Supramolecular assembly of a phosphole-based moiety into nanostructures dictated by alkyne-alkyne interactions: synthesis, characterization, and self-assembly behaviors†

Sammual Yu-Lut Leung, Sloane Evariste, Christophe Lescop, Muriel Hissler and Vivian Wing-Wah Yam* a

A new class of platinum(II) terpyridine complexes with a phosphole-derived bridging alkyne ligand have been prepared. The X-ray crystal structure of complex 2 has been determined, and reveals a polymeric zig-zag chain structure with the existence of π−π stacking interactions. The photophysical properties have also been studied, with MLCT/LLCT phosphorescence exhibited in degassed CH2Cl2, the energy of which is varied by the π-conjugation of the terpyridine ligands. The solvent-induced assembly of complex 1 has been studied. The incorporation of hydrophobic hydrocarbon chains has been shown to play an important role in assisting the formation of self-assembled nanostructures via Pt⋯Pt, π−π stacking and hydrophobic–hydrophobic interactions. It has been established that an isodesmic growth mechanism operates in polar media to give nanospheres, while fibrous networks originate from the self-assembly of the complexes in non-polar media, predominantly driven by π−π stacking interactions.

Introduction

Studies on metallosupramolecular π-conjugated amphiphiles and their use in the construction of supramolecular architectures have aroused increasing interest recently. This is not only because of their interesting spectroscopic and luminescence properties, but also their propensity to exhibit the unprecedented complexity of multiple non-covalent interactions such as hydrogen bonding, π−π stacking interactions, metal–metal interactions and hydrophobic–hydrophobic interactions, among others. Particularly, directional Pt⋯Pt interactions have emerged as a key driving force for providing precise control and tuning of the self-assembly of metallosupramolecular π-conjugated amphiphiles into designated molecular architectures with well-defined morphologies. The unique chromophore alkyneplatinum(II) terpyridine as well as platinum(II) complexes with other related cyclometalating pincer ligands that exhibit the formation of Pt⋯Pt and π−π stacking interactions have been demonstrated to undergo self-assembly into metallogels and liquid crystals, together with a variety of nanosize molecular architectures such as hairpin conformations, tubes, helical ribbons, rods, rings and plates etc. These studies have provided an in-depth understanding into metallosupramolecular π-conjugated amphiphiles, which have been found to play an important role in dictating the well-organized molecular packings that determine nanostructures in supramolecular chemistry.

Moreover, there has been a surge of interest in electronic materials derived from organo main group motifs due to the exhibition of unique structural and electronic properties. Among these organo main group molecule motifs, phospholes are one of the unique classes of compounds in electronic material applications. Indeed, not only have they aroused attention in the development of electronic devices, but also in the development of self-assembled materials. The development of self-assembled phosphate-based species has established interesting self-assembly behaviors via the subtle balance between the intermolecular π−π interactions of the phosphate backbones, electrostatic interactions as well as the thermal disorder in the hydrocarbon chains. Distinct from previously reported phosphate compounds, metal-containing phospholes are relatively less explored. Although there have been a few
examples in the literature of chloro- and alkynyl-gold(i) phosphole systems, which are confined to studies into their application in organic light-emitting diodes (OLEDs) and self-assembly behaviors, research into functionalization of the phosphole system with a platinum(n) metal center, which is well-known to exhibit interesting and rich luminescence and self-assembly behaviors, is essentially unexplored. Exploration in this area may lead to the discovery of unique coordination, supramolecular and material properties. Thus, in view of the rich self-assembly behaviors of square-planar d8 platinum(n) polypyridine systems and their propensity towards the formation of directional Pt⋯Pt interactions, it is anticipated that such directional and noncovalent interactions can act as additional driving forces for the self-assembly of metallosupramolecular -conjugated phosphole derivatives that may demonstrate interesting spectroscopic, luminescence and morphological properties. These may also provide important insight into the supramolecular assembly of the phosphole-based moiety driven by non-covalent Pt⋯Pt and π–π stacking interactions, which is distinct from previous studies on pure organic analogues and gold(i) systems. The utilization of such directional and non-covalent Pt⋯Pt interactions is anticipated to dictate the assembly behavior of the phosphole-based moiety in a controllable manner under various external stimuli.

Herein, we report the design and synthesis of dinuclear platinum(n) terpyridine complexes as well as that of a mononuclear complex containing a -conjugated phosphole-derived alkynyl ligand. To the best of our knowledge, this study represents the first example of platinum(n)-containing phosphole complexes, which can exhibit non-covalent metal–metal and π–π stacking interactions in the self-assembly and lead to morphological changes under different conditions. Its physicochemical properties of the complexes have been studied. It is also found that the dinuclear complexes with long hydrocarbon chains would self-assemble into spheres in DMSO, while they would aggregate into fibrous networks in diethyl ether. Temperature-dependent UV-vis absorption spectroscopic studies have provided an in-depth understanding into the nature of molecular self-assembly and the role of metal–metal interactions in the formation of these nanostructures, with the isodesmic growth mechanism established for their formation.

Results and discussion

Synthesis and characterization

The detailed synthetic route for the phosphole-based bridging alkynyl ligand is given in Scheme S1.† This class of bridging phosphole-containing ligand would steadily polymerize upon deprotection of the triisopropylsilyl (TIPS) groups. Thus, dinuclear alkynylplatinum(n) terpyridine complexes have been prepared by reacting a mixture of the TIPS-protected phosphole-containing bridging alkynyl ligand with the corresponding chloroplatinum(n) terpyridine precursor in degassed dichloromethane containing triethylamine and a catalytic amount of copper(i) iodide. The bridging phosphole-containing alkynyl ligand was deprotected in situ by the dropwise addition of tetra-n-butylammonium fluoride (TBAF) at 273 K. The reaction mixture was then stirred overnight at room temperature. Chromatography on silica gel using a dichloromethane-acetone mixture (10:1 v/v) as the eluent, followed by the diffusion of diethyl ether vapor into the CH2Cl2 solution, gave complexes 1 and 2 (Fig. 1) as dark red and red solids, respectively. The mononuclear platinum(n) complex with the bridging phosphole alkynyl ligand was isolated as a side-product in the preparation of complex 1. All complexes were found to be air-stable upon storage in the dark. These cationic complexes were found to be very soluble in common organic solvents such as acetonitrile and acetone, and soluble in dichloromethane and chloroform. In particular, complex 1 is barely soluble in diethyl ether, because of the four long hydrocarbon chains (–C18H37). All dinuclear complexes were characterized by 31P{1H} NMR and IR spectroscopy, FAB mass spectrometry and elemental analyses. The 31P{1H} NMR signal of 2 is situated at δ = +51.37 ppm, which is typical of the phosphole moiety. The IR spectra of complexes 1 and 2 exhibit characteristic C≡C absorptions at 2081–2086 cm⁻¹, which are in accordance with the presence of the alkynyl ligand in a terminal σ-coordination mode. In addition, the crystal structure of complex 2 has been successfully determined by X-ray crystallography.

X-ray crystal structure

The coordination geometry at the platinum centre of the alkynylplatinum(n) terpyridine complex is distorted square-planar with the N(1)–Pt(1)–N(3), N(1)–Pt(1)–N(2) and N(2)–Pt(1)–N(3) bond angles being 160.7°, 80.2° and 80.8°, respectively. The steric demand of the tridentate terpyridine ligand distorts the
square-planar geometry from the ideal values of 90° and 180°. The square-planar coordination promotes planarization of the terpyridine moieties, which facilitates the formation of supramolecular architectures. A perspective drawing of complex 2 is shown in Fig. 2a. The crystal structure determination data are given in the Experimental section and in Table S1.† The selected bond lengths and angles of complex 2 are summarized in Table 1. The bond lengths of Pt(1)–C(12) and C(11)–C(12) are 1.981 and 1.223 Å, respectively, which are similar to those observed in the previously reported alkynylplatinum(II) terpyridine and buzimpy systems.‡ Furthermore, the phosphorus center adopts a distorted tetrahedral geometry with a small endocyclic C(1)–P(1)–C(8) angle in the range of 92.7°. The endocyclic P–C bonds, P(1)–C(1) and P(1)–C(8), are found to have lengths of 1.836 and 1.806 Å, respectively, which are comparable to those observed in the previously reported phosphole systems.‡ The crystal packing diagram shows that the dicationic complexes are stacked in a head-to-tail configuration. Interestingly, the crystal packing of complex 2 displays zig-zag chains that are held together mainly through π–π stacking interactions (Fig. 2b). This is because the interplanar distance between the two platinum(u) terpyridine moieties of the dimeric structure is found to be 3.47 Å, which suggests weak to almost insignificant Pt···Pt interactions in the polymeric zigzag chain structures.

Photophysical studies
The red to orange CH2Cl2 solutions of complexes 1 and 2, together with the mononuclear complex, were examined using UV-vis absorption and emission spectroscopy. Their photophysical data are summarized in Table 2. In the UV-vis absorption spectra, high-energy absorption bands at ca. 313–412 nm, together with low-energy absorption bands at ca. 512–565 nm, are observed. The former are assigned as intense intraligand (IL) [π → π*] transitions of the terpyridine and alkynyl ligands, as supported by the UV-vis absorption spectra of the organic counterparts (Fig. S1†), while the latter are assigned as metal-to-ligand charge-transfer (MLCT) [dσ(Pt) → π*(tpy)] transitions mixed with alkynyl-to-terpyridine ligand-to-ligand charge transfer (LLCT) [π(C≡CR) → π*(tpy)] character (Fig. 3a).† In general, the dinuclear complexes exhibit more red-shifted MLCT/LLCT absorption bands (533–565 nm) than the mononuclear complex (512 nm). Notably, complex 1 displays the most red-shifted MLCT/LLCT absorption band due to the more π-conjugated and electron-deficient terpyridine ligand, thus lowering the π*(tpy) orbital energy (LUMO), leading to

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Selected bond distances and angles, with estimated standard deviations shown in parentheses, for complex 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bond angles (deg)</td>
<td>N1–Pt1–N3</td>
</tr>
<tr>
<td>N1–Pt1–N2</td>
<td>160.7(4)</td>
</tr>
<tr>
<td>N2–Pt1–N3</td>
<td>80.2(3)</td>
</tr>
<tr>
<td>S1–Pt1–C1</td>
<td>115.8(5)</td>
</tr>
<tr>
<td>S1–P1–C8</td>
<td>120.1(4)</td>
</tr>
<tr>
<td>Bond distances (Å)</td>
<td>Pt1–C12</td>
</tr>
<tr>
<td>C11–C12</td>
<td>1.981(10)</td>
</tr>
<tr>
<td>P1–S1</td>
<td>1.951(5)</td>
</tr>
</tbody>
</table>

Fig. 2  (a) Perspective drawing of the complex cation of 2. Hydrogen atoms, counter-ions and solvent molecules have been omitted for clarity. Thermal ellipsoids are shown at the 30% probability level. (b) Crystal packing diagram of the complex cations of 2.
Table 2  Photophysical data of complexes 1, 2 and of the mononuclear complex

<table>
<thead>
<tr>
<th>Complex</th>
<th>Absorption $\lambda$/nm [ε/dm$^3$ mol$^{-1}$ cm$^{-1}$]</th>
<th>Medium (T[K])</th>
<th>Emission $\lambda$ [nm] ($\tau_s$ [μs])</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>339 (28 990), 419 sh (13 350), 565 (11 530)</td>
<td>CH$_2$Cl$_2$ (298)</td>
<td>714 (&lt;0.1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Solid (298)</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Solid (77)</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Glass (77)$^a$</td>
<td>—</td>
</tr>
<tr>
<td>2</td>
<td>342 (26 340), 414 (11 410), 533 (14 100)</td>
<td>CH$_2$Cl$_2$ (298)</td>
<td>701 (&lt;0.1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Solid (298)</td>
<td>700 (&lt;0.1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Solid (77)</td>
<td>702 (&lt;0.1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Glass (77)$^a$</td>
<td>633, 698 sh (&lt;0.1)</td>
</tr>
<tr>
<td>Mononuclear complex</td>
<td>325 (24 210), 415 sh (11 410), 512 (6720)</td>
<td>CH$_2$Cl$_2$ (298)</td>
<td>596 (&lt;0.1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Solid (298)</td>
<td>582 (&lt;0.1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Solid (77)</td>
<td>583 (&lt;0.1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Glass (77)$^a$</td>
<td>574, 620 sh (&lt;0.1)</td>
</tr>
</tbody>
</table>

$^a$ Non-emissive. $^b$ Measured in n-butyronitrile.

Fig. 3  (a) UV-vis absorption and (b) emission spectra ([Pt] = 1 × 10$^{-5}$ M) of (——) 1, (——) 2 and (——) mononuclear complex in CH$_2$Cl$_2$ at 298 K. (The asterisk indicates the instrumental artifact.)

a lower MLCT/LLCT absorption energy. In the emission spectra, weak luminescence at ca. 596–714 nm in degassed CH$_2$Cl$_2$ was observed upon excitation at $\lambda \approx 360$ nm (Fig. 3b). The structureless emission bands are derived from an excited state which is predominantly 3MLCT [dπ(Pt) → π* (tpy)] in origin, with mixing of 3LLCT [π(C≡CR) → π*(tpy)] character (vide infra). Concomitant with the findings from the UV-vis absorption spectra, the mononuclear complex exhibits the highest energy 3MLCT/3LLCT emission bands (596 nm) and complex 1 displays the lowest emission energy (714 nm).

Complex 1 is found to be non-emissive at room and low temperatures in the solid state as well as in glass matrices at 77 K, while complex 2 and the mononuclear complex exhibit structureless emission bands at 582–700 nm in the solid state at 298 K and 77 K, which can be assigned as being derived from 3MLCT/3LLCT origin. In the glass state, they are found to exhibit vibronically structured emission bands at ca. 574–633 nm with a shoulder at ca. 620–698 nm. The vibrational progression spacings, ranging from 1292 to 1471 cm$^{-1}$, are in line with the C=C and C=N vibrational modes of the tridentate terpyridine ligands.

To provide further insight into the nature of the excited states for this class of complexes, nanosecond transient absorption (TA) measurements were carried out on complex 2 at room temperature in degassed CH$_2$Cl$_2$. The nanosecond TA spectrum of complex 2 recorded immediately after laser flash excitation at 355 nm is shown in Fig. S2.$^\dagger$ Upon laser excitation, the TA spectrum displayed a high-energy absorption band at 368 nm and a low-energy absorption band at 610–657 nm, as shown in Fig. S2.$^\dagger$ With reference to previous TA studies on the related alkynylplatinum(n) terpyridine system, the absorption bands were tentatively assigned as the radical anion absorption of the tridentate terpyridine ligand that results from the absorption of the 3MLCT [dπ(Pt) → π*(tpy)]/3LLCT [π(C≡CR) → π*(tpy)] excited state. The absorption signals for complex 2 were found to exhibit monoexponential decay with a decay time constant of about 20.0 ns. Bleaching was also observed in the spectrum at approximately 650 nm, which was ascribed to the depletion of the ground state upon MLCT/LLCT absorption.

Self-assembly study

The hydrophobic dinuclear complex 1 with long hydrocarbon chains had its self-assembly behavior further examined in solvents with different polarities. In different CH$_2$Cl$_2$ solution concentrations, the plot of the apparent absorbance at 620 nm is found to show good agreement with Beer’s law in the concentration range of 11–180 μM (Fig. S3$^\dagger$). This suggests that there is no significant intermolecular aggregation of the complex molecules in CH$_2$Cl$_2$. Notably, this complex would exhibit strong solvatochromism upon increasing the polarity of the solvent. The solution color of complex 1 is found to change from red to purple in high concentrations of DMSO, which is accompanied by a significant drop in the high-energy absorption band at ca. 350 nm, together with a new absorption shoulder at ca. 620 nm with clear isosbestic points in the UV-vis absorption spectra, as depicted in Fig. 4a. The drop in the absorbance in the high-energy region (ca. 350 nm), which is a result of the hypochromicity effect, indicates that complex 1 is driven to undergo self-assembly through π–π stacking interactions with increasing DMSO concentration. On the other hand, the emergence of the absorption shoulders (ca. 620 nm) in pure DMSO has been further monitored at various concentrations.
shoulder is assigned as metal

terpyridine moieties upon increasing the solvent

stacking and metal

relationship of Beer

UV-vis absorption spectra of $z$

K ($[Pt] \approx 1 \times 10^{-5}$ M). The apparent absorbance values have been obtained by correcting to a 1 cm path length equivalence.

(Fig. S4,† 102–445 μM), which shows a deviation from the linear relationship of Beer’s law. Therefore, the low-energy absorption shoulder is assigned as metal–metal-to-ligand charge-transfer (MMLCT) transitions, resulting from the formation of $\pi$-$\pi$

stacking and metal–metal interactions, which are a result of the intermolecular self-assembly of the hydrophobic alkynylplatinum

(II) terpyridine moieties upon increasing the solvent polarity.

To further investigate the self-assembly mechanism of complex 1, temperature-dependent UV-vis absorption studies were conducted in pure DMSO. Upon decreasing the temperature from 360 K in DMSO ([Pt] = 218 μM), emergence of the MMLCT absorption band (620 nm) is observed with depletion of the MLCT/LLCT band at ca. 530 nm (Fig. 4b), thus indicating the formation of self-assembled aggregates of complex 1 with the formation of $\pi$-$\pi$

stacking and metal–metal interactions. Interestingly, the cooling curve of complex 1, obtained by plotting the fraction of aggregated species ($\alpha_{agg}$) against temperature at 620 nm, is clearly sigmoidal (Fig. 4b, inset), which is indicative of an isodesmic self-assembly process. The thermodynamic parameters associated with the self-assembly process of complex 1 in DMSO (Table 3) have been determined by fitting $\alpha_{agg}$ against temperature at 620 nm in accordance with the isodesmic model. In particular, the enthalpy and entropy changes are determined to be $\Delta H = -138 \pm 5$ kJ mol$^{-1}$ and $\Delta S = -428 \pm 2$ J mol$^{-1}$ K$^{-1}$, respectively. The negative enthalpy and entropy values indicate that the self-assembly process is enthalpy driven. Therefore, the origin of the self-assembly behavior can be attributed to the synergistic participation of $\pi$-$\pi$ and Pt····Pt interactions in the self-assembly of complex 1 in its aggregated form, thus giving rise to the changes observed in the UV-vis absorption spectra. It is

worth noting that the phosphole moiety does play an important role in influencing the self-assembly mechanism. Recently, a series of dinuclear alkynylplatinum(II) terpyridine complexes containing the $\pi$-conjugated, planar and linear oligo(phenylene ethynylene) (OPE) moiety have been reported to undergo self-assembly through the mechanism of cooperative supramolecular polymerization in DMSO to give nanostructured tubular aggregates.$^{2}$ In the present system, the mechanism of self-assembly behavior is found to adopt an isodesmic polymerization with the presence of the phosphole moiety. This distinct feature could be attributed to the difference in the molecular design of the bridging alkynyl ligands, in which the OPE unit is $\pi$-conjugated and linear while the phosphole unit is bent-shaped with the phosphorus center adopting a tetrahedral geometry. The linear nature of the OPE alkynyl ligands might facilitate lamellar packing to give straight nanotubes via cooperative supramolecular polymerization. In contrast, the tetrahedral geometry at the phosphorus center of the phosphole moiety together with the bent nature of the bridging alkynyl ligands favors the formation of spheres with curved surfaces in the self-assembly process. Complex 1 essentially undergoes the isodesmic polymerization mechanism, gradually undergoing self-assembly into spherical aggregates in the polar DMSO medium. Therefore, the incorporation of the phosphole moiety in the alkynylplatinum(II) terpyridine complexes can provide features that are distinct from those reported previously.$^{3}$ In addition, the phosphorus centre in phosphole can be readily further functionalized for modification of the bridge. This can allow one to explore the influence of the nature and the geometry of the spacers on the supramolecular assembly of the metal complex systems.

Apart from the self-assembly induced by polar solvents, studies on the addition of non-polar solvents have also been performed. By increasing the diethyl ether content in CH$_2$Cl$_2$, complex 1 exhibits similar changes in the UV-vis absorption spectra, such as a drop in the absorbance in the high-energy region (ca. 350 nm) and the emergence of a MMLCT absorption shoulder (Fig. 5). This indicates that complex 1 undergoes self-assembly, driven by the formation of Pt····Pt interactions. However, in comparison with the self-assembly induced by polar solvents, the growth of the red-shifted MMLCT absorption band in the presence of increasing diethyl ether content is less significant. This may possibly be due to the presence of four hydrophobic hydrocarbon chains (–C$_{18}$H$_{37}$), which leads to

![UV-vis absorption spectra of 1 in CH$_2$Cl$_2$ at 298 K upon increasing the diethyl ether content from 10 to 100% ($[Pt] \approx 1 \times 10^{-5}$ M).](image-url)

Table 3  Thermodynamic parameters describing the self-assembly of complex 1

| C/μM | $\Delta H$/kJ mol$^{-1}$ | $T_m$/°K | $K_S \times 10^{-5}$/M$^{-1}$ | $\Delta S$/J mol$^{-1}$ K$^{-1}$ | $D_P$/$a$
|-----|-----------------|--------|-----------------|-----------------|-------
| 218 | $-138 \pm 5$ | 322.5 | 24.2 | $-428 \pm 2$ | 23.5 |

$^a$ Values determined at 298 K.
better solvation and a lesser extent of metal–metal interaction formation in non-polar solvents. Nevertheless, the hypochromicity effect in the high-energy region indicates the presence of intermolecular self-assembly of the complexes with the formation of π–π stacking interactions.

**Morphological studies**

The distinctive self-assembly behavior of complex 1 in environments of different polarities prompted us to study the self-assembled morphologies using electronic microscopy. In DMSO solution, complex 1 is found to self-assemble into spherical aggregates with diameters of ca. 20–50 nm on carbon-coated copper grids, as demonstrated in the TEM and SEM images depicted in Fig. 6a and b, respectively. Energy-dispersive X-ray (EDX) analysis into the spherical aggregates was conducted and the results are shown in Fig. S5,† which indicates the presence of platinum. Similar spherical architectures on silica wafers can also be observed in the AFM image shown in Fig. 6c, which shows heights of ca. 12 nm. However, the spherical aggregates become flattened on the silica wafers, possibly due to the strong interaction between the soft material and the surface of the substrate. By considering the hydrocarbon chains to be in an all-trans configuration, the molecular length of complex 1 is much smaller than the dimensions of the spherical aggregates. Thus, these supramolecular structures are constructed from multiple layers of the complexes with interdigitated hydrocarbon chains in order to minimize the contact of the polar DMSO media. Together with the observations from the UV-vis absorption spectroscopic study, it is believed that complex 1 undergoes stepwise self-assembly into spherical aggregates with a single equilibrium constant of \( K_c = 24.2 \times 10^5 \) M\(^{-1}\), driven by Pt···Pt, π–π stacking and hydrophobic–hydrophobic interactions in DMSO media.

In contrast to its aggregation behavior observed in polar solvents, in diethyl ether complex 1 aggregates into a network of irregular fibrous structures with widths in the range of ca. 20–50 nm and lengths of a few hundreds of μm, as shown in the TEM and SEM images (Fig. 7a and b respectively). This discrepancy in size was further evidenced by DLS measurements conducted in two different solvents, in which the hydrodynamic diameter of the fibrous networks is much larger in diethyl ether (Fig. S6†). The four hydrophobic hydrocarbon chains help the doubly-positive charged complexes become solvated in non-polar solvents. It is believed that the fibrous networks originate from the self-assembly of complex 1, predominantly driven by π–π stacking interactions.

**Conclusion**

To conclude, a new class of platinum(II) terpyridine complexes with a phosphole-based bridging alkynyl ligand have been synthesized through in situ protection of the TIPS groups by TBAF, followed by a catalytic reaction. X-ray crystallography analysis demonstrated that the structure of complex 2 exhibits polymeric zigzag chain structures. The photophysical properties of the complexes have also been studied. They exhibit \(^3\)MLCT/\(^3\)LLCT phosphorescence in degassed CH\(_2\)Cl\(_2\) with emission energies that are varied by the π-conjugation of the terpyridine ligands. Furthermore a solvent-induced aggregation study on complex 1 was also carried out. The incorporation of hydrophobic hydrocarbon chains is thought to play an important role in governing the formation of the nanostructure. The formation of spheres is stabilized by Pt···Pt, π–π stacking and hydrophobic–hydrophobic interactions via the isodesmic mechanism in polar media, while fibrous networks originate from the self-assembly of the complexes in non-polar media, predominantly driven by π–π stacking interactions.

**Experimental section**

**Materials and reagents**

1,7-Diiodooctadiyne was prepared as described in the literature.\(^{14}\) [Pt(R-tpy)Cl][OTf], where R-tpy = \( 4'-(3,5\)-bis(octadecylxyloxy)phenyl)-2,2′:6′,2″-terpyridine,\(^{19}\) and [(Bu₄tpy)PtCl][OTf]\(^{20}\)
were synthesized according to modifications of previously reported procedures. Pd(PPh₃)₂Cl₂, ethynyltrimethylsilane, I₂, (trisopropylsilyl)acetylene and TBAF (1.0 M) were obtained from Aldrich Chem. Co. Ltd., 1,7-Octadecyln (Alfa Aesar Chem. Co. Ltd.), K₂PtCl₄ (Strem Chemicals), triethylamine (Apollo Scientific Ltd) and CuI (Aeros Chemical Co. Ltd.) were purchased from the corresponding chemical suppliers. All other reagents and solvents were of analytical grade and were used as received.

Synthesis and characterization of the phosphole-based alkylnyl ligand

All reactions were carried out under an inert atmosphere of argon using standard Schlenk techniques.

Synthesis of tetrayne. To a solution of 1,8-diiodoocta-1,7-diyne (4.06 g, 11.2 mmol) and trisopropylacetylene (5.00 g, 27.4 mmol) in degassed pipetidine (45 mL) was added CuI (760 mg, 4 mmol) at 273 K. The mixture was stirred for 3 h at room temperature. This mixture was poured into a saturated aqueous solution of NH₄Cl, and the aqueous phase was extracted three times with CH₂Cl₂. The combined organic extracts were washed with brine, dried over MgSO₄, and then filtered through Celite. The filtrate was evaporated under reduced pressure. The crude residue was purified by column chromatography on silica gel (100% heptane, Rf = 0.26) to afford 1,12-bis(triisopropylsilyl)dodeca-1,3,9,11-tetrayne as a beige solid. Yield: 5.1 g (99%). ¹H NMR (400 MHz, CDCl₃, 298 K): δ = 2.33 (m, 4H, C≡CCH₂), 1.66 (m, 4H, C≡CCH₂CH₂), 1.08 (s, 42H, Si(CH₃)₃). ¹³C NMR (101 MHz, CDCl₃, 298 K): δ = 90.1 (s, C≡C), 80.6 (s, C≡C), 78.1 (s, C≡C), 66.5 (s, C≡C), 27.3 (s, C≡CCH₂), 19.0 (s, C≡CCH₂CH₂), 18.7 (s, Si–CH–(CH₃)₃), 11.4 (s, Si–CH). HR-MS (EI): m/z found 607.3377 [M + H]+; C₃₆H₅₆Si₂PS calcd 607.33789.

Elemental analysis calcd (%) for C₃₆H₅₆Si₂PS: C 71.23, H 9.13, S 5.28; found: C 71.45, H 9.01, S 5.24.

Synthesis of the phosphole-based alkylnyl ligand. To an Et₂O solution (30 mL) containing tetranyne (1.1 g, 2.42 mmol) and Ti(OiPr)₄ (0.790 mL, 2.66 mmol) was added dropwise, at 195 K, i-PrMgCl (1.85 M in Et₂O, 2.6 mL, 4.84 mmol). The reaction mixture was stirred for 1 h 30 at 273 K and for 18 h at room temperature. A saturated aqueous NH₄Cl solution was then added and the pale brown mixture was stirred for 20 h at room temperature. The solution was then filtered and evaporated under reduced pressure. The crude product was dissolved in chloroform and washed twice with saturated NaCl solution. The organic layer was dried over anhydrous MgSO₄ and evaporated under reduced pressure. The product was further purified by precipitation of the chloroform solution in hexane twice, and the product was obtained as a yellow powder.

Synthesis of 1. To a solution of phosphole-based alkylnyl ligand (50 mg, 0.08 mmol) and [Pt(tpy)Cl]OTf (172 mg, 0.16 mmol), where R-tpy = 4’-(3,5-bis(octadecyloxy)phenyl)-2,2’:6’,2’’-terpyridine, in degassed dichloromethane (30 mL) containing triethylamine (5 mL) and a catalytic amount of CuI was added dropwise tetra-n-butylammonium fluoride (TBAF) (0.1 M, 0.40 mmol) at 0 °C. The solution was stirred overnight at room temperature. After removing the solvent, the reaction mixture was purified by column chromatography on silica gel using a chloroform-acetone mixture (10:1 v/v) as the eluent, followed by the diffusion of diethyl ether vapor into a CH₂Cl₂ solution of the complex to give 1 as red crystals. Yield: 20 mg (32%).

¹H NMR (400 MHz, DMSO-D₆, 350 K): δ = 9.01 (d, J = 7.6 Hz, 4H, terpy), 8.87 (s, 4H, terpy), 8.81 (d, J = 7.6 Hz, 4H, terpy), 8.60 (d, J = 10.8 Hz, 4H, terpy), 8.43 (s, J = 7.6 Hz, 4H, terpy), 7.89 (m, 1H, Ph), 7.75 (t, J = 7.6 Hz, 4H, terpy), 7.62 (dd, 2H, Ph), 7.54 (m, 2H, Ph), 7.19 (s, 2H, terpy), 4.01 (t, 8H, –OCH₂–), 2.92 (m, 4H, –CH₂–), 2.20 (m, 4H, –CH₂–), 2.83 (m, 4H, –C₆H₄), 1.23 (s, 66H, –C₆H₄). Positive FAB-MS: ion clusters at m/z 1633.5 [M + OTf]⁺, 742.4 [M – 2OTf]²⁺. FTIR: 2081 cm⁻¹ ν(C=C). Elemental analyses calcd (%) for...
C_{14}H_{18}F_{6}N_{6}O_{16}PPt_{2}S_{3} CH_{2}Cl_{2}: C 58.79, H 6.91, N 3.05; found: C 58.88, H 7.07, N 2.89.

**Synthesis of 2.** Yield: 152 mg (60%). 1H NMR (400 MHz, CDCl$_3$, 298 K, relative to Me$_4$Si); $\delta = 9.17$ (d, $\text{J} = 7.6$ Hz, 4H, terpy), 8.45 (s, 4H, terpy), 8.37 (d, $\text{J} = 2.0$ Hz, 4H, terpy), 8.12 (dd, 2H, Ph), 7.54 (m, 7H, terpy and Ph), 2.92 (m, 4H, $\text{CH}_{2}$), 2.20 (m, 4H, $\text{CH}_{2}$), 1.57 (s, 18H, $\text{CH}_{12}$), 1.52 (s, 36H, $\text{CH}_{12}$), $^{31}$P ($\text{1H}$ NMR (161 MHz, CDCl$_3$, 298 K, relative to 85% H$_3$PO$_4$): $\delta = 51.37$. Positive FAB-MS: ion clusters at $m/z$ 1633.5 [M – OTf$^-$], 742.4 [M – 2OTf$^-$]. FTIR: 2086 cm$^{-1}$ ($\text{C=O}$). Elemental analyses calculated (%) for C$_{74}$H$_{83}$F$_6$N$_6$O$_6$PPt$_2$S$_3$: C 58.79, H 6.91, N 3.05; found: C 58.88, H 7.07, N 2.89.

**Synthesis of mononuclear alkenylplatinum(n) terpyridine complex**

$^{1}$H NMR (400 MHz, CDCl$_3$, 298 K, relative to Me$_4$Si); $\delta = 9.05$ (d, $\text{J} = 7.2$ Hz, 2H, terpy), 8.76 (s, 2H, terpy), 8.68 (s, 2H, terpy), 7.99 (t, $\text{J} = 7.7$ Hz, 2H, terpy), 7.88 (dd, $\text{J} = 13.7$ Hz, $\text{J} = 7.6$ Hz, 2H, $\text{CH}_{2}$), 7.42 (m, 5H, terpy and $\text{CH}_{2}$), 7.08 (s, 2H, terpy), 6.45 (s, 1H, terpy), 4.08 (m, 4H, $\text{CH}_{2}$), 2.73 (m, 4H, $\text{CH}_{2}$), 2.65 (m, 4H, $\text{CH}_{2}$), 2.83 (m, 4H, $\text{CH}_{2}$), 1.23 (s, 66H, $\text{CH}_{12}$), 1.15 (s, 90H, $\text{CH}_{12}$), $^{13}$P $^{[1]}$H NMR (161 MHz, CDCl$_3$, 298 K, relative to 85% H$_3$PO$_4$): $\delta = 51.55$. Positive ESI-MS: ion clusters at $m/z$ 1334.7 [M – OTf$^-$].

**Physical measurements and instrumentation**

$^{1}$H and $^{31}$P $^{[1]}$H NMR spectra were recorded using a Bruker DPX-400 Fourier transform NMR spectrometer at 400 MHz ($^{1}$H NMR) and 161 MHz ($^{31}$P NMR) with chemical shifts recorded relative to tetramethylsilane (Me$_4$Si) and 85% phosphoric acid (H$_3$PO$_4$), respectively. High-resolution mass spectra for the ligands were obtained using Varian MAT 311, Waters Q-TOF 2 or ZabSpec TOF Micromass Instruments at the CRMPO, University of Rennes 1. Elemental analyses of the metal complexes were carried out by the slow diffusion of vapors of pentane into dichloromethane solutions, as very thin and small dark crystalline blocks that were in most cases not single crystals. After several attempts, a small single crystal that presented a satisfactory but weak diffraction pattern was obtained and used for the X-ray diffraction studies. Single-crystal data collection was carried out using an APEX II Bruker-AXS instrument (Centre de Diffractionométrie, Université de Rennes 1). The reflections were indexed, and the Lorenz polarization corrected and integrated using the DENZO program of the KappaCCD software package. The data-merging process was performed using the SCALEPACK program. Structure determinations were performed by direct methods using the SIR97 solving program, which revealed all the non-hydrogen atoms. The SHELXL program was used to refine the structures using the full-matrix least squares method based on $\chi^2$. All of the non-hydrogen atoms were refined with anisotropic displacement parameters. The hydrogen atoms were included in idealized positions and refined with isotropic displacement parameters (as well as two non-H atoms that could not be refined with anisotropic displacement parameters). In the studied crystal lattices of 2, dichloromethane solvent molecules were found in addition to the cationic coordination complexes and their counter-ions. These solvent molecules in most cases have a strong tendency to leave the bulk crystal via evaporation once the crystals are removed from their mother liquor, a process that induces rapid degradation of the single-crystal integrity of the crystals investigated. In order to slow down this process, the single crystal was always coated in paratone oil once removed from the mother liquor, and mounted at a low temperature of 150 K as quickly as possible on the diffractometer goniometer, and X-ray data collection was carried out at 150 K. Nevertheless, even by applying such a procedure, the mounted single crystal most probably partially lost its included dichloromethane solvent molecules, which induced partial degradation of the bulk single crystal integrity and resulted in the collection of...
a weak diffraction data set. The included dichloromethane solvent molecules and triflate counter-ions were found to be highly disordered, and modeling of the disorder of these solvent molecules was not possible, which led to rather high anisotropic displacement parameters for some of their atoms. Therefore a ‘SQUEEZE’ treatment was carried out in order to remove the scattering contribution of these molecules that could not be satisfactorily modeled. In addition, some of the carbon atoms of the six tert-butyl groups present a high ADP max/min ratio and the large Hirshfeld difference is a result of the disordered arrangement of these tert-butyl groups in the crystal structure. Nevertheless, correct modeling of this disorder was not possible, which led to unstable refinement cycles together with an unjustified increase in the number of modeled parameters. Consequently, the final R1 and wR2 factors were quite high both before and after the ‘SQUEEZE’ procedure was applied, thus reflecting the unfavorable parameters. The crystal data and structure refinement at 150 K for 2 before and after this ‘SQUEEZE’ treatment are given in Table S1. Atomic scattering factors for all of the atoms were taken from the International Tables for X-ray Crystallography. The CCDC reference number 1506200 contains the supplementary crystallographic data for 2.

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References


