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Introduction

The ever increasing demand to save energy and the global awareness of the need to efficiently use fossil fuels have sparked the deployment of cleaner energy alternatives¹ and provoked the utilization of freely available natural feedstocks such as CO₂.² Extensive effort has also been made to attain sustainability in lighting technology³ by using emissive complexes as the primary building blocks for electroluminescent devices.^{3c} Furthermore, the motivation for this emerging technology also rests on the industrial demand for the fabrication

Bidentate NHC^pyrozolate ligands in luminescent platinum(II) complexes†

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A bidentate C^N donor set derived from an N-heterocyclic carbene (NHC) precursor linked to a trifluoromethyl (CF₃) functionalized pyrazole ring is described for the first time. The ligands have been employed to prepare four new phosphorescent complexes by the coordination of platinum(II) centres bearing cyclometalated phenyl-pyridine/triazole-pyridine chelates. The electronic and steric environments of these complexes were tuned through the incorporation of suitable substituents in the phenyl-pyridine/triazolepyridine ligands, wherein the position of the phenyl-ring substituent (a CF₃ group) also directs the selective adoption of either a *trans* or a *cis* configuration between the C_{NHC} and the C_{phenyl} donor atoms. Molecular structures obtained by X-ray diffraction for three of the complexes confirm a distorted squareplanar configuration around the platinum centre, and DFT calculations show that the substituents have a significant influence on the energies of the frontier orbitals. Moreover, a platinum(II) complex featuring the new bidentate NHC^pyrazolate ligand and a bulky adamantyl functionalized pyridine-triazole luminophore was observed to be highly emissive and exhibiting a sky-blue luminescence ($\lambda_{Em} = 470$ nm) with photoluminescence quantum yields as high as 50% in doped PMMA matrices. A complete photophysical investigation of all of the complexes in solution as well as in the solid state is herein reported.

> of efficient flat panel displays.⁴ The use of transition metal complexes as triplet emitters is advantageous, as they can deliver four-fold enhanced efficiencies, as compared to organic fluorophores.⁵ Electroluminescent complexes based on iridium,⁶ ruthenium,⁷ rhenium,⁸ osmium,⁹ platinum,¹⁰ gold,¹¹ copper,¹² and zinc¹³ have been previously well documented. Ligand design plays a pivotal role in determining the stabilities and luminescence efficiencies of these complexes. For example, symmetric NCN type benzene bis(pyrazolates)^{10d} or unsymmetric NNN¹⁴ type pincer ligands constituting rigid ligand frameworks exhibit rich photoluminescence upon coordination to platinum(II) or iridium(III) centers. Transition metal complexes bearing N-heterocyclic carbene (NHC) ligands are vastly applied in organometallic catalysis,¹⁵ as their strong σ donor properties and large ligand cone angles facilitate the stabilization of the transition states, preventing the degradation of the involved catalytic species. These aspects also render them comparable to their phosphine counterparts.¹⁶ More recently, several research groups^{6a,c,10h,i,17} have dedicated extensive efforts to obtain photoactive complexes bearing NHC ligands as functional units. The strong ligand field of NHC ligands assists in destabilizing the otherwise thermally accessible yet non-emissive metal-centred d-d states in their complexes. Moreover, the coordination of bidentate, tridentate or tetradentate-NHCs or mixed NHCs to the platinum atom renders a rigid coordination sphere that minimizes



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[†]Electronic supplementary information (ESI) available: Description of the synthetic procedure for compound **6**; crystallographic refinement parameters and CIF files of all the crystal structures reported herein; ambient temperature *vs.* 77 K excitation spectra of all the complexes; solid state neat film emission spectra of all the complexes; a list of the selected molecular orbital energies [eV] for all the complexes and HOMO–LUMO energy gaps; isodensity surface plots of the selected orbitals for all the complexes; cyclic voltammetry plots. CCDC 981765–981767. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c4dt03651d

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the vibronic deactivation of excited states and consequently affords high photoluminescence quantum yields (PLQY).⁴

Deep blue emission was observed in homoleptic platinum(II) complexes featuring bis(NHC) ligands.¹⁸ Also complexes bearing cyclometallated pincer bis(carbene) ligands with phenylene linkers (^RCCC^R)¹⁹ or pyridyl linkers (^RCNC^R)²⁰ exhibit blue emission along with vapochromic or aquachromic responses. Few rare examples of green-blue emitting heteroleptic Pt-NHC complexes that feature cyclometalation via the N-wingtip are known.²¹ Iridium(III) or platinum(II) based phosphors derived from $P^{\wedge}N^{22}$ or $P^{\wedge}O^{23}$ donor sets also emit blue light.²² In the search for neutral platinum-based phosphors with green-blue emission, ligands bearing ring-fluorinated aromatic moieties should be avoided since they can easily degrade, and problems also arise when readily oxidizable phosphine donors are employed. Thus we have developed a mono-anionic bidentate NHC^pyrazolate ligand system to coordinate the platinum(II) center via a C^N donor set. By combining suitably functionalized phenyl-pyridine/pyridinetriazole luminophores with a NHC^pyrazolate ligand, four new heteroleptic platinum phosphors were synthesized. The structural, photophysical and electrochemical characteristics of these complexes are evaluated along with TD-DFT calculations.

Results and discussion

Synthesis and structural characterization

The ligand precursor (NHC^pyrazolate) H_2 ·Br (7) was obtained by reacting stoichiometric amounts of 1-methylbenzimidazole and the bromomethyl pyrazole derivative (6). Compound 6 was previously synthesized by a modified five step procedure (ESI[†]).²⁴ Schematically, the reaction of an adequate diketone derivative (1) and hydrazine hydrate in ethanol yielded an unsymmetrically substituted pyrazole (2). Oxidation of the methyl function of the pyrazole derivative into the respective carboxylic acid (3) was achieved *via* a standard procedure using KMnO₄ as the oxidant. Esterification of the carboxylic acid using H_2SO_4 -methanol resulted in compound 4. The ester derivative was further transformed into the corresponding alcohol (5) by reduction with LiAlH₄ in tetrahydrofuran (thf), which was subsequently brominated using an excess of PBr₃ with the formation of (6) (Scheme 1).²²

The phenyl-pyridine ligand precursors were obtained *via* the Suzuki coupling of the corresponding aryl-boronic acids and 2-bromopyridine in the presence of catalytic amounts of $[Pd(PPh_3)_4]$.²⁵ The pyridine-triazole ligand precursor (8) was obtained *via* the modification of our previous protocol in which a bulky adamantyl group replaces the fluorinated aryl substituent.²⁶ Adequate cyclometallated phenyl-pyridine-Pt(II) precursors were synthesized according to literature procedures.²⁷ Alternatively, the (pyridine-triazole)–Pt(II) precursor [12] was obtained by heating equal stoichiometric amounts of compounds 8 and $[K_2PtCl_4]$ in a 2-ethoxyethanol-water mixture for 12 h at 85 °C.²⁶ Heteroleptic Pt(II) complexes [9], [10], [11] and [13] were obtained by reacting the ligand precur-



sor 7 and the corresponding (μ -Cl) cyclometallated Pt(π) precursor or the (pyridine-triazole)–Pt(π) precursor [12] under reflux in methanol using triethylamine as the base (Scheme 2).

All the complexes have been characterized using ¹H NMR, ¹³C NMR, ¹⁹F NMR, and high resolution mass spectrometry. In addition X-ray structural determination of complexes [9], [10], and [11] was performed on single crystals (see below). The 1 H NMR spectra of complexes [9], [10], [11] and [13] were lacking the characteristic benzimidazolium-C2-H resonances, along with the absence of pyrazole N-H resonances that were observed in the free ligand 7. In addition, chemically and magnetically equivalent CH₂ protons of the methylene bridges that link the pyrazole ring and the NHC precursor of the uncoordinated-bidentate ligand 7, were rendered diastereotopic upon complexation, due to the formation of a rigid metallocycle. The ¹H NMR signals of these diastereotopic protons were observed as two singlets at δ 5.43 and 5.30 ppm in [9], δ 5.41 and 5.39 ppm in [10] and δ 5.48 and 5.41 ppm in [11]. However only a doublet was observed at δ 5.35 ppm (J = 2.2 Hz) for [13]. The pyridine– $\alpha_{\rm H}$ protons of the phenyl pyridine or triazole pyridine co-ligands also tend to be more acidic in all complexes, as confirmed by the down-field resonances observed at δ 10.01 and 10.22 ppm in complexes [9] and [10] respectively, and at δ 9.73 ppm in [11] and [13]. Formation of the complexes was also confirmed by ¹³C NMR spectra, in which the carbenoid carbon resonances appeared at δ 168.7, 166.5, 166.0 and 174.3 ppm for the complexes [9], [10], [11] and [13], respectively. ¹³C NMR resonances of the methylene bridge $(-CH_2-)$ that links the pyrazole ring and the NHC donor of the coordinated NHC[^]pyrazolate ligand, were observed in the range of δ 42.6–44.3 ppm in all of the complexes. In addition, ¹⁹F NMR resonances of the pyrazole-CF3 substituent were observed at δ -60.52 and -60.89 ppm in complexes [9] and



Scheme 2 Synthesis of Pt(II)-NHC complexes bearing phenyl-pyridine/triazole-pyridine chromophores.



Fig. 1 Structures obtained by X-ray diffractometry of [9] (a), [10] (b), and [11] (c) shown as 50% displacement ellipsoids (hydrogen atoms omitted for clarity).

[13], respectively. Complexes, [10] and [11] bearing two additional CF₃ substituents on the phenyl ring, revealed in the ¹⁹F NMR spectra three resonances at δ –57.36, –60.49, –63.16 ppm and δ –58.63, –60.61, –62.85 ppm, respectively.

Single crystals of complexes [9], [10] and [11] were obtained by the slow diffusion of hexane vapors into their dichloromethane solutions at ambient temperature. The molecular structures of [9], [10] and [11] are shown in Fig. 1 and selected bonding parameters are listed in Table 1.

The phenyl-pyridine–N and the carbenoid carbon of [9] adopt a *cis*-configuration, as opposed to [10] and [11] that display a *trans* configuration in order to avoid a steric repulsion between the CF₃ groups of the pyrazole ring and the

 Table 1
 Some relevant bonding parameters of complexes [9], [10] and [11]

Bond lengths (Å)/angles (°)	[9]	[10]	[11]
Pt-C _{NHC}	2.045(8)	1.981(3)	1.967(4)
Pt-C _{Ph}	2.054(8)	1.991(3)	2.028(4)
Pt-N _{Pyrazole}	2.021(7)	2.100(3)	2.073(3)
Pt-N _{Pvridine}	2.026(7)	2.061(3)	2.073(3)
C _{NHC} -Pt-N _{Pyrazole}	84.8(3)	85.04(13)	83.37(15)
C _{NHC} -Pt-C _{Ph}	173.0(3)	96.82(14)	100.05(16)
C _{Ph} -Pt-N _{Pvrazole}	97.9(3)	175.05(13)	175.08(15)
N _{Pvridine} -Pt-N _{Pvrazole}	171.0(3)	98.62(12)	96.33(15)
N _{Pvridine} -Pt-C _{Ph}	81.0(3)	79.86(13)	80.78(16)
(N–C–N)NHC	107.0(7)	107.0(3)	106.6(4)

phenyl-pyridine-ligand. Due to the resulting *trans* labilizing effect of NHC ligands, the Pt–N_{pyridine} bond distances in [10] and [11] are elongated as compared to that of [9]. The longer Pt–C_{NHC} bond distance observed in [9] (2.045(8) Å), as compared to those of [10] (1.981(4) Å) or [11] (1.967(4) Å), can be attributed to the *trans* disposed phenyl-*C* donor.

The bite angles of the bidentate NHC^pyrazolate ligands coordinated to the platinum centers (84.8(3)°, 85.04(13)° and 83.37(15)° in [9], [10] and [11], respectively) indicate a distorted square-planar configuration. Also in [9], the *trans* angles between the pyridyl fragment of the phenyl pyridine ligand and the pyrazole fragment of the NHC-pyrazolate ligand clearly deviate from linearity (171.0(3)°). Similarly, in [10] and [11] the C_{ph} and the N_{pyrazole} donor atoms occurring in a *trans* configuration also feature platinum-centered *trans*-bond angles (C_{Ph} -Pt-N_{pyrazole}) of 175.05(13)° and 175.08(15)°, respectively. The inter-ligand bond angles (N_{pyrazole}-Pt-N_{pyridine}) in [10] and [11] were 98.62(12)° and 96.33(15)°, respectively.

Complex [13] also adopts a *trans* configuration between the $N_{pyridyl}$ atom and the C_{NHC} atom of the two bidentate ligands in order to avoid the steric repulsion between the bulky adamantyl group and the CF₃ substituent. As single crystals suitable for X-ray studies could not be obtained for complex [13], the bond connectivity and the geometry were established with the aid of 2D NMR spectroscopy (HMBC, HSQC and NOESY). The absence of polarization transfer within the NHC methyl wingtip and the pyridyl- $\alpha_{\rm H}$ protons of complex [13] in the NOESY spectra confirms a *trans* configuration between these fragments.

Absorption spectroscopy and frontier orbitals

Absorption spectra of all complexes were measured in dichloromethane at ambient temperature and are shown in Fig. 2. In the figure are also shown the relevant electronic transitions calculated using time dependent density functional theory (TD-DFT). In addition quantum-chemical calculations were performed to predict the nature and energies of the main transitions. Isodensity surface plots of the highest occupied (HOMOs) and lowest unoccupied molecular orbitals (LUMOs), along with the energies of selected molecular frontier orbitals, are listed in Tables S1 and S2 (ESI†). The computed vertical transitions under vacuum, their oscillator strengths and the expansion coefficients of the corresponding monoelectronic excitations are listed in Table S3.†

TD-DFT calculations reveal that the $S_0 \rightarrow S_1$ transitions for all complexes can be mainly described as HOMO \rightarrow LUMO excitations, whereas higher energy transitions involve admixtures of HOMO–*n* and LUMO+*m* excitations. Complex [9] exhibits a platinum-centred HOMO orbital with significant contributions from the phenyl fragment of the cyclometalated-C[^]N ligand, whereas the LUMO (π^*) appears largely delocalized over the pyridyl fragment of the cyclometalated C[^]N chelate with a small contribution from the phenyl ring as well as from the metal center. Hence, we reasonably ascribe the relatively weak band at 382 nm (HOMO \rightarrow LUMO excitation) to an electronic transition with a mixed intraligand charge-transfer (¹ILCT) and metal-to-ligand charge-transfer (¹MLCT) charac-



Fig. 2 Absorption spectra and calculated transitions for complexes [9], [10], [11], [13] depicted in (a), (b), (c), and (d), respectively. The dashed lines represent the spin-forbidden S_0-T_1 transitions.

ter.²⁸ The moderately intense absorption peak at a relatively high energy (338 nm) is due to a HOMO–1 \rightarrow LUMO excitation, whereas the highest intensity absorption peak at 282 nm is due to an admixture of HOMO–5 \rightarrow LUMO and HOMO–1 \rightarrow LUMO+1 excitations. The orbital diagrams of both HOMO–1 and HOMO–5 orbitals show electron delocalization over the phenyl ring and the pyrazole/benzimidazole fragments of the heteroleptic complex. As a sizeable participation of platinum-d π orbitals is also observed in the HOMO –5, HOMO–1 and LUMO orbitals, we ascribe these electronic transitions peaking at 338 nm and 282 nm to mixed metal-

Table 2Electrochemical data of complexes [9], [10], [11], and [13]. Thepotentials are given vs. SCE (saturated calomel electrode) a

Complex	$E_{\rm pa} \left({\rm HOMO}^b \right)$	$E_{\rm pc}$ (LUMO ^b)	$\Delta E_{\mathrm{HOMO-LUMO}}\left(\mathrm{eV}\right)$
[9]	1.33 V (-5.92 eV)	-2.12 V (-2.97 eV)	2.95
[10]	1.64 V (-6.17 eV)	-1.62 V (-3.31 eV)	2.86
[11]	1.69 V (-6.29 eV)	-1.91 V (-3.05 eV)	3.24
[13]	1.65 V (-6.31 eV)	-1.97 V (-3.10 eV)	3.21

^{*a*} 1 mM of each complex in a dichloromethane solution in the presence of 0.1 M TBAPF₆, glassy carbon 3 mm, $\nu = 0.1 \text{ V s}^{-1}$. ^{*b*} The HOMO and LUMO energies were calculated using the relation: $E_{\text{HOMO/LUMO}} = -(E_{\text{onset ox./red.}} \nu s. \text{ Fc/Fc}^+ + 4.8) (eV).^{29}$

perturbed intra-ligand (¹ILCT)/ligand-to-ligand charge transfer (¹LLCT) transitions.

The frontier orbitals of complexes [10] and [11] display similar iso-surfaces. Analogously to [9], the LUMO remains primarily localized on the pyridyl fragment. However, the HOMO is spread over the phenyl rings of the cyclometalated C^N ligand and the pyrazole fragment of the chelating NHC[^]pyrazolate ligand. The orbital analysis indicates that the low energy absorption bands at 408 nm for [10] and at 381 nm for [11] can be ascribed to HOMO \rightarrow LUMO transitions, whereas the intense absorption bands at higher energies (around 300 nm) are assigned to mixed metal-perturbed ¹ILCT/¹LLCT transitions arising from HOMO-4 \rightarrow LUMO for [10] and HOMO-1 \rightarrow LUMO/HOMO \rightarrow LUMO+1 excitations for [11].

The HOMO of complex [13] is mainly localized on the triazole moiety. Analogously to the other complexes described herein, the LUMO appears pyridine-centered (π^*), but with a significant contribution from the Pt-d π -orbitals. The lowenergy absorption band peaking at 369 nm involves a HOMO \rightarrow LUMO excitation and is ascribed to a mixed ¹MLCT/¹ILCT transition. The absorption bands at 306 nm and 282 nm are due to the HOMO-1 \rightarrow LUMO and HOMO \rightarrow LUMO+2 excitations, respectively, and can be ascribed to metal-perturbed ¹LLCT transitions.



Fig. 3 Emission spectra of complexes [9] (a), [10] (b), [11] (c), [13] (d) at ambient temperature (solid lines) and 77 K (dashed lines). Excitation wavelengths: 357 nm, 354 nm, 336 nm and 308 nm, respectively.

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All of the complexes were further characterized by cyclic voltammetry (CV) and differential pulse voltammetry (DPV) (Table 2 and Fig. S6[†]). Each compound displays an irreversible oxidation wave in the range between +1.33 and +1.69 V that can be attributed to HOMO-centred oxidation processes that clearly involve the metal centre. In particular a relatively facile oxidative process was observed for [9] (+1.33 V), in comparison to those of [10] (+1.64 V) and [11] (+1.69 V). The observed positive shift in the oxidation potential of [10] and [11] can be attributed to the presence of two additional electron withdrawing CF₃ groups in the bidentate ligand. In the range of -2.12 V to -1.62 V, complexes [9], [11] and [13] display irreversible reduction processes, whereas complex [10] reveals a quasireversible behaviour ($\Delta E_p = 0.11$ mV). Such reduction processes are pyridine-centered,^{10b} as also suggested by the calculated unoccupied frontier orbitals. The reduction potentials of [10] and [11] are, as expected, less negative than that of [9], due to the presence of the electron withdrawing CF₃ groups on the ligands and are -1.62 V and -1.91 V, respectively (see Table 2). However, it is clear that the emissive excited states



Fig. 4 Emission spectra of [9] (solid line), [10] (long dashed line), [11] (dotted line) and [13] (short dashed) in 10 wt% PMMA films. Excitation wavelengths: 357 nm, 354 nm, 336 nm and 308 nm, respectively.

Table 3	Photophy	sical data	in	solution
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cannot be described as pure HOMO-LUMO excitations, as also indicated by the TD-DFT calculations (*vide supra*).

Photophysical characterization

Emission spectra of complexes [9], [10], [11], [13] at ambient temperature and at 77 K are shown in Fig. 3 (the corresponding excitation spectra are shown in Fig. S1–S4†). Thin film emission spectra of [9], [10], [11], [13] are shown in Fig. 4 (10 wt% doped into PMMA) and Fig. S5† (neat films). Table 3 summarizes all relevant photophysical data in solution and in glassy matrices at 77 K, whereas Table 4 includes the data for the films.

In fluid dichloromethane at room temperature, [9] and [11] display comparable luminescence spectra. Complex [9] emits with a clear vibrational progression typical for predominantly ligand-centered states, with a maximum at 487 nm. The emission profile of complex [11] has a less pronounced band at 489 nm and a maximum at 519 nm. A remarkable characteristic of complex [10] is its featureless and broad emission, which appears with a large bathochromic shift as compared to [9] and [11]. This smaller energy gap of [10], if compared with complex [11], is a result of the CF₃-substitution pattern, as the *m*,*m*'-substitution lowers the π^* -orbital energy of the C^NN ligand to a larger extent than for the *o*,*p*-pattern. Indeed, the calculated MO energies reveal that the position of the CF3 substituents mostly affects the LUMO without significantly altering the HOMO of the complexes. The bluest emission is observed for complex [13] bearing the pyridine-triazole luminophore. Its emission spectrum in dichloromethane peaks at 476 nm and at 505 nm. The sensitivity of the long excited state lifetimes towards oxygen is indicative of emissions from the triplet manifolds.

In glassy matrices of 2-MeTHF at 77 K the emission spectra of [9] and [11] display enhanced vibrational progressions which are indicative of a predominant LC character. In contrast, the emission spectrum of [10] remains essentially unaffected. A hypsochromic shift with a pronounced vibrational progression was observed for complex [13] at 77 K,

Complex	$ \begin{array}{c} \lambda_{\rm abs} \stackrel{a,b}{=} [{\rm nm}] \\ (\varepsilon \times 10^{-3} / \\ [{\rm M}^{-1} \ {\rm cm}^{-1}]) \end{array} $	$\lambda_{\mathrm{em}}{}^{a,c}$ (rt) [nm]	$\lambda_{\mathrm{em}}{}^{d}$ (77 K) [nm]	$\Phi_{ m em}{}^{a,e}$ (×100, aer.)	$\Phi_{ m em}{}^{a,e}$ (×100, deaer.)	$ au^{a}$ (rt, aer.) [µs]	$ au^{a}$ (rt, deaer.) [µs]	τ ^c (77 K) [μs]	$\begin{array}{c} k_{\rm r} \times 10^{-4} \\ [\rm s^{-1}] \end{array}$	$\frac{k_{\rm nr} \times 10^{-5}}{[\rm s^{-1}]}$
[9]	282 (28.4) 338 (7.5) 282 (2.7)	487 518	482 518	0.2	0.7	0.004 (33%) 0.070 (67%)	0.9 (45%) 8.2 (55%)	12.2	0.14	2.0
[10]	382(2.7) 288(29.5) 354(6.1) 400(2.7)	547	508 547	1.0	1.5	0.41	1.0 (77%) 12.4 (23%)	11.9	0.41	2.7
[11]	408 (2.7) 286 (22.6) 338 (8.2) 281 (2.1)	489 519	481 515	5.0	8.0	0.1	1.3 (70%) 16.4 (30%)	20.7	1.4	1.6
[13]	381 (2.1) 282 (11.3) 306 (8.4) 369 (1.8)	476 505	461 492	1.5	7.2	0.36	2.6	19.8	2.7	3.5

^{*a*} In a dichloromethane solution. ^{*b*} The abbreviation "sh" denotes a shoulder. ^{*c*} λ_{exc} = 282 nm ([9]), 354 nm ([10]), 284 nm ([11]), 285 nm [13]. ^{*d*} In a 2-MeTHF glassy matrix. ^{*e*} Emission quantum yields were measured with an integrating sphere.

Table 4 Photophysical data in films

Complex	$\lambda_{\mathrm{em}}{}^{a,b}$ (neat film) [nm]	λ _{em} ^{<i>a</i>} (10 wt% in PMMA) [nm]	$\Phi_{ m em}{}^{a,c}$ (×100, neat film)	$\Phi_{ m em}{}^{a,c}$ (×100, 10 wt% in PMMA)	τ (neat film) [µs]	τ (10 wt% in PMMA) [µs]	$k_{\rm r} \times 10^{-4} d [{ m s}^{-1}]$	$\frac{k_{\rm nr} \times 10^{-4 d}}{\left[{\rm s}^{-1}\right]}$
[9]	491 527	489 524	3.4	34.0	0.7 (65%) 12.8 (35%)	14.1	2.4	4.6
[10]	518 543	509 537	10.0	26.7	9.9	4.0 (38%) 15.9 (63%)	2.3	6.4
[11]	494 521	490 518	3.2	10.0	2.6(41%) 11.1(59%)	3.1 (16%) 16.1 (84%)	0.7	6.4
[13]	474 501	470 498 537 (sh)	11.0	49.0	3.7 (54%) 14.6 (46%)	18.8	2.6	2.7

 $^{a}\lambda_{\text{exc}}$ = 357 nm ([9]), 354 nm ([10]), 336 nm ([11]), 308 nm [13]. b The abbreviation "sh" denotes a shoulder. c Quantum yields were measured in an integrating sphere system. d 10 wt% in PMMA.

and is most likely caused by a loss of MLCT character in a frozen environment.

The photoluminescence quantum yields in solutions are relatively low for all complexes, even under deaerated conditions, which is due to the solvent-related vibronic deactivation of the excited states. Indeed, a significant enhancement of the solid state PLQY is observed in 10 wt% doped PMMA films, where complex [13] reaches up to 49% (Table 4).

Conclusions

In summary, we have synthesized a new bidentate NHC^pyrazolate ligand precursor featuring a methylene spacer between the benzimidazol-2-ylidene and the pyrazolate moieties. It was used to obtain four new luminescent platinum(II) complexes along with either phenyl-pyridine or pyridine-triazole luminophores. A full structural, electrochemical and photophysical characterization was carried out, and the results were interpreted with the aid of TD-DFT calculations. The emission of the complexes was in the sky blue/green region of the electromagnetic spectrum. Amongst all the complexes reported herein, the one featuring a pyridine-triazole luminophore along with the bidentate NHC^pyrazolate ligand exhibited a more blue phosphorescence with a high photoluminescence quantum yield in the solid state. An additional tweaking of the pyridine-triazole chromophore (for instance, replacing the adamantyl moiety by a CF₃ group) could further shift the emission into the blue region, as the HOMO would be stabilized and the HOMO-LUMO gap would be increased, along with the energy of the emissive triplet state. Thus, the new bidentate building block introduced herein constitutes a convenient tool in the design of phosphorescent metal complexes for optoelectronic applications if combined with a suitable luminophore.

Experimental

Synthetic methods and characterization

All reagents were of analytical grade and used as received. Solvents were purified according to standard procedures. All reactions were performed under an atmosphere of nitrogen using standard Schlenk techniques unless specified otherwise. Column chromatography was performed using silica gel 60 (particle size 63–200 mm or 230–400 mesh, Merck). NMR spectra were recorded either on a Bruker AV 300/400 or an Agilent DD2 600 MHZ spectrometer. All chemical shifts are given in ppm and referenced to the residual solvent peaks. ¹⁹F NMR are referenced to CFCl₃ (0.00 ppm) as an internal standard. The signals are abbreviated as follows: s, singlet; d, doublet; t, triplet; q, quartet; m, multiplet. All coupling constants (*J*) are given in hertz (Hz). High resolution mass spectra (HRMS) of all the complexes were measured at the department of Organic Chemistry, University of Muenster in a Bruker Daltronics or Micro TOF with loop injection. Elemental analyses were recorded at the University of Münster, Germany or at the University of Milano, Italy.

Photophyscis

Absorption spectra were recorded on a Varian Cary 5000 double-beam UV-Vis-NIR spectrometer and the baseline corrected. Steady-state emission spectra were recorded on a Horiba Jobin-Yvon IBH FL-322 Fluorolog 3 spectrometer equipped with a 450 W xenon-arc lamp, double grating excitation, and emission monochromator (2.1 nm mm⁻¹ of dispersion), and a Hamamatsu R928 photomultiplier tube or a TBX-4-X single photon-counting detector. Emission and excitation spectra were corrected for source intensity (lamp and grating) by using standard correction curves. Time-resolved measurements were performed using the time-correlated single-photon counting (TCSPC) option on the Fluorolog 3 equipment. NanoLEDs (295 nm; full width at half maximum, 1 ns) with repetition rates between 10 kHz and 1 MHz were used to excite the sample. The excitation sources were mounted directly on the sample chamber at 90° to a doublegrating emission monochromator (2.1 nm mm^{-1} of dispersion, 1200 per groove) and collected by using a TBX-4-X single photon counting detector. The photons collected at the detector were correlated by means of time to amplitude converter to excitation pulse. Signals were collected with the aid of an IBH Data-station Hub photon counting module and data analysis was performed by using DAS6 program (Horiba Jobin- Yvon IBH). The quality of the fit was assessed by minimizing the reduced χ^2 function and visually inspecting the weighed

residuals. Luminescence quantum yields were measured with a Hamamatsu Photonics absolute PL quantum yield measurement system (C9920-02) equipped with a L9799-01 CW Xenon light source (150 W), monochromator, C7473 photonic multichannel analyser, integrating sphere and employing U6039-05 PLQY measurement software (Hamamatsu Photonics Ltd, Shizuoka, Japan). All solvents used were of spectrometric grade. Deaerated samples were prepared by the freeze–pump–thawtechnique.

Computational details

Geometries were optimized by means of density functional theory (DFT). The parameter-free hybrid functional Perdew-Burke–Ernzerhof³⁰ was employed along with the standard valence double- ζ polarized basis set 6-31G(d,p)³¹ for C, H, F and N. For Pt, the Stuttgart_Dresden (SDD) effective core potentials were employed along with the corresponding valence triple- ζ basis set. No imaginary frequency was obtained (NImag = 0). Simulated electronic absorption spectra were recorded on the optimized geometry at S₀ by means of time-dependent density functional theory (TD-DFT) calculations.³² All the calculations were performed under vacuum conditions using the Gaussian 09 program package.³³

Cyclic voltammetry

All photoactive complexes reported herein are characterized by cyclic voltammetry (CV) and differential pulse voltammetry (DPV) in a dichloromethane solution and in the presence of tetra-^{*n*} butylammonium hexafluorophosphate (0.1 M TBAPF₆) as a supporting electrolyte. The concentration of all the samples was 1 mM in dichloromethane (Sigma-Aldrich, Chromasolv® Plus, 99.9%). TBAPF₆ (electrochemical grade, 99%, Fluka) was used as the supporting electrolyte. The solvent and the supporting electrolyte were used as received without any further purification. For the electrochemical experiments, a CHI750C electrochemical workstation (CH Instruments, Inc., Austin, TX) was used. Electrochemical experiments were performed in a glass cell under an argon atmosphere. To minimize the Ohmic drop between the working and the reference electrodes, a feedback correction was employed. A 3 mm diameter glassy carbon disk electrode (CH Instruments, Inc., Austin, TX) was employed as the working electrode. The working electrode was polished with a 0.05 µm diamond suspension (Metadi Supreme Diamond Suspension, Buehler) before each measurement. A platinum wire served as the counter electrode, while a silver wire was employed as the quasi-reference electrode (Ag-QRE), which was separated from the cathode by a glass frit (Vycor). The reference electrode was calibrated at the end of each experiment against the ferrocene/ ferricenium couple (Fc/Fc⁺), whose formal potential against the saturated calomel electrode (SCE) was 0.46 V in dichloromethane; all potential values are reported against the SCE.

X-Ray diffraction

Data sets were collected with a Nonius KappaCCD diffractometer. Programs used: data collection, COLLECT (Nonius B.V., 1998); data reduction Denzo-SMN;³⁴ absorption correction;³⁵ structure solution SHELXS-97;³⁶ structure refinement SHELXL-97³⁷ and graphics, XP (BrukerAXS, 2000). Thermal ellipsoids are shown with 50% probability. *R*-Values are given for the observed reflections, and wR^2 values are given for all reflections (see ESI† for crystallographic data).

Syntheses

Compound 7. 1-Methyl benzimidazole (235.1)mg, 1.78 mmol) was added to a solution of compound 6 (408 mg, 1.78 mmol) in dichloromethane (20 mL) and heated under reflux for 4 h. The precipitation of a colourless solid was observed upon completion of this time. The dichloromethane solution was further concentrated to half the volume ($\approx 10 \text{ mL}$) to ensure complete precipitation. The resulting precipitate was filtered and washed again with dry diethyl ether (20 mL \times 3) using a filter-cannula under an atmosphere of N₂ to avoid contact with moisture. Upon drying the colourless solids in vacuo, an analytically pure form of the ligand precursor 7 was isolated as a hygroscopic powder. Yield: 686 mg (88.2%). ¹H NMR (300 MHz, DMSO- d_6): δ 14.33 (s, 1H, Pyrazole NH), 9.93 (s, 1H, C_2 -H), 8.05 (d, J = 3.66 Hz, 2H, benzimidazole-H), 7.70 (d, J = 3.66 Hz, 2H, benzimidazole-H), 6.94 (s, 1H, pyrazole-H), 5.95 (s, 2H, methylene bridge), 4.11 (s, 3H, CH_3). ¹³C ${}^{1}H$ NMR (101 MHz, CD₂Cl₂): δ 154.3, 143.3, 137.0, 131.9, 130.5, 127.4, 126.8, 126.6, 113.8, 113.3, 104.3, 48.3 (-(CH₂)-), 33.5 (CH₃). ¹⁹F NMR (282 MHz, DMSO- d_6): δ -60.42. MS (ESI HRMS) m/z (%): 281.1010 (100) [7]⁺, (calcd for [7]⁺ 281.1009).

Compound 8 (2-(3-(adamantan-1-yl)-1*H*-1,2,4-triazol-5-yl)**pyridine**). This ligand precursor was prepared according to a previously known procedure using adamantanecarbonylchloride.²⁶ Yield: 1.3 g (91.0%). ¹H NMR (300 MHz, CDCl₃) δ 8.69 (d, *J* = 4.0 Hz, 1H), 8.20 (d, *J* = 7.9 Hz, 1H), 7.81 (t, *J* = 7.0 Hz, 1H), 7.41–7.29 (m, 1H), 2.12 (s, 9H), 1.81 (s, 6H). MS (ESI HRMS) (*m*/*z*): 281.1762 [8 + H]⁺ (calcd for [8 + H]⁺ 281.1761), 303.1580 [8 + Na]⁺ (calcd for [8 + Na]⁺ 303.1580).

Compound [9] (R = H). A mixture of the ligand precursor 7 (100 mg, 0.27 mmol), platinum(II)-phenylpyridine dimer (R = H) (106 mg, 0.14 mmol) and triethylamine (109 mg, 1.08 mmol) was dissolved in methanol (10 mL) in a flamedried Schlenk flask. The reaction mixture was heated under refluxion for 6 h. After this time, the solvent was removed in vacuo and the product was suspended in dichloromethane and washed with water (1:1, v:v). The organic phase was separated and dried over MgSO4 and dichloromethane was removed in vacuo. A further purification of the crude product was done on a silica gel column using a mixture of dichloromethane-methanol (10:1, v:v) as an eluent to obtain [9] as pale yellow powder. Yield: 46 mg (27.1%). ¹H NMR (300 MHz, CD_2Cl_2): δ 10.01 (ddd, J = 5.8 Hz, 1.6 Hz, 0.8 Hz, 1H), 8.53 (m, 1H), 8.31 (m, 1H), 7.86 (m, 2H), 7.65 (dd, J = 7.7 Hz, 1.2 Hz, 1H), 7.60 (m, 1H), 7.44 (m, 2H), 7.34 (td, J = 7.4 Hz, 1.3 Hz, 1H), 7.18 (td, J = 7.4 Hz, 1.4 Hz, 1H), 7.01 (ddd, J = 7.4 Hz, 5.9 Hz, 1.7 Hz, 1H), 6.49 (s, 1H), 5.43 (s, 1H), 5.30 (s, 1H), 3.91 (s, 3H). ¹³C{¹H} NMR (101 MHz, CD₂Cl₂): δ 168.7, 157.7, 151.9, 150.2, 146.8, 142.9, 139.9, 138.4, 137.2, 135.6, 130.4, 129.3,

127.3, 124.2, 123.9, 123.0, 122.7, 120.9, 119.0, 111.6, 110.5, 101.4 (d, J = 2.1 Hz), 44.3 (-(CH_2)-), 36.4 (CH_3). ¹⁹F NMR (282 MHz, CD_2Cl_2): δ -60.52. MS (ESI HRMS) m/z (%): 629.12201 (100) [9 + H]⁺ (calcd for [9 + H]⁺ 629.1235). Anal. Calcd (for $C_{24}H_{19}F_3N_5Pt\cdotH_2O\cdothexanes$): C, 49.31; H, 4.41; N, 9.58. Found: C, 49.56; H, 4.48; N, 9.65.

Compound [10] ($R = o_{,p}$ -CF₃). A mixture of the ligand precursor 7 (100 mg, 0.27 mmol), platinum(II)-phenyl pyridine dimer (R = o, p-CF₃) (145 mg, 0.14 mmol) and triethylamine (109 mg, 1.08 mmol) was dissolved in methanol (10 mL) in a flame-dried Schlenk flask and heated under refluxion for 6 h. The subsequent work-up procedure was analogous to that of compound [9]. A further purification of the crude product was done by silica gel column chromatography using a mixture of dichloromethane-methanol (v:v, 10:0.5) to obtain [10] a crystalline yellow solid. Yield: 92 mg (44.4%). ¹H NMR (400 MHz, CD_2Cl_2): δ 10.22 (ddd, J = 5.7 Hz, 1.7 Hz, 0.7 Hz, 1H), 8.29 (d, J = 8.5 Hz, 1H), 8.06 (ddd, J = 8.5 Hz, 7.5 Hz, 1.8 Hz, 1H), 7.79 (m, 1H), 7.65 (m, 2H), 7.53 (m, 2H), 7.45 (m, 2H), 6.57 (s, 1H), 5.41 (s, 1H), 5.39 (s, 1H), 3.93 (s, 3H). ¹³C{¹H} NMR (101 MHz, CD_2Cl_2) δ 166.5 (C_{NHC}), 161.4, 154.0, 152.0, 150.2, 148.6, 142.1, 141.7, 138.8, 136.8, 136.3, 135.4, 134.1, 132.8, 131.8, 128.7, 123.5, 123.4 (d, J = 10.4 Hz), 123.1, 122.4, 118.9, 110.4, 109.4, 100.4 (d, J = 1.8 Hz), 43.0 (-(CH_2)-), 34.8 (CH_3). ¹⁹F NMR (282 MHz, CD_2Cl_2): δ –57.36 (s, satellites were observed with J = 13.0 Hz), -60.49 (s, satellites were observed with J = 36.0 Hz), -63.16 (s). MS (ESI HRMS) m/z (%): 765.0988 (100) $[10 + H]^+$ (calcd for $[10 + H]^+$ 765.0985). Anal. Calcd (for $C_{26}H_{16}F_9N_5Pt$): C, 40.84; H, 2.11; N, 9.16. Found: C, 40.79; H, 2.26; N, 9.23.

Compound [11] (R = mm' CF₃). A mixture of the ligand precursor 7 (100 mg, 0.27 mmol), platinum(II)-phenylpyridine dimer (R = m, m'-CF₃) (145 mg, 0.14 mmol) and triethylamine (109 mg, 1.08 mmol) was dissolved in methanol (10 mL) in a flame-dried Schlenk flask and heated under refluxion for 6 h. A work-up procedure analogous to that of compound [9] was followed to obtain a crude sample. A further purification of the crude sample was done on a silica gel column using dichloromethane-methanol (10:0.5, v:v) as the eluent to obtain [11] as a pale yellow powder. Yield: 52 mg (25.2%). ¹H NMR (300 MHz, CD_2Cl_2): δ 9.73 (ddd, J = 5.7 Hz, 1.6 Hz, 0.8 Hz, 1H), 8.08 (t, J = 1.2 Hz, 1H), 8.00 (dd, J = 1.5 Hz, 0.8 Hz, 1H), 7.70 (s, 1H), 7.61 (d, J = 8.02 Hz, 1H), 7.47 (ddd, J = 7.3, 5.7, 1.5 Hz, 1H), 7.37 (m, 4H), 7.35 (m, 1H), 6.56 (s, 1H), 5.48 (s, 1H), 5.41 (s, 1H), 3.56 (s, 3H). ${}^{13}C{}^{1}H{}$ NMR (75 MHz, CD_2Cl_2): δ 166.0 (C_{NHC}), 164.5, 160.7, 152.4, 150.3, 143.6, 140.2, 138.0, 137.6, 135.0, 134.4, 133.9, 130.2, 126.9, 126.4, 124.3, 124.0, 123.6, 122.4, 121.8, 119.8, 113.3, 110.6, 101.6 (d, J = 2.1 Hz, pyrazole-C), 44.2 (-(CH₂)-), 35.4 (CH₃). ¹⁹F NMR (282 MHz, CD₂Cl₂): δ -62.85, -60.61, -58.63. MS (ESI HRMS) m/z (%): 765.0992 (100) $[11 + H]^+$ (calcd for $[11 + H]^+$ 765.0985). Anal. Calcd (for C₂₆H₁₆F₉N₅Pt): C, 40.84; H, 2.11; N, 9.16. Found: C, 40.57; H, 2.47; N, 8.33.

Compound [12] (potassium dichloro(pyridine-triazole^{Ad})platinate). To a solution of 2-(3-(adamantan-1-yl)-1*H*-1,2,4triazol-5-yl)pyridine (140 mg, 0.5 mmol) in methoxyethanol (25 mL) was added a solution of K_2 PtCl₄ (208 mg, 0.5 mmol) in water (10 mL) and the reaction mixture was stirred at 85 °C overnight. After cooling the reaction mixture to ambient temperature, the yellow precipitate was filtered off, washed with water, ethanol and diethylether and finally dried *in vacuo*. Yield: 185 mg (68.0%). ¹H NMR (300 MHz, DMF-*d*7) δ 9.55 (d, *J* = 5.8 Hz, 1H), 8.47 (t, *J* = 7.8 Hz, 1H), 8.25 (d, *J* = 7.9 Hz, 1H), 7.87 (td, *J* = 6.0 Hz, 2.9 Hz, 1H), 2.14 (s, 9H), 1.80 (s, 6H). ¹³C {¹H} NMR spectra of this precursor complex could not be obtained due to its poor solubility in DMF-*d*₇. MS (ESI HRMS, negative mode) *m*/*z* (%): 544.0641 (100) [**12**]⁻ (calcd for [**12**]⁻ 544.0646). Anal. Calcd (for C₂₄H₁₉F₃N₅Pt): C, 34.94; H, 3.28; N, 9.59. Found: C, 34.68; H, 3.37; N, 9.63.

Compound [13]. A mixture of ligand precursor 7 (60 mg, 0.16 mmol), potassium dichloro(pyridine-triazole^{Ad})platinate (97 mg, 0.16 mmol) and triethylamine (100 mg, 0.98 mmol) in methanol (10 mL) in a flame-dried Schlenk flask was heated under refluxion for 6 h. After completion of the reaction, the solvents were dried in vacuo and the product was suspended in dichloromethane and washed with water (1:1, v:v). The organic phase was separated and dried over MgSO4 and dichloromethane was removed in vacuo. A further purification was done using a preparative TLC with a mixture of dichloromethane-hexane (10:2, v:v) as the eluent to obtain [13] as a colourless powder. Yield: 28 mg (23.1%). ¹H NMR (600 MHz, CD_2Cl_2 : δ 9.73 (dt, J = 5.7 Hz, J = 1.2 Hz, 1H), 8.03–8.00 (m, 2H), 7.60-7.57 (m, 2H), 7.49-7.40 (m, 3H), 6.65 (s, 1H), 5.35 (d, J = 2.2 Hz, 2H), 4.21 (s, 3H), 2.04 (s, 6H), 1.79 (s, 3H), 1.27 (s, 6H). ¹³C{¹H} NMR (151 MHz, CD₂Cl₂): δ 174.3 (C_{NHC}), 163.6 (triazole-C), 151.6 (pyridyl-C), 151.3 (pyridyl-C), 141.9 (s), 140.9 (pyrazole-C), 140.0 (pyridyl-C) 135.3 (benzimidazole-C), 135.1 (pyrazole-C), 134.2 (triazole-C), 133.7 (benzimidazole-C), 124.7 (benzimidazole-C), 124.4 (benzimidazole-C), 124.1 (pyridyl-C), 120.3 (pyridyl-C), 112.0 (benzimidazole-C), 110.6 (benzimidazole-*C*), 101.8 (d, J = 2.1 Hz, pyrazole-*C*), 44.0 (adamantyl-*C*), 42.6 (-(CH₂)-), 37.5 (CH₃), 36.7 (adamantyl-C), 29.4 (adamantyl-C). ¹⁹F NMR (564 MHz, CD_2Cl_2): δ –60.89. MS (ESI HRMS) m/z (%): 754.2194 (100) $[13 + H]^+$ (calcd for $[13 + H]^+$ 754.2101).

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