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Dual luminescence and infrared circularly polarized luminescence up to 900 nm with platinum complexes bearing a helical donor–acceptor ligand†

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Chiral molecular materials able to emit circularly polarized luminescence (CPL) have attracted considerable interest in the last few decades, due to the potential of CP-light in a wide range of applications. While CP luminescent molecules with blue, green, and yellow emissions are now well-reported, NIR CPL from organic and organometallic compounds lags behind due to the dual challenge of promoting radiative deexcitation of the excited state in this low energy region while assuring a significant magnetic dipole transition moment, a prerequisite for generating CPL. Based on a versatile axially chiral arylisoquinoline ligand, we report the synthesis and chiroptical properties of chiral donor–acceptor platinum(II) complexes displaying CPL that extends up to almost 900 nm. Interestingly, these emitters show both fluorescence and phosphorescence emissions in solution, with intensities depending on the charge-transfer character of the organic ligand. Experimental and theoretical investigations show that this feature strongly impacts the intersystem crossing event between the singlet and triplet excited states of these complexes and the related phosphorescence lifetime. The effect is less important regarding the CPL, and most complexes show luminescence dissymmetry factors with values up to ca. 2×10^{-3} around 800 nm.

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Introduction

Molecular compounds able to emit in the near infrared region (NIR) of the electromagnetic spectrum attract intense interest due to their potential applications, ranging from optoelectronics to bioimaging. During the last decade, the development of chiral emitters able to display circularly polarized luminescence (CPL) has motivated chemists working in this area to develop NIR CPL compounds. Indeed, CP-light displays potential advantages over

its non-polarized counterpart in the aforementioned applications, such as in organic light-emitting diodes (OLEDs), CP light detectors, as well as in chiral sensing and bioimaging.^{1–7}

Accordingly, the design of NIR CPL emitters has emerged with lanthanides and chromium complexes constituting the first compounds reaching high intensity of CPL thanks to their magnetically allowed and electrically forbidden transitions.^{1,8–12} In a complementary strategy, chiral molecular organic and organometallic complexes have also been investigated due to their readily tuneable optoelectronic properties, simple processability, and often higher photoluminescence quantum yields (PLQYs). The latter aspect, associated with the high molar extinction coefficient of the related electric dipole-allowed transitions, provide this class of chiral complexes with potentially high CPL brightness.^{13,14} These features make these dyes valuable candidates for CPL applications,^{15–22} with promising examples of blue, green, and yellow CPL emitters.²³ However, reaching the far red, and even the NIR remains a considerable challenge for both organic and organometallic derivatives,²⁴ since the competition with nonradiative deactivation pathways of the excited state is the first aspect to overcome, commonly underlined by the so-called ‘energy gap law’. In addition, the needed electronic and structural factors for reaching this low-energy emission, *i.e.*, a delocalized

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➤ Previously reported molecular NIR CPL emitters with CPL beyond 750 nm

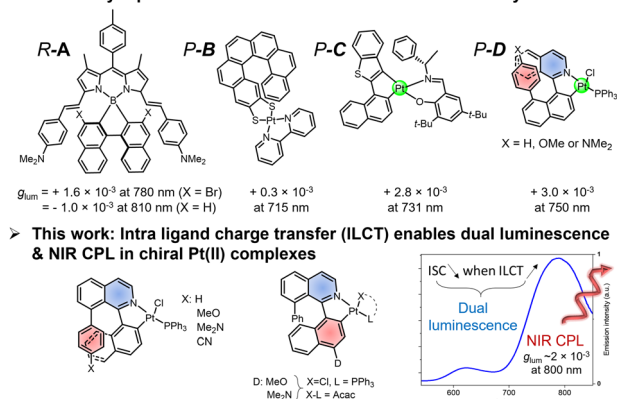


Fig. 1 Top: Chemical structures of the molecular CPL emitters displaying CPL maxima beyond 750 nm in solution. Bottom: Chemical structures of the investigated helical donor–acceptor platinum(II) complexes in this report.

π -conjugated system with often a rigid and planar molecular backbone, are usually hardly compatible with a significant magnetic (m) dipole transition moment, a prerequisite for chiroptical properties such as CPL.

While elegant designs of helical organic and organometallic systems have shown CPL in the red and far-red region,^{25–34} only a few examples have currently achieved a CPL over 750 nm (Fig. 1).^{35,36} For these, the obtained luminescence dissymmetry factor, g_{lum} , remains rather modest with values rarely larger than 3×10^{-3} . The investigated molecular designs are either based on an extended π -conjugated chromophoric unit based on BODIPY (A, Fig. 1) or rely on platinum complexes incorporating chiral ligands such as carbohelicene.^{36–43} Indeed, Pt complexes have been found to be promising to generate NIR phosphorescence, with cyclometalated phosphorescent d⁸ metal complexes extensively studied for the development of high-performance far red and NIR-OLEDs.^{36,42,44}

Following this line of research, we recently reported a new family of chiral donor–acceptor platinum(II) complexes involving an organic helical ligand possessing a significant intra-molecular charge-transfer (ICT) character.⁴⁵ This feature affords a dual luminescence behaviour with a fluorescence emission arising from the ligand and a NIR phosphorescence that can be modulated by the ICT character of the helical ligand. Interestingly, these complexes exhibit NIR CP phosphorescence with a g_{lum} of 3×10^{-3} , a significant value in this low energy region. While these results highlight interesting aspects regarding the design of NIR CPL emitters, such “helical donor–acceptor platinum complex design” remains limited to a few examples only, hampering a clear rationalization of the dual luminescence process and its implication on the possibility to reach NIR CPL. Accordingly, we decided to synthesize a complete family of chiral platinum complexes to explore the structural and electronic impact of the electron donating groups on the dual fluorescence/phosphorescence emissions as well as on the associated CPL signals.

We report here on the synthesis, photophysical, and chiroptical properties of a new family of NIR CPL helical

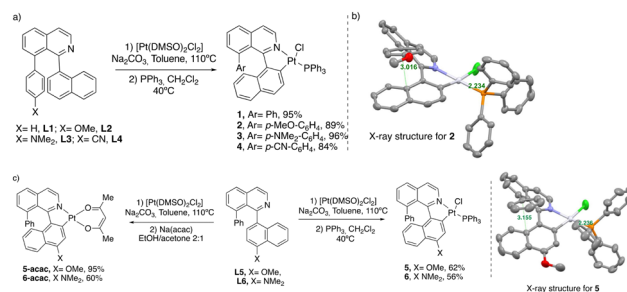
donor–acceptor platinum complexes based on axially chiral arylisoquinoline ligands. Playing with the functionalization of either the isoquinoline or the naphthyl unit of these molecular systems strongly impacts both the photophysical and chiroptical properties, by notably promoting dual luminescence emission and NIR CPL extending up to 900 nm. Experimental and theoretical investigations revealed that this effect is related to a subtle interplay between the ILCT transition localized on the ligand and the MLCT one involving the metallic centre, with a dual emission process becoming more favoured when electron donor substituents are attached to the naphthyl core. Such an impact is however less pronounced on the chiroptical properties and NIR CPL can be detected for most of the complexes with a promising g_{lum} of up to $ca. 2 \times 10^{-3}$ around 800 nm.

Results and discussion

Synthesis and characterization

Based on our previous results⁴⁵ and with the idea of developing a new family of CPL active Pt-based complexes with NIR-luminescence, we decided to modify the structure of the N,C-ligands by introducing aryl moieties with different electron richness to control the balance between the ILCT and MLCT transitions and tune the resulting emission process. Initially, the synthesis of the [Pt(N,C)(PPh₃)Cl] complexes **1–4** containing an aryl substituent at the 8-position of isoquinoline was carried out (Scheme 1a). Starting from the free ligands **L1–L2** (see the ESI†), a cyclometallation step followed by a ligand replacement (DMSO \rightarrow PPh₃), affords the desired complexes **1–4** in excellent yields.^{40,46–48} These complexes were completely characterized by NMR and HRMS (see the ESI†) and were found to be stable to oxygen and moisture, being purified by flash chromatography on silica gel. The X-ray structure for (\pm)-**2** (Scheme 1b) clearly shows the expected helicoidal disposition of the ligand (aryl...aryl distance of 3.016 Å) and the *trans* configuration between the phosphine moiety and the nitrogen atom (Pt–P distance of 2.234 Å) which fits with the solution-phase observation ($J_{Pt,P} = 4454$ Hz).

Along the same lines, we decided to analyse the effect of changing the naphthalene ring on complex **1** by more electron rich substituents by introducing an electron donating groups like –OMe and –NMe₂ at the 4-position of the naphthalene



Scheme 1 (a) Synthesis of the Pt-complexes **1–4**. (b) X-ray structure of **2**; (c) synthesis of the Pt-complexes **5–6** and their corresponding acetylacetonate (acac), with the corresponding X-ray structure of (\pm)-**5**.



fragment (Scheme 1c). Following the same synthetic strategy, the corresponding complexes **5–6** were obtained in modest 56–62% yields as orange to purple solids. Finally, we decided to study the effect of replacing the 3D dimensional bulky triphenylphosphine ligand with a bidentate anionic ligand such as acac, *i.e.*, acetylacetonate, with the aim of increasing the planarity of the complexes, and therefore, to increase $\pi \cdots \pi$ and Pt \cdots Pt intermolecular interactions. Following a similar strategy, ligands **L5–L6** were used in the cyclometalation step to give the corresponding [Pt(N,C)(DMSO)Cl] intermediates which, after reaction with Na(acac), afforded the final **5-acac–6-acac** complexes in 60–95% yields as pink to red solids.

The racemic complexes were resolved using semi-preparative chiral HPLC separation to give the enantiopure compounds with ee values up to 99% (see the ESI† for details). Their absolute configuration was assigned by comparison of their ECD responses with those of previously reported complexes⁴⁷ and further confirmed by theoretical calculations (see later). All products were characterized by ¹H and ¹³C NMR spectroscopy and HRMS (see the ESI†).

Ground- and excited-state photophysical properties

In line with our recent study on Pt-based complexes with NIR-CPL,⁴⁵ we investigate here the structural and electronic

parameters influencing the photophysical and chiroptical properties of these new chiral compounds, namely: (1) the impact of the donor position on the π -helical ligand and the resulting chiroptical properties for both the ground and excited states and (2) the magnitude of the intra ligand charge transfer and its interaction with the metal to ligand one, influencing the dual emission process and the efficiency of the NIR phosphorescence emission (*vide infra*). As depicted in Fig. 2, the absorption spectra of complexes **1–4** display similar profiles with two intense bands around 350 and 390 nm ($\epsilon \sim 10 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$) that are assigned to ligand $\pi \rightarrow \pi^*$ transitions and a lowest energy broad signal between 420 and 530 nm ($\epsilon \sim 4\text{--}7 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$) that involves both the metal and the ligand, *i.e.*, that can be characterized as a mixture of metal-to-ligand (ML) and intra-ligand (IL) charge-transfer excitations (for comparison, the UV-vis spectra of the corresponding organic ligands are depicted in Fig. S1 (ESI†)). This assignment is supported by theoretical calculations, see the electron density difference (EDD) plots in Fig. 2 and Table S1 (ESI†).

Despite the difference in the donating/accepting electron ability of the functional groups at the 8-position of the isoquinoline fragment in these systems, the lowest absorption band shows only a minor shift, indicating a small effect of the substitution on the excited-state energies. This is easily

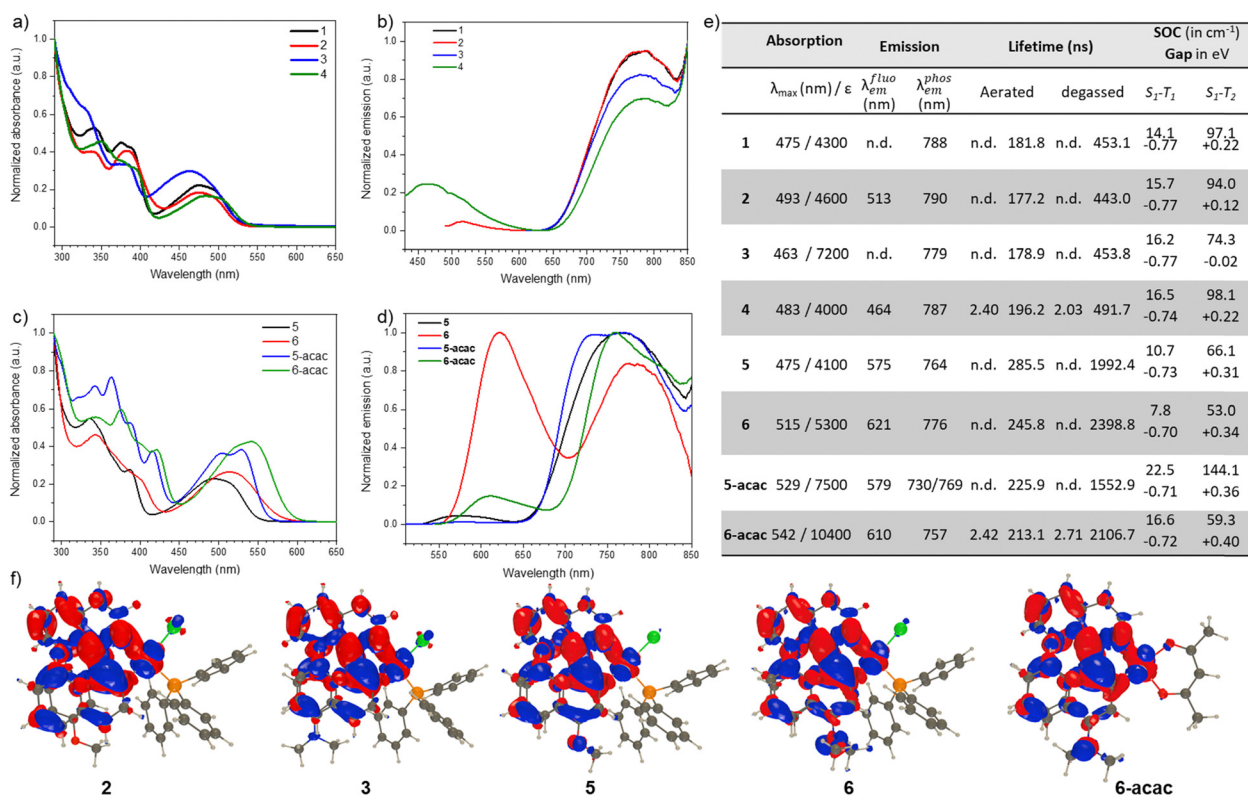


Fig. 2 (a) Normalized absorption and (b) emission spectra of complexes **1**, **2**, **3**, and **4** in toluene at 298 K; (c) normalized absorption and (d) emission spectra of complexes of complexes **5**, **5-acac**, **6**, and **6-acac** in toluene at 298 K; (e) photophysical data of the platinum complexes recorded in toluene at 298 K, under aerated and degassed conditions, with the corresponding lifetimes for the fluorescence and phosphorescence emissions (n.d.: not detected); S_1-T_1 and S_1-T_2 gaps (in eV) computed on the optimal S_1 geometries together with the associated SOC (in cm⁻¹); (f) electron density difference plots for the lowest absorption of complexes **2**, **3**, **5**, **6** and **6-acac**. The blue (red) lobes correspond to region of decrease (increase) of density upon excitation. Contour threshold: 0.001 au (see the ESI† for details).

understandable by the EDD plots that all show almost no involvement of the substituent at 8-position in the lowest excited state. This can look surprising, especially for **3** that bears a potent NMe₂ group. However, the expected IL-CT in which this group acts as a strong donor is actually the second state (Fig. 2), and it is slightly blue-shifted as compared to the S₀-S₁ transition (Table S1, ESI†).

Consistent with the above, the same is found in the emission spectra, which show a broad phosphorescence emission at 780 nm for all complexes, with similar lifetimes of *ca.* 500 ns and low photoluminescence quantum yields (PLQY, >1%, Fig. 2). Under aerated conditions, a second emission band at higher energy is additionally distinguishable for complexes **2** and **4**, albeit with very low intensity (Fig. 2). Reminiscent of what we observed in our previous study,⁴⁵ we attribute this behaviour to a dual emission process. Further experimental evidence of both fluorescence and phosphorescence emissions has been obtained by recording the luminescence response of the complexes under air atmosphere, which shows only a decrease in the NIR signal due to the quenching of the emitting triplet state by molecular oxygen (Fig. S3–S5, ESI†). The corresponding nanosecond lifetime of this luminescence signal confirms the fluorescence nature of this emission, which has also been reported for platinum complexes exhibiting a weak electronic interaction between the d orbitals of the metal and the π -orbitals of the organic ligand.^{49–52} Furthermore, theory predicts phosphorescence in the NIR spectral region whereas fluorescence should appear at much shorter wavelengths (Table S1, ESI†), confirming the experimental assignment. In Fig. 2 and Table S2 (ESI†), we report computed data for the ISC process. The two most interesting molecules are likely **2** and **3**. In both, one notices large S₁-T₁ gaps of -0.77 eV associated with SOC of *ca.* 16 cm⁻¹, that is, large gaps and moderate SOC for Pt-bearing systems. In **2**, the S₁-T₂ gap is +0.12 eV, but this small uphill process is likely available, especially since the associated SOC is 94 cm⁻¹, *i.e.*, it is likely that some ISC takes place towards the second triplet state due to thermal effects. In **3**, this state is almost perfectly aligned with S₁ (-0.02 eV gap) and associated with a large SOC of 74 cm⁻¹ as well, indicating a very efficient ISC, consistent with the absence of fluorescence in the experiment.

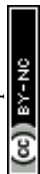
In contrast, modifying the position of the electron donor group on the naphthyl unit, as in complexes **5** and **6**, induces a noticeable change for both the absorption and emission properties. Indeed, while the absorption spectra of these complexes show a similar intensity as their isoquinoline substituted analogues, they undergo a redshift of *ca.* 50 nm for the lowest energy band with a noticeable difference between the methoxy and dimethyl amino substituents, indicating a stronger impact of these functional groups on the excited-state properties (Fig. 2). The donating groups of **5** and **6** indeed play an active role in the EDD plots (Fig. 2 and Fig. S34, S35, ESI†) contrasting with dyes **2** and **3**. Under aerated conditions, the emission spectra of complexes **5** and **6** shows a phosphorescence band at *ca.* 770 nm with the same profile as that for complexes **2** and **3**, with also a higher energy emission band centred at 575 and 620 nm for **5** and **6**, respectively, much higher in intensity than

the ones hardly observed in the case of complexes **2** and **3**. Interestingly, increasing the electric dipole moment of the substituent in complex **6** with the dimethyl amino group impacts significantly the intensity of the dual emission process since the fluorescence/phosphorescence ratio becomes higher than for complex **5**; the fluorescence of **6** remaining distinguishable even when the measurement is performed under degassed conditions, which is not the case for complex **5** (Fig. S3–S5, ESI†). This suggests that the radiative deexcitation of the singlet excited-state, in competition with ISC, is favoured when it acquired higher ILCT character. This aspect can be also evidenced by the lifetimes of the fluorescence emission which is longer for complex **6** in comparison with complex **5**, respectively (Fig. 2). Additional insights into these experimental outcomes are provided by theory (Table S1, ESI†), which first predicts the fluorescence (phosphorescence) at 523 and 544 (780) nm for **5** and **6**, respectively, confirming the experimental assignments, although the theoretical emission of **6** is too blue shifted. Second, for **5** the S₁-T₁ and S₁-T₂ gaps (on the optimal S₁ geometry) are, respectively, -0.73 eV and +0.31 eV. Even given the error of theory, it therefore suggests that ISC to the second triplet state is difficultly achievable, and only the S₁-T₁ channel is open. For that transition, we compute a rather small SOC matrix element of 10.7 cm⁻¹. In **6**, the gaps are similar, -0.70 and +0.34 eV, but the SOC is significantly smaller at 7.8 cm⁻¹. These values are qualitatively consistent with the clear observation of fluorescence in **5**–**6**, and its absence in **3**, which present notably different ISC features (*vide supra*).

Increasing the electron donor nature of the ancillary ligand in complexes **5-acac** and **6-acac**, by replacing the triphenyl phosphine and chloride ones with an acac unit, redshifts their absorption spectra which suggest a slight increase of the role of the metal in the lowest energy band. A small change can indeed be seen in Fig. 2, although the global topology of the transition is preserved. The acac ligand also impacts the phosphorescence emission with a hypsochromic shift when compared to their PPh₃/Cl analogue complexes ($\Delta\lambda_{em}$ = 39 and 19 nm for **5-acac** and **6-acac**, respectively, Fig. 2). This is also associated with a longer lifetime of the emission, also in line with more pronounced MLCT character of the triplet excited-state.

Ground- and excited-state chiroptical properties

The impact of the substituent position and the type of ancillary ligands on the chiroptical properties was also investigated both in the ground and excited states of the complexes using ECD and CPL characterization techniques. As depicted in Fig. 3, the ECD spectra of all compounds display obvious similar responses at the high energy region with a bisignate signal at *ca.* 270 nm, which is reminiscent of the binaphthyl ECD-type signature.⁵³ This characteristic signal is also found in previously reported ECD spectra of related platinum complexes,²² as well as in the simulated ECD spectra of complexes **1**, **3**, **5** and **6** (Fig. S36–S39, ESI†) and allow the absolute configuration of these new derivatives to be assigned: the ECD spectrum with the positive \rightarrow negative couplet corresponds to the *M*-enantiomers while the mirror-image signal is for the *P*-enantiomers.



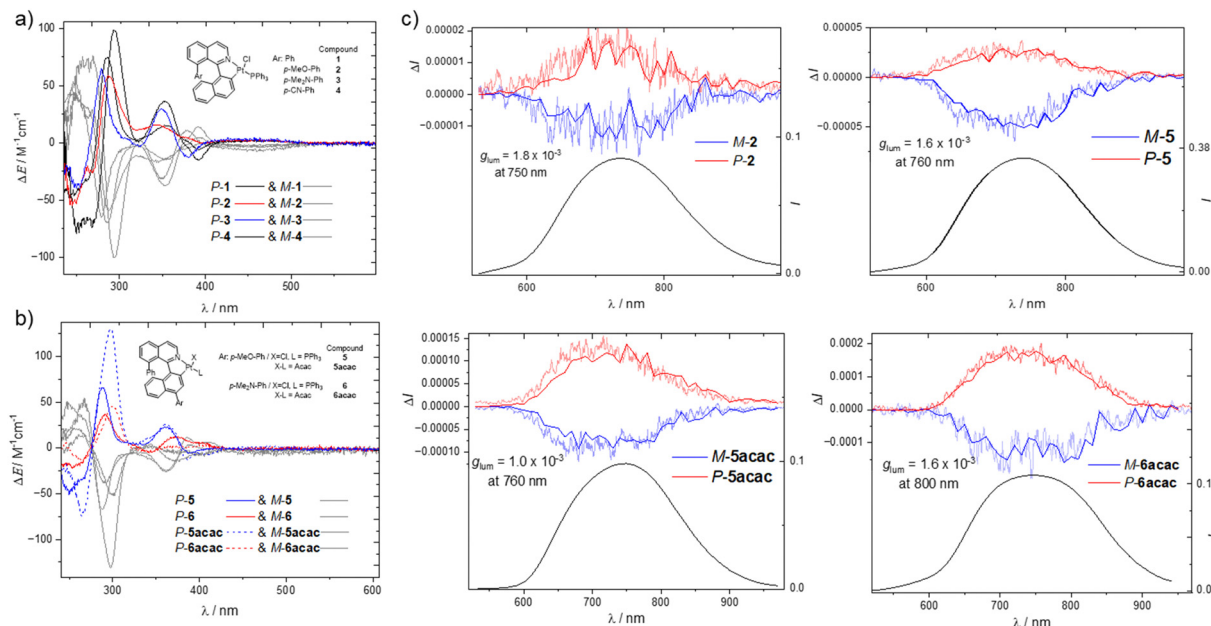


Fig. 3 (a) ECD of complexes **1**, **2**, **3** and **4**, and (b) of complexes **5**, **6**, **5acac** and **6acac** (right) recorded in dichloromethane at 298 K ($[c] \sim 10^{-5}$ M); (c) CPL of complexes **2** and **5** (top), and **5acac** and **6acac** (bottom) recorded in toluene solution at 298 K ($[c] \sim 10^{-5}$ M). For CPL measurements, the faded spectra are the ones directly measured, and the spectra in solid lines are corrected using a calibration curve for the detector.

In addition to this couplet, complexes **1–3** display very similar ECD responses with a set of three signals at 355, 392 and a broader one between 400 and 500 nm, respectively, positive, negative and positive. The corresponding intensity of these bands decreases when going to lower energy, ranging from up to 25 to $2 \text{ M}^{-1} \text{ cm}^{-1}$, resulting in similar absorption dissymmetry factors, g_{abs} , of $\text{ca. } 6.0 \times 10^{-4}$ at 460 nm.

Changing the position of the donor group on the naphthalene unit in complexes **5** and **6** does not significantly affect the couplet signal at 270 nm, but clearly impacts the intensity of the lower energy bands in a similar way for both methoxy and dimethyl amino groups, which evidences the role of the donor group in these optical transitions. For instance, the positive signal at 340 nm for compound **2** almost disappears in the case of compound **5** concomitantly to an intensity increase of the signal at 370 nm and an almost complete vanishing of the lowest energy broad band, as illustrated by the obtained zero value of g_{abs} at 460 nm. Finally, the replacement of the triphenylphosphine and chloro ligands by the acac one in complexes **5-acac** and **6-acac** does not tune the overall shape of the obtained ECD spectra but rather impacts the intensity of the involved bands, in a similar manner for both compounds. As an illustration, **5-acac** shows a strong increase in the band at 300 nm (from $\text{ca. } 60$ to $110 \text{ M}^{-1} \text{ cm}^{-1}$ when comparing complexes **5** and **5-acac**, respectively), together with a less pronounced but clearly distinguishable increase of the low energy signal, resulting in g_{abs} of $\text{ca. } 5.0 \times 10^{-4}$ at 460 nm.

In line with the NIR luminescence obtained for these new Pt complexes, all of these complexes display CP phosphorescence with mirror-image responses that extend lower than 900 nm, as depicted in Fig. 3 for complexes **2**, **5**, **5-acac**, and **6-acac**. The chiral platinum complexes exhibited almost constant g_{lum}

values between 650 and 900 nm with values of $\text{ca. } 1.0\text{--}2.0 \times 10^{-3}$ (Fig. S32, ESI[†]), showing a limited influence of both the electron donor group and the ancillary ligands on the intensity of the CPL. Among the complexes, only **6** shows a different behavior than the other systems, with a weak dual CPL at 570 and 760 nm, opposite in sign (Fig. S33, ESI[†]), presumably linked to the significant dual emission observed for this compound.

Accordingly, a first CP fluorescence can be observed, arising mainly from ILCT transitions, followed by a lowest energy CP phosphorescence involving MLCT, and presumably explaining the sign inversion observed between these two responses. The introduction of the acac ligands in **5-acac** and **6-acac** does not significantly modulate the CPL intensity, and the overall CPL brightness (B_{CPL}) of these complexes falls in the range of $10^{-2} \text{ M}^{-1} \text{ cm}^{-1}$.^{13,14} These values are rather weak in the context of molecular CPL emitters,^{2,13,54} which is mainly attributed to the low PLQY of the complexes. However, these results represent an interesting example of NIR CPL, among the most redshifted reported to date for chiral luminophores based on organic and platinum complexes.

Conclusions

In conclusion, we extended the family of NIR CPL molecular emitters by designing innovative chiral platinum complexes displaying phosphorescence around 750–800 nm. Through molecular engineering around a common platinum arylisoquinolines core, we show that the presence of donor groups on both the organic ligand and the metallic ion strongly impacts the interplay between the ILCT and the MLCT transitions,



promoting a significant dual emission process involving a fluorescence signal between 500 and 700 nm. The later process appears firmly linked to the position of the electron donor substituents on the structure and is favoured when methoxy and amino-donor groups are on the naphthyl unit in complexes **5** and **6**. Such behavior was linked to singlet–triplet excited-state energy gaps and SOC elements, hence to the ISC, for which methoxy and amino donor groups tend to maintain large S_1 – T_1 gaps (> -0.77 eV) associated with rather small SOC (< 10 cm $^{-1}$). In comparison, when these donor groups are on the isoquinoline moiety or when an acac ligand is introduced on the metallic ion, the ISC efficiency increases, ultimately favouring phosphorescence emission. While these combined experimental and theoretical investigations reveal the crucial role played by the extended π -helical donor acceptor ligand on the unpolarized emission of these new complexes, these factors seem less crucial for the chiroptical properties. Indeed, this family of chiral molecular emitters afforded NIR CPL at wavelength lower than 900 nm, with dissymmetry factor up to *ca.* 2.0×10^{-3} , which is a significant value for this low energy region. We hope that this investigation may offer new opportunities to design innovative and efficient NIR CPL emitters.

Data availability

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

Conflicts of interest

There are no conflicts to declare.

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