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Photoisomerization in Aminoazobenzene-Substituted 
Ruthenium (II) tris(bipyridine) Complexes : 
Influence of the Conjugation Pathway

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Abstract

Transition metal complexes containing stimuli-responsive systems attract extensive attention for potential applications in optical devices, photonic memory, photosensing, as well as luminescence imaging. Among them, photochromic metal complexes offer the possibility to combine the specific properties of the metal centre and the optical response of the photochromic group. Here we report on the synthesis, the electrochemical properties and the photophysical characterization of a series of donor-acceptor azobenzene derivatives that possess bipyridine groups connected to a 4-dialkylaminoazobenzene moiety through various linkers. DFT and TD-DFT calculations are performed to complement experimental findings and contribute to their interpretation. Position and nature of the linker (ethynyl, triazolyl, none) are engineered and shown to induce different electronic coupling between donor and acceptor, in ligands and complexes. This in turn leads to strong modulations in terms of photoisomerization of the ligands and complexes.
Introduction

Multi-mode photo-functional materials are currently attracting significant attention in relation with the emergence of molecular electronics. Light-driven real-time information-transmitting systems may be achieved by using photochromic switches\textsuperscript{[1]} and this issue raises growing interest in the field of materials chemistry.\textsuperscript{[2]} In particular, azobenzene derivatives are known to undergo efficient \textit{trans}-to-\textit{cis} isomerization upon irradiation with UV or blue light, which can be reversed either thermally or photochemically.\textsuperscript{[3]} Thanks to the simplicity of this moiety, the ease in molecular structure modification, their optical properties and photo-fatigue-resistance, azobenzene-based chromophores form a very attractive class of molecules. They are currently widely investigated for various applications.\textsuperscript{[3-4]} In order to develop azobenzene-based photoswitches that can exchange from the \textit{trans} to the \textit{cis} states under visible light, various strategies have been implemented. Among these, the most popular relies on the incorporation of electron-withdrawing or -donating substituents in the $\pi$-system of azo-chromophore, shifting the $\pi-\pi^*$ transition to the visible region. This route has already been explored in terms of dyes, polymers and materials processing methods.\textsuperscript{[5]} An alternative approach to red-shift the activation wavelength consists in gathering a transition metal group and an azobenzene moiety through a suitable linker. In this way, the specific properties of the metal centre and the optical response of the organic group are combined in a single entity.\textsuperscript{[6]} For instance, this has already been implemented by linking the azo unit to a polypyridine and coordinating the resulting ligand to a transition metal atom.\textsuperscript{[7]} Some of these complexes show photochromism\textsuperscript{[7c]} or luminescence switching upon irradiation.\textsuperscript{[7a, 8]} Even if the isomerization is assumed to take place from intraligand excited states
localized on the azobenzene unit, the choice of the metal centre is known to drive the photochromic properties of the azobenzene unit.\textsuperscript{[7c]} In fact, the photoisomerization was reported to be suppressed in azobenzene-containing polypyridine Ru(II) complexes, presumably due to an energy transfer from the reactive azobenzene-localized intraligand (IL) state to low-lying MLCT states.\textsuperscript{[9]} The location of the azobenzene unit in the bipyridine ligands was also established as an alternative way to influence the photochromatic properties of the azobenzene unit in such complexes.\textsuperscript{[10]}

In this paper, we report on the synthesis, the characterization and the photophysical properties of a series of azobenzene-containing bipyridines (L\textsubscript{1-3}) and their respective monometallic ruthenium complexes (Scheme 1). These ruthenium (II) heteroleptic complexes, Ru(bpy)\textsubscript{2}(L\textsubscript{1-3})(PF\textsubscript{6})\textsubscript{2} (abbreviated as RuL\textsubscript{1-3}) differ by the nature of the azobenzene-containing bipyridine ligands L in which both the linker and its position were modified. L\textsubscript{1} and L\textsubscript{2} exhibit a donor/conjugated bridge/acceptor (D-\pi-A) structure, but difference stems from the position, nature and length of the conjugated bridge. While for L\textsubscript{1} the azo unit is directly linked to position 4 of the bipyridine, it is linked to the bipyridine by an ethynyl bridge at position 5 in L\textsubscript{2}. L\textsubscript{3} is similar to L\textsubscript{2} except for the linker that contains a triazole moiety. The effect of the ligand types on the photophysical properties (absorption, emission and photoisomerization) of the ligands (L\textsubscript{1-3}) and complexes RuL\textsubscript{1-3} are investigated experimentally and theoretically, and compared to the simple models L\textsubscript{4}\textsuperscript{[11]} (Scheme 1) and Ru(bpy)\textsubscript{3}(PF\textsubscript{6})\textsubscript{2}. 
Results and Discussion

Synthesis. A specific strategy was considered for the synthesis of the azo-bipyridine ligand L1 which is different from compounds L2-3 as the azo unit is directly fixed to one of the pyridine cycles, in position 4 (Scheme 2; for experimental details, see SI).

L1 was prepared via a Negishi cross-coupling reaction between the (E)-4-((2-bromopyridin-4-yl)diazenyl)-N,N-dibutylaniline 2 and a pyridyl zinc reagent 4, obtained from the 2-bromopyridine 3\textsuperscript{[12]} (Scheme 2). The (E)-4-((2-bromopyridin-4-yl)diazenyl)-N,N-dibutylaniline 2 was previously obtained in high yield (83%), by reacting N,N-dibutyl aniline with the appropriate diazonium salt formed by the reaction of aqueous tetrafluoroboric acid\textsuperscript{[13]} with 4-amino-2-bromopyridine 1. The second step consists of a lithium-halogen exchange by reaction of tert-butyllithium with 2-bromopyridine at -78 °C. After transmetallation to Zn, a CHCl\textsubscript{3} solution of Pd\textsubscript{2}(dba)\textsubscript{3}, t-Bu\textsubscript{3}PH·BF\textsubscript{4} and 2 were added, and the reaction was refluxed until completion. L1 was obtained in high yield (89%) after aqueous workup in the presence of ethylenediaminetetraacetic acid (EDTA) and purification by column chromatography. The syntheses of the ligands L2-3, both substituted via an ethynyl (L2) or a triazolyl (L3) bridge in position 5 (Scheme 3), result from different catalysed reactions from 5-(ethynyl)-2,2'-bipyridine 5\textsuperscript{[14]} (Scheme 3). L2 was synthesized by the Pd-catalysed Sonogashira cross coupling reaction\textsuperscript{[15]} of 5 with 4-(N,N-dibutylamino)-4'-iodoazobenzene 6, which was previously prepared by reaction of N,N-dibutylaniline with 4-iodobenzene diazonium chloride. L3 was obtained from 5 and 6, in the presence of sodium azide, via a Huisgen-type azide–alkyne cycloaddition\textsuperscript{[16]} to give the 1, 4-disubstituted 1, 2, 3-triazole ligand L3. The \textsuperscript{1}H NMR signals for the bipyridine protons of L2 coincide with values reported for other 5'-monosubstituted 2,2'bipyridines.\textsuperscript{[12]} Shifts for the H\textsuperscript{4} and H\textsuperscript{6} of the pyridine linked to the triazolyl bridge,
in L3, are significantly downfield shifted [H4': 8.39 (L3) and 7.98 (L2); H6': 9.20 (L3) and 8.85 ppm (L2) respectively], while the CH of the triazole is observed at 8.36 ppm, confirming the formation of L3. The Ru(bpy)2(L1-3)(PF6)2 complexes were prepared by the reaction of commercial cis-(bpy)2RuCl2.2H2O, in presence of one equivalent azo-bipyridine ligands L1-3, in refluxing ethanol/water (8:2). These complexes were obtained in good yields (72 to 89%) after silica gel column chromatography, using a mixture of acetonitrile and saturated aqueous KNO3 as eluent, and were fully characterized by 1H and 13C NMR spectroscopy as well as Zabspec-TOF MS and elemental analysis.

**Electrochemical Properties.** Figure 1 shows typical cyclic voltammograms (CVs) of L1, RuL1 and Ru(bpy)32+ in acetonitrile (see also Figure S1). All the ligands globally display comparable electrochemical features (Table 1). The same holds true for their corresponding ruthenium-based complexes. A quasi-reversible oxidation process is clearly observed for the ligands which display a redox potential (Eox) in the 0.9 – 1.0 V range. These values are slightly more positive than that of the N,N’ dimethylaniline (0.81 V vs. SCE[17]) due to the electron withdrawing character of the diazo –N=N– group in para position, that should promote a better stabilization of the generated amine radical cation through an extended charge delocalization. The complexation with Ru2+ hardly affects the values of these redox potentials, which indicates a weak electronic interaction between the rims of each complex. This effect is also consistent with the quasi invariance of the second oxidation potential observed for all complexes assigned to the metal-centred oxidation. These potentials nicely match the values measured for Ru(bpy)32+ (Figure 1). In the cathodic region, a reversible wave should be attributed to the reduction of the azobenzene moiety of L1-3.[18] In all cases, the
reduction potential is more positive than that of the 4-diethylamino-azobenzene (–1.57 V vs. SCE\textsuperscript{[18a]}). Moreover, \(E_{\text{red}}\) shifts in the positive direction by about 0.15 V on going from L3 to L1. This suggests that the azobenzene moiety is sensitive to the electron withdrawing strength of its bipyridine-based substituent group. Interestingly, the presence of Ru\(^{2+}\) strongly amplifies this shift: the first reduction potential of RuL1 is about – 0.95 vs. SCE as compared to a value of – 1.29 vs. SCE for L1. Finally, all ruthenium-based complexes typically display three reversible ligand-centred (i.e. bipyridine) reduction waves at the low potential range. Even though such successive reduction processes occur at very close potential values, the measured half-wave potentials are globally comparable to those measured for Ru(bpy)_3\(^{2+}\) (Table 1), whereas the dialkylamino-azobenzene moieties, whose oxidation potential remains invariant, present a better reducibility with a noticeable increase of \(E_{\text{red}}\) on going from RuL3 to RuL1. Such a change of the first reduction potential should presumably impact the relative position of the LUMO of the complexes (vide infra).

**Photophysical Properties.** The photophysical data of the compounds are gathered in Table 2. Figure 2 shows the absorption spectra of all chromophores in dichloromethane. In order to identify the position and the nature of the electronic transitions which are present within the longest wavelength absorption bands, DFT and TD-DFT calculations were carried out both for the ligands and their respective complexes.

All the ligands exhibit intense structureless bands in the blue visible region (Figure 2), with \(\varepsilon_{\text{MAX}}\) in the 28500 – 35000 M\(^{-1}\) cm\(^{-1}\) range. TD-DFT calculations predict the presence of two low-lying \(n\pi^*\) and \(\pi\pi^*\) states and indicate that their energy gap is below 0.13 eV for all ligands (Table 2). In particular, the TD-DFT results reveal that the symmetry forbidden \(n\pi^*\) transition mainly arising from HOMO-2 \(\rightarrow\) LUMO
contributions lies close to the ππ* one, for all three ligands L1-3. In other words, the observed structureless bands comprise two electronic transitions with distinctive symmetries: 

1. A S0 → S1 transition which has a very weak intensity due to its nπ* nature
2. A S0 → S2 transition which is a strongly allowed ππ* transition with a charge transfer (CT) character involving electronic delocalization all along the azobenzene backbone.[11, 18a, 19] Due to the electron withdrawing ability of the bipyridine fragment which leads to a better energy stabilization of the S0 → 1ππ* transition, the longest wavelength band of L2 is red-shifted by about 890 cm⁻¹ as compared to L4 (Figure S2). The energy diagram of frontier orbitals (Figure 3) indicates that the ππ* transition of L2 mainly corresponds to a HOMO-LUMO transition which exhibits a long range electronic delocalization all along the entire structure with a CT character from the aminoazobenzene to bipyridine moieties. Such CT character has been quantified by the amount of transferred charge (qCT) and the charge transfer distance (dCT) (Table S1). This is consistent with the fact that the ππ* transition is strongly sensitive to 4,4'-azobenzene substitution effects and shifts to low energy region when increasing the strength of donor and/or acceptor.[19-20]

The same electronic feature can be observed for L1. The fact that the frontier orbitals are not localized on either the donor (amino) or acceptor (bipyridine) moieties is consistent with the idea that the conjugated azo bridge allows for effective conjugation within this ligand. On the other hand, L1, with the azobenzene group directly bonded to the 4 position, shows stronger absorption band, at low energy, than that observed for 4-phenylazopyridine [310 nm (ε = 23360 L.mol⁻¹.cm⁻¹), 450 nm (ε = 265 L.mol⁻¹.cm⁻¹)],[21] in agreement with the expected greater push-pull character of L1.
In the case of L3, the HOMO (π-type) and LUMO (π*-type) are mainly localized on the photochromic part of the ligand with a very weak participation of the adjacent pyridine; such a disruption in the conjugation path should be presumably due to the presence of the triazole acting as a site-separating moiety.\[22\] Interestingly, comparable qCT and dCT are maintained in L3 (Table S1).

Complexation with Ru$^{2+}$ clearly enhances the electron withdrawing ability of the acceptor group (Figure S3).\[23\] A stabilization of the ππ* state due to a strong enhancement of its CT character is observed (Table 2). In turn, the longest wavelength absorption band observed for the free ligands shifts to the red region and increases in intensity upon complexation (Figure 2). Interestingly, bathochromic shift and hyperchromic effect both depend on the nature of the linker and on the positioning of the 2, 2'-bipyridine unit (i.e. position 4 vs. 5). A direct link of the bipyridine to the aminoazobenzene subunit (i.e. ligand L1) leads to the larger effects with a band red-shift by ca 4300 cm$^{-1}$ and a 1.7-fold increase in intensity. Such important spectral changes contrast to those observed for RuL3 where the band hardly shifts by ca 90 cm$^{-1}$ with a 1.2-fold intensity increase. Finally, the absorption band of RuL2 displays an intermediate behavior with a bathochromic shift of ca 650 cm$^{-1}$ and a 1.4-fold intensity increase. TD-DFT calculations reproduce with a good agreement such metal-induced spectral shifts ($\Delta$E < 0.2 eV), even if the calculated S$_1$ energy of RuL1 seems significantly overestimated (Table S2). In fact, a good correlation ($R^2$>0.979) is obtained when plotting the theoretical bathochromic shifts of the complexes compared to their related ligands vs. those determined experimentally.

The lowest-energy transition of RuL1 is dominated by a HOMO-LUMO ligand-centred transition (Figure S3) with a strong intramolecular CT character (Table S1) from the 4-dialkylaminoazobenzene to the ‘fused’ bipyridine chelating group. Interestingly,
similar electronic effects are observed for the $S_0 \rightarrow S_1$ transition of RuL2 which leads to a larger charge delocalization within the entire L2-subunit (Figure 4), leading to a dramatic increase of the charge transfer distance $d_{CT}$ from 2.5 to 4.3 Å (Table S1). However, in the case of RuL3, the $S_0 \rightarrow S_1$ transition which displays a mixed $\pi\pi^*$ and $n\pi^*$ character is only localized on the 4-dialkylaminoazobenzene subunit (Figure 4). As for L3, this confirms that the triazole linker between the azobenzene and bipyridine units leads to a strong disruption of the electronic conjugation\textsuperscript{[22]}. Moreover, our calculations exclude any metallic character within the lowest energy transition of RuL3 whereas a weak weighted contribution of the metal cation (< 5%) is noted within the LUMOs of RuL1 and RuL2. These findings suggest that the ruthenium tris(bipyridine) subunit is electronically decoupled with respect to its photochromic fragment. Such an assumption is also consistent with the unchanged redox properties of the metal complex site (vide supra). Moreover, similar electronic decoupling is suggested when comparing the absorption spectra of RuL1-3 with that of the Ru(bpy)$_3^{2+}$ model system (Figure 2). For RuL1-3, the high energy side of the spectrum (i.e. < 300 nm) displays the same spectral feature than that observed for Ru(bpy)$_3^{2+}$: i) a structured band at 240 nm corresponding to a first set of Metal-to-Ligand Charge Transfer (MLCT) transitions\textsuperscript{[24]} ii) a second intense band located at 290 nm which should be assigned to the ligand-centred (LC) transitions\textsuperscript{[24]} (bipyridine-centred transitions). The substitution effect on one of the three bipyridine ligands induces a slight decrease of the LC band with a larger effect for RuL3 (i.e. ~ 20 % decrease). The lowest-energy band of Ru(bpy)$_3^{2+}$ located at 455 nm corresponds to a second set of $S_0 \rightarrow ^1$MLCT transitions. This characteristic band is clearly observed at the blue edge of the charge transfer band of RuL1. However, on
going from RuL2 to RuL3, the MLCT band is progressively masked due to a gradual overlap with the lowest-energy band which shifts to the blue region.

Figure 5 shows the phosphorescence and the excitation spectra of the complexes in glassy matrix of ethanol. According to the excitation spectra recorded at the maximum luminescence wavelength, the MLCT band is dominating the excitation spectrum and coincides to that of Ru(bpy)₃²⁺. Therefore, the energy of the S₀ → ¹MLCT transitions remains invariant whatever the structural changes on complexes. It is worth noting that the lowest-energy band is not observed within the excitation spectrum of RuL1 and is hardly visible for the other complexes. This suggests that the efficiency of the intersystem crossing (ISC) toward the ³MLCT state is much higher upon excitation to the high-energy ¹MLCT state as compared to an excitation to the ¹ππ* one. This will have strong consequences on the photoisomerization properties of the complexes (vide infra). The phosphorescence spectra of all complexes are located in the same spectral range which yields to triplet energies of 2.15, 2.02 and 2.10 eV for RuL1, RuL2 and RuL3, respectively. Moreover, their luminescence lifetimes are 11, 10 and 8 µs, respectively, in agreement with that of Ru(bpy)₃²⁺ (~ 5 µs[18b, 24]). The lowest triplet state is the ³MLCT which would thus exclude any triplet-triplet energy transfer process from the excited ruthenium tris(bipyridine) subunit to the azo photochromic moiety.

Hence, coordination to Ru²⁺ clearly impacts the charge transfer character of the lowest-energy transitions which are mainly localized on the conjugated photochromic subunit (Figure 4, Table S1). Even though the metal coordination sphere globally maintains its electronic integrity, important changes will be observed in the efficiency of the photochromic switches.
Photoisomerization Reaction and Mechanism. Table 3 reports the values of the \( \text{trans} \rightarrow \text{cis} \) photoisomerization quantum yields for all the chromophores in various solvents as well as the rate constants of the thermal \( \text{cis} \rightarrow \text{trans} \) isomerization reactions. Figure 6 illustrates the changes of the absorption spectra of L2 and its complex during the isomerization reactions. Considering the simulated UV-visible spectra of both \( \text{trans} \) and \( \text{cis} \) isomers (Figure S4) and corresponding data (Table S3), we can first tentatively assign experimental changes of absorption spectra occurring after the irradiation inducing partial conversion of \( \text{trans} \) to \( \text{cis} \) conformation. As expected, calculations indicate that the \( \text{trans-cis} \) isomerisation induces a slight red-shift (ca 0.1 eV) and a significant increase of oscillator strength for the \( n\pi^* \) transitions mainly located on the azobenzene moiety (Figure S5). Indeed, the \( \text{cis} \) L1-3 ligands exhibit their first bright low-energy transition at 445 nm, 479 nm and 467 nm, respectively. For instance, changes in the UV-vis absorption spectrum of L2, which exhibits experimentally the best conversion ratio (Table 3), are qualitatively well explained by TD-DFT calculations. As can be seen from the observed spectrum of L2 under irradiation (Figure 6), the decrease of the intensity of the low-energy band indicates the partial conversion of the \( \text{trans} \) to the \( \text{cis} \) form. Simultaneously, the intensity of the absorption bands in the UV range increases upon irradiation. These experimental observations are consistent with our theoretical results, namely the decrease of the intensity of the calculated low-energy absorption band when going from \( \text{trans} \) to \( \text{cis} \) isomers (Figure S4). A weak increase is observed in the \( \text{trans-cis} \) energy differences for the simulated complexes RuL1-3 as compared to the ligands. The highest calculated \( \text{trans-cis} \) energy difference is found for RuL1. By contrast, the calculated \( \text{trans-cis} \) energy difference of RuL3 is almost comparable to the value obtained for the ligand (L3), indicating that the complexation with ruthenium should
only slightly affect the thermodynamics of the trans-cis isomerism. This is a direct consequence of the site isolation character of the triazole moiety (vide infra).

Figure 6 shows that upon continuous irradiation at 436 nm, the longest wavelength absorption band of the L2 strongly decreases concomitantly with the growth of new bands in the 270–375 nm range and above 490 nm. Three isosbestic points are observed at 256 nm, 385 nm and 512 nm. At the photostationary state, the intensity of the main absorption band collapsed by a factor ~ 4 while the low energy band of RuL2 only decreased by a factor 0.75. For this complex, a slight increase of the absorption bands is also observed in the 270-380 nm range with the presence of an isosbestic point at 401 nm. The trans-to-cis photoconversion yield is relatively high for L2 with a total conversion of ca 83 % in methylcyclohexane (MCH) which only decreases by a factor 1.2 in dichloromethane (DCM). For the other ligands, the nature of the solvent strongly influences the photoconversion efficiency, especially for L1 whose conversion yield drops from 69 to 0 % on going from MCH to DCM. In DCM the relative proportion of L2 cis isomers is divided by a factor ~ 8 with respect to that in MCH or THF. Such important changes globally parallel those observed for the rates of thermal cis-to-trans isomerization (Table 3). For instance, the rates of L1 and L3 vary over more than two orders of magnitude between MCH and DCM.

For the ruthenium complexes, the trans-to-cis photoisomerization quantum yields (when measurable) are globally divided by a factor 2.5 with respect to those of the ligands. As previously indicated, the metal-induced stabilization of the \(^{1}\pi\pi^*\)(CT) promotes a \(^{1}\pi\pi^*\) – \(^{1}\pi\pi^*\) state inversion (Table 2). The photoisomerization dynamics will be thereby strongly modified. For instance, the dramatic decrease of the S\(_1\) state energy observed for RuL1 (Ru directly involved in the conjugation pathway) probably favours the internal conversion processes according to the energy gap law.
Photoinduced intramolecular electron transfers (PeT) should be considered as minor deactivation channels for all the complexes. The free energy ($\Delta G_{eT}$) for such a PeT can be estimated from the Rehm–Weller equation$^{[25]}$: 

$$\Delta G_{eT} = E_{ox} - E_{red} - E_{00}$$

neglecting the Coulomb part of the stabilization energy. According to the redox properties of the complexes (Table 1) and estimating the singlet state energy ($E_{00}$) from the onset of the absorption CT band in DCM, $\Delta G_{eT}$ is estimated to be close to 0 eV for RuL1 and RuL2 and becomes hardly negative for RuL3 (−0.13 eV).

Noteworthy, the irradiation of the complexes at 436 nm corresponds to a direct excitation into their $^1$MLCT states. This should open a new deactivation channel to the $^3$MLCT state as confirmed by the phosphorescence fingerprints of the complexes (Figure 5). Indeed, the intersystem crossing to the triplet MLCT state occurs rapidly in less ~ 300 fs$^{[26]}$ for Ru(bpy)$_3^{2+}$ and with a $\Phi_{ISC}$ close to unity.$^{[24]}$ Interestingly, the phosphorescence properties of the complexes in deoxygenated THF are hardly affected by the trans-to-cis photoisomerization (see Figure S6)$^{[9c, 27]}$. This observation is in line with the fact that the lowest triplet state is the $^3$MLCT which excludes any triplet-triplet energy transfer to the azo photochromic moiety. This suggests that the significant decrease of $\Phi_{t\rightarrow c}$ observed for RuL1-3 presumably stems from this competing deactivation channel to the $^3$MLCT state as depicted in Scheme 4. By contrast to other azobenzene-conjugated Ru$^{2+}$ complexes where the photoisomerization properties are completely switched off,$^{[28]}$ the photochromic part of RuL2 and RuL3 remains relatively photoactivable despite the competition of intersystem crossing. This indicates that both competing processes should exhibit comparable rate constants.

**Conclusion.**
The present results demonstrate the importance of the position and nature of the linker in azobenzene containing polypyrroline systems and their effects on the electronic coupling between donor and acceptor, in both ligands and complexes. This in turn leads to strong modulations of the photoisomerization properties of these species. An efficient electronic communication along the whole molecular backbone with extended conjugation, which involves the Ru cation, disfavours the photoisomerization reaction. In addition, the very low activation barrier along the thermal cis → trans isomerization path prevents any photostationary cis / trans equilibrium at room temperature. By contrast, an extension of molecular backbone from the bipyridine to the azobenzene with an ethynyl or a triazolyl linker results in photoswitchable complexes where the electronic communication along the azo containing ligand is weakly influenced upon coordination. Thus, our approach allows to suggest a route to provide polypyrroline Ru(II) complexes where the specific properties of the metal centre and the optical response of the organic group can be gathered.

**Experimental Section**

**General synthetic procedure for Ru(bpy)$_2$(L1-3)(PF$_6$)$_2$ complexes.** Cis-dichloro, bis-(2,2'-bipyridine)-Ruthenium(II) (0.1 mmol) and corresponding substituted bipyridine L (0.1 mmol) were dissolved in 10 mL of a deoxygenated mixture of ethanol/water (8:2) and heated at reflux for 3 h. After this time the mixture was cooled down to room temperature, 30 mL of water and 10 mL of a KPF$_6$ saturated aqueous
solution were added. The precipitate was filtered, washed with water and diethyl ether and the red solid was dried under vacuum.

**Ru(bpy)\(_2\)(L1)(PF\(_6\))\(_2\)** (RuL1). The pure product was obtained after column chromatography [SiO\(_2\), CH\(_3\)CN/H\(_2\)O/ sat. KNO\(_3\) (95:5:0.5)], as a red solid (97 mg, 89%). \(^1\)H NMR (CD\(_2\)Cl\(_2\), 400 MHz): \(\delta = 8.70\) (s, 1H), 8.55-8.45 (m, 5H), 8.15-8.05 (m, 5H), 7.96 (d, \(J = 9.2\) Hz, 2H), 7.84 (d, \(J = 5.4\) Hz, 1H), 7.80-7.65 (m, 6H), 7.55-7.45 (m, 5H), 6.79 (d, \(J = 9.2\) Hz, 2H), 3.47 (m, 4H), 1.69 (m, 4H), 1.44 (m, 4H), 1.02 ppm (t, \(J = 7.3\) Hz, 6H). \(^{13}\)C \(\{^1\)H\} NMR (CD\(_2\)Cl\(_2\), 100 MHz): \(\delta = 158.3, 156.8, 156.3, 156.1, 156.0, 155.9, 152.5, 151.0, 150.6, 150.5, 150.3, 142.6, 137.3, 127.4, 127.3, 127.2, 123.6, 123.6, 123.5, 123.3, 119.4, 115.1, 111.0, 50.5, 28.7, 19.4, 12.9 ppm. MS (Zabspec-TOF): \(m/z\) 400.6425 \([M^{++}\] \), 946.2477 \([M^{++} + PF_6^{-}\] \); elemental analysis calcd (%) for C\(_{44}\)H\(_{45}\)F\(_{12}\)N\(_9\)P\(_2\)Ru, H\(_2\)O: C, 47.66; H, 4.27; N, 11.37; found: C, 47.92; H, 4.23; N, 11.43.

**Ru(bpy)\(_2\)(L2)(PF\(_6\))\(_2\)** (RuL2). The pure product was obtained after column chromatography [SiO\(_2\), CH\(_3\)CN/H\(_2\)O/ sat. KNO\(_3\) (95:5:0.5)], as a red solid (86 mg, 72%). \(^1\)H NMR (CD\(_2\)Cl\(_2\), 500 MHz): \(\delta = 8.55-8.45\) (m, 6H), 8.19 (dd, \(J = 8.5, 2\) Hz, 1H), 8.15-8.05 (m, 5H), 7.85 (d, \(J = 9.3\) Hz, 2H), 7.82 (d, \(J = 8.6\) Hz, 2H), 7.80-7.70 (m, 6H), 7.60 (d, \(J = 8.6\) Hz, 2H), 7.55-7.45 (m, 6H), 6.74 (d, \(J = 9.3\) Hz, 2H), 3.41 (m, 4H), 1.66 (m, 4H), 1.42 (m, 4H), 1.01 ppm (t, \(J = 7.4\) Hz, 6H). \(^{13}\)C \(\{^1\)H\} NMR (CD\(_2\)Cl\(_2\), 125 MHz): \(\delta = 156.8, 156.6, 156.2, 155.4, 152.7, 152.4, 151.4, 151.3, 151.3, 143.0, 140.1, 138.3, 138.3, 132.7, 128.6, 128.2, 128.2, 128.1, 125.6, 124.8, 124.7, 124.4, 124.3, 123.9, 122.2, 121.0, 111.2, 98.1, 84.8, 50.9, 29.4, 20.2, 13.7 ppm. MS (Zabspec-TOF): \(m/z\) 450.6578 \([M^{++}\] \), 1046.2829 \([M^{++} + PF_6^{-}\] \); elemental analysis calcd
(%) for C₅₂H₄₉N₉F₁₂P₂Ru, H₂O: C, 51.66; H, 4.25; N, 10.43; found: C, 51.58; H, 3.99; N, 10.12.

**Ru(bpy)₂(L₃)(PF₆)₂ (RuL₃).** The pure product was obtained after column chromatography [SiO₂, CH₃CN/H₂O/ sat. KNO₃ (95:5:0.5)] as a red solid (89 mg, 72%). ¹H NMR (CD₂Cl₂, 400 MHz): δ = 8.71 (s, 1H), 8.60 (d, J = 8.3 Hz, 1H), 8.56 (d, J = 8.3 Hz, 1H), 8.32 (s, 1H), 8.14 (t, J = 8.2 Hz, 1H), 8.08 (m, 4H), 7.98 (d, J = 8.7 Hz, 2H), 7.92 (d, J = 9.0 Hz, 2H), 7.90 (m, 1H), 7.84 (m, 1H), 7.83 (d, J = 8.7 Hz, 2H), 7.75 (m, 3H), 7.49 (m, 1H), 7.42 (m, 4H), 6.81 (d, J = 9.0 Hz, 2H), 3.47-3.38 (m, 4H), 1.63 (m, 4H), 1.40 (m, 4H), 0.98 ppm (t, J = 7.3 Hz, 6H). ¹³C {¹H} NMR (CD₃CN, 400 MHz): δ = 156.8, 156.7, 156.7, 156.4, 155.8, 152.9, 151.7, 151.6, 151.5, 151.4, 151.1, 147.4, 142.4, 142.3, 137.7, 137.6, 137.5, 137.5, 136.1, 133.5, 130.0, 127.3, 127.3, 127.3, 125.1, 124.3, 124.1, 124.0, 124.0, 124.0, 122.8, 121.1, 120.9, 111.0, 50.2, 28.8, 19.6, 12.9 ppm. MS (Zabspec-TOF): m/z: 466.6282 [M⁺], 1078.2177 [M++] + PF₆⁻]; elemental analysis calcd (%) for C₅₂H₅₀F₁₂N₁₂P₂Ru, H₂O: C, 49.88; H, 4.19; N, 13.42; found: C, 49.61; H, 4.13; N, 13.58.

More experimental and computational details are available in the Supporting Information.

**Acknowledgements**

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References.


FIGURES AND TABLES CAPTIONS

Scheme 1  Molecular structures of the metal complexes and corresponding ligands.

Scheme 2  Reaction scheme for the synthesis of L1.

Scheme 3  Reaction scheme for the synthesis of L2 and L3.

Scheme 4  Simplified photophysical schemes relative to D-\(\pi\)-A ligands and their metal complexes when excited into their S\(_2\)(\(\pi\pi^*\)) and \(^1\)MLCT states, respectively.

Figure 1  Cyclic voltammograms of L1, RuL1 and Ru(bpy)\(_3^{2+}\)·2PF\(_6^-\) (mM range in acetonitrile + NBu\(_4\)PF\(_6\) at 0.1M). \(v = 200\) mV s\(^{-1}\).

Figure 2  Absorption spectra of the free ligands and their respective Ru\(^{2+}\) complexes in dichloromethane (Dotted lines: absorption spectrum of RuL complexes; full lines: L ligands; dash-dotted lines: Ru(bpy)\(_2^{2+}\), 2PF\(_6^-\)).

Figure 3  Frontier molecular orbitals (MO) diagrams of L1-3 ligands with main transitions.

Figure 4  Density differences plots (\(\Delta \rho(r) = \rho^{S_1}(r) - \rho^{S_0}(r))\) of RuL1-3 between S\(_1\) - S\(_0\) states (light/yellow colour = increase; dark/blue colour = decrease of electron density; isovalue 0.03 au).

Figure 5  Phosphorescence (squares) and excitation (circles) spectra of the complexes in glassy matrix of ethanol (77K). (Dash-dotted lines: phosphorescence and excitation spectra of Ru(bpy)\(_3^{2+}\) in the same conditions).

Figure 6  Evolution of the absorption spectra of L2 in MCH and RuL2 in THF during the isomerization reactions (i.e. \(h\nu\) or \(\Delta\)), dotted lines: stationary states. Insets: (a) Plots of the maximum absorbance at 436 nm as function of the irradiation time (b) Plots of the maximum absorbance at 436 nm during the cis \(\rightarrow\) trans thermal reaction (sample is kept in the dark). Least square fit using a simple first-order equation (i.e. \(f(t) = A + B(1 - e^{-k_{th}t})\) with as \(A\), \(B\) and \(k_{th}\) as variables).

Table 1  Redox potentials of the compounds in acetonitrile.

Table 2  Absorption properties of the compounds and corresponding TD-DFT data (solvent: dichloromethane).

Table 3  Isomerization properties of compounds in various solvents (\(\lambda_{irr}: 436\) nm).
Scheme 1
Scheme 2
Scheme 3
Scheme 4
Figure 1

$E / V$ vs. SCE
Figure 2
Figure 3
**RuL1**: $S_1 - S_0 (\pi-\pi^*)$

**RuL2**: $S_1 - S_0 (\pi-\pi^*)$

**RuL3**: $S_1 - S_0 (n-\pi^* + \pi-\pi^*)$
Figure 5
Figure 6
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<th>$E_{ox}$ V vs. SCE</th>
<th>$E_{red}$ V vs. SCE</th>
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<td>L1</td>
<td>0.95</td>
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<td>RuL1</td>
<td>1.00, 1.34</td>
<td>0.95, -1.31, -1.49, -1.72</td>
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<td>0.89, 1.29</td>
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<td>-1.35, -1.54, -1.79</td>
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(i) Irreversible wave.
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<th>Type</th>
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<td>L1</td>
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<tr>
<td>278 (54.2), 459 (30.3)</td>
<td>440 0.0 n-π' H₂ → L (0.91)</td>
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<tr>
<td></td>
<td>421 1.3 π-π' H → L (0.93)</td>
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<tr>
<td>RuL1</td>
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<tr>
<td>292 (72.3), 465 (21.3), 572 (50.9)</td>
<td>471 1.7 π-π' H → L (0.90)</td>
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<tr>
<td></td>
<td>446 0.0 n-π' H₄ → L (0.83)</td>
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<tr>
<td>L2</td>
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<td>343 (24.8), 472 (28.2)</td>
<td>447 2.2 π-π' H → L (0.85)</td>
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<td>435 0.0 n-π' H₂ → L (0.73)</td>
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<td>RuL2</td>
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<td>290 (67.3), 358 (25.3), 493 (42)</td>
<td>465 2.3 π-π' H → L (0.57), H → L₃ (0.30)</td>
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<td>439 0.0 H₅ → L₃ (0.55), H₅ → L (0.30)</td>
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<td>L3</td>
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<td>303 (27.3), 460 (35.1)</td>
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<td>291 (63.4), 320 (35.5), 464 (43.5)</td>
<td>437 0.8 n-π' + π-π' H₅ → L₃ (0.42), H → L₃ (0.34)</td>
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<td>424 1.0 π-π' H → L₃ (0.43), H₅ → L₃ (0.34)</td>
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Table 2
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<td>$\tau_{1/2}^c$ / s</td>
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$^a$no change of the absorption spectrum, $^b$first-order reaction rate, $^c\tau_{1/2} = \ln 2 / k_{th}$, $^d$insoluble.
Graphical Abstract
Engineering of azobenzene-containing bipyridine ligands leads to strong modulations of the photoisomerization properties of their ruthenium heteroleptic complexes: an electronic communication involving the ruthenium cation disfavours the photoisomerization; by contrast, extending the molecular backbone from the bipyridine to the azobenzene with ethynyl or triazolyl linkers results in photoswitchable complexes.

Short title if needed:
Photoswitchable azobenzene-ruthenium complexes
Keywords

Photochromism; Azo compounds; Coordination chemistry; Ruthenium; Density functional calculations