**. Thus replacement of the chloride ligand by the 4-nitrophenylethynyl ligand upon progressing from **3** to **4** significantly improves the third-order NLO activity, in spite of the concomitant increase of the metal-centered oxidation potential.

Likewise, the TPA associated with the first MLCT band of 4 is significantly larger than that associated with the first MLCT band of 3. As previously discussed, the first MLCT states are different in each compound and strongly polarized transitions such as that of 4 have already been shown to give rise to stronger twophoton absorption (TPA) than less polarized ones.^[55] A closer look at the data reveals that the apparent TPA value associated with the first MLCT excited state of **3** ($\pi^*_{\text{Fluorene}} \leftarrow d_{\text{Ru}} + \pi_{\text{C}=\text{C}}$) is clearly lower than that reported for 1 in CH₂Cl₂, although this corresponds to a similar MLCT.^[11] Taken at face value, these data would indicate that Fe(II) is superior to Ru(II) for this particular property when ligated with a 2-ethynylfluorenyl ligand. However, the photochemical changes in CH₂Cl₂ previously noted for 1 during Z-scan measurements might perhaps have led to some overestimation of the measured TPA values in that particular solvent. In that respect, TPA values comparable to those of 15 were found for 1 when measured in THF (Table 3). Comparison of the second TPA peaks of 4 and 15, which possess comparable metal-to-fluorene ($\pi^*_{Flu} \leftarrow d_M$) charge transfer character (M = Fe(II) and Ru(II)) exhibit nearly similar cross-sections. Interpretation of the changes in absolute values of these TPA cross sections between Fe(II) and Ru(II) must therefore be made with caution until more data are available.

Finally, spectroelectrochemical investigations conducted on 3-4 (Figure 3 and ESI) revealed that oxidation gives rise (reversibly) to a Ru(III) parent with an inverted polarization of the metalalkynyl-fluorene π -manifold. As a result, an intense and broad LMCT band appears in the 850-1050 nm range while the MLCT band observed at lowest energy for the Ru(II) precursor disappears. While TPA (associated with a positive sign of γ_{im}) takes place in this spectral range for the Ru(II) complex, SA (associated with a negative sign of $\gamma_{\rm m}$) is now observed for the Ru(III) isomer (Table 3). As previously shown for related monoand bis- Ru(II) alkynyl complexes,[41] the LMCT states usually give rise to SA when excited. Thus, likewise to what had been previously stated for the redox-active analogues 1 and 15,[11] oxidation of 3-4 leads to a "switch" of the sign of the imaginary part of the cubic NLO coefficient (m) near 900 nm,[56] itself associated with a dramatic change of the nonlinear absorption properties in this spectral range.^[7a]

Conclusions

We have reported in this contribution the synthesis and characterization of four new Ru(II) and Fe(II) alkynyl complexes (**2-4** and **5**[PF₆]) containing 2-fluorenyl groups. Spectrochemical investigations reveal that the lowest lying (MLCT) singlet excited states of these compounds is essentially non-emissive. In line

with previous investigations on the related organoiron analogue 1, the weak luminescence that can be detected for 3-4, when excited around 300 nm, possibly originates from a higher lying excited state presumably centered on the fluorenyl group. As such, the luminescence quantum yields of these compounds remain too weak to allow the determination of their TPA crosssections by two-photon induced fluorescence (TPEF). Finally, we show that both 3 and 4 exhibit significant third-order NLO responses in the near-IR range. Similar to 1, the related (airsensitive) mononuclear organoiron complex previously studied, these air-stable compounds behave as two-photon absorbers in the 850-1050 nm region, 4 being significantly more active than 1 and 3. Furthermore, in this particular spectral range, these complexes should also give rise to a dramatic change in their NLO response upon oxidation, an observation emphazising their interest for innovative developments based on the redoxswitching of their third-order NLO properties.

Experimental Section

General. All manipulations were carried out under an inert argon atmosphere. Solvents and reagents were used as follows: MeOH, distilled from Mg/I2; THF, Et2O and n-pentane, distilled from Na/benzophenone; CH2Cl2, distilled from CaH2 and purged with Ar, opened/stored under Ar. High-field NMR spectroscopy experiments were performed on multinuclear Bruker 500 MHz, 300 MHz or 200 MHz instruments (AVANCE 500, AM300WB and 200DPX). Chemical shifts are given in parts per million relative to tetramethylsilane (TMS) for ¹H and ¹³C NMR spectra and external H₃PO₄ for ³¹P NMR spectra. Transmittance-FTIR spectra were recorded using a Bruker /IFS28 spectrometer (400-4000 cm⁻¹). UV-Visible spectra were recorded using a Cary 5000 spectrometer. Fluorescence spectra were recorded using an Edinburgh Instrument FLS 920 fluorimeter equipped with a 450 W Xenon lamp and a Peltier-cooled Hamamatsu R928P photomultiplier tube in photon-counting mode. MS analyses were performed at the "Centre Regional de Mesures Physiques de l'Ouest" (CRMPO, University of Rennes) on a high resolution MS/MS ZABSpec TOF Micromass Spectrometer. Elemental analyses were performed at the "Centre Regional de Mesures Physiques de l'Ouest" (C.R.M.P.O., University of Rennes). The solid-state structures (X-ray) were resolved at the "Centre de Diffractométrie X" (UMR CNRS 6226, University of Rennes).

Unless specified, all reagents were of commercial grade. Petroleum spirit refer to the 40-60 °C boiling range fraction. *cis*-Ru(κ^2 -dppe)₂Cl₂ (**8**),^[13a] *trans*-Ru(κ^2 -dppe)₂[C≡C(4-C₆H₄NO₂)](Cl) (**9**),^[15a] Fe(η^5 -C₅Me₅)(CO)₂(Cl) (**6**),^[57] Fe(η^5 -C₅Me₅)(κ^2 -dppe)[C≡C(4-C₅H₄N)] (**10**)^[20], 2-alkynyl-9,9-dibutyl-9*H*-fluorene (**7**)^[11] and 2-bromometyl-9*H*-fluorene (**11**)^[58] were prepared analogously to published procedures.

Synthesis of compound 1: A solution of the fluorene derivative 4 (61 mg, 0.11 mmol) and TCNE (14 mg, 0.11 mmol) in CH₂Cl₂ (1.1 mL) was stirred at 20 °C for 15 h. The reaction mixture was concentrated under reduced pressure to give the title compound (75 mg, 100%) as a dark red solid. MP: 108-111 °C. R_f: 0.24 [Petroleum ether/Et₂O (9:1)]. ¹H **NMR** (400 MHz, CDCl₃) δ = 7.83 (1H, d, J = 1.5 Hz, H_{Flu}), 7.72 (1H, d, J = 8.1 Hz, H_{Flu}), 7.71-7.67 (1H, m, H_{Flu}), 7.61 (2H, d, J = 9.3 Hz, H_{Ar}), 7.46 (1H, dd, J = 1.9 and 8.1 Hz, H_{Flu}), 7.39-7.28 (7H, m, H_{Ar} and 3H_{Flu}), 7.21-7.11 (6H, m, 2 H_{Ar}), 6.86 (2H, d, J = 9.3 Hz, H_{Ar}), 2.03-1.85 (4H, m, H_{Bu}), 1.05-0.92 (4H, m, H_{Bu}), 0.57 (6H, t, J = 7.3 Hz, H_{Bu}), 0.54-0.44 (4H, m, $H_{\text{Bu}}\text{)}.$ $^{13}\text{C}\{^{1}\text{H}\}$ NMR (100 MHz, CDCl_3) δ = 168.9, 164.7, 153.8, 152.6, 152.3, 148.2, 144.7, 139.0, 132.1, 130.3, 130.2, 129.9, 129.4, 127.6, 127.1, 126.8, 124.5, 123.4, 122.0, 121.4, 121.0, 118.2, 113.8, 112.8, 112.8, 111.9, 85.1, 78.6, 55.8, 39.9, 26.1, 23.0, 13.9. **IR** (KBr, cm⁻¹): v = 3061, 2928 (m, C_{Ar}-H), 2219 (m, C=N), 1607 (m, C=C), 1586 (s, C=C_{Ar}). Raman (neat, cm⁻¹): v = 3066 (vw, C_{Ar}-H), 2221 (s, C=N), 1609 (s, C=C), 1520 (vs, C=C_{Ar}). **HRMS**: calculated for C₄₇H₄₀N₅ [M+H]⁺ 674.32837, found 674.3278, calculated for C₄₇H₃₉N₅ M^{+.} 673.32055, found 673.3197. **UV-vis** (CH₂Cl₂): λ_{max} (log ϵ) = 274 (4.40), 340 (4.20), 414 (4.53), 489 (4.40).

Synthesis of $Fe(\eta^5-C_5Me_5)(CO)_2[C=C(2-C_{21}H_{25})]$ (2). In a Schlenk tube under argon, Cul (37 mg, 0.19 mmol, 0.3 eq.) were dissolved in triethylamine (30 mL). After 10 min of stirring, a solution of 2-ethynyl-9,9dibutyl-9H-fluorene (7; 194 mg, 0.64 mmol, 1 eq.) in THF (30 mL) was added. The mixture was stirred for 30 min and then a solution of Fe(η^5 -C5Me5)(CO)2(Cl) (6; 200 mg, 0.70 mmol, 1.1 eq.) in THF (30 mL) was added. The reaction mixture was stirred overnight in the dark. After removal of the solvent, the product was extracted with CH₂Cl₂ and filtered through an alumina plug and the filtrate taken to dryness. The compound was dissolved in diethylether, precipitated with n-pentane, and subsequent washing with cold methanol and cold n-pentane (-50 °C) afforded the title compound as a yellow solid (120 mg, 0.22 mmol, 31 %). Anal. Calc for C₃₅H₄₀O₂Fe•½CH₂Cl₂: C, 72.15, H 6.99; Found: C, 71.69, H 7.44. MS (ESI) m/z 571.2270 [M+Na]⁺ calc. for C₃₅H₄₀O₂FeNa: 571.2270 ([M+Na]⁺⁺), 543.2324 [M-CO+Na] calc. for $C_{34}H_{40}OFeNa$: 543.2321 ([M+Na]+•), 515.2365 [M-2CO+Na] calc. for C₃₃H₄₀FeNa: 515.2372 ([M-2CO+Na]+•). ¹H NMR (200 MHz, CDCl₃): δ = 7.78 (s, 1H, H_{flu}), 7.74 (d, 1H, H_{flu}), 7.59-7.75 (m, 2H, H_{flu}), 7.17 (m, 3H, H_{flu}), 1.93 (t, 4H, CH_{2-Bu}), 1.47 (m, 15H, C₅(CH₃)₅), 0.90 (m, 8H, CH_{2-Bu}), 0.70 (m, 8H, CH_{2-Bu}), 0.50 (s, 6H, CH₃). ¹³C{¹H} NMR (125 MHz, C₆D₆): δ = 216.3 (s, CO), 151.2 (s, C_{flu}), 151.6 (s, C_{flu}), 151.3 (s, C_{flu}), 142.4 (s, C_{flu}), 138.9 (s, Cflu), 131.6 (s, CHflu), 129.5 (s, Cflu), 127.5 (s, CHflu), 127.2 (s, CHflu), 126.3 (s, CH_{flu}), 123.4 (s, CH_{flu}), 120.3 (s, CH_{flu}), 120.2 (s, CH_{flu}), 115.1 (s, FeC=C), 105.1 (s, FeC=C), 97.4 (s, C₅(CH₃)₅), 55.6 (s, C_{flu}), 41.2 (s, CH_{2/Bu}), 26.9 (s, CH_{2/Bu}), 23.9 (s, CH_{2/Bu}), 14.4 (s, CH_{2/Bu}), 10.4 (s, C₅(CH₃)₅). IR (KBr/CH₂Cl₂, cm⁻¹): v = 2095/2093 (m, C≡C), 2010/2017 (s, C=O), 1960/1967 (s, C=O).

Synthesis of trans-Ru(x²-dppe)₂[C=C(2-C₂₁H₂₅)](CI) (3). A mixture of cis-Ru(k²-dppe)₂Cl₂ (8; 400 mg, 0.41 mmol, 1 eq), 2-ethynyl-9,9-dibutyl-9H-fluorene (7; 150 mg, 0.50 mmol, 1.2 eq) and NaPF₆ (90 mg, 0.5 mmol, 1.2 eq) was strirred 12 h at 35 °C. After filtration and concentration, the solution was poured into Et₂O. The precipitate was filtered and washed several times with Et₂O. The compound was then dissolved in CH₂Cl₂ and Et₃N (3 mL). Subsequently, the volatiles were removed and the product was eluted through a basic alumina plug using CH₂Cl₂ as eluent. After evaporation of the solvent, the title complex was obtained as a yellow solid (313 mg, 0.25 mmol, 61 %). Anal. Calc for C₇₅H₇₃ClRuP₄: C, 72.95; H, 5.96; Found: C, 72.87; H, 6.06. MS (ESI) m/z 1240.4 ([M- $Cl+CH_{3}CN]^{+\bullet},\ 100\ \%),\ calc.\ for\ C_{75}H_{73}P_{4}Ru\bullet CH_{3}CN:\ 1240.4\ ([M-$ CI+CH₃CN]^{+•}). ³¹P{¹H} NMR (121 MHz, CDCI₃): $\delta = 50.2$ (s, P_{dppe}). ¹H **NMR** (300 MHz, CDCl₃): δ = 7.51-7.45 (m, 9H, H_{Ar}), 7.41 (d, 1H, ³J_{H,H} = 8.1 Hz, H_{Ar}), 7.22 (m, 11H, H_{Ar}), 7.13 (m, 8H, H_{Ar}), 6.92 (m, 16H, H_{Ar}), 6.59 (m, 2H, H_{Ar}), 2.66 (m, 8H, CH_{2/dppe}), 1.89 (m, 4H, CH_{2/Bu}), 1.13 (m, 4H, CH_{2/Bu}), 0.67 (m, 10H, CH_{2/Bu}+ CH₃). ¹³C{¹H} NMR (125 MHz, CDCl₃): δ = 151.1 (s, C_{flu}), 150.5 (s, C_{flu}), 142.5 (s, C_{flu}), 137.3 & 136.4 (2 × m, CPh/dppe), 136.7 (s, Cfiu), 135.1 (2 × s, CHPh/dppe), 130.0 (s, Cfiu), 129.5 $(2 \times m, CH_{Ph/dppe}), 129.4 (s, CH_{flu}), 127.8 \& 127.7 (2 \times s, CH_{Ph/dppe}), 127.4$ (s, CH_{flu}), 126.5 (s, CH_{flu}), 125.0 (quin, ${}^{2}J_{C,P} = ca.$ 16 Hz, Ru-C=C), 124.8 (s, CH_{flu}), 123.4 (s, CH_{flu}), 119.6 (s, CH_{flu}), 119.5 (s, CH_{flu}), 115.6 (m, Ru-C=C), 55.4 (s, Cflu), 41.4 (s, CH_{2-flu}), 31.4 (m, CH_{2/dppe}), 26.8 (s, CH_{2/flu}), 24.0 (s, CH_{2/flu}), 14.7 (s, CH₃). IR (KBr/CH₂Cl₂, cm⁻¹): v = 2065/2066 (s, C≡C).

Synthesis of $Ru(\kappa^2-dppe)_2[C=C(2-C_{21}H_{25})][C=C(4-C_6H_4NO_2)]$ (4). A mixture of *trans*-Ru(κ^2 -dppe)_2[C=C(4-C_6H_4NO_2)](CI) (9; 200 mg, 0.18 mmol, 1 eq.), 2-ethynyl-9,9-dibutyl-9*H*-fluorene (7; 73 mg, 0.24 mmol, 1.3 eq.), NaPF₆ (47 mg, 0.28 mmol, 1.5 eq.) and 5 drops of Et₃N were stirred 12 h at reflux. The solvents were removed and the compound was filtered through a deactivated alumina plug and eluted with a petroleum spirit/dichloromethane mixture (1:2 to 2:1). After removal of the volatiles, the product was washed with petroleum spirit (3 x 20 mL) to afford the

title compound as a red solid (218 mg, 0.16 mmol, 90 %). Anal. Calc for C83H77NO2P4Ru: C, 74.09; H, 5.77; N, 1.04, Found: C, 73.96; H, 5.89; N, 1.10. MS (ESI) m/z 1384.7 ([M+K]**, 100 %), 1368.7 ([M+Na]**, 60 %), calc. for C₆₇H₆₈FeP₂K: 1384.4. ³¹P{¹H} NMR (121 MHz, CDCl₃): δ = 53.2 (s, P_{dppe}). ¹H NMR (400 MHz, CD₂Cl₂): δ = 8.01 (d, 2H, ³J_{H,H} = 9.0 Hz, H_{Ar}), 7.71 (m, 8H, H_{Ar}), 7.67 (m, 1H, ${}^{3}J_{H,H} \approx 7.5$ Hz, H_{flu}), 7.56 (dd, 1H, ${}^{3}J_{H,H} = 7.8$ Hz, ${}^{5}J_{H,H} = 0.4$ Hz, H_{flu}), 7.40-7.33 (m, 10H, H_{Ar}), 7.30 (dd, 1H, ${}^{3}J_{H,H}$ = 7.3 Hz, ${}^{4}J_{H,H}$ = 1.2 Hz, H_{flu}), 7.27-7.21 (m, 8H, H_{Ar}), 7.06-6.97 (m, 16H, H_{Ar}), 6.92 (dd, 1H, ${}^{3}J_{H,H}$ = 7.8 Hz, ${}^{4}J_{H,H}$ = 1.2 Hz, H_{flu}), 6.90 (s, 1H, *H*_{flu}), 6.67 (d, 2H, ³*J*_{H,H} = 9.0 Hz, *H*_{Ar}), 2.71 (m, 8H, C*H*_{2/dppe}), 1.93 (m, 4H, CH_{2/Bu}), 1.22 (m, 4H, CH_{2/Bu}), 0.79 (t, 6H, ³J_{H,H} = 7.3 Hz, CH_{3/Bu}), 0.69 (m, 4H, CH_{2/Bu}). ¹³C{¹H} NMR (125 MHz, CDCl₃): δ = 155.0 (quin, ²J_{C,P} = ca. 15 Hz, Ru-C=C), 151.4 (s, Cflu), 150.7 (s, Cflu), 143.3 (s, CAr), 142.4 (s, $C_{flu}),~138.1$ (s, $C_{Ar}),~137.4$ & 137.2 (2 \times m + s, $C_{Ph/dppe}\text{+}C_{flu}),~135.1$ & 134.6 (2 \times s, CH_{Ph/dppe}), 130.6 (CH_{Ar}), 129.9 (s, C_{flu}), 129.5 (2 \times m, CH_{Ph/dppe}), 129.2 (s, CH_{flu}), 127.9 (2 × s, CH_{Ph/dppe}), 127.4 (s, CH_{flu}), 126.7 (s, CH_{flu}), 124.8 (s, CH_{flu}), 124.1 (s, CH_{Ph}), 123.3 (s, CH_{flu}), 120.1 (m, Ru-C=C), 119.7 (s, 2CH_{flu}), 119.6 (m, Ru-C=C), 55.4 (s, C_{flu}), 41.4 (s, CH_{2/flu}), 32.1 (m, CH_{2/dppe}), 26.8 (s, CH_{2/flu}), 24.0 (s, CH_{2/flu}), 14.7 (s, CH₃), Ru-C=C (C_a) not detected. IR (KBr/CH₂Cl₂, cm⁻¹): v = 2070 (sh), 2051/2046 (s, C=C), 1579/1581 (s, NO₂[v_{as}]), 1324/1323 (s, NO₂[v_{sym}]).

Synthesis of [Fe(η⁵-C₅Me₅)(κ²-dppe)[C=C(4-C₅H₄N{CH₂(2-C₁₃H₉)})]] [PF₆] (5[PF₆]). In a Schlenk tube under argon, $Fe(\eta^5-C_5Me_5)(\kappa^2$ dppe)[C=C(4-C₅H₄N)}] (**10**; 300 mg, 0.43 mmol, 1 eg) and 2bromomethylfluorene (11; 110 mg, 0.42 mmol, 0.98 eq) were stirred for 12 h in THF at room temperature. To this mixture, potassium hexafluorophosphate (88 mg, 0.48 mmol, 1.1 eq) was added and the reaction was stirred for a further 5 h. The solvent was then removed and the deep blue product was extracted with dichloromethane. After removal of the solvent, the compound was precipitated in a mixture of toluene and *n*-pentane. The solid was then washed with n-pentane (2 x 50 mL) to afford the title complex as a deep blue solid (337 mg, 0.33 mmol, 79 %). Anal. Calc for C₅₇H₅₄F₆FeNP₃•1/₂CH₂Cl₂: C, 65.26; H, 5.24; N, 1.32; Found C, 65.72; H, 5.44; N, 1.34. MS (ESI) m/z 870.3088 (M+•) calc. for C₅₇H₅₄FeNP₂: 870.3081 (M+*). ³¹P{¹H} NMR (81 MHz, CDCl₃): δ = 98.1 (s, P_{dppe}), -142.9 (sept, ${}^{1}J_{P,F} = 713$ Hz, PF_{6}). ${}^{1}H$ NMR (200 MHz, CDCl₃): $\delta =$ $\Box \Box J_{H,H} = 6.6 \text{ Hz}, \Box \Box H, H_{Py} \Box \Box \Box \Box \Box \Box m, 3H, H_{flu}, 7.57$ (m, 8H, $H_{Ph/dppe}$ + H_{flu}), 7.49-7.20 (m, 16H, $H_{Ph/dppe}$), 6.59 (d, 2H, ${}^{3}J_{H,H}$ = 6.6 Hz, 2H, H_{Py}), 5.30 (s, 2H, CH₂), 3.95 (s, 2H, CH₂), 2.63 (m, 2H, $CH_{2/dppe}),\ 2.13\ (m,\ 2H,\ CH_{2/dppe}),\ 1.43\ (s,\ 15H,\ C_5(CH_3)_5).\ ^{13}\text{C}\{^1H\}\ \text{NMR}$ (100 MHz, CD₂Cl₂): δ = 144.7 (s, C_{flu}), 143.7 (s, C_{flu}), 143.0 (s, C_{flu}), 140.5 (s, C_{fiu}), 139.7 (s, CH_{Py}), 137.0-135, (2 × m; $C_{Ph/dppe}$), 133.6 & 133.3 (2 × m, CH_{Ph/dppe}), 131.8 (s, CH_{flu}), 130.7 (t, ²J_{C,P} = ca. 6.4 Hz, C_{Py}), 130.0 & 129.6 (2 × s, CHPh/dppe), 127.9 & 127.9 (s, CHPh/dppe), 127.5 (s, CH_{flu}), 127.3 (s, C_{flu}), 126.9 (s, CH_{flu}), 126.4 (s, CH_{Py}), 125.3 (s, CH_{flu}), 125.2 (s, CH_{flu}), 120.6 (s, CH_{flu}), 120.2 (s, CH_{flu}), 91.6 (s, $C_5(CH_3)_5$), 62.2 (s, NCH2), 36.8 (s, CH2/flu), 30.3 (m, CH2/dppe), 9.8 (s, C5(CH3)5, [Ru-C=C (C_α), Ru-C=C (C_β) not detected]. **IR** (KBr/CH₂Cl₂, cm⁻¹): v = 1987/1988 (s, C=C), 1619/1620 (s, Py), 842/847 (s, P-F).

Luminescence measurements. The samples used to make the solutions were freshly recrystallized or thoroughly washed with cooled ether / pentane to remove any organic impurity prior to the measurements. Luminescence measurements in solution were performed in dilute deoxygenated solutions (except in the case of compound **12**[PF₆]) contained in air-tight quartz cells of 1 cm pathlength (*ca.* 10^{-6} M, optical density < 0.1) at room temperature (298 K). Fully corrected excitation and emission spectra were obtained with an optical density at $\lambda_{exc} \leq 0.1$ to minimize internal absorption. Luminescence quantum yields were measured according to literature procedures.^[59] UV-vis absorption spectra used for the calculation of the luminescence quantum yields were recorded using a double beam Jasco V-570 spectrometer.

Spectroelectrochemistry. Solution UV-Vis-NIR spectra of the oxidized species were obtained at 243 K or 298 K by electrogeneration on a platinum mesh electrode in a 0.05 mm optically transparent thin-layer

electrochemical (OTTLE) cell,^[60] using a silver wire as pseudo-reference and a platinum wire as counter-electrode. Solutions were made up with 0.20 M [*n*-Bu₄N][PF₆] in dry and deoxygenated CH₂Cl₂, and were kept under an atmosphere of pure nitrogen.

Z-scan Measurements. Third-order nonlinear optical properties were investigated as previously described,^[61] but with some modifications. The laser system consisted of a Quantronix Integra-C3.5F pumping a Quantronix Palitra-FS optical parametric amplifier, tuneable over a wavelength range from 500 nm to 2000 nm. The output wavelength was confirmed by use of an Ocean Optics USB2000+ spectrometer (500-1000 nm) or an Ocean Optics NIR-Quest spectrometer (1000-1800 nm). The system delivered 130 fs pulse with a 1 kHz repetition rate. Colored glass filters and a Thorlabs polarizing filter were used to remove unwanted wavelengths and the power adjusted by use of neutral density filters, attenuating it to the µJ/pulse range to obtain nonlinear phase shifts between 0.2 to 1.3 rad. The focal length of the lens used in the experiment was 75 mm, which gave 25-40 µm beam waists resulting in Rayleigh lengths sufficiently longer than that of the sample thickness that a "thin-sample" assumption was justified. Solutions of compounds in "as received" CH2Cl2 (see text), deoxygenated and distilled CH2Cl2, or deoxygenated and distilled THF, of 0.1 w/w% concentration in 1 mm glass cells were analyzed. Samples travelled down the Z-axis on a Thorlabs motorized stage between -20 and 20 mm (where 0 was the laser focus). Data were collected by three Thorlabs photodiodes, 500-900 nm with Si based detectors, 900-1300 nm with InGaAs detectors and 1300-2000 nm with amplified InGaAs detectors, monitoring the laser input, the open-aperture signal and the closed-aperture signal. Data from the detectors were fed into three channels of a Tektronix digital oscilloscope, collected with a custom LabVIEW program, and fitted with theoretical traces with a program that used equations derived by Sheik-Bahae et al.[62] A sample of the solvent was run at each wavelength to account for solvent and cell contribution to the Z-scan signals and the light intensity was determined from a Z-scan run on a 3 mm fused silica plate; the real and imaginary components of the second hyperpolarizability (γ) of the materials were calculated assuming additivity of the nonlinear contributions of the solvent and the solute and the applicability of the Lorentz local field approximation. The values of the imaginary parts of γ were also converted into values of the two-photon absorption cross-sections σ_2 .

Crystallography. Data collection of crystals of **4** and **12**[PF₆] were performed on an Oxford Diffraction Xcalibur Saphir 3 diffractometer or an image plate APEXII Bruker-AXS spectrometer at 150 K, with graphite monochromatized MoK_a radiation. The cell parameters (ESI) were obtained with Denzo and Scalepack.^[63] The data collection^[64] provided reflections from which the independent reflections were obtained after data reduction using Denzo and Scalepack.^[63] The structures were solved with SIR-97 which revealed the non-hydrogen atoms.^[65] After anisotropic refinement, the remaining atoms were found in Fourier difference maps. The complete structures were then refined with SHELXL97^[66] by the full-matrix least-square technique. Atomic scattering factors were taken from the literature.^[67]

Computational Details. Calculations were performed using the Amsterdam Density Functional (ADF) program,^[68] version ADF2012. Scalar relativistic effects were treated via the Zeroth-Order Regular Approximation (ZORA) method.^[69] In all calculations, all-electron Triple Zeta plus Polarization (TZP) Slater orbital basis sets were used for all atoms. Geometry optimizations were undertaken with suitable symmetry constraints using the exchange-correlation (XC) functional and the Generalized Gradient Approximation (GGA) proposed by Becke and Perdew (BP).^[70] The model systems were simplified by using 1,2-bis(dimethylphosphino)methane (dmpm) ligand instead of 1,2-bis(diphenylphosphino) ethane (dppe) and hydrogen instead of methyl in the butyl groups. UV-Vis spectra were calculated using the Statistical Average of Orbital Potentials (SAOP) functional^[71] with the same TZP basis sets.

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Entry for the Table of Contents

FULL PAPER



The linear and third-order nonlinear optical properties of a series of new 2ethynylfluorene complexes are investigated and compared to those of known Fe(Cp*)(dppe) analogues. Among these, the bis-alkynyl Ru(II) derivatives give rise to remarkably large and potentially redox-switchable apparent two-photon absorption in the 700-750 nm wavelength range. Floriane Malvolti, Cedric Rouxel, Guillaume Grelaud, Loic Toupet, Thierry Roisnel, Adam Barlow, Fazira I. Abdul Razak, Robert Stranger, Marie P. Cifuentes, Mark G. Humphrey*, Olivier Mongin, Mireille Blanchard-Desce, Christine Paul-Roth and Frédéric Paul*

Page No. – Page No.

Iron and Ruthenium Alkynyl Complexes with 2-Fluorenyl Groups: Some Linear and Nonlinear Optical Absorption Properties