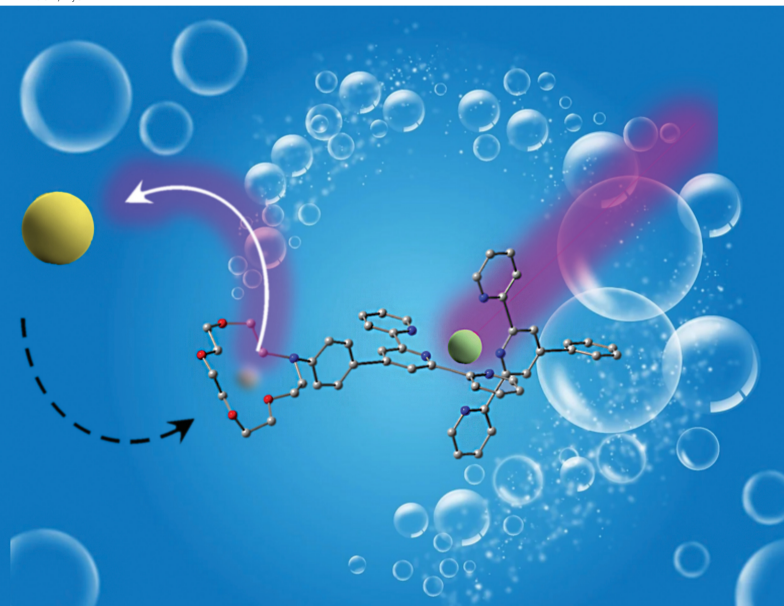




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Cedric Mongin, Stephane Aloise *et al.*Photoejection-recapture of the Ca²⁺ cation studied by time resolved spectroscopy and TDDFT calculations: the case study of an azacrown-iridium(III) complex





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Photoejection-recapture of the Ca²⁺ cation studied by time resolved spectroscopy and TDDFT calculations: the case study of an azacrowniridium(III) complex†

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In this study, we examined the photophysical properties of an azacrown-iridium(iii) complex while focusing on its interactions with calcium ions (Ca2+). We explored the dynamic processes within the complex combining time-dependent density functional theory (TDDFT) calculations and time-resolved spectroscopies. In the presence of Ca²⁺, the complex exhibits significant shifts in absorption and emission profiles, from 494 nm to 375 nm, aligning with theoretical predictions. Notably, we observed the ultrafast photo-ejection of Ca²⁺ within 70 femtoseconds, followed by its recapture in 250 nanoseconds, revealing a 10-million-fold timescale difference between the two phenomena. These behaviors confirm the established photophysical properties of polypyridyl iridium(III) complexes and their intrinsic sensitivity to their surrounding environment. Our comprehensive kinetic analysis highlights the azacrown moiety's competitive binding and photo-release capabilities, suggesting its potential for practical sensing applications. The versatile properties of these iridium(III) complexes offer promising prospects for their application as stimuli-responsive materials and in advanced optoelectronic devices, targeted imaging, and biomedical ion sensors and delivery systems

chemical strategies.

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1. Introduction

Complexation and precise spatiotemporal release of bioactive compounds are becoming increasingly vital in the fields of supramolecular chemistry, life science, and nanomedicine. A significant focus has been directed towards the development of chemical cages for metal ions, particularly calcium ions (Ca²⁺), due to their biological significance.2 In the pursuit of achieving precise control over bioactive cations, the use of light is appealing because of its non-invasive nature and its ability to be delivered with high spatial and temporal precision. Among all the possible chemical strategies, one solution involves using photo-induced charge transfer (PCT) molecules combined with

a complexing agent, typically an aza-crown compound:

PCT induces positive polarization of the heteroatom of the

macrocycle, leading to coulombic repulsion between the heteroatom and the metal cation initially present in the cavity. Such a system has been proposed using various compounds, including azacrown-substituted [(bpy)Re(CO)₃L]⁺ complexes³⁻⁵ and azacrown-substituted merocyanine. 6-9 In 2016, we investigated a novel system, azacrown-substituted pyridinum betaine (PyB-Aza) by combining stationary spectroscopy, ultrafast absorption spectroscopy and DFT calculations. Focusing on PyB-Aza, the first aim was to assess the competitive complexation of the imidazole bridge vs. the macrocycle. In acetonitrile, it was found by absorption and emission that the imidazole moiety binds efficiently through lateral electrostatic interaction to form a (1,1) metal-ligand complex with $\log K = 6.8$. From a photochemical point of view, the rate of the photo-release process, assessed by tracking the transient stimulated emission band was found to be less than 200 fs, the fastest photorelease characteristic time reported so far. Furthermore, it has been shown that this rate of photorelease is proportional to the complexation constant. Finally, no evidence was found for a photoinduced translocation from an imidazole bridge to crown ether. Currently, our group is interested in exploring novel

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The study of iridium(III) complexes' photophysics represents a cutting-edge topic within the field of photoluminescent materials. 10-13 These complexes, renowned for their exceptional luminescent characteristics, are pivotal contenders for a diverse range of applications, spanning optoelectronic devices to bioimaging agents. 14-16 Upon further examination, iridium(III) complexes can be classified into two primary categories: cyclometalated complexes and coordination complexes, including polypyridyl complexes. Cyclometalated iridium(III) compounds present at least one characteristic carbon-iridium bond and exhibit strong absorption in the visible region arising from intense metal-toligand charge transfer (MLCT) singlet state transition and high emissivity due to their lowest energy excited states being a blend of strong ³MLCT and triplet ligand-centered (³LC) π - π * states, rather than metal centered (MC) non-radiative d-d states. 11,12 Among phosphorescent compounds, cyclometalated iridium(III) complexes emit from triplet excited states at room temperature with exceptional quantum yields up to unity and radiative rates in the microsecond regime.¹⁷

In contrast, polypyridyl complexes often exhibit ligand-toligand charge transfer (LLCT) and intra-ligand charge transfer (ILCT), which influence their photophysical behavior and result in weaker or even no MLCT transitions. 12,18-21 Consequently, these complexes are often less emissive and display weaker phosphorescence at room temperature. However, due to the significant alteration of the polarization in the excited state, they are highly susceptible to minor alterations in their immediate surroundings, rendering them promising candidates for ion chelation and photo-ejection monitoring.^{20,22}

This work will concentrate on the photophysical properties of polypyridyl complexes, with a particular emphasis on heteroleptic bisterpyridine iridium(III) complexes. These complexes are distinguished by their molar extinction coefficients, which are typically lower than those of cyclometalated complexes. Thus, the oscillator strength of the transitions is a critical parameter, influencing their photophysical properties.

The absorption and emission properties of these complexes are largely influenced by the nature of the ligands. The strong absorption bands around 450 to 500 nm, corresponding to the ¹ILCT state, and the emission band above 500 nm, arising from the intersystem crossing (ISC) to the ³ILCT, are sensitive to changes in solvents and pH.11,12 This makes them excellent candidates for use in sensors and other applications where changes in environmental conditions need to be monitored. They also exhibit emission in the visible region involving the lowlying ³ILCT state. Furthermore, strong wavelength-dependent photoluminescence has been reported, as well as dual phosphorescence.^{23,24} The latter phenomenon is associated with the formation of a specific ion pair, which results in the iridium center and a counter ion, particularly PF6-, being in close proximity in organic solvents such as acetonitrile.²³ This leads to a decrease in the excited state energy, which in turn causes the emission from the ³ILCT to shift towards the red end of the spectrum. The emission of the ion pair is similar to that observed in the solid state, where the crystalline structure allows for the specific interaction to occur. This study has explored the

Scheme 1 Structure of molecule 1

photophysics of these systems using time-resolved spectroscopies, characterizing very short ¹ILCT lifetimes (approximately 100 fs),²⁵ dynamics dependent on substituents,26 and solvent-dependent ISC 1 ILCT \rightarrow 3 ILCT (70 fs-3 ps). 27 It is also noteworthy that the use of iridium(III) cyclometalated complexes substituted with metallic cation chelating moieties has been previously investigated. This research has demonstrated a notable alteration of the phosphorescent properties, both in the absence of a metallic cation (weak 3ILCT emissive states) and in the presence of a metallic cation (strong ³LLCT/³MLCT emissive state).²⁸

In this paper, we are investigating a new class of iridium(III) terpyridine complex, molecule 1 (Scheme 1), that comprises an azacrown moiety dedicated to trap and photo-release calcium cations. This molecule has been studied using a combination of stationary and ultrafast spectroscopies together with (TD-)DFT calculations.

Results and discussion

Synthesis

The synthesis of molecule 1 is outlined in Scheme 2. Terpyridine 4 was synthesized from 2 and 3 according to a method from the literature via the Paal Knorr reaction in 70% yield.²⁹ The synthesis of the iridium complex was achieved using a twostep method described by Collin et al. 30 IrAzaTpyCl3 was obtained in 40% yield by a reaction of IrCl₃ and 4 in ethylene glycol at 160 °C. The formation of the final complex was achieved by heating complex 5 in the presence of 6, followed by anion exchange with PF₆⁻. The purities of the complexes were carefully checked by 1H NMR spectroscopy and highresolution mass spectrometry.

2.2. Photophysical framework and complexation scheme

Absorption and complexation properties. The absorption spectra of 1 in CH₃CN are presented in Fig. 1 with three features in the UV part and a strong absorbance band in the visible range peaking at 494 nm responsible for the red color of the solution. Upon addition of the Ca²⁺ cation (perchlorate calcium solution), the solution color turns gradually to yellow. Indeed, due to the complexation reaction the visible band vanishes giving way to a new band peaking at 375 nm assigned to the 1-Ca2+ complex. Note that a large excess of calcium perchlorate has been tested but only one complexation constant is required during multicurve fitting (see the details in the ESI†).

Scheme 2 Synthesis of the iridium complex 1

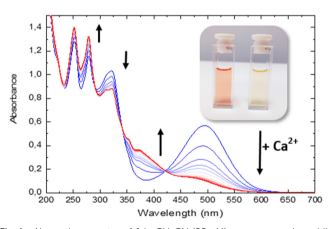


Fig. 1 Absorption spectra of ${\bf 1}$ in CH $_3$ CN (28 μ M) upon successive addition of calcium perchlorate solution from 0 to 0.29 M. The inset shows a picture of the solution of 1 before (blue) and after adding a large excess of Ca2+ ion (red).

The complexation constant $\log K = 2.4 \pm 0.002$ (K = 250) is assigned to a (1:1) stoichiometry complex as evidenced by the JOB plot presented in Fig. S1 in the ESI.† Note that this latter value is in accordance with various aza-crown-Ca²⁺ complexes found in the literature for PCT based fluorescent sensors inducing cation photorelease.31,32

Theoretical insights. As seen in Fig. 2, the TDDFT calculations were able to reproduce both 1 and 1-Ca²⁺ experimental

absorption bands at 494 nm (calc. 540 nm) and at 375 nm (calc. 389 nm) respectively. For molecule 1, the calculated maximum absorption wavelength (540 nm) does not correspond to the $S_1 \leftarrow S_0$ electronic transition (LLCT transition with an oscillator strength close to zero, see Fig. S2 in the ESI†) but rather to a $S_2 \leftarrow S_0$ electronic excitation arising from a HOMO \rightarrow LUMO+1 transition. A close inspection of the variation of electronic density shows a depletion from the phenyl-azacrown moiety toward the terpyrydine (tpy) chromophore which corresponds to an ILCT transition. Note that according to our computational strategy, the analysis of charge transfer (CT) descriptors indicates a large CT dipole moment (μ_{CT} = 23 D) along with a large CT distance ($D_{\text{CT}} = 6.3 \text{ Å}$).

For 1-Ca²⁺ (with the cation inside the crown ether), the maximum absorption band is now related to the S1 electronic state which arises from a similar aza → tpy ILCT transition with CT descriptor values close to those of molecule 1 ($\mu_{\rm CT}$ = 21.6 D, D_{CT} = 4.6 Å). These descriptors suggest that 1-Ca²⁺ is a good candidate for cation photoejection since there is a depletion of electronic density on the azacrown complexation site after excitation. In addition to this, we considered the evolution of the distances of Ca²⁺ in the azacrown upon relaxation on the first singlet excited state (Table S1, ESI†). One can notice a preejection process of Ca2+ from the azacrown with (i) a distance between the aza-nitrogen and Ca2+ stretched by more than 1 Å and (ii) an average distance between the oxygens and Ca²⁺ increased by 0.05 Å.

Paper

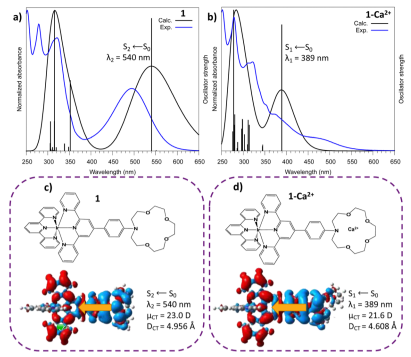


Fig. 2 Theoretical excitation energies and convoluted absorption spectra (convolution: FWHM = 0.25 eV) (a) for 1 and (b) the $1-Ca^{2+}$ complex. The variations of the electronic density upon excitation ($\Delta \rho \vec{lr}$), red: gain of electronic density, blue: depletion of electronic density) are shown for the most important transition related to the calculated bands peaking at 540 nm for 1 (c) and at 389 nm for $1-Ca^{2+}$ (d). The charge transfer dipole moment \vec{u}_{CT} is also represented (golden arrow) and the charge transfer parameters are given

Finally, as anticipated with transient laser experiments, for molecule 1, an excitation near 350 nm coincides with the $S_6 \leftarrow S_0$ transition which presents an ILCT character (calculated value: 352 nm; see Fig. 2 and Fig. S2, Table S2, ESI†). In contrast, for complex 1-Ca2+, there is no transition with a strong oscillator strength in this region (Fig. 2 and Table S3, ESI†). In summary, according to theoretical predictions, UV excitation of 1-Ca2+ will only induce a $S_1 \leftarrow S_0$ electronic excitation and an instantaneous Aza \rightarrow tpy ILCT process, leaving a positive charge on the crown nitrogen, inducing the pre-ejection of Ca²⁺ into the bulk. The corresponding ILCT state of molecule 1, namely S2, should then be populated.

Emission properties. Emission spectra of 1 and 1- Ca^{2+} are presented in Fig. 3 for 310, 355 and 425 nm excitations

(together with absorption spectra). Three features are noticed according to the excitation wavelength: (i) a narrow emission band E1 near 350 nm with one vibronic feature (1500 cm⁻¹ spacing) and poorly sensitive to the presence of Ca²⁺; (ii) a broad emission band E2 peaking at 550 nm with two vibronic transitions (1647 and 1570 cm⁻¹ spacing) and not sensitive to the presence of Ca²⁺; and (iii) a broad band E3 peaking at 605 nm without Ca²⁺ that vanishes after the addition of calcium perchlorate. All three features have already been reported in the literature. The band E1 is the well-known fluorescence band of the iridium complex related to the ILCT process that can be observed for heteroleptic³³ or homoleptic iridium complexes.²¹ The lifetime of the ¹ILCT state – responsible for the E1 emission

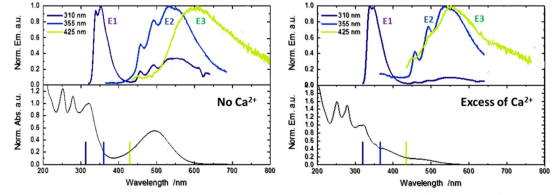


Fig. 3 Absorption and emission spectra upon 310, 355 and 425 nm excitation for 1 in CH₃CN (left) and 1-Ca²⁺ (excess of Ca²⁺) in CH₃CN (right) (10 μM). The E3 band was missing after addition of calcium perchlorate.

band - was determined using the TCSPC technique under 310 nm laser excitation. A mono-exponential decay was found with a 2.37(9) ns lifetime in accordance with literature values. In the presence of Ca²⁺, one observes a slight increase of the latter value up to 2.82(2) ns which suggests a possible electrostatic effect.

By comparison with the literature, with its typical vibronic structure, the E2 band is assigned at first glance to the triplet phosphorescence originating from the ³LLCT state for homoleptic iridium complexes.²⁴ However, in our case, for molecule 1, TDDFT calculations provide evidence that T₁(³ILCT) is the lowest in energy while the triplet T₂(³LLCT) is quasi degenerated with both the singlet state populated after photo-ejection, namely the S₂(ILCT) and S₁(LLCT) states (Fig. S2, ESI†). Therefore, the E2 band could be attributed to delayed emission involving LLCT states. Finally, the unusual E3 band that arises for 425 nm light excitation is due to ion-pair formation between the metallic center Ir³⁺ and the counter anion (PF₆)⁻.²³ With anticipation, this latter state can be important to rationalize the upcoming transient spectroscopy experiments.

2.3. Ultrafast dynamics and cation ejection

In order to depict the full picture of molecule 1 with and without Ca²⁺ cations, we will focus on transient spectroscopy with laser excitation near 350 nm to ensure that triplet states are surely populated knowing that cation ejection can occur at singlet or triplet manifolds. Two set-ups were employed corresponding to two distinct ranges - femtosecond/picosecond and nanosecond/microsecond regimes. The overall results are reported in a Jablonski diagram at the end of the article.

Microsecond regime. As seen in Fig. 3, with the two overlapping contributions E2 (delayed fluorescence) and E3 (ionpair), it is interesting to decipher the photophysics of those two emissive states. In Fig. 4, under 355 nm excitation, in the luminescent mode, one is able to identify the temporal decay of two distinct spectroscopic signatures: the vibronic structure is related to delayed emission near 450 nm while the redshift

long tail is related to the ion-pair near 575 nm. Because of the double contribution, every wavelength is better fitted by a biexponential function. However, the E2 emission is predominant near 450 nm while near 575 nm the ion-pair state emission becomes preponderant. In this way, we can assign a lifetime of 2.1 µs for S₁/T₂(LLCT) states and 6.1 µs for the ionpair state. Note that unlike a regular triplet state, the ion-pair state is only partially quenched by oxygen (6.1 $\mu s \rightarrow 1.5 \mu s$ with O₂) which suggests a non negligeable triplet character with a singlet-triplet mixed character possible. However, an in-depth study of the ion pair photophysics is not in the scope of this paper. Still in luminescent mode, the addition of Ca²⁺ drastically suppresses the two strong bands for two different reasons: the splitting of the S1-T2 energy gap for the delayed emission band on the one hand, and the electrostatic screening effect with the perchlorate anion for the ion pair band on the other hand.

According to the strategy of Moore et al., the recapture of the Ca²⁺ cation can be highlighted with laser flash photolysis with a regular absorption configuration. The transient data have to be compared with the difference in the stationary spectra with and without excess of the cation (the absorption curve with the Ca²⁺ cation is subtracted from the absorption curve without Ca²⁺). 4,31 Both μ-second TA data after 355 nm laser excitation and difference spectra are presented in Fig. 4. Clearly, the transient band perfectly matches the difference spectra, providing evidence that this band is associated with the photoproduct, namely molecule 1 in its excited states following Ca²⁺ photo-release. The kinetics at 500 nm is fitted with a monoexponential function with a 250 ns characteristic time that represents the mean time for the Ca²⁺ recapture.

Femtosecond regime. The results of femtosecond transient spectroscopy for a laser excitation at 350 nm are presented in Fig. 5. Initially, we will analyze the photophysics of molecule 1 without Ca^{2+} . In the first temporal window [-0.25; 0.16 ps], one can observe the rise of the signal along a negatively centered

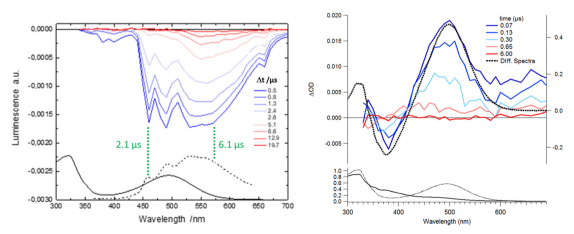


Fig. 4 (left) Transient emission traces obtained with laser flash photolysis in luminescent mode for 1 in CH₃CN (200 μM) with 355 nm laser excitation. Biexponential fits have been performed at two chosen wavelengths (green dashed line) and the predominant characteristic time is indicated. (right) Transient absorption for $\mathbf{1}$ - \mathbf{Ca}^{2+} (excess of \mathbf{Ca}^{2+}) in CH₃CN with 355 nm laser excitation. The kinetics at a selected wavelength are better fitted with a mono-exponential function

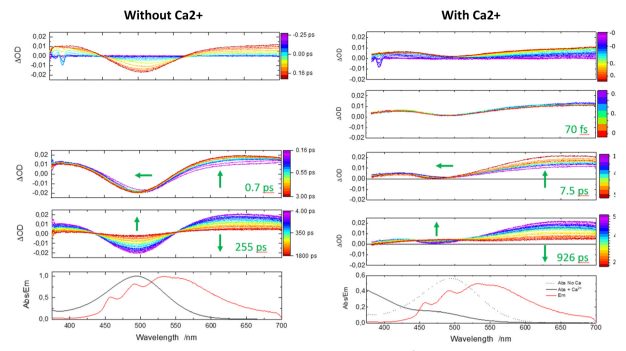


Fig. 5 Femtosecond TA for a solution of 1 in CH₃CN without (left) and with excess (right) of Ca²⁺ cations under 350 nm laser excitation presented in different temporal windows from -0.25 ps to 2000 ps. Global fitting has been undertaken at selected wavelengths (max of transient bands, bleaching bands...) and the characteristic times are indicated. The first panel represents the rise of the TA signal. The last panels are stationary absorption and emission curves for the 350 nm excitation

bleaching band at 490 nm, with two positive transient bands below 450 nm and above 550 nm. With respect to TDDFT calculations, S₆(ILCT) is populated upon 350 nm excitation but S₂(ILCT) has a lower energy (and the S1 state obtains the LLCT character, Fig. S2, ESI†). Providing that $S_6 \rightarrow S_2$ internal conversion occurs at an ultrafast scale well below our temporal resolution, the two positive transient bands are probably related to the S₂(ILCT) state absorption. The second temporal window [0.16-3 ps] is characterized by a small rise in the red edge transient band together with a slight blue shift of the bleaching band with a 700 fs characteristic time. Taking into consideration the participation of the $T_1(ILCT)$ (see the above discussion), this ultrafast event is logically attributed to the ISC processes $S_2(ILCT) \rightarrow T_1(ILCT)$. The question about the second route $S_2(ILCT) \rightarrow S_1(LLCT) \rightarrow T_1(LLCT)$ is still open but this through space mechanism is more likely to occur upon a bimolecular process, i.e., in a longer time range. Finally, in the third temporal window [4-1800 ps], one can observe the simultaneous decrease of the positive bands along with the total disappearance of the bleaching band both with a characteristic time of 255 ps (very negligible residues are present after 2 ns, probably related to ion-pair species). This final process is safely attributed to $T_1(ILCT) \rightarrow S_0$.

Now, one can analyze the femtosecond transient data taking into account the presence of the Ca²⁺ ion. In comparison, the first panel shows immediately a similar rise in positive transient bands related to S₁(ILCT) (originating from 1-Ca²⁺ species) except that the negative bleaching band is strongly reduced in accordance with stationary spectroscopy. Then, a

novel discrete process, a slight decrease of intensity at the red spectral edge, is reported on a second temporal window [0.45; 0.9 ps] with a characteristic time of 70 fs. Because the complexation constant is somewhat weak (K = 250), one does not expect a drastic decrease of the excited state absorption, and for that reason, we assign that process to the photo-release of the Ca²⁺ cation. That assignment is consistent with the 100 fs photo-release reported for betaine pyridinium molecules.³⁴ The following temporal window [1; 50 ps] logically displays the expected result, that is to say, ISC from S₁(ILCT) to T₁(ILCT) but with a characteristic time of 7.5 ps. This drastic increase by a factor of 10 compared to molecule 1 alone can be rationalized by an electrostatic effect induced by the "loose complex", i.e. Ca²⁺ is still in the neighboring molecule.⁸ Indeed, the energy levels (not computed by TDDFT) should be influenced by the presence of the cation. Following the analogy with the previous case, the next temporal window [50; 2000 ps] concerns the decay to the ground state, $T_1(ILCT) \rightarrow S_0$. The characteristic lifetime of 926 ps is also multiplied by a factor of 3 rationalized in terms of the energy gap law for the ISC processes (Fig. 6).35

3. Conclusion

The results of our investigation on the photophysical properties of an iridium(III) polypyridyl heteroleptic complex, namely an azacrown-iridium(III) complex, have elucidated several crucial aspects of its response to photoexcitation and interactions with Ca²⁺ ions. The integration of TD-DFT calculations and time-

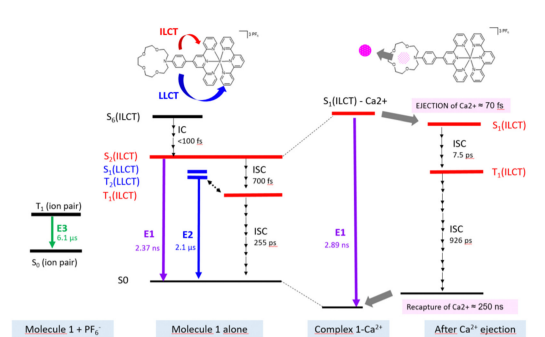


Fig. 6 Proposed Jablonski diagram for molecule 1 and complex 1-Ca²⁺ according to our experiment and theoretical calculations

resolved spectroscopies has allowed us to characterize the intricate dynamics governing this complex. Our results indicate that the introduction of Ca²⁺ significantly alters the absorption and emission profiles, with the complex displaying a notable shift from 494 nm to 375 nm upon metal binding, consistent with the calculated transitions. In addition, the theoretical results as a whole are very consistent with the spectra obtained from experimental studies, thus demonstrating the relevance of this method in the selection of a molecule to be synthesized. Furthermore, we documented the photo-ejection of Ca²⁺, occurring at a remarkable 70 fs, and its subsequent recapture at 250 ns, and these phenomena aligned with existing literature on ultrafast cation dynamics in comparable systems. These results are among the few to show both photo-ejection and recapture of the cation and feature a unique 10 million time factor between the two processes, allowing temporal isolation of the two phenomena and potential applications in fields where ion control, both temporal and spatial, is critical.

Comparisons with earlier studies show that this complex exhibits typical behaviors of polypyridyl iridium(III) complexes, such as strong intra-ligand charge transfer (ILCT) and ligandto-ligand charge transfer electronic transitions as well as sensitivity to environmental changes through, for example, ion-pair formation. These findings are consistent with established knowledge of the photophysics of iridium complexes, particularly regarding the influence of ligand environments on charge transfer states and emission properties. The distinction in our results lies in the detailed kinetic analysis and the exploration of interaction dynamics, which have not been thoroughly examined in prior studies. This filled a significant gap, particularly in understanding the excited states' landscape and their role in facilitating efficient ion transfer processes. The competitive binding and photo-release capabilities of the azacrown

moiety in our system are also corroborated by previous reports on similar aza-crown compounds, cementing our understanding of their potential in sensor, biolabeling or optoelectronic applications.

Looking ahead, the versatile photophysical properties of these complexes present exciting opportunities for the development of advanced biosensors, molecular machines and stimuli-responsive materials. The precise control over cation binding and release demonstrated here could be leveraged to design sophisticated ion sensors and delivery systems with potential for applications in biomedical fields. Additionally, further exploration into the fine-tuning of ligand environments and photophysical responses assisted by preliminary TD-DFT calculations may yield new insights into the manipulation of charge transfer processes for various technological applications. The promising results showcased in this study lay the groundwork for future developments in the use of iridium(III) complexes for excited state cation transfer for innovative applications in dynamic and responsive material systems.

4. Experimental

4.1. Synthesis

Materials and methods. All starting materials and solvents required for the synthesis were purchased from Sigma-Aldrich, TCI or Carlo Erba, and used as supplied without any further purification. ¹H NMR and ¹³C NMR spectra were recorded on a JEOL-FT RMN 400 MHz. The deuterated solvents used are specified for each spectrum and have been produced by Euriso-top[®]. The chemical shift (d) is expressed in parts per million (ppm) using tetramethylsilane (TMS) as the reference, and the coupling constant (J) is given in Hertz (Hz). For

spectral description, these abbreviations were used: singlet (s), doublet (d), triplet (t), and multiplet (m). All NMR spectra are available in the ESI† (Fig. S6-S10).

HRMS have been obtained by the mass spectrometry service from the Institut de Chimie des Substances Naturelles (ICSN), at the research center from Gif sur Yvette.

Synthesis of compound 4. N-(4-Formylphenyl)aza-15-crown-5 2 (2 g, 6.18 mmol) and 2-acetyl pyridine 3 (1.5 g, 12.36 mmol) were dissolved in 70 mL of ethanol and stirred at room temperature under an argon atmosphere for 20 min, and 1.25 g of KOH and 40 mL of aq NH₃ solution were added to the mixture and left under stirring for 18 H. The obtained pale precipitate was filtered off and rinsed with acetone to afford 2.3 g of the desired product in 70% yield. ¹H NMR (CDCl₃, 400 MHz,) δ /ppm: 8.73 (d, 2H, J 4.6 Hz), 8.68 (s, 2H), 8.66 (d, 2H, J 7.8 Hz), 7.8-7.90 (m, 4H), 7.34-7.26 (m, 2H), 6.7 (d, 2H, J 7.8 Hz), 3.7 (m, 20 H) ¹³C NMR (CDCl₃, 100 MHz) 156.8, 155.8, 149.9, 149.3, 149.0, 148.50, 137.0, 136.7, 128.4, 128.3, 125.1, 123.9, 123.5, 121.6, 121.5, 121.3, 117.6, 117.3, 111.8, 111.6, 71.4, 70.4, 70.3, 70.2, 68.6, 52.7, HRMS (TOF MS ESI) m/z calculated for [M+H]⁺ C₃₁ H₃₅ N₄ O₄: 527.2663, found 527.2666.

Synthesis of compound 5. 4 (66 mg, 0.28 mmol) and IrCl₃ (104 mg, 0.28 mmol) were heated at 160 °C in degassed ethylene glycol (5 mL), under argon and in the dark, for 4 hours, during which a red precipitate formed. It was filtered off after cooling to room temperature and was washed with EtOH, H₂O, and Et₂O to give pure 5 in a 40% yield (60 mg) as an orange-red solid. 1H NMR (400 MHz, DMSO-d₆,80 $^{\circ}$ C): δ /ppm: 9.21 (d, 2H, J 5.4 Hz); 8.88 (s, 2H); 8.78 (m, 2H); 8.21 (m, 2H); 8.17 (m, 2H); 7.89 (m, 2H) 6.88 (d, 2H, J 5.4 Hz); 3.8-3.5 (m, 20H). ¹³C NMR (DMSO-d₆, 80 °C, 100 MHz) 160.2, 157.4, 153.5, 150. 7, 140.4, 129.8, 128.4, 1253, 119.0, 112.4, 71.0, 70.3, 69.9, 68.7, 52.4 HRMS (MALDI TOF) m/z calculated for $[M+H]^{\dagger}$ C₃₁H₃₄Cl₃IrN₄O₄ 824.12690 found 824.12410.

Synthesis of compound 1. 5 (100 mg, 0.12 mmol) and terpyridine 6 (28 mg, 0.12 mmol) were heated at 190 °C in degassed ethylene glycol (10 mL), under argon and in the dark, for 25 min, during which a red precipitate formed. 10 mL of a saturated NH₄PF₆ solution was added to the mixture, it was filtered off after cooling to room temperature and was washed with H_2O , EtOH/ H_2O (50/50 v/v) to give pure 1 in a 28% yield (48 mg) as a red solid. ¹H NMR (400 MHz, acetone-d6): δ /ppm: 9.41 (s, 2H); 9.22 (d, 2H, H6, I 5.4 Hz); 9.13 (d, 2H, H6, I 5.4 Hz); 8.94 (m, 3H); 8.85 (m, 2H); 8.38 (m, 4H); 8.28 (m, 2H); 8.13 (m, 2H), 7.65 (m, 4H), 7.08 (d, 2H, J 5.6 Hz); 3.8-3.5 (m, 20H). MS (MALDI TOF) m/z calculated for $C_{46}H_{45}F_{12}IrN_7O_4P_2$ 1242.25, found 1242.25, HRMS (MALDI TOF) calculated for [M] C₄₆H₄₅IrN₇O₄ 952.3162 found 952.3157.

4.2. Calculations

All the calculations were performed using the Gaussian 16 package³⁶ relying on (TD-)DFT to determine the ground-state (GS, S_0) and excited-state (ES, S_n and T_n) properties, respectively. We have used (i) the 6-311+G(d) basis set for the H, C, N, O, and Ca atoms, and (ii) the LANL2DZ³⁷ relativistic pseudopotential and the corresponding basis set for Ir. The geometry

optimizations and the calculation of the excited state properties have been carried out with the PBE038 hybrid exchange-correlation functional. Previous works have demonstrated the performances of global hybrid functionals³⁹ and, more specifically, PBE0, to investigate the nature, dynamics, and reactivity of the (charge transfer) excited states of iridium(III) complexes. 40,41

We calculated and analyzed frequency vibrations to guarantee that we obtained true minima on the potential energy surfaces. For all TD-DFT calculations, 25 excited states are considered. The "stick spectra" are convoluted with a Gaussian function (full width at half maximum FWHM = 0.25 eV). The computed vertical excitation wavelengths λ with the highest oscillator strength f are compared to the experimental maximum absorption wavelength. One should note that these values do not necessarily coincide, and a more refined computational model should consider the vibronic structure of the electronic absorption spectra.

All calculations were performed by considering the polarizable continuum model42 (IEF-PCM) in the equilibrium limit using the linear-response (LR)PCM scheme to quantify the impact of the acetonitrile solvent (ACN).

The Breit-Pauli spin-orbit matrix elements (SOCMEs) are calculated using PySOC, 43 a program interface based on MolSOC⁴⁴ and implemented within the MOMAP package.⁴⁵ At that stage, the level of calculation is exactly the same as the one considered in the TD-DFT calculations (solvent included).

To analyze the CT character of the different excited states, we have used three different descriptors. First, the variation of electronic density $\Delta \rho(\vec{r})$ is calculated on a grid of points using the cubegen utility provided by the Gaussian package:

$$\Delta \rho(\vec{r}) = \rho^{ES}(\vec{r}) - \rho^{GS}(\vec{r})$$

where $\rho^{GS}(\vec{r})$ and $\rho^{ES}(\vec{r})$ are the electron density of the ground state and the considered excited states, respectively.

Secondly, the density derived indexes namely the D^{CT} index and the norm of the dipole moment corresponding to the CT observed upon excitation μ_{CT} are computed using the DctVia-Cube program.46

4.3. Spectroscopies

The stationary absorption spectra were measured with a CARY 100bio absorption spectrometer while emission spectra were recorded with a Fluoromax3 spectrometer. The TCSPC were measured with FT200 apparatus (PicoQuant) using femtosecond laser pulses at 310 nm.

Our nanosecond transient absorption spectroscopy setup has been described elsewhere. 47 Briefly, a third harmonic pulse (355 nm) of a Nd:YAG laser with ca. 1 mJ output energy and 5 ns pulse width was used as the excitation light and a pulsed Xe lamp was utilized as the probe light. Transient absorption spectra were obtained from the transient absorption decays recorded at various wavelengths by sampling the absorbance changes for different given delay-times. Note that a "luminescent" mode is available by switching off the Xe lamp.

Femtosecond time-resolved experiments were performed using a pump-probe spectrometer⁴⁸ based on a Ti:sapphire

laser system (Coherent Mira-900-D oscillator and Libra-S regenerative amplifier) delivering 800 nm, 1 mJ and 90 fs pulses with a repetition rate of 1 kHz. The pump pulses adjusted and tuned at 320 nm were generated by frequency quadrupling the output of a Quantronix Palitra OPA pumped at 800 nm and the energy at the sample was about $2 \mu \text{J}$ (0.2 mJ cm⁻²). Note that, we chose to excite the molecule far from the maximum of absorption localized near 270 nm to avoid a short-time 2-photon absorption signal coming from the solvent. The probe pulses were obtained by focusing 1 µJ, 800 nm pulses into a 1 mm thick CaF₂ plate to generate a white light continuum. The pumpprobe polarization configuration was set at a magic angle (54.7°). Transient absorbance was obtained by comparing the signal and reference spectra with and without pump pulses for different delay-times. The delay-time between the pump and probe was varied up to 0.5 ns using an optical delay line. Sample solutions ($\sim 10^{-3}$ - 10^{-5} mol dm⁻³, OD = 1 at the pump wavelength) were circulated in a cell equipped with a 200 µm thick CaF2 entrance window and a 2 mm optical path. All the transient spectra presented in this paper are GVD corrected according to the typical extrapolation method. 49 The characteristic times deduced from kinetics were obtained using a global fitting approach, and the data being fitted with the results of a multiexponential function convolved with a Gaussian pulse mimicking the pump-probe cross-correlation function (FWHM \sim 180 fs).

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts of interest to declare.

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