Dalton Transactions

COMMUNICATION



View Article Online

Check for updates

Cite this: DOI: 10.1039/c8dt04823a

Received 6th December 2018, Accepted 7th December 2018 DOI: 10.1039/c8dt04823a Four different emissions from a Pt(Bodipy) (PEt₃)₂(S-Pyrene) dyad⁺

Peter Irmler,^a Franciska S. Gogesch,^a Christopher B. Larsen,^b Oliver S. Wenger ^b and Rainer F. Winter ^b *^a

The Pt(bodipy)-(mercaptopyrene) dyad BPtSPyr shows four different emissions: intense near-infrared phosphorescence ($\Phi_{\rm ph}$ up to 15%) from a charge-transfer state pyrS⁺⁺-Pt-BDP⁺⁻, additional fluorescence and phosphorescence emissions from the ${}^{1}\pi\pi^{*}$ and ${}^{3}\pi\pi^{*}$ states of the bodipy ligand at r.t., and phosphorescence from the pyrene ${}^{3}\pi\pi^{*}$ and the bodipy ${}^{3}\pi\pi^{*}$ states in a glassy matrix at 77 K.

In accordance with Kasha's rule, the vast majority of compounds emits solely from their lowest excited electronic state.¹ Dual fluorescence and phosphorescence emissions may, however, be observed if intersystem crossing (ISC) occurs at a competitive timescale to fluorescence and if the rate constant for phosphorescence emission is increased by a proximal heavy atom with a large spin-orbit coupling constant.²⁻⁵ By capitalizing on this remote heavy-atom effect we and others have recently observed dual fluorescence and phosphorescence emissions at r.t. from metal complexes with a σ -bonded⁶⁻¹¹ or remotely attached bodipy (BDP) ligand.^{12,13} Multiple emissions from a single compound can also result from inefficient Förster resonance energy transfer in dyads with two electronically decoupled, orthogonally aligned chromophores.¹⁴ Intriguing examples are multiply emissive Ir(ppy)₂(Q) complexes with two cyclometalated phenylpyridine (ppy) and a 2-hydroxyquinoline ligand Q, which show ligand-localized fluorescence from ${}^{1}n\pi^{*}$ and phosphorescence from ${}^{3}MLCT$ (MLCT = metal-ligand charge-transfer) states of the different ligands.¹⁵ We report here on the new *meso*-Pt bodipy complex BPtSPyr (Scheme 1), which combines the above principles and presents four different emissions owing to the coexistence of energetically close-lying ligand-localized and charge-separated excited states.

BPtSPyr was synthesized from *trans*-Pt(bodipy)I(PEt₃)₂ (**BPtI**)⁷ by halogenide abstraction with AgOTf and subsequent addition to a solution of deprotonated 1-mercaptopyrene (pyrS). The same protocol was also used to prepare the complex **MesPtSPyr**, where the BDP chromophore is replaced by mesityl (Mes, Mes = 2,4,6-trimethylphenyl) from newly synthesized *trans*-PtBr(Mes)(PEt₃)₂ (for synthetic details and NMR spectroscopic characterization see ESI and Fig. S1–S18[†] therein).

The electronic absorption spectrum of **MesPtSPyr** is dominated by the $\pi\pi^*$ band of the mercaptopyrene ligand at 431 nm (Fig. 1). Upon excitation into this band, **MesPtSPyr** exhibits dual fluorescence and phosphorescence emissions, at r.t. and at 77 K. Both emissions emanate from the mercaptopyrene ligand as follows from the typical shapes of the emissions and their lifetimes (Fig. S19–S25, ESI† and Table 1).^{5,16,17} ISC in **MesPtSPyr** is obviously triggered by the heavy-metal effect of the attached Pt ion, similar to what has been observed in Pt complexes with a σ -bonded pyrenyl ligand.^{5,16,17}

BPtSPyr reveals a three-band absorption pattern comprising of the structured pyrS-based $\pi\pi^*$ band at 350–425 nm, the bodipy (BDP) $\pi\pi^*$ absorption at 470 nm, and a partially overlapping, weaker pyrS-to-BDP charge-transfer (PB-CT) band at *ca.*



Scheme 1 Synthesis of BPtSPyr and MesPtSPyr.

^aFachbereich Chemie, Universität Konstanz, Universitätsstraße 10,

D-78457 Konstanz, Germany. E-mail: rainer.winter@uni-konstanz.de

^bDepartment of Chemistry, University of Basel, St.-Johanns-Ring 19, CH-4056 Basel, Switzerland

[†]Electronic supplementary information (ESI) available. See DOI: 10.1039/ c8dt04823a



Fig. 1 UV/Vis spectra of BPtSPyr, MesPtSPyr and BPtI in toluene.

Table 1 Emission data of MesPtSPyr and BPtSPyr

Solvent	$\lambda_{ m exc}{}^{a}$		$\lambda_{\mathrm{fl}}{}^{b}$	$\lambda_{\rm ph}^{\ \ b}$ (assignment)	${\Phi_{\mathrm{ph}}}^{c}$	$ au_{ m ph}{}^d$
MesPtSPyr						
Toluene	430		460^{e}	664 (³ pyrS)	< 0.01	$448 \pm 6 \ \mu s$
$Me-THF^{f}$	430		450	658 (³ pyrS)	n.d.	$1.04 \pm 0.06 \text{ ms}$
BPtSPyr						
Toluene	520			724 (³ PB-CT)	0.08	$6.7 \pm 0.4 \ \mu s$
	470		484	724 (³ PB-CT)	0.15	$6.7 \pm 0.5 \ \mu s$
	420			724 (³ PB-CT)	0.15	$6.4 \pm 0.3 \ \mu s$
THF	520			784 (³ PB-CT)	n.d.	n.d.
	470		484^g	635 (³ BDP)	n.d.	78.0 \pm 0.4 μ s,
				784 (³ PB-CT)		51.4 ± 0.8 ns
	420			784 (³ PB-CT)	n.d.	50.4 ± 0.3 ns
Acetone	520			861 (³ PB-CT)	n.d.	n.d.
	470		484	$635 (^{3}BDP)$	n.d.	$49.9\pm0.3~\mu s$
				861 (³ PB-CT)		$2.44\pm0.02~\mathrm{ns}$
Me-THF ^J	530	Т		635 (³ BDP)		270 ± 1 us
	450	7	n.o.	$649 (^{3}$ pyrS)	n.d.	$279 \pm 1 \ \mu s$ 1 12 + 0 02 ms
	415	I		049 (Pyro)		1.12 ± 0.02 1115

^a Excitation wavelength in nm. ^b Wavelength of the fluorescence (fl) or phophorescence (ph) emissions in nm. ^c Phosphorescence quantum yield. Lifetime of the phosphorescence emissions. ${}^{\hat{e}} \tau_{\rm fl} = 1.113 \pm 0.005$ ns, $\Phi_{\rm ph} = 1.113 \pm 0.005$ 0.24. ^f Measured in a 2-MeTHF glass at 77 K. ^g $\tau_{\rm fl} = 3.70 \pm 0.2$ ns.

500 nm. The latter has no equivalent in the precursors BPtI and MesPtSPyr (Fig. 1, Table S1, ESI[†]). The above band assignments are supported by time-dependent density functional theory (TD-DFT) calculations (Fig. S26, ESI⁺). According to our calculations all transitions are ligand-based with only minor Pt contributions to the corresponding donor and acceptor orbitals (Table S2[†]). In line with a CT character, the PB-CT excitation is solvatochromic while the pyrS and BDP $\pi\pi^*$ absorptions are only modestly influenced by solvent polarity (Fig. S27, ESI†).

Importantly, and in contrast to MesPtSPyr, BPtSPyr exhibits four different emissions depending on excitation wavelength, solvent polarity and temperature (Fig. 2). As shown in Fig. 2a, excitation into the PB-CT absorption band at 520 nm leads to broad and unstructured NIR emission with a quantum yield of 8% in toluene. The emission shifts red with increasing solvent polarity, from 724 nm in toluene to 784 nm in THF and to 816 nm in acetone (Table 1). Its strong solvatochromism and the lifetime of



nalized intensity

600

500

700

 λ / nm

800

Fig. 2 Emission spectra of BPtSPyr in ca. 1 µM solutions of acetone, THF or toluene at r.t. on excitation at (a) 520 nm, (b) 450 nm and (c) 405 nm. (d) Emission spectra recorded in 2-MeTHF at 77 K at different excitation wavelengths.

900

500

600

700

900

800

6.7 µs in toluene (for lifetime decays see Fig. S28-S30, ESI†) indicate that this emission emanates from the ³PB-CT state. We note that heavy atom-free donor-substituted BDPs were also shown to undergo efficient ISC to the T1 state via an excited CT state; the latter systems are, however, non-phosphorescent.¹⁸⁻²⁰

Excitation into the BDP $\pi\pi^*$ absorption band at 470 nm (Fig. 2b and S31, ESI[†]) opens two further radiative pathways: BDP-based fluorescence at 484 nm and phosphorescence at 635 nm are observed in addition to the ³PB-CT emission. Changing the solvent from toluene to THF or acetone increases the energy separation between the BDP ${}^{3}\pi\pi^{*}$ and the ${}^{3}PB-CT$ states and allows for tuning of the emission colour from red in toluene (CIE coordinates 0.6052/0.3211 in toluene) to almost white in THF (CIE coordinates: 0.2209/0.3206) or acetone (CIE coordinates 0.3357/0.2901). Concomitantly with its redshift, the lifetime of the ³PB-CT emission decreases from 6.7 μs in toluene solution to only 2.4 ns in acetone (Fig. S30a, ESI[†]). This in turn increases the intensity ratio of the ${}^{1}\pi\pi^{*}/{}^{3}\pi\pi^{*}$ to the ³PB-CT emissions (Fig. 2b). We note that the BDP-based emissions of BPtSPyr are not due to contamination with the BPtI precursor (Fig. S32, ESI[†]). Triple emission upon excitation into the BDP $\pi\pi^*$ band at 470 nm indicates that ISC from the $\pi^*\pi^*$ to the ${}^{3}\pi\pi^{*}$ state of the BDP ligand as well as formation of the {}^{1}PB-CT state and subsequent ISC to the ³PB-CT state occur at similar rates as fluorescence from the BDP ${}^{1}\pi\pi^{*}$ state. When the sample is excited into the pyrS $\pi\pi^*$ band at 415 nm, again solely ³PB-CT emission is observed, but with an even increased quantum yield of 15% (λ_{exc} = 420 nm) as compared to 8% for λ_{exc} = 520 nm (Fig. 2a and c), rendering BPtSPyr a powerful NIR emitter. All the above conclusions are fully supported by excitation vs. emission maps and excitation spectra (Fig. S33 and S34, ESI⁺).

Emission spectra recorded in a glassy Me-THF matrix at 77 K show only two features, irrespective of the excitation wave-

length: the ${}^{3}\pi\pi^{*}$ BDP emission with a maximum at 635 nm and a typical lifetime of 279 µs (Fig. S35, ESI[†]) as well as a structured emission enveloping the region in from 649 to 750 nm (Fig. 2d). This latter feature and its lifetime of 1.12 ms (Fig. S35, ESI[†]) are typical of a pyrene-based ${}^{3}\pi\pi^{*}$ emission.^{5,16,17} The ${}^{3}\pi\pi^{*}$ excited states of the BDP and pyrS ligands are hence both accessible from excitation into any band in the rigid matrix. This indicates that, at 77 K, population of the close-lying BDP and pyrS ${}^{3}\pi\pi^{*}$ from the ¹PB-CT state of **BPtSPyr** is more rapid than direct intersystem crossing to the ³PB-CT state and that deactivation to the ³PB-CT state is blocked under these conditions. One possible explanation is that the ¹PB-CT and ³PB-CT states possess very similar nuclear coordinates and the BDP and pyrS ${}^{3}\pi\pi^{*}$ potential wells have more easily accessible intersections with the ¹PB-CT potential well. Previous studies have similarly demonstrated direct singlet-totriplet electron-transfer by spin-orbit coupling.²¹⁻²³

The Jablonski diagram of Scheme 2 summarizes the processes following the different excitations of BPtSPvr. Electronic decoupling of the pyrS- and the BDP-localized ${}^{1}\pi\pi^{*}$ states, as indicated by the confinement of the HOMO to the pyrS and of the LUMO to the BDP ligand (Fig. S26, ESI[†]), explains the absence of pyrS to BDP energy transfer from the pyrene-based ${}^{1}\pi\pi^{*}$ state S₃ and of any BDP emission after excitation into the pyrS $\pi\pi^*$ band at 415 nm. The BDP-based ${}^{1}\pi\pi^*$ and ${}^{3}\pi\pi^*$ states S_2 and T_3 are thus exclusively populated by excitation into the BDP ${}^{1}\pi\pi^{*}$ band. In contrast, the 3 PB-CT state, which constitutes the energetically lowest excited triplet state T_1 , is accessible from all three excited singlet states, either via IC to the ¹PB-CT and subsequent ISC, or via ISC and subsequent electron transfer from the higher-lying pyrS-based T₂ and BDP-based T₃ states (note that MesPtSPyr also undergoes ISC from the pyrene-based ${}^{1}\pi\pi^{*}$ state). At 77 K, direct singlet to triplet electron-transfer from the ¹PB-CT state allows population of the pyrS- and BDP-based triplet states T2 and T3 (blue arrows in Scheme 2) and renders the pyrS-based phosphorescence emission observable, which, at r.t., is quenched by rapid decay into the lower-lying ³PB-CT state.

The proposed excited state cascade is further supported by transient absorption (TA) spectroscopy in THF. TA spectra recorded over the first 200 ns after excitation are very similar to each other, irrespective of whether the sample is excited at 410



Scheme 2 Qualitative Jablonski diagram.



Fig. 3 Transient absorption spectra of BPtSPyr at r.t. (a) in deaerated 20 μ M THF solution at different time delays after excitation at 470 nm; and (b) in a 20 μ M toluene solution directly after laser excitation at the given wavelength. All spectra are time-integrated over 200 ns.

(pyrS), 470 (BDP), or 532 nm (PB-CT absorption) (Fig. 3 and Fig. S36, ESI[†]). Three main features are observed: the bleached pyrS and BDP ground state absorption signals at ca. 400 nm or 470 nm, and an excited state absorption (ESA) signal at 550 nm, which we assign to the pyrS^{*+} radical cation (vide infra). Identical monoexponential kinetics with a common time constant of 50 ns were found for all three transients when excited at 532 nm (for single point kinetics for each TA feature see Fig. S37 and S38, ESI[†]). When excited into the BDP absorption at 470 nm the features at 415 nm and 550 nm still have a time constant of 50 ns, but the bleach recovery at 475 nm has much longer kinetics of 16.5 µs in THF. This is in accordance with the long-lived BDP ${}^{3}\pi\pi^{*}$ excited state in that solvent (red trace in Fig. S37a, ESI[†]). Transient absorption bleach recoveries and decays match the luminescence lifetimes in all cases. This corroborates that none of the observable emissions originate from impurities but represent inherent photoluminescence from BPtSPyr.

The common CT state after excitation should exhibit spectroscopic features similar to the oxidized and reduced forms of BPtSPyr. Cyclic voltammograms show a reversible oxidation at +0.18 V and a reduction at -1.97 V against the ferrocene/ferrocenium couple (Fig. S39 and Table S3, ESI⁺). This allowed us to experimentally probe the UV/Vis/NIR features of the oxidized and reduced forms of BPtSPyr by means of spectroelectrochemistry. Upon oxidation, the pyrS-based $\pi\pi^*$ absorption band bleaches while the BDP $\pi\pi^*$ band exhibits an only minor red-shift of 316 cm⁻¹ (Fig. 4). Based on literature data^{24,25} and the results of our TD-DFT calculations (Fig. S40 and Table S4, ESI[†]), the new absorption bands at ca. 520, 550, 750, and 1000 nm can be assigned to $\pi\pi^*$ transitions of the pyrS⁺⁺ radical cation. When **BPtSPyr** is reduced, the BDP $\pi\pi^*$ and the PB-CT absorption bands bleach while the pyrS-based $\pi\pi^*$ band experiences a moderate red-shift. The weak new band at 518 nm constitutes mainly a $\pi\pi^*$ transition of the reduced BDP ligand (Fig. S41 and S42, Table S5, ESI[†]); it is essentially identical in shape and energy to that observed for BPtI^{-.9} Calculated spin densities for the BPtSPyr radical cation and anion also agree with a pyrS-based oxidation and a BDP-based reduction (Fig. 5 and Table S6, ESI[†]). The results of our spectroelectrochemical experiments and our calculations thus confirm our assignment of the ESA signal to the pyrS^{*+} constituent as well as the character of the HOMO and LUMO of BPtSPyr.



Fig. 4 Changes of UV/Vis/NIR spectra of BPtSPyr on oxidation (THF, NBu_4PF_6 , 293 K).



Fig. 5 Calculated spin densities of (a) the radical cation and (b) the radical anion of BPtSPyr.

In summary, the **BPtSPyr** dyad combines several remarkable features. It constitutes an extremely rare case of a compound showing emissions from four different excited states and acts as a powerful NIR emitter. Its emission profiles and colour can be tuned by the excitation wavelength and solvent polarity from deep red to almost white. The crucial role of the pyrS⁺⁺-Pt-BDP⁺⁻ charge-transfer state within the excited state cascade was identified by TA spectroscopy, spectroelectrochemical methods and TD-DFT calculations.

Conflicts of interest

The authors declare no competing financial interests.

Acknowledgements

This work was supported by Deutsche Forschungsgemeinschaft (grant Wi1262/10-2) and the Swiss National Science Foundation (grant number 200021_176780).

Notes and references

- 1 M. Kasha, Discuss. Faraday Soc., 1950, 9, 14–19.
- 2 C.-W. Hsu, C.-C. Lin, M.-W. Chung, Y. Chi, G.-H. Lee, P.-T. Chou, C.-H. Chang and P.-Y. Chen, *J. Am. Chem. Soc.*, 2011, 133, 12085–12099.

- 3 C.-C. Hsu, C.-C. Lin, P.-T. Chou, C.-H. Lai, C.-W. Hsu, C.-H. Lin and Y. Chi, *J. Am. Chem. Soc.*, 2012, **134**, 7715-7724.
- 4 Y.-C. Chang, K.-C. Tang, H.-A. Pan, S.-H. Liu, I. O. Koshevoy, A. J. Karttunen, W.-Y. Hung, M.-H. Cheng and P.-T. Chou, *J. Phys. Chem. C*, 2013, **117**, 9623–9632.
- 5 W. Y. Heng, J. Hu and J. H. K. Yip, *Organometallics*, 2007, **26**, 6760–6768.
- 6 F. Geist, A. Jackel and R. F. Winter, *Inorg. Chem.*, 2015, 54, 10946–10957.
- 7 P. Irmler and R. F. Winter, *Dalton Trans.*, 2016, **45**, 10420–10434.
- 8 F. Geist, A. Jackel, P. Irmler, M. Linseis, S. Malzkuhn, M. Kuss-Petermann, O. S. Wenger and R. F. Winter, *Inorg. Chem.*, 2017, 56, 914–930.
- 9 P. Irmler and R. F. Winter, *Organometallics*, 2018, **37**, 235–253.
- 10 W. Wu, J. Zhao, H. Guo, J. Sun, S. Ji and Z. Wang, Chem. Eur. J., 2012, 18, 1961–1968.
- 11 F. Zhong, A. Karatay, L. Zhao, J. Zhao, C. He, C. Zhang, H. G. Yaglioglu, A. Elmali, B. Küçüköz and M. Hayvali, *Inorg. Chem.*, 2015, 54, 7803–7817.
- 12 J. Sun, F. Zhong, X. Yi and J. Zhao, *Inorg. Chem.*, 2013, 52, 6299–6310.
- 13 W. Wu, J. Sun, X. Cui and J. Zhao, *J. Mater. Chem. C*, 2013, 1, 4577–4589.
- 14 B. Ma, P. I. Djurovich, M. Yousufuddin, R. Bau and M. E. Thompson, J. Phys. Chem. C, 2008, 112, 8022–8031.
- 15 T. Tsuboi, D.-F. Huang, T. J. Chow and W. Huang, Opt. Mater., 2014, 36, 1734–1738.
- 16 J. Hu, J. H. K. Yip, D.-L. Ma, K.-Y. Wong and W.-H. Chung, Organometallics, 2009, 28, 51–59.
- 17 W. Wu, W. Wu, S. Ji, H. Guo and J. Zhao, Eur. J. Inorg. Chem., 2010, 2010, 4470–4482.
- 18 M. A. Filatov, S. Karuthedath, P. M. Polestshuk, S. Callaghan, K. J. Flanagan, M. Telitchko, T. Wiesner, F. Laquai and M. O. Senge, *Phys. Chem. Chem. Phys.*, 2018, 20, 8016–8031.
- 19 X.-F. Zhang and N. Feng, *Chem. Asian J.*, 2017, **12**, 2447–2456.
- 20 S. Ji, J. Ge, D. Escudero, Z. Wang, J. Zhao and D. Jacquemin, *J. Org. Chem.*, 2015, 80, 5958–5963.
- 21 M. R. Wasielewski, D. G. Johnson, W. A. Svec, K. M. Kersey and D. W. Minsek, *J. Am. Chem. Soc.*, 1988, **110**, 7219–7221.
- 22 S.-H. Lee, A. G. Larsen, K. Ohkubo, Z.-L. Cai, J. R. Reimers, S. Fukuzumi and M. J. Crossley, *Chem. Sci.*, 2012, 3, 257– 269.
- 23 T. Higashino, T. Yamada, M. Yamamoto, A. Furube,
 N. V. Tkachenko, T. Miura, Y. Kobori, R. Jono,
 K. Yamashita and H. Imahori, *Angew. Chem., Int. Ed.*, 2016, 55, 629–633.
- 24 T. Shida and S. Iwata, J. Am. Chem. Soc., 1973, 95, 3473-3483.
- 25 J. Maurer, M. Linseis, B. Sarkar, B. Schwederski, M. Niemeyer, W. Kaim, S. Záliš, C. Anson, M. Zabel and R. F. Winter, *J. Am. Chem. Soc.*, 2008, **130**, 259–268.