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Triplet state CPL active helicene-dithiolene platinum bipyridine complexes

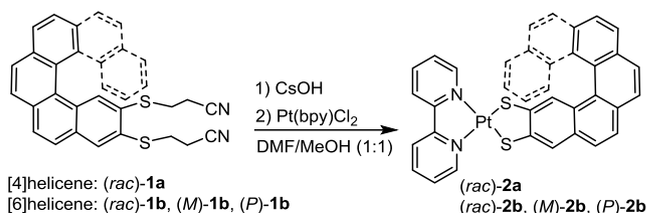
Thomas Biet,^a Thomas Cauchy,^a Qinchao Sun,^b Jie Ding,^b Andreas Hauser,^{*b} Patric Oulevey,^b Thomas Bürgi,^b Denis Jacquemin,^c Nicolas Vanthuyne,^d Jeanne Crassous^e and Narcis Avarvari^{*a}

Chiral metal dithiolene complexes represent a family of chiral precursors, which can give rise to molecular materials with properties resulting from the interplay of chirality with conductivity, magnetism, and photophysics. We describe herein the first examples of chiral metal diimine dithiolene complexes, by the use of a platinum(II) centre coordinated by 2,2'-bipyridine and helicene-dithiolene ligands. Straightforward synthesis of racemic and enantiopure complexes allows the preparation of luminescent Pt(bpy) [4] and [6]helicene compounds for which the solid-state structure was determined as well. TD-DFT calculations support the assignment of the low energy bands observed in the UV-vis absorption spectra as mixed metal-ligand-to-ligand charge transfer transitions and confirm that the emission band results from the T₁ excited state. Interestingly the enantiopure [6]helicene complexes show CPL activity at room temperature in acetonitrile solutions with anisotropy factors of 3×10⁻⁴.

Chiral metal dithiolene complexes represent an emerging family of molecular materials where chirality is expected to modulate properties such as conductivity, magnetism, luminescence, etc.¹ For example, differences in conductivity between diastereomeric pairs of anionic Ni(II) bis(dithiolene) complexes with chiral viologene type cations have been noticed,² while the first chiral single component conductors based on neutral Au(III) bis(dithiolene) complexes have been recently described.³ Besides, stable anionic, neutral or cationic species can be easily accessed in metal bis(dithiolene) complexes thanks to the “non-innocent” character of the dithiolene ligands, so that redox modulation of the chiroptical properties can be observed.⁴ Although square planar platinum diimine dithiolene complexes have been investigated over more than twenty years especially for their emission properties in solution,⁵⁻⁸ and more recently for photocatalytic water splitting,⁹ no chiral derivative of this heteroleptic family has been yet reported. For example, in Pt(II) 2,2'-bipyridine (bpy) arene-dithiolene complexes such as Pt(bpy)(tdt) (tdt = toluene-dithiolate) showing room temperature luminescence in solution arising from a MMLL'CT (mixed metal-ligand-to-ligand charge transfer) transition,^{5,6} chirality could be in principle introduced either on the diimine or the benzodithiolene fragments in order to influence the photophysical properties. One of the interests of such complexes relies on the possible observation of circularly polarized luminescence (CPL), which is the differential spontaneous emission of left and right-handed circularly polarized light.¹⁰ While chiral lanthanide complexes are generally the most intense CPL emitters,^{11,12} several examples of transition metal complexes have been reported as well.¹³⁻¹⁵ Among them those containing helicene based ligands

are particularly interesting,^{16,17} as helicenes^{18,19} and heterohelicenes²⁰ are well known non-planar conjugated molecules possessing strong chiroptical properties. Note that CPL active cationic dioxo, azaoxa and diaza [6]helicenes have been recently reported.²¹ This work is presenting out investigations on helical dithiolene platinum 2,2'-bipyridine complexes using the hitherto unknown helicene-dithiolate (hel-dt) ligands, by analogy with the achiral toluene-dithiolate (tdt) previously mentioned. We describe herein the synthesis and the structural characterization of Pt(bpy)([n]hel-dt) (n = 4, 6) complexes together with their chiroptical and photophysical properties supported by DFT calculations.

The racemic 2,3-dithiolate-[4] and [6]helicene ligands, generated *in situ* from the protected precursors (*rac*)-**1a** and (*rac*)-**1b** respectively, which have recently been used by some of us for the synthesis of TTF-helicenes,²² have been reacted with Pt(bpy)Cl₂ to generate the corresponding complexes. Thus, (*rac*)-Pt(bpy)([4]hel-dt) **2a** and (*rac*)-Pt(bpy)([6]hel-dt) **2b** have been isolated as dark purple crystalline solids after column chromatography (Scheme 1 and ESI).



Scheme 1 Synthesis of Pt(bpy)(hel-dt) complexes **2a-b**.

As the racemization barrier for [4]helicenes is generally very low,²³ the enantiopure forms have been prepared only for the [6]helicene dithiolene complexes **2b** starting from the precursors (*M*)-**1b** and (*P*)-**1b** separated by chiral HPLC (Fig. S1-S3, ESI). The racemic complexes **2a** and **2b** crystallize in the centrosymmetric space groups *C2/c* and *P-1* respectively, with both enantiomers (*M*) and (*P*) in the unit cell (Table S1, ESI). Worth noting are the helical curvatures (hc) defined by the dihedral angle between the terminal rings of the helicene skeleton amounting to 25.40° and 53.76° for (*rac*)-**2a** (Table S2, Fig. S4-S5, ESI) and (*rac*)-**2b** (Table S3, Fig. S6-S7, ESI), which are typical values for [4] and [6]helicenes, and the square planar coordination geometry of the platinum centres. DFT calculations in acetonitrile yield dihedral angles 28.06° and 41.68° for **2a** and **2b**, suggesting that the impact of crystal packing is stronger for the latter compound. The enantiomerically pure complex (*M*)-**2b** has been also analysed by single crystal X-ray diffraction thus allowing to confirm that it was obtained from the (–)-**1b** precursor. (*M*)-**2b** crystallized

in the orthorhombic system, non-centrosymmetric space group $P2_12_12_1$, with four independent molecules in the unit cell (Fig. 1).

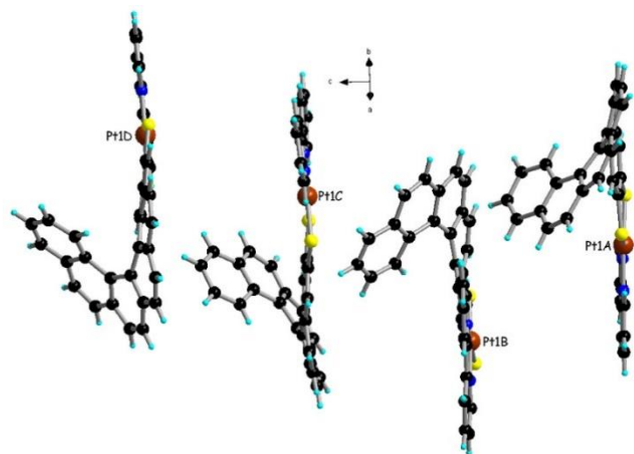


Fig. 1 The four independent molecules of complex in the solid state structure of (*M*)-**2b**.

The four molecules named as Pt1A – Pt1D slightly differ by the helical curvature values ranging from 58.58° (Pt1C) to 62.36° (Pt1D), while the Pt–S (2.24–2.25 Å) and Pt–N (2.05–2.07 Å) are in the normal range for such complexes (Table S4, ESI).²⁴ The packing of the molecules is very likely governed by the π - π stacking interactions occurring along the *c* direction (Fig. 1 and Fig. S8–S9, ESI).

The enantiomeric (*P*) and (*M*)-**2b** complexes represent the first chiral members of the platinum diimine dithiolenes family. As outlined above, Pt(diimine)(dithiolate) complexes are emissive in fluid or frozen solutions.^{5,6,25} We have therefore set out to measure in a first time the photophysical properties of the racemic **2a** and **2b**. In the low energy region the complex (*rac*)-**2a** has an absorption band from 450 to 650 nm with a maximum around 550 nm (18180 cm^{-1}) and an absorption coefficient $\epsilon \approx 6700\text{ M}^{-1}\text{cm}^{-1}$ (Fig. 2, top). For (*rac*)-**2b** the maximum of the corresponding band is around 562 nm and the absorption coefficient $\epsilon \approx 3640\text{ M}^{-1}\text{cm}^{-1}$ (Fig. 2, bottom). This low energy absorption band is typical for Pt(diimine)(dithiolate) complexes and has been assigned to a MMLL'CT transition, as the HOMO has metal/dithiolene character while the LUMO is developed over the unsaturated diimine ligand.⁶ The small redshift of the CT transition from **2a** to **2b** is caused by the slight change in the dithiolate ligand, with a more extended rigid π backbone in the latter, which makes the HOMO energy slightly higher by +0.03 eV in **2b** than in **2a**. Moreover, the absorption coefficient is smaller for **2b** likely because of the more distorted structure. TD-DFT reproduces the experimental trends with a vertical absorption at 556 nm ($f = 0.21$) for **2a** and 562 nm ($f = 0.19$) for **2b**. Both complexes are emissive in fluid solutions of CH_2Cl_2 when irradiated into the CT bands, showing low energy emission bands at 720 nm (**2a**) and 715 nm (**2b**) (Fig. 2 and Fig. S10, ESI, for (*rac*)-**2a** in CH_3CN). The more distorted structure of **2b** might be as well at the origin of the lower emission quantum yield for **2b** (0.15%) than for **2a** (0.19%) (Table S5, ESI). The perfect agreement between absorption and excitation spectra is proof that the luminescence indeed originates from the two compounds despite the low emission quantum yield. For **2b** a luminescence life-time of 124 ns was measured in

deoxygenated solution with pulsed excitation at 458 nm (decay curve shown in Figure S11, ESI). With the quantum efficiency of 0.15% this corresponds to a radiative lifetime of around 100 μs , indicating that the emission originates from the T1 state, as is generally the case for Pt(II) complexes.⁶

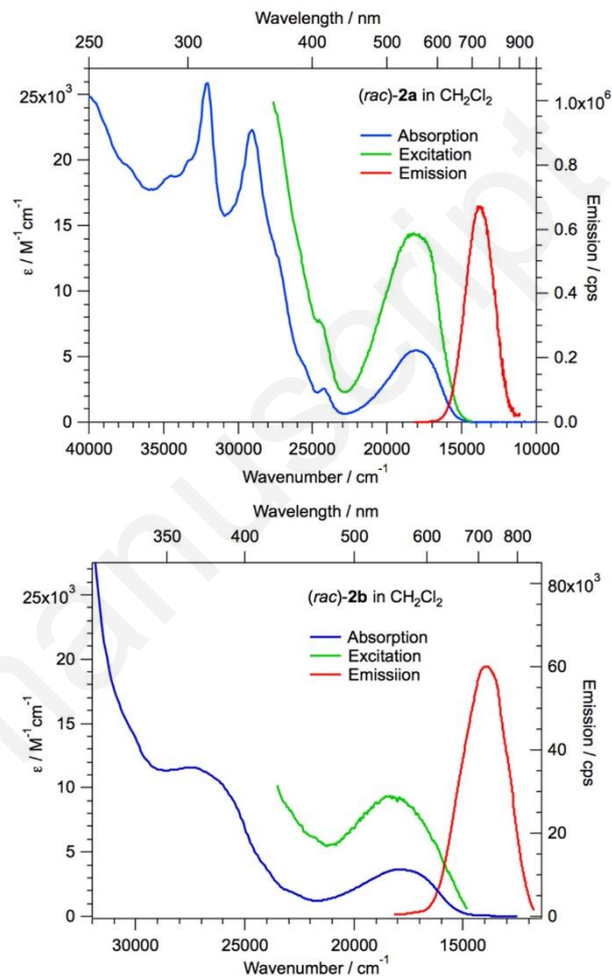


Fig. 2 Absorption, emission and excitation spectra of (*rac*)-**2a** (top) and (*rac*)-**2b** (bottom) in CH_2Cl_2 . Absorption spectra were measured at concentrations of $2.2 \times 10^{-5}\text{ M}$ for (*rac*)-**2a** and at $8 \times 10^{-5}\text{ M}$ for (*rac*)-**2b**. Emission spectra were measured using the same solutions degassed by nitrogen bubbling for 20 min with excitation at 525 nm for (*rac*)-**2a** and at 560 nm for (*rac*)-**2b**. Excitation spectra were measured at emission wavelengths of 720 nm for (*rac*)-**2a** and 715 nm for (*rac*)-**2b**.

To characterize the charge transfer and emission properties, DFT calculations have been performed on **2a** and **2b** (Fig. S12–S19, Tables S6–S8, ESI). For both compounds, optimized as (*M*) and (*P*) enantiomers respectively, the fully relaxed molecular geometries obtained by DFT are in line with those obtained by X-Ray diffraction. In Fig. 3, we represent the electron density difference (EDD) plots corresponding to the transition to the lowest singlet excited-state and the spin density of the lowest emissive triplet state. The EDD representation clearly shows that there is a strong CT from the dithiolene (donor, mostly in blue in Fig. 3) to the diimine (acceptor, mostly in red). The computed CT distance attains 4.0 Å in both **2a** and **2b**, which is a rather large value. Interestingly, one notices that the metal centre presents both positive and negative

density contributions, indicating that it also partially plays the role of an accepting unit. The implication of the metal in the EDD plots is also consistent with a possible intersystem crossing to the T1 state. The spin density of the lowest triplet state is unsurprisingly localized in exactly the same regions as the corresponding S1 state. The emission for the lowest triplet state was estimated in both the vertical and adiabatic approximations. In the former, DFT yields an emission at 802 nm and 815 nm for **2a** and **2b**, respectively, whereas in the latter that takes into account the difference of vibrational energies, we obtained 759 nm and 760 nm, for the two compounds. These latter values are in good correspondence with the experimental data, with an error smaller than 0.1 eV, confirming that the observed emission is indeed coming from T1. The fact that the vertical values indicate (incorrectly) a small redshift when going from **2a** to **2b**, whereas the adiabatic energies show essentially no shift, hints that the T1 geometrical relaxation is smaller in the latter compound than in the former.

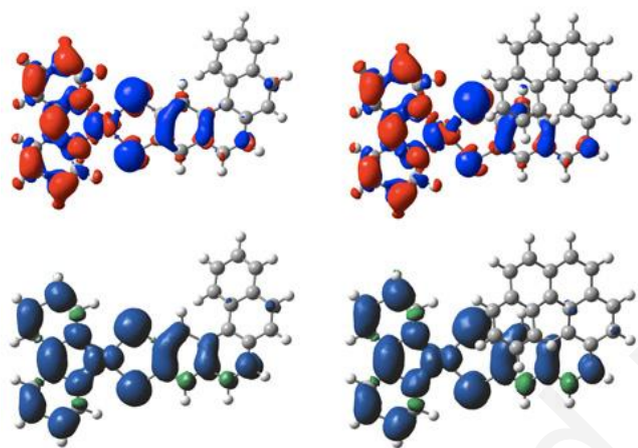


Fig. 3 Top: Density difference plots between the lowest S₁ and the S₀ state as determined by the TD-DFT on the optimal ground-state geometry. The blue (red) regions indicate regions of loss (gain) of density upon transition. Bottom: spin-density difference plots between the T₁ and S₀ states considering the T₁ state in its optimal geometry. (*M*)-**2a** and (*P*)-**2b** are displayed on the left and right hand-side, respectively. See the SI for computational details.

Investigation of the photophysical properties of both enantiomers of **2b** has been performed in acetonitrile solutions. Absorption spectra are shown in the top panel of Fig. 4, together with the emission spectrum of (*M*)-**2b**. Compared to the spectra in CH₂Cl₂, the maximum of the absorption is shifted to 530 nm, that is, by 30 nm to lower wavelength. The emission maximum in acetonitrile appears at 720 nm, that is, shifted to higher wavelength by 5 nm. The CD spectra of (*P*) and (*M*)-**2b** in acetonitrile, image mirror of each other, are shown in the bottom panel of Fig. 4. For the CD spectra, TD-DFT indeed yields a weakly positive contribution to the rotary strength for the transition to the lowest excited-state of (*P*)-**2b** and a weakly negative contribution to that strength for the lowest excited-state of (*M*)-**2b**, which is consistent with the experimental findings.

CPL, representing the differential emission between left and right circularly polarized light and characterized by the anisotropy factor $g_{em} = 2(I_L - I_R)/(I_L + I_R)$ at the maximum of the emission band, has been measured for solutions of (*M*) and (*P*)-**2b** in acetonitrile at room temperature (Fig. 4, bottom). As hypothesized through the introduction of the helicene backbone

in the dithiolenic ligand, the enantiomers of **2b** show CPL activity with an anisotropy factor of $\pm 3 \times 10^{-4}$, which is a typical value for organic, organometallic and coordination complexes in solution excepting the lanthanides.^{11,12} It should be mentioned however that this value of CPL anisotropy is for a compound with a luminescence quantum efficiency of only 0.15%, and which represents the first CPL active metal dithiolenic complex.

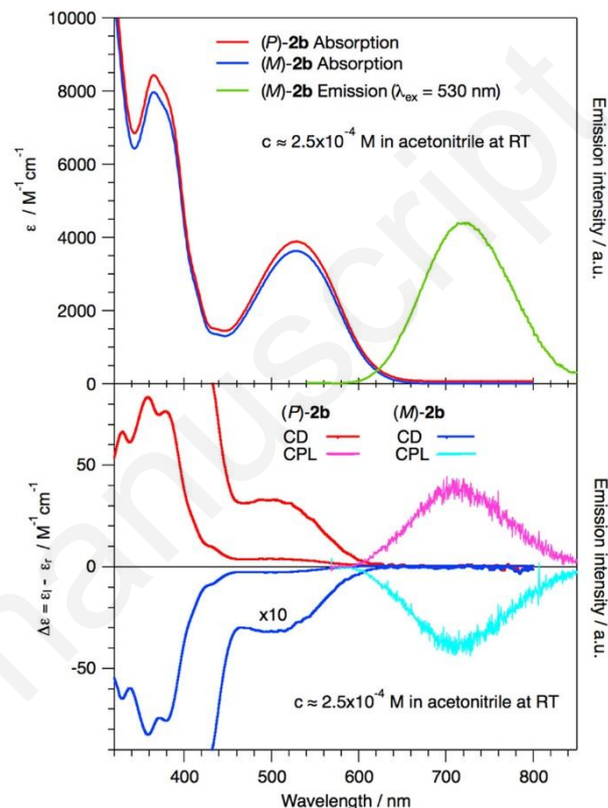


Fig. 4 Absorption spectra of (*P*)-**2b** and (*M*)-**2b** and emission spectrum of (*M*)-**2b** in acetonitrile (2.5×10^{-4} M) at $T = 298$ K, $\lambda_{ex} = 532$ nm (top); CD and CPL spectra of (*P*)-**2b** and (*M*)-**2b** (bottom).

In summary, the first helical Pt(diimine)(dithiolenic) complexes have been prepared through the introduction of [4] and [6]helicene backbones in the structures of the dithiolenic ligand. The solid state structures of the racemic complexes show the presence of both (*P*) and (*M*) enantiomers, with helical curvatures typical of [4] and [6]helicenes and packings controlled by π interactions. Enantiopure [6]helicene complexes have been prepared from the corresponding enantiopure precursors separated by chiral HPLC. The complexes are emissive in fluid solutions at room temperature when excited in the MMLL'CT band, the triplet state being responsible for the observed emission band centered at 715-720 nm. DFT calculations support the absorption, emission and CD properties of the enantiopure compounds. The conformationally stable [6]helicene enantiopure complexes show CPL activity. These results underline the interest of helical dithiolenic ligands as means to access chirality related combined properties in the derived complexes and open the way towards the preparation of other related compounds by the use of chiral bipyridines in combination with diverse helicene-dithiolenes in order to tune their photophysical properties.

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Notes and references

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Electronic Supplementary Information (ESI) available. Experimental details and characterization of compounds, X-ray data, CV, UV-vis, ECD, photophysical measurements and theoretical calculations details (PDF).

CCDC 1550054 ((*rac*)-**2a**), 1550055 ((*rac*)-**2b**), 1550056 ((*M*)-**2b**) contain the supplementary crystallographic data (cif files) for this paper. See DOI: 10.1039/c000000x/

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