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### Introduction

Phosphorescence is a distinctive photophysical property of transition-metal complexes, which has widespread applications in diverse areas. As it is derived from the 'forbidden' radiative relaxation of a triplet-excited state to the singlet ground state, it

## The interplay between fluorescence and phosphorescence with luminescent gold(I) and gold(III) complexes bearing heterocyclic arylacetylide ligands†

YAL SOCIETY<br>**CHEMISTRY** 

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The photophysical properties of a series of gold(i)  $[LAu(C=CR)]$  (L = PCy<sub>3</sub> (1a–4a), RNC (5a), NHC (6a)) and gold(III) complexes [Au(C^N^C)(C=CR)] (1b-4b) bearing heterocyclic arylacetylide ligands with narrow band-gap are compared. The luminescence of both series are derived from an intraligand transition localized on the arylacetylide ligand ( $\pi\pi*(C\equiv CR)$ ) but 1a–3a displayed prompt fluorescence  $(\tau_{PF}$  = 2.7–12.0 ns) while 1b–3b showed mainly phosphorescence  $(\tau_{Ph}$  = 104–205 µs). The experimentally determined intersystem crossing (ISC) rate constants ( $k_{\text{ISC}}$ ) are on the order of 10<sup>6</sup> to 10<sup>8</sup>  $s^{-1}$  for the gold(i) series (1a-3a) but 10<sup>10</sup> to 10<sup>11</sup>  $s^{-1}$  for the gold(iii) analogues (1b-3b). DFT/TDDFT calculations have been performed to help understand the difference in the  $k_{\text{ISC}}$  between the two series of complexes. Owing to the different oxidation states of the gold ion, the Au(I) complexes have linear coordination geometry while the Au(III) complexes are square planar. It was found from DFT/TDDFT calculations that due to this difference in coordination geometries, the energy gap between the singlet and triplet excited states ( $\Delta E_{ST}$ ) with effective spin–orbit coupling (SOC) for Au(i) systems is much larger than that for the Au(III) counterparts, thus resulting in the poor ISC efficiency for the former. Timeresolved spectroscopies revealed a minor contribution (<2.9%) of a long-lived delayed fluorescence (DF)  $(\tau_{DF} = 4.6 - 12.5 \text{ }\mu\text{s})$  to the total fluorescence in 1a–3a. Attempts have been made to elucidate the mechanism for the origins of the DF: the dependence of the DF intensity with the power of excitation light reveals that triplet-triplet annihilation (TTA) is the most probable mechanism for the DF of 1a while germinate electron–hole pair (GP) recombination accounts for the DF of 2a in 77 K glassy solution (MeOH/EtOH  $= 4:1$ ). Both 4a and 4b contain a BODIPY moiety at the acetylide ligand and display only  ${}^{1}$ IL( $\pi\pi^{*}$ ) fluorescence with negligible phosphorescence being observed. Computational analyses attributed this observation to the lack of low-lying triplet excited states that could have effective SOC with the  $S_1$  excited state. **EDGE ARTICLE**<br> **Solutions**<br> **Solution**<br> **Solution School Constrained School Constrained** 

> is featured by long emission lifetime (in  $\mu s$ ) and reduced emission energy compared to fluorescence commonly encountered in organic luminophores. Electronic transitions associated with a change of spin are prohibited by the spin-selection rule. However, transition-metal ions that have high atomic number and hence, large spin–orbit coupling constant  $(\xi)$ , usually lead to efficient spin–orbit coupling (SOC) that relaxes the spin selection rule. Fast intersystem crossing (ISC) in the sub-picosecond to picosecond time regime<sup>1</sup> leads to rapid depletion of a singlet excited state to a triplet excited state instead of fluorescence as fluorescence radiative lifetime is typically in the nanosecond range. Thus, in transition-metal complexes, phosphorescence normally prevails in their luminescence spectra.

> However, in recent years, there are an increasing number of reports on transition-metal complexes which display slow ISC

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rate with lifetimes ranging from hundreds of ps to ns. For instance, 2,5-bis(arylethynyl)rhodacyclopentadiene complexes  $(\xi_{\rm Rh} = 1260 \text{ cm}^{-1})^2$  were reported to display exclusively prompt fluorescence with high emission quantum yields of 0.3-0.7 and lifetimes of 1–3 ns, corresponding to ISC rate constants of  $\sim 10^8$  $s^{-1}$ .<sup>3</sup> In addition, transition-metal complexes containing fused aromatic systems such as perylene, perylene diimide, pyrene and tetracene also show ligand-dominated fluorescence (see Fig. 1).<sup>4</sup> Hence, it has become clear that the presence of heavy elements does not guarantee fast ISC rate; the molecular structure and the nature of the ligands may play more critical roles in determining the ISC rate.

Luminescent  $Au(i)$  complexes are well documented to display rich photophysical properties. Although Au(I) complexes generally display phosphorescence owing to the large SOC constant of Au(I) ion ( $\xi_{\text{Au}} \sim 5100 \text{ cm}^{-1}$ ),<sup>2</sup> ligand-centered fluorescence has also been reported in a number of  $gold(i)$ complexes. For example, as revealed by the luminescence of  $[TEE(AuPCy<sub>3</sub>)<sub>4</sub>]$  and  $[TEB(AuPCy<sub>3</sub>)<sub>3</sub>]$  (TEE = tetraethynylethene;  $TEB = 1,3,5$ -triethynylbenzene), subtle changes in the electronic structure of the bridging alkynyl ligand leads to intense phosphorescence ( $\Phi_{\text{em}} = 0.46$ ,  $\tau = 285 \text{ }\mu\text{s}$ ) in the latter but solely fluorescence ( $\Phi_{\text{em}} = 0.22$ ,  $\tau \le 0.05$  µs) in the former.<sup>5</sup> In both cases, the luminescence originates from the ligand-centered transition mainly localized on the bridging alkynyl ligands. Che and co-workers also reported a series of Au(I)-conjugated acetylides,  $[(Cy_3P)Au(C\equiv C-C_6H_4)_{n-1}(C\equiv CPh)]$   $(n \ge 2)$ , which display dual fluorescence (prompt and delayed) and phosphorescence.<sup>6</sup> Both the  $\Phi_{\text{em}}$  and ratio of fluorescence versus phosphorescence were found to depend on the conjugation length (number of repeating units  $n$ ) and the substitution pattern of arylacetylide ligands.

As a continuous effort to elucidate the ligand effects on the photophysics of luminescent gold complexes, heterocyclic arylacetylide ligands containing narrow band-gap moieties (benzothiadiazole (L1), coumarin (L2), naphthalimide (L3) and borondipyrromethene (referred to as Bodipy) (L4); Chart 1) were chosen in this study. A series of Au(I) heterocyclic arylacetylide complexes, 1a–4a, were synthesized. Tricyclohexylphosphine



Fig. 1 Selected examples of transition-metal complexes that display dominant fluorescence instead of phosphorescence. Auxiliary ligands coordinated to the metal ions are omitted.



Chart 1 Au(I) and Au(III) acetylide complexes studied in this work.

 $(PCy<sub>3</sub>)$  was used as the auxiliary ligand in these complexes because (1) it is optically transparent at wavelength >250 nm so that it is not involved in the emissive excited states in the UV-visible spectral region and (2) its steric bulkiness would prevent the gold ions from coming into close contact that could lead to low-lying excited states originated from metal–metal and  $\pi$ – $\pi$  interactions. It is worth mentioning that several recently reported Au(I) alkynyl complexes bearing similar benzothiadiazole,<sup>7</sup> coumarin<sup>8</sup> and naphthalimide<sup>9</sup> derivatives also show similar luminescence properties as our  $Au(1)$  complexes.<sup>10,11</sup> The effects of auxiliary ligands on the photophysical properties of  $Au(i)$  complexes were also studied by comparing 1a with two derivatives containing 2,6-dimethylphenyl isocyanide (RNC, 5a) and 1,3-dimethylimidazol-2-ylidene (NHC, 6a) instead of the phosphine auxiliary ligand, respectively.

The effects of the oxidation state of the metal ion on the photophysical behaviours of transition-metal complexes are relatively unexplored. Herein, an analogous series of  $Au(m)$ acetylides supported by the cyclometalated  $[C^N^C]$  ligand (1b–4b; HC $\textdegree N\textdegree CH = 2,6$ -diphenylpyridine) were also prepared and their photophysical properties were compared with those of the Au(I) counterparts. The photophysical properties of both  $Au(i)$  and  $Au(i)$  complexes were investigated by steadystate and time-resolved spectroscopic measurements. DFT/ TDDFT calculations were performed on the pairs (1a, 1b) and (4a, 4b) in order to understand the origin of the dramatic difference in ISC efficiencies between these  $Au(I)$  and  $Au(III)$ complexes.

### Results

#### Synthesis and characterization

The gold $(I)$  alkynyl complexes **1a–6a** were synthesized in 53–79% yields following the protocol of base deprotonation (NaOMe) of terminal alkynes and substitution of chloride ion of the corresponding  $Au(i)$  precursors.<sup>5a,6a,b,12</sup> As these complexes were observed to show signs of decomposition on  $SiO<sub>2</sub>$  column, column chromatography was not used for their purification. Analytically pure 1a–6a were obtained by recrystallization from  $CH_2Cl_2/h$  exane mixtures. The Au( $\text{m}$ ) complexes **1b–4b** were synthesized by copper-catalyzed Sonogashira coupling between terminal alkynes and [Au(C^N^C)Cl] using deoxygenated  $CH_2Cl_2$  as the solvent, with NEt<sub>3</sub> added to initiate the deprotonation of alkynes.<sup>13</sup> These complexes were purified by chromatography on SiO<sub>2</sub> column using dichloromethane and hexane as eluent. The yields were 51–84%.

All complexes have been characterized by  ${}^{1}\mathrm{H}$  and  ${}^{13}\mathrm{C}$  NMR, mass spectrometry (FAB+) and elemental analyses. Ligands L1–L4 were characterized by  $^1\mathrm{H}$  NMR and MS-EI. The complexes are stable in the solid state and in solution under ambient conditions. Complexes  $1a-6a$  are highly soluble in  $CH_2Cl_2$  and THF but are less soluble in alcoholic solvents such as MeOH. Complexes 1b–4b have lower solubility compared with their  $Au(i)$  counterparts. All of these gold complexes appear as yellow or orange solids except for 4a and 4b that are purplish red. The <sup>31</sup>P signals of **1a-4a** occur at *ca.*  $\delta$  56.3 as a singlet, characteristic of the  $31P$  signals of the Au–PC $y_3$  moiety that usually appear in the range of  $\delta$  56.0–58.0.<sup>6a,b,12</sup> In the <sup>13</sup>C NMR spectra, two doublets are observed at *ca.*  $\delta$  131.2–146.2 ( $\mathcal{C}_{JCP} \approx 130$  Hz) and 94.6–98.4 ( ${}^{3}J_{CP} \approx 24$  Hz) which can be assigned to the  $\alpha$  and  $\beta$ -acetylenic carbons.<sup>6a,b,12b</sup> In 6a, the carbene carbon ligated to gold occurs at  $\delta$  187.7.<sup>14</sup> Chemical Schene:<br> **Properties and characterization**<br> **Supplies and characterization**<br> **Supplies and characterization**<br>
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#### X-Ray crystallography

Crystals of 1a, 4a and 2b were obtained by layering hexane over concentrated CH<sub>2</sub>Cl<sub>2</sub> solutions. Their crystal data and selected bond lengths and angles are given in ESI.† Fig. 2 shows the



structures of 1a and 4a (top panel). The P1–Au1–C(acetylide) angles of 1a and 4a are 175.0(13) and 178.1(2)° and Au1–C $\equiv$ C angles are  $170.9(4)$  and  $177.6(5)^\circ$ , respectively, revealing slight deviation from linear coordination geometry. The Au1–C(acetylide) distances of 2.049(4) and 2.001(5)  $\AA$  and C=C distances of 1.146(7) and 1.191(8)  $\AA$  for **1a** and **4a**, respectively, are comparable with those of other reported  $\text{gold}(I)$  acetylide complexes.<sup>6a,b,12a,b</sup> The crystal packing diagrams of 1a and 4a are shown in ESI (Fig. S1†). In both cases, there are no short intermolecular contacts; the closest  $Au \cdots Au$  distances are 5.9715(5) and  $5.2726(4)$  Å for **1a** and **4a**, respectively.

The crystal structure of 2b (Fig. 2, bottom) shows a slightly distorted square-planar geometry with C1–Au1–C17 angle of 162.44(14)°. The Au1-C(acetylide) and C $\equiv$ C distances are 1.969(4) and 1.197(5)  $\AA$ , respectively. These parameters are similar to those found in related cyclometalated  $Au(m)$  arylacetylide complexes.<sup>13</sup> The torsional angle between the Au( $C^N$ <sup> $\sim$ </sup>C) and arylacetylide planes is approximately 72.6°. This non-planarity gives rise to negligible  $\pi-\pi$  stacking between molecules as shown in Fig. S2 in ESI.†

#### Electrochemical properties

The electrochemical properties of selected complexes, 1a–4a and 1b–4b, were investigated by cyclic voltammetry. The electrochemical data are summarized in Table 1. The cyclic voltammograms of the  $Au(1)$  complexes and their  $Au(III)$ counterparts are shown in Fig. S3, ESI.† Except for 1a and 2a, both classes of complexes display both irreversible oxidation  $(E_{pa} = +0.6$  to +1.4 V) and quasi-reversible/irreversible reduction waves ( $E_{\text{pc}} = -1.5$  to  $-1.8$  V) attributed to the redox process localized on the arylacetylides. For the pairs [2a, 2b] and [4a, 4b], the  $E_{pa}$  occur at relatively low potential of *ca.* +0.6 and +0.7 V, respectively, suggesting higher HOMO level of the conjugated coumarin and Bodipy than the other heterocyclic moieties. For the  $Au(III)$  complexes, other than the redox reactions occurring at the arylacetylide ligands at potentials similar to the  $Au(1)$  counterparts, there are also irreversible reduction waves at *ca.*  $-1.9$  to  $-2.0$  V attributable to reduction of the  $[C^NC]$  ligands.



<sup>*a*</sup> Values determined in CH<sub>2</sub>Cl<sub>2</sub> (C<sub>P<sub>2</sub>Fe<sup>+/0</sup></sup> occurs at  $E_{1/2} = +0.15-0.16$  V)</sub> at 298 K; values reported versus Ag/AgNO<sub>3</sub> reference electrode; electrolyte: 0.1 M  $nBu_4NPF_6$ ; scan rate = 100 mV s<sup>-1</sup>. <sup>b</sup> Cathodic peak potential  $(E_{\text{pc}})$  of irreversible wave. <sup>c</sup> Anodic peak potential of  $(E_{\text{pa}})$ irreversible wave.  ${}^{d}E_{1/2} = E_{\text{pa}} + E_{\text{pc}}$  of quasi-reversible wave.

#### Table 2 Photophysical data of  $1a-6a$ ,  $1b-4b$  and  $L1-L4^a$



<sup>a</sup> Data were obtained from steady-state measurements with degassed CH<sub>2</sub>Cl<sub>2</sub> solutions ( $2 \times 10^{-5}$  M) unless specified. Measurements with glassy solutions were performed in EtOH/MeOH  $(4:1)$  mixture at 77 K.  $^b$  Emission lifetimes of prompt fluorescence  $(\tau_{\rm PF})$  were determined by timecorrelated single photon counting (TCSPC) measurement.  $c$  Emission lifetimes of delayed fluorescence ( $\tau_{\rm DF}$ ) were obtained from fitting the decay of the time-resolved emission (TRE) as a mono-exponential decay in the delay time range of 0–40 ns and 1–46 µs, respectively. Measurements were performed in degassed CH<sub>2</sub>Cl<sub>2</sub> (5  $\times$  10<sup>-5</sup> M) solutions. <sup>d</sup> Emission quantum yields ( $\Phi_{\rm em}$ ) were obtained using quinine sulfate in degassed 0.5 M H<sub>2</sub>SO<sub>4</sub> ( $\Phi = 0.546$ ) as the standard unless specified.  $\Phi_{\text{em}}$  measured in steady state is the overall emission quantum yield, i.e.  $\Phi_{em} = \Phi_{pr} + \Phi_{DF}$  for 1a–3a and 5a–6a. <sup>e</sup> Obtained from time-resolved emission spectra. <sup>f</sup> Emission lifetime was not determined (n.d.) due to weak emission signal. <sup>g</sup> Determined from time-resolved emission spectra in degassed CH<sub>2</sub>Cl<sub>2</sub> ( $1 \times 10^{-5}$  M) solutions. <sup>h</sup> Determined from fs-TRF spectra. <sup>i</sup> Emission quantum yields ( $\Phi_{\rm cm}$ ) were obtained using [Ru(bpy)<sub>3</sub>][PF<sub>6</sub>]<sub>2</sub> in degassed acetonitrile as the standard ( $\Phi$  = 0.062).



Fig. 3 UV-vis absorption spectra (left) and emission spectra (right) of **1a–4a** (top) and **1b–4b** (bottom) in CH<sub>2</sub>Cl<sub>2</sub> at 298 K (2  $\times$  10<sup>-5</sup> M).

#### UV-vis absorption spectroscopy

All photophysical data of the gold complexes and the free ligands L1–L4 are listed in Table 2.

Electronic absorption of Au(I) complexes 1a–6a. Fig. 3 (top left) shows the absorption spectra of 1a–4a. The lowest energy absorption bands of 1a–4a are at 379, 410, 397 and 553 nm, respectively, and their molar absorptivities  $(\varepsilon)$  fall in the range 6.9  $\times$  10<sup>3</sup> to 4  $\times$  10<sup>4</sup> mol<sup>-1</sup> dm<sup>3</sup> cm<sup>-1</sup>. These lowest energy absorption bands have spectral features resembling those of L1–L4 (Fig. S4 in ESI†) and are attributable to the dipole-allowed intraligand transitions of the arylacetylide ligands  $(^1\pi\pi^*(C\text{=}CR))$ with some charge-transfer character. Similar assignments were also made for other Au(1) alkynyl complexes in the literature.<sup>6,7,8*a*</sup> Bathochromic shifts of <sup>1</sup>IL transitions of arylacetylides are observed upon coordination of the arylacetylides to the  $Au(i)$  ion and are ascribed to  $\pi$ -interaction between Au(I) 5d orbitals and the ligand  $\pi$ -orbitals (see MO surfaces in Fig. 9 and 10).

Replacing the neutral auxiliary ligand  $PCy_3$  in 1a with 2,6dimethylphenyl isocyanide (RNC, 5a) and 1,3-dimethylimidazol-2-ylidene (NHC, 6a) results in a slight change in  $\lambda_{\text{max}}$  of the lowest energy absorption band ( $\lambda_{\text{max}} = 371$  nm (5a, RNC) and 383 nm (6a, NHC) cf.  $\lambda_{\text{max}} = 379$  nm (1a, PCy<sub>3</sub>)) (Fig. S6 in ESI†).

Electronic absorption spectra of  $Au(m)$  complexes 1b-4b. Fig. 3 (bottom left) shows the UV-vis absorption spectra of the  $Au(m)$  complexes studied in the present work. The spectral features at  $\lambda > 300$  nm are similar to those of the Au(I) analogues and free ligands L1–L4. Complexes 1b–4b display low-energy absorption bands at  $\lambda_{\text{max}} = 381, 432, 395$  and 546 nm, respectively, and with  $\varepsilon$  values in the range of 1.38  $\times$  $10^4$  to  $4.17 \times 10^4$  mol<sup>-1</sup> dm<sup>3</sup> cm<sup>-1</sup>. These absorption bands are, like the Au(1) complexes, attributable to  $^1\pi\pi^*(C\!\!\equiv\!\!\mathbb{CR})$ transitions. Comparisons of the absorption spectra of 1a–4a and 1b–4b at  $\lambda \le 300$  nm revealed that the absorption bands in this spectral region are more intense in  $1b-4b$  with  $\varepsilon$  values of *ca.*  $4 \times 10^4$  mol<sup>-1</sup> dm<sup>3</sup> cm<sup>-1</sup> and these high energy absorption bands likely involve intraligand  ${}^1\pi\pi^*(C^\wedge N^\wedge C)$  transitions (Fig. S7 in ESI†).

#### Steady-state emission spectroscopy

All of the complexes are luminescent in degassed  $CH_2Cl_2$  at room temperature and in 77 K glassy solutions (EtOH : MeOH  $=$ 4 : 1) upon excitation at the corresponding lowest-energy absorption  $\lambda_{\text{max}}$ . As depicted in Table 2, there is a distinct difference between the two classes of complexes: the  $Au(i)$ complexes 1a–6a display predominantly fluorescence while the  $Au(m)$  complexes,  $1b-3b$ , exhibit exclusively weak phosphorescence. Complex 4b, on the other hand, shows fluorescence only.

Emission of 1a–4a and 5a–6a. In dichloromethane solutions, structureless emission bands are observed at  $\lambda_{\text{max}} = 467, 466,$ 439 and 553 nm for 1a–4a, respectively (Fig. 3, top right). The corresponding excitation spectra of 1a–4a can be found in the ESI (Fig. S8†). The emission quantum yields for 1a–3a are high  $(\Phi_{em} = 0.91, 0.70$  and 0.78, respectively). In the case of 4a, its emission quantum yield is low ( $\Phi_{em}$  = 0.04). Emission lifetimes of 1a–4a are in the nanosecond time regime: 0.8–12 ns. As the emissions of these Au(I) complexes resemble those of the corresponding free ligands L1–L4, they are attributable to  $1\pi\pi^*(C\equiv CR)$  excited states, with some charge transfer character, which probably arise from mixings of metal-to-ligand charge-transfer (MLCT) character. Solvent effects on the emissions of 1a–3a can be found in Fig. S10, ESI.† There is no discernible phosphorescence for 1a–4a under steady-state conditions in solutions at either room temperature or 77 K (Fig. S9, ESI†). Openies Science<br>
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Comparing the three  $Au(i)$  complexes bearing the benzothiadiazole moiety, the emission energies ( $\lambda_{\text{max}} = 467$ , 456 and 476 nm for  $1a$  (PCy<sub>3</sub>),  $5a$  (RNC) and  $6a$  (NHC), respectively (Fig. S11 in ESI†)) and emission lifetimes ( $\tau_{PF} \sim 11-14$  ns) are similar, indicating that the auxiliary ligand plays an insignificant role in modification of the electronic structures of the excited states.

**Emission of 1b–4b.** Emission spectra of the  $Au(\text{III})$  complexes are depicted in Fig. 3 (bottom right). Contrary to the  $Au(1)$ analogues where the emission profiles are structureless, the emission spectra of complexes 1b–3b are vibronically structured with  $\lambda_{\text{max}}$  at 630, 592 and 603 nm and quantum yields of 0.003, 0.01 and 0.04, respectively. The emission lifetimes are of hundreds of microseconds ( $\sim$ 100 µs for 1b and 2b;  $\sim$ 200 µs for 3b). Taking into account the large Stokes shifts (between 6300 and 10 400  $\text{cm}^{-1}$ ), structured emission profiles, and long emission lifetimes, the emissions of 1b–3b could be attributed to  $\sqrt[3]{\pi \pi^*}$ (C $\equiv$ CR) excited states with negligible mixings of MLCT and LLCT character (LLCT = ligand-to-ligand charge transfer). Solvent effects on the emissions of 1b can be found in Fig. S12, ESI.† On the contrary, 4b shows emission with a small Stokes shift of 920  $\text{cm}^{-1}$  and emission lifetime of only 2.1 ns. Thus, the emission of 4b is derived from fluorescence with  $1\pi\pi$ \*(C=CBodipy) parentage.

#### Time-resolved spectroscopies for the  $gold(i)$  complexes

Nanosecond time-resolved emission (ns-TRE) spectra of the Au(1) complexes in degassed CH<sub>2</sub>Cl<sub>2</sub> solutions at 298 K (5  $\times$  10<sup>-5</sup> M) are measured at different time delays and are presented in Fig. 4 (1a) and ESI (2a–3a, 5a–6a; Fig. S13†). There are two



Fig. 4 ns-TRE spectra of 1a recorded from (a) 0–42 ns and (b) after a time delay of 1  $\mu$ s in degassed CH<sub>2</sub>Cl<sub>2</sub> (5  $\times$  10<sup>-5</sup> M) at 298 K. Inset shows the emission kinetic decay trace. Decay time constants were fitted as mono-exponential decay ( $\lambda_{\text{exc}} = 355$  nm).

components in the emission decay: a major component which decays within nanoseconds  $(\tau_1 = 12.0 \text{ (1a)}, 2.7 \text{ (2a)}, 2.8 \text{ (3a)}, 11.0)$ (5a) and 14.0 ns (6a)) and a minor component with microsecond decay lifetime ( $\tau_2$  = 11.9 (1a), 4.6 (2a), 2.6, 12.5 (3a), 6.8 (5a) and 7.4  $\mu$ s (6a)). For each of these Au(I) complexes, both decay components have identical emission profile and peak energy and so, the short-lived one  $(\tau_1)$  is assigned to be prompt fluorescence (PF) while the long-lived one  $(\tau_2)$  is delayed fluorescence (DF) of  $^1\pi\pi^*(C\!\!\equiv\!\!\mathbb{CR})$  character. In the case of  $4a$ , only PF  $(\tau_{PF} = 0.8 \text{ ns})$  is observed. The proportion of DF and PF constituting the total emission of 1a–3a have been estimated (Table 3): the intensity of DF is minute (<3%) when compared with that of PF  $(>97%)$ . It is noted that delayed fluorescence in the microsecond time regime is indicative of the emission generated from a long-lived excited state. This is further supported by nanosecond transient absorption (ns-TA) measurements that reveal the presence of long-lived absorbing species in the microsecond timescale (vide infra).

Table 3 Proportion of PF and DF constituting the fluorescence of  $1a-3a^c$ 

Complex	% PF	% DF
1a	99.2	0.81
2a 3a	99.9 97.1	0.1 2.9

 $a$  % PF and % DF are estimated by integrating the emission intensity of degassed CH<sub>2</sub>Cl<sub>2</sub> ( $5 \times 10^{-5}$  M) in the spectral region of  $\lambda = 350-$ 700 nm over the time range:  $0-500$  ns and  $800$  ns to 999  $\mu$ s, respectively ( $\lambda_{\rm exc}$  = 355 nm).

Weak phosphorescence bands were observed for 1a–3a under different conditions. For dilute CH<sub>2</sub>Cl<sub>2</sub> solutions (1  $\times$ 10-<sup>5</sup> M) at 298 K, dominant emissions were observed in the spectral region of 440-470 nm, which correspond to fluorescence (Fig.  $S14$ , left panel in ESI†). In addition, weak emission peaks at ca. 600 nm become discernible for 2a and 3a and the lifetimes measured are 13.6 and 61.9 µs, respectively (Fig. S13, right panel in ESI†). Cooling to 77 K gives more resolved phosphorescence bands with vibrational progression spacings of 1300-1400  $\text{cm}^{-1}$  for all three complexes (Fig. 5 and 6). For 1a, contrary to the ns-TRE spectra recorded in degassed CH<sub>2</sub>Cl<sub>2</sub> at room temperature (Fig. 4 (5  $\times$  10<sup>-5</sup> M); Fig. S14, ESI†  $(1 \times 10^{-5}$  M)) where only DF could be observed over the time range 1–46 ms, in 77 K glassy solution, phosphorescence at 630 nm is dominant and the weak DF at 467 nm vanishes after  $80 \mu s$  (Fig. 5a). The phosphorescence band decays with firstorder kinetics at  $\tau_{\text{phos}} = 109 \,\mu s$ . Similarly, the low-temperature ns-TRE spectra of 3a is dominated by phosphorescence at 609 nm and DF vanishes after 200  $\mu$ s (Fig. 5b). The phosphorescence band also decays mono-exponentially with  $\tau_{\text{phos}} =$ 530  $\mu$ s. The photodynamics of 2a at 77 K, however, is different from that of 1a and 3a: both DF and phosphorescence of 2a are of comparable intensities initially  $(\sim 1 \,\mu s)$  in the 77 K ns-TRE spectra (Fig. 6); in addition, DF and phosphorescence do not follow first-order kinetics but decay according to the power law  $(I \propto t^{-1})$  in the time interval 1 µs to 1.2 ms (inset of Fig. 6). The thermally induced Stokes shifts ( $\Delta E_{\rm s} = E_{00} (77 \text{ K}) - E_{00} (298 \text{ K})$ ), being  $\sim$ 0 (2a) and  $\sim$ 107 cm<sup>-1</sup> (3a), are small, thus supporting that the phosphorescence bands are originated from <sup>3</sup>IL.<sup>15</sup> Moreover, as the emission energies and profiles of the lowenergy bands are similar to those of the steady-state emission Edge Article<br>
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Fig. 5 ns-TRE spectra of (a) 1a and (b) 3a in 77 K glassy solution (EtOH/ MeOH = 4 : 1) recorded at different time intervals.  $\lambda_{\text{exc}} = 355$  nm; integration time: 80 and 200 µs for 1a and 3a, respectively. Insets of (a) and (b) show the kinetic decay traces at the specified wavelengths with the estimated phosphorescence lifetime  $(\tau_{\text{phos}})$ .



Fig. 6 ns-TRE spectra of 2a in 77 K glassy solution (EtOH/MeOH  $=$ 4 : 1) recorded at different time intervals.  $\lambda_{\text{exc}} = 355$  nm; integration time: 200 µs. Inset shows the log-log plot of emission intensity of DF ( $\lambda_{\text{DF}}$  = 500 nm) and phosphorescence ( $\lambda_{\text{phos}}$  = 596 nm) of 2a in 77 K glassy solution against time; both decay according to a power law:  $I \propto t^{-1}$ .

spectra of the  $Au(m)$  analogues, the low-energy emission bands of 1a–3a are assigned to be from phosphorescence decay of the  $\sqrt[3]{\pi \pi^*}$ (C $\equiv$ CR) excited state.

Nanosecond transient absorption (ns-TA) difference spectra of 1a–3a (Fig. 7) and 5a–6a (Fig. S15 in ESI†) have been recorded in deoxygenated  $CH_2Cl_2$  at a gate delay of 1 µs after excitation at  $\lambda = 355$  nm. The ns-TA spectra are characterized by an intense positive signal due to excited-state absorption (ESA) within the spectral range 400–700 nm. The decay time constants of the lowest-energy ESA ( $\tau_{\text{ESA}}$ ) are 20.3 (1a), 39.2; 304 (2a), and 13.3;  $70.9 \,\mu s$  (3a) (insets of Fig. 7). Changing the auxiliary ligand from  $PCy_3$  (1a) to RNC (5a) and NHC (6a) results in negligible changes in the ns-TA spectra and  $\tau_{ESA}$  (Fig. 7 vs. S15†), suggesting that auxiliary ligand has little effect on the photophysics of the gold(I) arylacetylide complexes.

#### Time-resolved spectroscopies for the gold $(m)$  complexes

ns-TRE and ns-TA difference spectra of 1b–3b have been recorded in degassed  $CH_2Cl_2$  solutions at 298 K at a gate delay of 1 µs. The

ns-TRE spectra of 1b–3b (Fig. S16 in ESI†) have the same emission profiles and peak positions as the corresponding steadystate phosphorescence spectra and exhibit single exponential decay lifetimes of 20.8 (1b), 9.7 (2b) and 18.3  $\mu$ s (3b). For the ns-TA difference spectra of 1b–3b (Fig. 8, bottom panel), a broad positive ESA band was observed in the spectral region 450–800 nm; this ESA signal follows first-order kinetics with lifetimes determined to be 23.8 (1b), 13.6 (2b) and 25.4  $\mu$ s (3b), in reasonable agreement with the phosphorescence decay lifetimes determined from their respective ns-TRE spectra, thus indicating that the broad ESA is derived from  $T_1 \rightarrow T_n$  absorption.

To probe the early excited state dynamics of the  $\text{gold(m)}$ complexes, in particular the events associated with ISC, femtosecond time-resolved fluorescence (fs-TRF) and transient absorption difference spectra (fs-TA) of 1b–3b have been recorded. Fig. 8 depicts the fs-TRF (top panel) and fs-TA spectra (middle panel) of complexes  $1b-3b$  in CH<sub>2</sub>Cl<sub>2</sub> solution at various time intervals after 400 nm excitation at 298 K. Promptly  $({\leq}2$  ps) after photo-excitation, an unstructured fluorescence band peaking at 461 (1b), 473 (2b) and 459 nm (3b) appears and decays completely within 100 ps. As the TRF emission peaks and profiles closely resemble those of their  $Au(i)$  analogues, 1a-3a, these TRF spectra are suggested to be originated from the  $1\pi\pi$ <sup>\*</sup>(C $\equiv$ CR) excited state. Fitting of the kinetic traces at their peaking wavelengths reveals that bi-exponential functions are required for 1b–3b with  $\tau_1$  and  $\tau_2$  being 1.28 and 13.8 ps for 1b, 0.95 and 9.04 ps for 2b, and 0.74 and 5.22 ps for 3b.

In the fs-TA of 1b–3b (Fig. 8, middle panel), all three complexes displayed similar spectral transformations: the initially formed  $(\sim1.4-2.5 \text{ ps})$  excited state absorption peaking at  $\sim$ 490 nm (ESA1) decays with a concomitant growth of a broad band covering a spectral region 450–800 nm (ESA2) and is fully developed within 40 ps and persists up to 2.7 ns (the longest time recorded in the fs measurements). Clear isosbestic points could be observed at  $\sim$ 500 nm (1b), 530 nm (2b) and  $\sim$ 500 and 700 nm (3b) during the temporal evolution. Such kind of spectral conversion points to a precursor–successor relationship between ESA1 and ESA2. Kinetic analyses at representative wavelengths of these TA spectra reveals that ESA1 of 1b and 3b decay bi-exponentially with  $\tau_1$  and  $\tau_2$  being 0.80 and 13.2 ps for 1b and 0.63 and 3.49 ps for 3b, respectively, whereas ESA1 of 2b decays with a single exponential time constant of  $\tau_2 = 8.38$  ps.



Fig. 7 Nanosecond transient absorption (ns-TA) difference spectra of  $1$ a–3a recorded at selected decay times in degassed CH<sub>2</sub>Cl<sub>2</sub> (5  $\times$  10<sup>-5</sup> M) at 298 K. Insets show the ESA kinetic decay trace at the specified wavelengths; decay lifetimes were fitted as mono-exponential decay for 1a and bi-exponential decays for 2a and 3a. ( $\lambda_{\rm exc} = 355$  nm; integration time: 200 ns).



Fig. 8 (Top) fs-TRF spectra and (middle) fs-TA difference spectra of 1b–3b in CH<sub>2</sub>Cl<sub>2</sub> (5  $\times$  10<sup>-5</sup> M) at 298 K ( $\lambda_{\rm exc}$  = 400 nm; 120 fs fwhm). Arrows indicate the spectral evolution. (Bottom) ns-TA difference spectra of 1b-3b in degassed CH<sub>2</sub>Cl<sub>2</sub> (laser  $\lambda_{\text{exc}}$ : 355 nm). Insets show the kinetic time profiles and the decay time constants at the specified wavelengths.

ESA2, on the other hand, grows with first-order kinetics for all three complexes **1b–3b** with time constants  $\tau_{ESA2} = 9.93$  (**1b**), 4.64 (2b) and 5.73 ps (3b). Given the similar decay time constants between the fs-TRF and ESA1 in fs-TA of 1b–3b, the spectral dynamics for both time-resolved spectra should be originated from the same  $S_1$  excited state, namely, the  $n^4\pi\pi^*(C\equiv CR)$  excited state as revealed in the fs-TRF. On the other hand, comparing the ESA2 in fs-TA spectra at the longest time recorded with the corresponding ns-TA spectra for each  $Au(m)$  complex (Fig. 8, bottom panel), the two spectra are similar, indicating that ESA2 is derived from  $T_1 \rightarrow T_n$  absorption. Because there is a precursor–successor relationship between the ESA1 (S<sub>1</sub>  $\rightarrow$  S<sub>n</sub> absorption) and ESA2 (T<sub>1</sub>  $\rightarrow$  T<sub>n</sub> absorption),  $\tau_2$  of ESA1 is assigned to ISC from the S<sub>1</sub> excited state to a receiving triplet excited state, which then internally converted to the  $T_1$  excited state with an ultrafast time scale. Thus,  $\tau_{\text{ISC}} = 13.2 \text{ ps } (\mathbf{1b})$ , 8.38 ps (2b) and 3.49 ps (3b). The short  $\tau_1$  = 0.80/1.28 (1b), 0.95 ps (2b) and 0.63/0.74 ps (3b) of ESA1/TRF may likely correspond to the  $S_1$  vibrational relaxation.

#### Intersystem crossing rate

The spectroscopically determined intersystem crossing rate constants ( $k_{\text{ISC}}$ ) and the corresponding time constants ( $\tau_{\text{ISC}}$ ) for both  $Au(i)$  and  $Au(m)$  complexes studied herein are tabulated in Table 4. For 1a–3a and 5a–6a, assuming that the major non-





 $k_{\text{ISC}}$  for 1a–3a, 5a–6a are calculated according to eqn (1).  $b \tau_{\text{ISC}} = 1/k_{\text{ISC}}$  of 1b–3b is estimated from  $\tau_2$  obtained from the fs-TA spectra of ESA1.

radiative decay of the S<sub>1</sub> excited state is ISC, *i.e.*  $k_{\text{nr}} \approx k_{\text{ISC}}$ , an upper bound approximation of the  $k_{\text{ISC}}$  values could be obtained by eqn (1):

$$
\Phi_{\rm em} = \frac{k_{\rm r}}{k_{\rm r} + k_{\rm ISC}}\tag{1}
$$

The estimated  $k_{\text{ISC}}$  for the gold(i) complexes are 7.5  $\times$  10<sup>6</sup> to  $1.1 \times 10^8 \text{ s}^{-1}$  and the intersystem crossing time constants ( $\tau_{\text{ISC}}$ ) are 9.0-133 ns. These  $\tau_{\text{ISC}}$  are much larger than those of many phosphorescent transition-metal complexes ( $\tau_{\text{ISC}}$  in the femtosecond to picosecond timescale). For 1b-3b, the  $\tau_{\text{ISC}}$  values are more than three orders of magnitude faster than their  $\text{gold}(1)$ analogues; these ISC rates, nevertheless, are comparable to other transition-metal complexes where  $S_1 \rightarrow T_1$  ISC is mediated by a higher-lying  $T_n$  triplet excited state.<sup>16,17</sup>

### Computational study

The different luminescence behaviors between the  $Au(1)$  and  $Au(m)$  systems were investigated by DFT/TDDFT calculations. The pair (1a, 1b) was chosen as a representative example to examine why the Au(I) complexes studied herein display only fluorescence while the  $Au(III)$  counterparts exhibit exclusively phosphorescence. As the Bodipy-functionalized complexes give fluorescence for both  $Au(i)$  and  $Au(m)$  complexes, DFT/TDDFT calculations were also performed on the pair, (4a, 4b). To save computational time, the cyclohexyl groups of the phosphine ligands in 1a and 4a were replaced by methyl groups.

#### Calculations on 1a and 1b

The frontier MO diagrams of 1a and 1b are shown in Fig. 9. The HOMO and LUMO for both complexes 1a and 1b are predominantly localized on the arylacetylide ligand. For 1b, a considerable contribution (17%) from the C $\gamma$ N $\gamma$ C moiety to the LUMO is also noted. The energy gap between HOMO and H-1 in 1a is approximately  $0.88$  eV. For  $1a$ , the H $-1$  is comprised of the antibonding combinations of Au(d<sub>xy</sub>) and  $\pi$ (C $\equiv$ C) orbitals with little

involvement of the phosphine ligand. For  $\bf 1b$ ,  $\bf H-1$  is composed of Au(d<sub>xz</sub>) and the  $\pi$ (C^N^C) orbitals; the energy gap between HOMO and  $H-1$  in 1**b** is only 0.2 eV. The  $H-2$  of 1**b** is made up of an antibonding combination of the Au(d<sub>xy</sub>),  $\pi$ (C $\equiv$ C) and  $\sigma$ (C $\wedge$ N $\wedge$ C) orbitals, with a HOMO/H $-2$  orbital energy gap of only  $\sim$ 0.4 eV. Clearly, the cyclometalated  $[C^{\wedge}N^{\wedge}C]$  ligand has a role in destabilizing the  $Au(d)$  orbitals. Therefore, the HOMO and  $H-1/H-2$ energy gaps in 1b are much smaller than that in 1a.

The energies of the singlet and triplet excited states and the associated nature and composition for 1a and 1b at their respective optimized singlet ground state geometries are obtained by TDDFT and are shown in Table S5 and S6 in ESI.† For 1a, there is only one triplet excited state  $(T_1)$  which is more than 10 000  ${\rm cm^{-1}}$  below S<sub>1</sub>. In addition, both S<sub>1</sub> and T<sub>1</sub> excited states are of the same parentage and are derived from HOMO  $\rightarrow$ LUMO transition  $(\sim)90\%$  and thus, there would be no effective SOC between them. The triplet excited states above  $S_1$  were also considered; the closest lying  $T_m$  excited state with efficient SOC is when  $m = 4$ , which is derived from a H $-1$  to LUMO transition (90% H-1  $\rightarrow$  L). However, the energy separation  $\Delta E(S_1-T_4)$  is  $-3180$  cm<sup>-1</sup>, which is too large to be overcome by thermal activation. Openical Science<br>
Clearing the S, excited state is EC, i.e.  $k_{\text{av}} \sim k_{\text{av}}$ , and modelling contribution of the Au(a) = 10 December 2016. Downloaded by each<br>
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On the other hand, for 1b, there are four triplet excited states which are lower-lying than  $S_1$ , of which the closest-lying  $T_4$ excited state is only  $\sim$ 70 cm<sup>-1</sup> below the S<sub>1</sub> excited state. Thus, thermal energy at room temperature assists facile ISC, even though SOC is small between the  $S_1$  and  $T_4$  excited states ( $|S_1|$ )  $H_{\text{SOC}}|T_4\rangle|^2 \sim 1.5 \text{ cm}^{-2}$ ). In addition, among the triplet excited states above  $S_1$ , there is a close-lying  $T_5$  excited state derived from the H-2  $\rightarrow$  LUMO transition (79%) which lies only 390 cm<sup>-1</sup> above the  $S_1$  excited state and ISC from  $S_1$  to  $T_5$  could be thermally activated. Besides, owing to the different orientations of the d-orbitals in HOMO and H $-2$ , the  $S_1$  and T<sub>5</sub> excited states could have effective SOC ( $|\ll_1|H_{\text{SOC}}|T_5>|^2 \sim 4.1 \times 10^3 \text{ cm}^{-2}$ ).

#### Calculations on 4a and 4b

The Frontier MOs for 4a and 4b are shown in Fig. 10. Relative to 1a and 1b, the HOMO is destabilized and the LUMO is stabilized



Fig. 9 Frontier MOs of 1a and 1b at the optimized  $S_0$  geometries. Orbital energies are also given in eV.



Fig. 10 Frontier MOs of 4a and 4b at their optimized  $S_0$  geometries. Orbital energies are also given in eV.

for 4a and 4b. Even for 4b, which contains a cyclometalated  $[C^{\wedge}N^{\wedge}C]$  ligand, the LUMO is predominantly localized on the Bodipy-functionalised arylacetylide ligand. The HOMO is composed of an antibonding combination of the  $Au(d_{vz})$  and  $\pi$ (C $\equiv$ CBodipy) orbitals. The energies and compositions of the singlet and triplet excited states of 4a and 4b at their respective optimized singlet ground state geometries were obtained by TDDFT and are collected in Tables S7 and S8 in ESI.† For this pair, (4a, 4b), the two triplet excited states,  $T_1$  and  $T_2$ , are more than 2000  $\mathrm{cm}^{-1}$  below the  $\mathrm{S}_1$  excited state and are all composed of Au $(d_{yz})$  orbitals. As SOC between the coupling singlet and triplet excited states would be ineffective with d-orbitals of the same orientation, ISC from  $S_1$  to  $T_2$  (or  $T_1$ ) for 4a and 4b would be sluggish.

For  $4a$ , the  $T_3$  excited state is the closest-lying triplet excited state that could have effective SOC with the  $S_1$  excited state due to a minor contribution of the  $H-5 \rightarrow LUMO$  transition to the T<sub>3</sub> excited state (H-5 is composed of the Au( $d_z^2$ ) orbital); however,  $\Delta E(S_1 - T_3)$  is  $-2955$  cm<sup>-1</sup> (negative sign indicates that  $T_3$  lies above  $S_1$ ) which is much larger than the thermal energy. For  $4b$ , the  $T_3$  excited state is also the closest-lying triplet excited state that could have effective SOC with the  $S_1$  excited state due to a minor contribution of  $H-1 \rightarrow LUMO$  transition in the  $T_3$ excited state (the d-orbitals of the  $Au(m)$  ion at the HOMO and H-1 of 4b are of different orientations, Fig. 10). However, the singlet-triplet gap,  $\Delta E(S_1 - T_3) = -1192$  cm<sup>-1</sup>, is also much larger than the thermal energy. Thus, the pair (4a, 4b) is expected to have slow ISC rates, when taking into consideration both the singlet–triplet energy gaps and SOC.

### **Discussion**

#### General remarks on the photophysical properties

The emissions of the  $Au(i)$  complexes, 1a-6a are attributable to  ${}^{1}$ IL  $\pi\pi^{*}$ (C $\equiv$ CR) excited states. Pronounced red shifts in emission  $\lambda_{\text{max}}$  of arylacetylides can be observed upon their coordination to Au(I) ion (e.g.  $\lambda_{\text{max}} = 412 \text{ nm}$  (L1) vs. 467 nm (1a)). Considering the complexes 1a, 5a and 6a, which have different neutral auxiliary ligands (phosphine (PCy<sub>3</sub>, 1a), isocyanide (RNC, 5a) and N-heterocyclic carbene (NHC, 6a)) but the same acetylide ligand with a benzothiadiazole moiety, the lowest energy emission  $\lambda_{em}$  red shifts with the auxiliary ligand from 456 nm (RNC) to 467 nm (PCy<sub>3</sub>) to 476 nm (NHC). A rationalization would be that the NHC, being the strongest  $\sigma$ -donor ligand among the auxiliary ligands in the three complexes, destabilizes the Au(d) orbital to the greatest extent. From DFT calculations, the HOMO is comprised of a Au(d) orbital and  $\pi$ (C $\equiv$ CR) (Fig. 9). Thus, the more electron-donating the auxiliary ligand, the more destabilized the HOMO, and hence, the smaller the HOMO–LUMO gap and the  ${}^1\pi\pi^*(C\text{+CER})$  energy. A similar trend in the lowest energy absorption  $\lambda_{\text{abs}}$  can also be observed on changing the auxiliary ligand from RNC (371 nm) to  $PCy_3$  (379 nm) and NHC (383 nm).

Most of the reported luminescent cyclometalated  $Au(III)$ complexes display phosphorescence that comes from the  $^3\pi\pi^*$ IL excited state localized on the cyclometalated ligands.<sup>13</sup> For the Au( $\text{III}$ ) complexes studied herein, 1b-3b, the lowest-

energy triplet excited states are of  $\pi \pi * (C \equiv CR)$  in nature, with  $\lambda_{\text{max}} = 630, 592$  and 603 nm respectively. These complexes, however, exhibit rather weak phosphorescence, with  $\Phi_{\rm em}$  values in the range of 0.003–0.04. It is noted that in 77 K glassy solutions, the phosphorescence lifetimes are signicantly increased compared with those obtained in degassed  $CH_2Cl_2$  at RT (e.g. 205  $\mu$ s at RT to 2.2 ms at 77 K for 3b). Since low temperature and rigid glassy matrix can impede structural distortion, the lifetimes obtained at 77 K could reflect the intrinsic radiative lifetime of the complexes. The especially long emission lifetimes can reflect the predominant localization of the emitting  $T_1$ excited state on the arylacetylide ligand, *i.e.*  $\sqrt[3]{\pi \pi^*}$  (C=CR) with little participation of the metal ion. This is also corroborated by the small thermally induced Stokes shifts ( $\Delta E_{\rm s}$  =  $E_{00}$  (77 K) –  $E_{00}$ (298 K)) of less than 600  $cm^{-1}$  (Table 2).

#### Intersystem crossing in  $gold(i)$  and  $gold(m)$  complexes

ISC is usually fast in transition-metal complexes with time constants ( $\tau_{\text{ISC}}$ ) in the fs to ps time regimes. In the literature, there are numerous examples which show ultrafast ISC,  $1,18-25$  e.g.  $[M(bpy)_3]^{2+}$   $(M = Ru \text{ or } Fe, \tau_{ISC} = 30 \text{ fs})$ ,<sup>19*a*,b</sup>  $[Re(L)(CO)_3(bpy)]$  $(\tau_{\text{ISC}} = 100-140 \text{ fs})$ ,<sup>20</sup> [Ir(piq)<sub>3</sub>] (piq: 1-phenylisoquinoline;  $\tau_{\text{ISC}}$  = 70 fs),<sup>21b</sup> [Pt(PBu<sub>3</sub>)<sub>2</sub>(C=CPh)<sub>2</sub>] ( $\tau_{\text{ISC}}$  = 70 fs),<sup>22</sup> and [Cy<sub>3</sub>PAu(2-naphthyl)] ( $\tau_{\text{ISC}} = 230 \text{ fs}$ )<sup>23</sup> etc. These  $\tau_{\text{ISC}}$  correspond to rates of intersystem crossing  $(k_{\text{ISC}})$  in the range of  $10^{12}$  to  $10^{13}$  $s^{-1}$ . The fast  $k_{\rm{ISC}}$  in transition-metal complexes is traditionally attributed to a large spin–orbit coupling (SOC) constant inherited from the heavy metal atom. However, there are increasing number of reports revealing slow ISC rate ( $k_{\text{ISC}} \sim 10^8$ )  $s^{-1}$ ) in spite of the presence of heavy transition metal, such as the cases of Rh( $i$ )- and Ir( $m$ )-bis(arylethynyl)cyclopentadiene,<sup>3</sup> Au(I)-pyrene,<sup>4b</sup> Pt(II)-perylene/<sup>4c</sup> tetracene,<sup>4e</sup> and Pd(II)-perylene diimide;<sup>4f</sup> all these complexes contain highly conjugated ligand systems and display ligand-dominated  $\frac{1}{2}\pi\pi^*$  fluorescence. There are also cases where comparable  $k_{\text{ISC}}$  and  $k_{\text{r}}$  of  $S_1 \rightarrow S_0$ leads to the observation of dual fluorescence-phosphorescence under steady-state condition,  $e.g.,$  [Pt(L)(acac)] and [Ir(L)(acac)]  $(L = 2$ -(oligothienyl)pyridine);<sup>26</sup>  $[Os(L)(CO)<sub>3</sub>X]$  (L = 8-quinolinolate<sup>27</sup> or isoquinoline-triazole),<sup>28</sup> and  $[Bu_4N]_4[Pt_2(\mu P_2O_5(BF_2)_2)_4$ <sup>29</sup> etc. Edge Article<br>
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> In this work, the luminescence behaviour of the  $Au(1)$  and  $Au(m)$  complexes are drastically different, even though they have the same metal and arylacetylide ligands. Ligand-dominated fluorescence has been observed with the Au(I) complexes, 1a-3a and 5a–6a, with  $k_{\text{ISC}}$  estimated to range from 7.5  $\times$  10<sup>6</sup> to  $1.1 \times 10^8$  s<sup>-1</sup>. The Au(III) complexes **1b-3b**, on the other hand, display phosphorescence, with  $k_{\text{ISC}}$  estimated to be larger than  $10^{10}$  s<sup>-1</sup>. The major difference between the two series of gold complexes is the oxidation state of Au ion, that dictates the coordination geometry, *i.e.* a linear geometry for the  $Au(1)$ complexes,  $1a-6a$ , and a square-planar geometry for Au(III) complexes, **1b–4b**. The coordination geometry has a significant impact on the relative energies of the frontier orbitals (specifically, the d-orbital energies) and hence the relative energies of the singlet and triplet excited states, which subsequently affect the  $k_{\text{ISC}}$ .

The two factors that determine the  $k_{\text{ISC}}$  are (1) the SOC matrix element  $\langle S_n|H_{\text{SOC}}|T_m\rangle$ , and (2) the energy gap ( $\Delta E_{\text{ST}}$ ) between the coupling singlet  $(S_n)$  and triplet  $(T_m)$  excited states. The larger the  $H_{\text{SOC}}$  and the smaller the energy gap ( $\Delta E_{\text{ST}}$ ), the faster will be  $k_{\text{ISC}}$ . For effective SOC, this requires the metal d-orbitals of the coupling singlet and triplet excited states to have different orientations. For example, if  $S_n$  is derived from a metal-to-ligand charge transfer (MLCT) excited state where  $Au(d_{xz})$  orbital is involved,  $H_{\text{SOC}}$  would be zero if the triplet excited state is also an MLCT state that involves  $Au(d_{xz})$  orbital because of symmetry reasons.

The pair (1a, 1b) has been chosen as a representative example to illustrate the different photophysical properties exhibited by the  $Au(i)$  and  $Au(m)$  arylacetylide complexes studied in this work. From the DFT/TDDFT calculations, it is revealed that owing to the inherent linear coordination geometry of the  $Au(i)$  complex, the dorbitals of the gold $(i)$  ion is mainly destabilized by the arylacetylide ligand (Fig. 9 and Table S9 in ESI†). On the other hand, as  $Au(m)$  complexes are assumed to have a square-planar four-coordinated geometry, thus, in addition to the antibonding interactions with the arylacetylide ligand, the d-orbitals of  $\text{gold}(m)$  ion could also be destabilized by the cyclometalated  $[C^NC]$  ligand (both  $\pi$ -type, *e.g.* H $-1$ , and  $\sigma$ -type, *e.g.* H $-2$  in 1**b**; Fig. 9 and Table S9 in ESI†); these latter interactions result in smaller d-orbital splittings in the Au( $\text{m}$ ) series than the Au( $\text{i}$ ) series. In effect, S<sub>1</sub> and  $\mathrm{S}_2$  excited states are  ${\sim}4200$   $\mathrm{cm}^{-1}$  apart for  $\bf{1}$ a while the analogous splitting (between  $S_1$  and  $S_3$  excited states) is only  $\sim$ 1300  $\rm cm^{-1}$  for **1b.** As S<sub>2</sub> of **1a** is derived from <sup>1</sup>[Au(d<sub>xy</sub>)  $\rightarrow \pi^*(C\equiv CR)]/$ <sup>1</sup>[ $\pi(C\equiv C)$ ]  $\rightarrow \pi^*(C\equiv CR)]$  (<sup>1</sup>MLCT/<sup>1</sup>ILCT) and S<sub>3</sub> of **1b** from <sup>1</sup>[Au(d<sub>xy</sub>)  $\rightarrow$  $\pi^*(C\equiv CR)]/{}^{1}[\pi(C\equiv C)\rightarrow \pi^*(C\equiv CR)]/{}^{1}[\pi(C^{\wedge}N^{\wedge}C)\rightarrow \pi^*(C\equiv CR)]$ (<sup>1</sup>MLCT/<sup>1</sup>ILCT/<sup>1</sup>LLCT), *i.e.*, both are of charge-transfer type excited states, the singlet–triplet energy gaps for this type of transitions are small ( $\Delta E(S_2-T_4) \sim 1000$  cm<sup>-1</sup> for 1a and  $\Delta E(S_3-T_5) \sim 900$  cm<sup>-1</sup> for 1b)  $(T_4$  (1a) and  $T_5$  (1b) are the triplet counterpart of  $S_2$  (1a) and  $S_3$  (1b) respectively). As depicted in Fig. 11, the  $S_1/T_5$  energy gap for 1b is small but the  $S_1/T_4$  energy gap for 1a is large. In other words, the oxidation state of the gold ion affects the coordination geometry of the complex, which in turn change the interactions between the metal d-orbitals and ligand orbitals, giving rise to different d-orbital splitting and subsequently the singlet–triplet splitting  $(\Delta E_{ST})$  of the two coupling excited states in the gold complexes. Openical Science<br>
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Moreover, DFT/TDDFT calculations also revealed that there is a triplet excited state  $(T_4)$  almost isoenergetic with the  $\mathrm{S}_1$  excited state (<70  $\mathrm{cm}^{-1}$  below the  $\mathrm{S}_1$  excited state) in  $\bf{1b}$ such that even though the SOC between  $S_1$  and  $T_4$  is small due to the similar d-orbital orientations involved in both excited states, thermal energy could promote facile ISC. With 1a, the closest triplet excited state  $(T_2)$  to the  $S_1$  excited state is more than 500  $\mathrm{cm}^{-1}$  above the  $\mathrm{S}_1$  excited state, which is more than twice the thermal energy at room temperature and SOC is also small between these two excited states as the d-orbitals involved are also of the same orientations. Thus, taken together both the SOC and  $\Delta E_{\textrm{ST}}$ , 1b should have a much faster  $k_{\text{ISC}}$  than 1a.

On the other hand, for the Bodipy-functionalized complexes, 4a and 4b, only  ${}^1\pi\pi^*(C\text{ }\equiv$ CBodipy) fluorescence

with no long-lived species are observed under ns-TRE and ns-TA measurements. The photophysical behavior of the Bodipyfunctionalized complexes can be attributed to the intrinsically small band-gap of the Bodipy moiety. Due to the highly conjugated structure of Bodipy, the HOMO is much destabilized and there is a wide orbital energy gap between the HOMO and other occupied MOs, even in the case of 4b which contains a [C^N^C] ligand. As a result, the  $HOMO/H-x$  orbital energy gap is the largest among the four arylacetylide ligands studied herein  $(H-x)$  is the other occupied orbitals lower in energy than the HOMO;  $x = 1, 2, ...$ ). In effect, the closest  $T_m$  excited state that could have effective SOC with  $S_1$  is more than 1000  $cm^{-1}$  above the S<sub>1</sub> excited state. With such a large  $\Delta E(S_1-T_m)$ , thermal energy would be insufficient to promote ISC. Therefore, similar to the scenario in the case of 1a (Fig. 11, left), ISC is sluggish for  $Au(II)$  and  $Au(III)$  arylacetylide complexes bearing Bodipy.

#### Mechanism for the generation of delayed fluorescence

From the ns-TRE measurements of 1a–3a, DF contributes to the total fluorescence, though only a minute proportion  $($ <3%, Table 3). In general, the mechanism of DF could be inferred from the dependence of the DF intensity  $(I_{\text{DF}})$  with the power of excitation light.<sup>30</sup> According to Bässler, a quadratic dependence of excitation power with the DF intensity indicates that the mechanism of the DF is TTA with dominant phosphorescence.<sup>30</sup><sup>b</sup> On the other hand, a linear dependence of DF intensity with excitation power could be due to three possible mechanisms: TTA with dominant delayed-fluorescence, TADF, and GP-recombination. As depicted in Fig. 12, the plot of  $I_{\text{DF}}$  against excitation intensity in double-logarithm scale gave a slope of 1.71  $\approx$  2 for 1a; this nearly quadratic dependence is most consistent with the TTA mechanism with dominant phosphorescence. However, for 2a and 3a (slope  $=$ 0.916 and 1.10), both display nearly linear dependence between  $I_{\text{DF}}$  and excitation intensity. Therefore, it is not possible to confirm the mechanism for DF in the case of 2a and 3a by solely considering the excitation power dependence.

Time-dependence of  $I_{\text{DF}}$  and phosphorescence intensity  $(I_{\text{P}})$ could also give hints to the DF mechanism.<sup>30,31</sup> For TTA with dominant DF, phosphorescence intensity decays with a power law,  $I_{\text{P}} \propto t^{-1}$  while  $I_{\text{DF}}$  is approximately constant at short time and  $I_{\text{DF}} \propto t^{-2}$  at longer time. For the GP-recombination mechanism, both DF and phosphorescence decay in accordance with the power law,  $I_{\text{DF}} \propto t^{-1}$ , at both short and long times.<sup>30a</sup> In the case of 2a in 77 K glassy solution, both DF and phosphorescence decayed according to the power law:  $I \propto t^{-1}$ over the time intervals investigated  $(1 \mu s)$  to 1.2 ms) (Fig. 6, inset), suggesting that the DF mechanism under this condition is most likely the GP-recombination mechanism. As for 3a, there is no power law decay relation with both DF and phosphorescence and so it seems unlikely that GP-recombination is the mechanism for the generation of DF in 3a. There is still not enough information to conclude on the DF mechanism for 3a.



Fig. 11 Illustration of the low-lying singlet and triplet excited states of Au(I) (left) and Au(III) (right) complexes that accounts for the different photophysical behaviour of the Au(i) and Au(iii) complexes investigated in this work. S<sub>1</sub> and T<sub>1</sub> for both complexes are derived from HOMO  $\rightarrow$ LUMO transitions; S<sub>2</sub> and T<sub>4</sub> excited states of **1a** are derived from  $^{1.3}$ [H-1  $\rightarrow$  LUMO] transitions while S<sub>3</sub> and T<sub>5</sub> excited states of **1b** are derived from  $^{1,3}$ [H $-2 \rightarrow$  LUMO] transitions. The d-orbitals involved in the T<sub>2</sub> of 1a and T<sub>4</sub> of 1b have the same orientations as their respective S<sub>1</sub> excited state (see Tables S5, S6 and S9 in ESI†). The wavy blue arrows indicate internal conversion (IC) from the T<sub>5</sub> to T<sub>1</sub> excited state. F = fluorescence and  $P =$  phosphorescence.



Fig. 12 Dependence of delayed fluorescence of  $1a-3a$  in  $5 \times 10^{-5}$  M degassed  $CH<sub>2</sub>Cl<sub>2</sub>$  with excitation power intensity. Emission intensity measured after a time delay of 1 µs. (Laser  $\lambda_{\text{exc}} = 355$  nm; 0.2–11 mJ per pulse; diameter  $= 8$  mm, integration time: 800  $\mu$ s.)

### Conclusion

A series of gold complexes with different oxidation states,  $\text{gold}(I)$  complexes  $\text{[LAu(C=CR)]}$  and  $\text{gold}(II)$  complexes  $[Au(C^NC^C)(C\equiv CR)]$  bearing the same heterocyclic arylacetylides with narrow bandgap were synthesized and characterized. The photophysical behaviors with the gold ion in different oxidation states are strikingly different: fluorescence dominates the luminescence of the  $Au(r)$  complexes while phosphorescence takes over in the  $Au(m)$  complexes. Detailed computational studies by DFT/TDDFT have accounted for these phenomena as a result of different coordination environments inherited from the gold ion in a particular oxidation state: a linear coordination geometry for  $Au(1)$  and a square-planar coordination geometry for  $Au(m)$ . This difference in

coordination geometry subtly affects the energy separation between the coupling singlet and triplet excited states, leading to smaller  $\Delta E_{ST}$  of the Au(III) complexes than the Au(I) complexes and hence, larger  $k_{\text{ISC}}$  in the Au(III) complexes than the Au(I) complexes. For the complexes bearing Bodipy-functionalized acetylide ligand, they only display prompt fluorescence. Computational analyses revealed that, due to the especially narrow bandgap of Bodipy, the  $\Delta E_{ST}$  is still large even in the  $Au(m)$  complex so that  $k_{\text{ISC}}$  could not compete with fluorescence radiative decay. Additionally, the mechanisms for the generation of DF in  $Au(i)$  complexes have been explored. To the best of our knowledge, this is the first report which systematically studies the effects of the metal ion oxidation state on the photophysical behaviours of transition-metal complexes.

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