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Self-assembling a neutral platinum(II) complex into highly emitting microcrystalline fibers through metallophilic interactions†

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The solvent-assisted self-assembly of blue-emitting neutral platinum(II) complexes into micrometer-long and highly crystalline fibers has been achieved. The aggregates show highly efficient (quantum yield up to 74%) polarized yellow-orange light emission, as consequence of their high degree of supramolecular order imparted by weak non-covalent intermolecular (metal···metal and π-π) interactions.

Supramolecular organometallic nanostructures have attracted a great deal of attention in the last few years because of the possibility to fabricate functional materials with superior properties compared to the bulk. In particular, square-planar Pt(II) and Pd(II) complexes with protruding filled dz2 orbitals have been known for a long time to show a high tendency towards stacking, through weak non-covalent metal···metal and/or π-π ligand–ligand interactions. By means of bottom-up approach, it has been shown that luminescent Pt(II) complexes are able to form either homo-2 or hetero-metallic3 supramolecular architectures as liquid crystals,4 nanowires,5 nanotubes,6 nanosheets,7 and metallogelators,8a,9a with very appealing (electro)optical,9–11 sensing,12 and semiconducting properties.7,13

To date, metal complexes forming long-range ordered soft structures showing polarized light emission and photoluminescence quantum yield (PLQY) exceeding 0.5 are very rare and typically require aid of orienting scaffolds and special techniques.13 This is despite the great potential applications that such self-assembled functional architectures might have in optoelectronic devices such as organic light emitting diodes (OLEDs),9 field effect transistors (FETs),7,13 and organic light-emitting FETs (OLEFETs).11 Noteworthy, linearly polarized materials with uniaxial molecular orientation find great application for improving light extraction in such devices,9 and as active materials in 3D OLED displays.14

In this communication, we report on the synthesis and characterization of a neutral Pt(II) complex, namely Pt-CF3tz-pyC5, containing a formally dianionic N-donor tridentate chromophoric ligand. Such complex, because of a square-planar geometry, displays great tendency to aggregate in bright polarized-light emitting fibers. The compound belongs to the class of highly luminescent Pt(II) complexes recently reported by our group,15 bearing N^N^N azoles-based ligands and the chemical structure depicted in Fig. 1. The one-pot synthetic procedure employs PtCl2(DMSO)2 (DMSO = dimethylsulfoxide) as the Pt(II) precursor, diisopropylethylamine as the base, 4-amylpyridine as the ancillary ligand and 2,6-bis(3-(trifluoromethyl)-1H-1,2,4-triazol-5-yl)pyridine (pyC5-CF3-tzH2), as the tridentate ligand in a 3:1 2-methoxyethanol:H2O solvent mixture at 83°C. The final pure compound was obtained by column chromatography and fully characterized by 1H, 13C and 19F NMR and high-resolution mass spectrometry. Interestingly, the complex is extremely soluble in halogenated solvents and ethers. The experimental details are given in the Electronic Supplementary Information (ESI).

Geometrical structure optimization by means of density functional theory (DFT) has been performed for the compound Pt-
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The complex adopts a distorted square-planar arrangement around the metal atom with C₂ point group symmetry. The computed bond distances are 2.027, 1.998 and 2.052 Å for Pt–N(1), Pt–N(4) and Pt–N(5), respectively, which nicely agree with other similar Pt(II) complexes.\(^{15}\) The atoms labelling and a more complete list of geometrical parameters can be found in Fig. S1 and Table S1 of the ESI, respectively. In order to gain insights into the frontier orbitals and the redox processes of the complex Pt-CF₃tz-pyC₅, cyclic voltammetry (CV) measurements were performed in CHCl₃ as the solvent and tetrabutylammonium hexafluorophosphate (TBAPF₆) as the supporting electrolyte (see ESI for further details and Fig. S2). The cyclic voltammogram displays a reversible reduction process at \(-1.83\) V (vs Fe⁺[Fe]), which is mainly located on the pyridlic moiety of the tridentate ligand, where the lowest unoccupied molecular orbitals, i.e. LUMO, is located. Such assignment is also supported by the frontier orbital analysis carried out on the S₀ DFT-optimized geometry. Further computational results are given in Table S2, and the isodensity surface plots are depicted in Fig. S3 of the ESI. No oxidation process is visible in the employed observation window.

As shown in Fig. 1, in dilute (concentration 5×10⁻⁵ M) CHCl₃ solution at room temperature, the absorption spectrum displays intense bands in the UV region (\(\lambda_{\text{abs}} = 254\) nm, \(\varepsilon = 2.3 \times 10^4\) M⁻¹ cm⁻¹; \(\lambda_{\text{max}} = 302\) nm, \(\varepsilon = 1.5 \times 10^4\) M⁻¹ cm⁻¹; \(\lambda_{\text{max}} = 337\) nm, \(\varepsilon = 2.6 \times 10^4\) M⁻¹ cm⁻¹). These transitions are mainly attributed to the intraligand (IL) and metal-perturbed interligand charge transfer (1ILCT) states.

**Fig. 2.** a) Emission spectra obtained for the complex Pt-CF₃tz-pyC₅ in CHCl₃ (black trace) at concentration of 5×10⁻⁵ M, self-assembled fibers obtained from acetone at concentration of 4 mg mL⁻¹ (red trace), fibers obtained from CHCl₃ at concentration of 4 mg mL⁻¹ (blue trace), upon excitation at \(\lambda_{\text{exc}} = 300\) nm. b) Fluorescence microscopy image of the fibers obtained from acetone under irradiation at \(\lambda_{\text{exc}} = 400-440\) nm. Scale bar = 100 μm.

At lower energy a broad band in the region 350–450 nm (\(\lambda_{\text{exc}} = 0.7 \times 10^5\) M⁻¹ cm⁻¹) is assigned to the HOMO \(\rightarrow\) LUMO (\(f = 0.01, 3.211\) eV, 386 nm), and ascribed to the admixture of spin-allowed metal-to-ligand charge transfer (MLCT) and intraligand (IL) transitions. These assignments are further corroborated by the similarity with already reported Pt(II) complexes,\(^{15}\) and by time-dependent DFT (TD-DFT) calculation in CHCl₃ at PBE1PBE and SDD + 6-31g(d,p) level. A more complete list of electronic transitions computed both in vacuum and CHCl₃ solvent is given in Table S3 of the ESI.

**Fig. 2a** shows the emission spectrum in degassed CHCl₃ solution of the complex. A structured blue luminescence is observed, peaking at \(\lambda_{\text{em}} = 463, 493, 525\) and 570 (shoulder) nm, with vibronical progression of 1236–1504 cm⁻¹ attributable to the tridentate intraligand vibrational modes. Such moderate (PLQY = 2%) luminescence is characterized by a very fast radiative decay, with a multi-exponential excited-state lifetime, being \(\tau_1 = 10\) ns (48%) and \(\tau_2 = 309\) ns (52%). The bi-exponential decay is due to the equilibrium between the emissive 3MLCT/ILC and a thermally populated non-radiative state. Hence, such emission can be attributed to the T₁ \(\rightarrow\) S₀ radiative transition with mainly metal-perturbed LIL character, and the highest energy peak observed at 298 K (463 nm, 2.68 eV) agrees with the theoretical S₀ \(\rightarrow\) T₁ (HOMO \(\rightarrow\) LUMO) vertical excitation process computed at 432 (2.87 eV) nm in CHCl₃.

Interestingly, the complex exhibits very different photophysical behaviour in the solid state. In the attempt to crystallize the compound to obtain good quality single-crystal suitable for X-ray analysis, we have isolated yellow bright emissive microcrystalline fibers. The luminescent fibers, Fig. 2b, can be obtained when an acetone solution of the Pt-CF₃tz-pyC₅ complex is slowly concentrated by evaporation (see ESI for details). Upon photo-irradiation with UV light, such discrete fibers showed intense and featureless emission centred at \(\lambda_{\text{em,max}} = 559\) nm with PLQY as high as 0.74 (see Fig. 2a). Such great increase of the emission quantum yield is accompanied by a concomitant prolongation of the mono-exponential decay of the excited-state (\(\tau = 355\) ns). Furthermore, the employed solvent plays also an important role in the photophysical properties of the self-assembled structures. Indeed, CHCl₃ solution at concentration of 4 mg mL⁻¹ yields fibers with slightly broader emission spectra peaking at \(\lambda_{\text{em,max}} = 563\) and with PLQY of 0.58.

The sizeable bathochromic shift, the featureless emission profile, and the high emission quantum yield are already strong indications that the emissive state of the fibers is different from the luminescent level of the complex in solution. Furthermore, the outcome is not a merely rigidochromic effect since the nature of the emission is different.

It is well known that the square planar geometry of the compound can favour aggregates formation leading to Pt-Pt interactions, through the dz² orbitals of the metals that rise in energy and become the HOMO orbitals of the assemblies. As a consequence the nature of the lowest electronic transitions can change from ligand centered to metal-metal ligand charge transfer, MLCT. For our system the strong coupling between the two platinum units is observed by the appearance of a lower energy band in the excitation spectrum of the fiber (\(\lambda = 501\) nm), as displayed in Fig. S4 (ESI). The transition involved has been ascribed as 1MLCT. Excitation in this band, as well as in any other absorption leads to the yellow emission attributed to the corresponding 3MLCT. Furthermore, such solvophobic metallophilic interactions are accounted to play a major role into the nanoscale organization and molecular propagation for the self-assembly of the Pt-CF₃tz-pyC₅ complex. As a result of this long-range order, light emission perpendicularly polarized with respect to the fiber growth axis is unambiguously observed, as shown in Fig. 5 and Movie S1 of the ESI.

Morphological and structural characterization of these Pt(II) aggregates has been performed by both scanning (SEM) and transmission electron microscopy (TEM) techniques, Fig. 3, as well as small-angle X-ray scattering, (SAXS). As shown in the SEM images, the sample of the emissive soft-structures prepared from...
fibers prepared by drop-casting a 4 mg mL\(^{-1}\) acetone solution, which
information on the packing and crystallinity of the assembly.
nanofibrils. SEM analysis, however, could not give us any
result of a self-assembly process of thinner nanostructures such as
Fig. 3. SEM images taken on a sample of fibers of complex Pt-CF\(_3\)tz-
pyC\(_5\) obtained from acetone at concentration of 4 mg mL\(^{-1}\) (a) and zoom-
in image (b); c) SAED pattern from a few fibres as shown in the corresponding shadow image (d).
nanofibrils. SEM analysis, however, could not give us any information on the packing and crystallinity of the assembly.
Additionally, SAXS/WAXS experiments were performed on fibers prepared by drop-casting a 4 mg mL\(^{-1}\) acetone solution, which is the same experimental condition to obtain the fibers investigated by means of spectroscopic techniques. The scattering pattern, which displays sharp peaks, confirmed the high degree of long-range order within the self-assembled structures, as shown in Fig. S6 of the ESI. Noteworthy, it is possible to detect a scattering peak at 3.28 Å that can be most likely ascribed to the distance corresponding to \(d_2^2\pi\cdots d_2^2\pi\) intermolecular interaction leading to the strong bathochromic shift of the emission into the self-assembled fibers (see Fig. S7 of the ESI).
Finally, to explore the crystal structure of the Pt(II) aggregates prepared from acetone, selected area electron diffraction (SAED) patterns were acquired corresponding to a local region with a few fibers present. In order to avoid damage of the crystal structure as a consequence of the electron beam, very low dose rate was employed for the analysis (see ESI for details). The morphology and orientation of the fibers contributing to electron diffraction were obtained looking at the shadow image in the central disk of a largely defocused diffraction pattern. Fig. 3c-d show a representative diffraction pattern and the corresponding shadow image. In the diffraction pattern, two pairs of spots close to the central beam correspond to a lattice period of 1.33 nm. The diffraction vectors are perpendicular to the two groups of fibers numbered in the shadow image, respectively. This indicates a periodic self-assembly of the Pt complex chains perpendicular to the fiber long axis. In addition, strongly streaked diffraction spots corresponding to a lattice period of 3.53 Å were observed parallel to the molecular propagation vector. This lattice period is different to that obtained by X-ray scattering analysis and is most likely due to different sample preparation required for the TEM analysis (lower concentration and fast evaporation rate). However, this periodicity can be attributed to the long-range order of the complex along the fiber axis, while the streaking of these spots perpendicular to such axis indicates a shift disorder of the stacked complexes along the fibre long axis.
Conclusions
The synthesis and characterization of a novel blue emitting platinum(II) complexes based on \(\text{N}^\circ\text{N}^\circ\text{N}^\circ\) chromophoric ligand is reported. The compound shows a remarkable tendency to spontaneously assemble into highly ordered structures, such as fibers, through metal···metal and \(\pi\cdots\pi\) interactions. The assemblies efficiently emit polarized yellow light, as shown by fluorescent microscopy, with the remarkably high PLQY value of 74%, which might find application in polarized light-emitting devices and 3D OLED displays. To gain insight into the structural properties of the self-assembled structures SAED and SAXS/WAXS experiments were performed, which displayed an intermolecular distance of 3.28 Å attributable to the \(d_2^2\pi\cdots d_2^2\pi\) interactions along the growth axis of the fibers.

Notes and references
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