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# Unveiling the intersystem crossing dynamics in N-annulated perylene bisimides†

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The efficient population of the triplet excited states in heavy metal-free organic chromophores has been one of the long-standing research problems to molecular photochemists. The negligible spin-orbit coupling matrix elements in the purely organic chromophores and the large singlet-triplet energy gap  $(\Delta E_{S-T})$  pose a hurdle for ultrafast intersystem crossing (ISC). Herein we report the unprecedented population of triplet manifold in a series of nitrogen-annulated perylene bisimide chromophores (NPBI and Br-NPBI). NPBI is found to have a moderate fluorescence quantum yield ( $\Phi_{\rm f}$  = 68  $\pm$  5%), whereas Br-NPBI showcased a low fluorescence quantum yield ( $\Phi_{\rm f}$  = 2.0  $\pm$  0.6%) in toluene. The femtosecond transient absorption measurements of Br-NPBI revealed ultrafast ISC ( $k_{\rm ISC}=1.97\times10^{10}~{\rm s}^{-1}$ ) from the initially populated singlet excited state to the long-lived triplet excited states. The triplet quantum yields ( $\Phi_{\top}$  = 95.2  $\pm$  4.6% for Br-NPBI,  $\Phi_{T}$  = 18.7  $\pm$  2.3% for NPBI) calculated from nanosecond transient absorption spectroscopy measurements showed the enhancement in triplet population upon bromine substitution. The quantum chemical calculations revealed the explicit role of nitrogen annulation in tuning the excited state energy levels to favor the ISC. The near degeneracy between the singlet and triplet excited states observed in NPBI and Br-NPBI ( $\Delta E_{S-T}=-0.01$  eV for NPBI,  $\Delta E_{S-T}=0.03$  eV for Br-NPBI) facilitates the spin flipping in the molecules. Nitrogen annulation emerges as a design strategy to open up the ISC pathway and the rate of which can be further enhanced by the substitution of a heavier element.

## Introduction

Intersystem crossing (ISC), a non-radiative transition involving spin flipping, is the quintessential process for populating the triplet manifold in chromophoric systems.<sup>1,2</sup> The efficient population and harvest of triplet excitons find diverse applications in organic light-emitting diodes, 3-5 organic photovoltaics, <sup>6-9</sup> photodynamic therapy, <sup>10,11</sup> photocatalysis, <sup>12,13</sup> etc. The long diffusion lengths of the triplet excitons have been exploited in optoelectronic devices to enhance the photocurrent as it increases the possibility of charge separation at donor-acceptor interfaces.8 The generation of reactive singlet oxygen via triplet sensitization from chromophores finds

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applications in photodynamic therapy. 11,14 As a result, chromophores exhibiting efficient ISC are in great demand.

Perylene bisimides (PBIs) are a class of highly sought-after n-type organic semiconductor materials for optoelectronic applications owing to their excellent electron affinity and mobility. 15,16 The large molar absorption coefficient, high stability, easier tunability of electronic properties, etc., are attractive features of PBIs. 16,17 A vital role in controlling the rate of ISC process in organic materials is played by spin-orbit coupling (SOC). 1,18 In addition to SOC, the reduction in the singlet-triplet energy gap is important for enhancing the rate of intersystem crossing.<sup>2</sup> Several strategies have been reported till date to generate triplet excitons, ranging from the conventional heavy atom effect,  $^{19}$  vibronic coupling,  $^{18,20}$  core twist,  $^{21-23}$  multiexciton generating singlet fission,  $^{24-28}$  and charge recombination.29-35 Seminal works by Castellano and coworkers<sup>36</sup> and Würthner and co-workers<sup>37,38</sup> have shown that incorporating transition metals to rylene bisimides enhances SOC-driven ISC. The large nuclear charge associated with transition metals imparts large SOC, thereby enhancing the rate of ISC.2 The cytotoxicity associated with chromophores encompassing heavy metals proposes the need for purely organic molecular materials exhibiting an enhanced triplet population. As an alternative strategy, non-metallic elements,

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including heavier halogens<sup>39–42</sup> and chalcogens<sup>43–47</sup> have been appended to chromophores to enhance the ISC rates. An unusual halogen bond-directed heavy atom effect was observed by Kim and co-workers in phosphorescent crystals of bromine-substituted aromatic carbonyls.<sup>48</sup>

The continuous efforts from our group in populating the triplet manifold have been fruitful in generating near unity triplet quantum yield in eight bromine atoms substituted PBI driven by the synergistic effect of core-twist and spin-orbit coupling.<sup>49</sup> Herein, we report the observation of an ultrafast triplet population in a single bromine atom substituted nitrogen-annulated PBI (Br-NPBI) chromophore. Nitrogenannulated PBI (NPBI) is found to have a moderately large fluorescence quantum yield ( $\Phi_{\rm f}$  = 68  $\pm$  5%). Upon single bromine substitution at the bay position of NPBI, the fluorescence quantum yield remarkably decreases, rendering ultrafast ISC in Br-NPBI ( $\Phi_{\rm f}$  = 2.0  $\pm$  0.6%). On the contrary, the unsubstituted PBI and mono-brominated Br-PBI exhibits strong fluorescence ( $\Phi_{\rm f} \sim 100\%$  for PBI and  $\Phi_{\rm f}$  = 97% for Br-PBI). The transient absorption measurements in the femtosecond and nano-second time regimes explicitly confirm the population of triplet excited states in Br-NPBI and NPBI. The quantum chemical calculations support the observation of triplet formation in Br-NPBI and NPBI, where efficient ISC occurs between the nearly degenerate S1 and T2 excited states  $(\Delta E_{\rm S-T} = -0.01 \text{ eV for NPBI}, \Delta E_{\rm S-T} = 0.03 \text{ eV for Br-NPBI})$ . The high singlet-triplet energy gap forbids the ISC pathway in PBI and Br-PBI. The present work sheds light on the research quest to achieve efficient triplet exciton formation in organic materials for advanced optoelectronic applications.

## **Experimental**

#### Photophysical measurements

Steady-state and time-resolved optical measurements of the compounds were carried out in standard quartz cuvettes having path length of 1 cm for solutions in dried and distilled solvents. Electronic absorption spectra were recorded on Shimadzu UV-3600 UV-Vis-NIR, while emission and gated emission spectra were recorded on Horiba Jobin Yvon Fluorolog 3 spectrometer. The pulse width of the flash lamp for gated measurement is 3  $\mu s$ .

#### Estimating triplet quantum yield $(\Phi_T)$

To estimate the triplet quantum yield ( $\Phi_T$ ) of Br-NPBI/NPBI, the process of triplet energy transfer to  $\beta$ -carotene was used with  $[Ru(bpy)_3]^{2+}$  as the reference ( $\Phi_T \sim 1$ ) as reported in the literature. A known, equal volume of  $\beta$ -carotene in CHCl<sub>3</sub> was added to the optically matched solutions (0.1–0.2 absorbance at 532 nm) of  $[Ru(bpy)_3]^{2+}$  in methanol and Br-NPBI/NPBI in toluene. The energy transfer from  $[Ru(bpy)_3]^{2+}$  or Br-NPBI/NPBI populates the  $\beta$ -carotene triplet, which is monitored at 530 nm. The  $\Phi_T$  values of the PBI derivatives were calculated according to the following equation where super-scripts "Sam" and "Ref" refers to the sample Br-NPBI/NPBI and reference

 $[Ru(bpy)_3]^{2+}$  respectively.  $k_{obs}$  is the pseudo-first-order rate constant for the growth of the β-carotene triplet, and  $k_0$  is the rate constant for the decay of the donor triplets (in the absence of β-carotene) in solutions that contained  $[Ru(bpy)_3]^{2+}$  or Br-NPBI/NPBI with same absorbance.

$$\boldsymbol{\varPhi}_{\mathrm{T}}^{\mathrm{Sam}} = \boldsymbol{\varPhi}_{\mathrm{T}}^{\mathrm{Ref}} \times \frac{\Delta A^{\mathrm{Sam}}}{\Delta A^{\mathrm{Ref}}} \times \frac{k_{\mathrm{obs}}^{\mathrm{Sam}}}{k_{\mathrm{obs}}^{\mathrm{Sam}} - k_{0}^{\mathrm{Sam}}} \times \frac{k_{\mathrm{obs}}^{\mathrm{Ref}} - k_{0}^{\mathrm{Ref}}}{k_{\mathrm{obs}}^{\mathrm{Ref}}}$$

## Singlet oxygen ( ${}^{1}O_{2}$ ) quantum yields ( $\Phi_{\Lambda}$ )

The singlet oxygen quantum yield ( $^{1}O_{2}$ ) measurements were carried out using Rose Bengal (RB) in methanol as a standard having a  $^{1}O_{2}$  quantum yield of  $0.80.^{53,54}$  1,3-Diphenylisobenzofuran (DPBF) in de-aerated dimethylformamide (DMF) solvent acts as the  $^{1}O_{2}$  scavenger, having absorbance around 1.0. The Br-NPBI (photosensitizer), was added into the cuvette (OD  $\sim 0.2$ –0.3) containing DPBF. The cuvette was photoirradiated using of 532 nm monochromatic light, and the absorption spectra was recorded in every 2 s interval. The slope of absorbance maxima of DPBF at 414 nm with irradiation time was plotted. Singlet oxygen quantum yield ( $\Phi_{\Delta}^{\rm Sam}$ ) was calculated according to a modified equation.

$$\Phi_{\Delta}^{\text{Sam}} = \Phi_{\Delta}^{\text{Ref}} \times \frac{k(\text{Sam})}{k(\text{Ref})} \times \frac{F(\text{Ref})}{F(\text{Samp})}$$

where k is the slope of the difference in the change in the absorbance of DPBF (at 414 nm) with irradiation time and F is the absorption correction factor, which is given by  $F = 1-10^{\text{OD}}$  (OD at the irradiation wavelength).

#### Femtosecond transient absorption measurements

A Spectra-Physics Mai Tai SP mode-locked laser (800 nm, 86 MHz) was used as a seed for Spectra-Physics Spitfire Ace regenerative amplifier (1 kHz, 5.5 mJ). Using TOPAS, a portion of the amplified 800 nm output beam was converted into a 470 nm pump pulse. An 800 nm residual pulse passing through the ExciPro pump-probe spectrometer optical delay line (3.5 ns) was used in the generation of a white light continuum using calcium fluoride crystal. White light continuum passes through a beam splitter to produce the probe and the reference pulse. The femtosecond transient absorption spectra of the samples were recorded using a dual diode array detector, having a detection window and optical delay of 200 nm and 3.5 ns, respectively. Sample solutions were prepared in a 1.2 mm path length rotating cuvette. Determination of instrument response function (IRF) is essential for the accurate deconvolution of the spectra. IRF was determined by a two-photon absorption experiment using 10% benzene in a methanol solution and was found to be approximately 110 fs at 530 nm. Samples were excited at 470 nm with 100 fs pulses, and the use of a neutral density filter controlled the incident flux of the sample.

#### Nanosecond transient absorption measurements

Nano-second laser transient absorption experiments of the samples were carried out in an Applied Photophysics Model LKS-60 laser kinetic spectrometer. Nitrogen-purged solution of samples were excited using the second harmonic (532 nm, pulse duration  $\sim 8$  ns) of a Quanta Ray INDI-40–10 series pulsed Nd:YAG laser. Oxygen-purged experiments confirm the presence of the triplet excited state through nanosecond tran-

### Global analysis

sient absorption studies.

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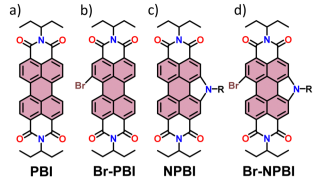
Glotaran software is used for the global analysis of fsTA and nsTA data of the samples. The method evaluates the group velocity dispersion of the white light continuum and instrument response function (IRF), which allows accurate estimation of decay time constants and dispersion-compensated spectra. Global analysis of fsTA data carried out in a sequential model resulted in evolution-associated spectrum (EAS) with decay time constants.<sup>55</sup>

#### Computational details

The ground state geometries of the Br-NPBI, NPBI, Br-PBI, and PBI were optimized using density functional theory (DFT) at CAM-B3LYP56/def2-SVP level of theory. Singlet excited states were computed using the time-dependent DFT (TD-DFT).56 Using the Tamm-Dancoff approximation (TDA) instead of TD-DFT the agreement to the experimental results deteriorates significantly (Table S6, ESI†). For triplet states, however, the Tamm-Dancoff approximation (TDA) were employed since it provides more reliable energies for such states.<sup>57</sup> For all calculations the range-separated CAM-B3LYP functional in combination with def2-SVP and aug-cc-pVDZ<sup>58</sup> basis sets were used. To obtain information about the shapes of the corresponding potential energy curves the S<sub>1</sub> as well as the T<sub>2</sub> equilibrium geometries were also determined. The spin-orbit coupling matrix elements between the spin singlet and triplet excited states were computed at the CAM-B3LYP/aug-cc-pVDZ levels of theory using the PySOC package. 59,60 All the quantum chemical calculations were carried out using Gaussian 16 package. 61

## Results and discussion

In this work, a series of perylene bisimide derivatives (PBI, Br-PBI, NPBI, and Br-NPBI) were synthesized and characterized following previously reported procedures (Fig. 1 and Scheme S1, ESI†).62,63 The geometries of the derivatives in the ground state were optimized using DFT at CAM-B3LYP/def2-SVP levels of theory (Fig. S1, ESI†). Unsubstituted PBI exhibited a planar structure in the ground state, whereas optimized geometry of Br-PBI revealed that the substitution of bromine atom on the bay position of PBI imposed a core twist in the chromophore with a twist angle of 20.6°. The optimized geometry of NPBI showcased a planar scaffold with a bowed geometry. The 5membered heterocyclic ring formed, with a carbon-nitrogen bond length of 1.40 Å, imposed the inward bend in the core. The substitution of the bromine atom in the bay position of NPBI resulted in a negligible twist in the core, resulting in planar Br-NPBI. The insignificant core twist in Br-NPBI resulted



**Fig. 1** Molecular structures of (a) PBI, (b) Br-PBI, (c) NPBI, and (d) Br-NPBI. R in the chemical structure denotes n-hexyl chain.

from the reduced steric repulsion between the bromine and hydrogen atoms in the bay position supported by bowed geometry of the molecule (Fig. S1, ESI†).

The steady-state optical measurements of PBI derivatives were performed in toluene at room temperature. The photophysical properties of PBI, Br-PBI, and NPBI, observed from the steady-state optical measurements, were consistent with the previous reports. The UV-Vis absorption spectra of PBI, Br-PBI, NPBI, and Br-NPBI show distinct vibronic features as a result of the strong coupling between the vinyl stretching modes of the PBI core ( $v_{C=C} \sim 1400~\text{cm}^{-1}$ ), and the  $S_0 \rightarrow S_1$  electronic transition (Fig. 2 and Fig. S2, ESI†). The UV-visible absorption spectrum of Br-NPBI showed two distinct absorption bands centered at 526 nm and 491 nm, corresponding to the 0-0, and 0-1 vibronic transitions, respectively. NPBI exhibited two distinct absorption bands at 527 nm and 492 nm

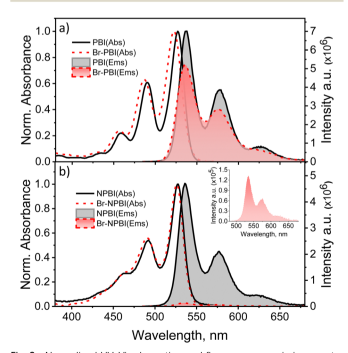


Fig. 2 Normalized UV-Vis absorption and fluorescence emission spectra of PBI and Br-PBI (top) and NPBI and Br-NPBI (bottom) in toluene. The enlarged fluorescence emission spectrum of Br-NPBI is shown in the inset.

**Table 1** Photoexcited properties of PBI, Br-PBI, NPBI, and Br-NPBI.  $\lambda_{abs}$ ; absorption maximum,  $\lambda_f$ ; emission maximum

	$\lambda_{\rm abs}~({\rm nm~eV}^{-1})$	$\lambda_f\left(nm\right)$	$\Phi_{\mathrm{f}}\left(\% ight)$	$\Phi_{\mathrm{T}}$ (%)
PBI	527/2.35	538	$\sim 100^a$	_
Br-PBI	522/2.38	535	97 <sup>a</sup>	_
NPBI	527/2.35	536	$68 \pm 5^{a}$	18.7 $\pm$ 2.3
Br-NPBI	526/2.36	535	$2.0\pm0.6$	$95.2 \pm 4.6$

<sup>&</sup>lt;sup>a</sup> Fluorescence quantum yield was obtained from previously published results63,64

corresponding to the 0-0, and 0-1 vibronic transitions, respectively. The absorption spectrum of Br-PBI showed three distinct vibronic bands at 522, 487, and 457 nm, whereas the unsubstituted PBI showcased vibronic bands centered at 527, 490, and 459 nm (Table 1).

The fluorescence emission spectra of Br-NPBI, NPBI, Br-PBI, and PBI revealed mirror images of the corresponding absorption spectra. Br-NPBI showed emission bands at 535 nm and 571 nm, whereas NPBI has emission bands centered at around 536 and 577 nm. The unsubstituted PBI has emission bands centered at 538, 577, and 627 nm, and Br-PBI has emission bands centered at 535, 575, and 628 nm (Fig. 2 and Fig. S3, ESI†). The fluorescence quantum yields ( $\Phi_f$ ) of the PBI derivatives were measured in toluene at room temperature. NPBI showed fluorescence quantum yield,  $\Phi_{\rm f}$  = 68  $\pm$  5%, compared to  $\Phi_{\rm f} \sim 100\%$  for PBI and  $\Phi_{\rm f}$  = 97% for Br-PBI. Solvent-dependent

fluorescence quantum yield measurements of Br-NPBI and NPBI have been carried out in acetone and acetonitrile solvents (Table S1, ESI†). Surprisingly, bromine substitution on NPBI reduced the fluorescence quantum yield to 2.0  $\pm$  0.6% (Table 1). The observed reduction in the fluorescence quantum yield can be attributed to competitive non-radiative processes occurring from the S1 state.

The heavy atom effect induced by bromine atom substitution is known to enhance the ISC rate. 49,64,65 However, a single bromine substitution on the PBI core has negligible effect on the triplet population.<sup>22</sup> The SOC induced by the single bromine atom on PBI is inadequate for the enhancement of the ISC rate, as evident from the fluorescence quantum yield (Table 1). The reduced fluorescence in NPBI and the remarkably low fluorescence in its mono-brominated derivative, Br-NPBI, suggests the possibility of an enhanced triplet population in the molecules. Vauthey and coworkers reported ultrafast ISC in bromine and nitrogen-substituted naphthalene bisimide chromophore.42 The rate of the ISC can be theoretically determined using the Fermi golden rule where the rate constant of ISC depends on the SOC.66

To understand the competitive decay pathways from the singlet excited state, femtosecond transient absorption (fsTA) measurements were carried out in toluene at room temperature. The fsTA spectra of NPBI were adapted from ref. 63. The fsTA spectra of NPBI (Fig. 3a) after photoexcitation at 470 nm

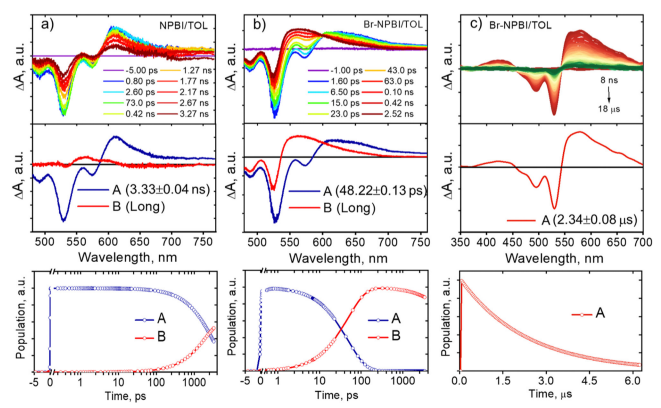


Fig. 3 (Top) Femtosecond transient absorption spectra of (a) NPBI (adapted from ref. 63) and (b) Br-NPBI in toluene. Nanosecond transient absorption spectra of (c) Br-NPBI in toluene. (Middle) EAS reconstructed from global analysis. (Bottom) The relative population profiles of the excited states fitted usina kinetic model

show negative ground state bleach (GSB) ranging from 480 nm to 557 nm. The band at 530 nm have contributions from both GSB and stimulated emission (SE). The band at 574 nm corresponds to higher vibronic bands of SE. The positive excited state absorption (ESA) lies in the region from 587 nm to 655 nm. In the initial delay times, NPBI shows ESA spectral features centered at 609 nm. As the delay time increases, the spectral signal decays with the slow evolution of a weak ESA feature centered at 570 nm. To elucidate the kinetics of the spectral evolution, a global analysis of the fsTA spectra was performed. The evolution-associated spectrum (EAS) of NPBI, was acquired by fitting using a two-component sequential kinetic model (A  $\rightarrow$  B  $\rightarrow$  GS). The two components can be attributed to two distinct excited state transitions,  $S_1 \rightarrow S_n$  and  $T_1 \rightarrow T_n$ . The initially populated singlet state spectral signature decays with the gradual evolution of spectral features corresponding to the triplet manifold. The ESA of the newly formed transient species centered at 570 nm can be attributed to the  $T_1 \rightarrow T_n$  transition. The fsTA spectra of Br-NPBI (Fig. 3b) after photoexcitation at 470 nm showed similar ESA and GSB compared to NPBI in the initial delay time. The positive spectral signature ranging from 613 nm to 627 nm attributed to the initially populated singlet state decay in ultrafast timescale  $(\tau_{A\to B} = 48.22 \pm 0.13 \text{ ps})$ , leading to the population of a new state with ESA centered at 570 nm. The decay of the newly formed species is not complete within the experimental time window. The de-convolution of the fsTA spectra was carried out through global analysis, extracting the EAS. The EAS of Br-NPBI was fitted into two components using a sequential kinetic model (A  $\rightarrow$  B  $\rightarrow$  GS). The first component (A) of the EAS is attributed to the absorption from the first singlet excited state to higher singlet states. The spectral signature associated with this transition decays in ultrafast timescale with  $(k_{\rm ISC})$  with the evolution of the second EAS component. The second component (B) can be attributed to the absorption from the first

Nanosecond transient absorption (nsTA) measurements of NPBI and Br-NPBI in toluene at room temperature were carried out to understand the dynamics of the excited state in the longer delay times. Photoexcitation at 532 nm of nitrogenpurged solution of NPBI resulted in a positive peak centered at around 570 nm and a negative peak corresponding to the ground state bleach ranging from 470 nm to 545 nm (Fig. S4, ESI†). The positive spectral signature at 570 nm was quenched upon oxygen-purging of the NPBI solution, confirming the triplet population (Fig. S5, ESI†). The decay time constant of the triplet excited state of NPBI was found to be 1.42  $\pm$  0.23  $\mu s$ (Fig. S4, ESI†) in the degassed solution with a quantum yield  $(\Phi_{\rm T})$  of 18.7  $\pm$  2.3% (Fig. S6, ESI†). nsTA measurements of nitrogen-purged Br-NPBI (Fig. 3c) solution in toluene showed an intense positive peak centered at around 570 nm after photoexcitation at 532 nm. The nsTA spectra showed ground state bleach in the region from 460 nm to 540 nm. The formation of a triplet excited state in Br-NPBI is confirmed by the depletion in the spectral peak intensity at 570 nm after purging the solution with oxygen (Fig. S7, ESI†). The decay time

triplet excited state to higher triplet states.

constant of the long-lived triplet excited state was found to be  $\tau_{\rm T}=2.34\pm0.08~\mu {\rm s}$ . The triplet quantum yield of Br-NPBI in toluene was estimated to be  $\Phi_{\rm T}=95.2\pm4.6\%$  using a triplet-triplet energy transfer mechanism (Fig. S8, ESI†). The singlet oxygen quantum yield ( $\Phi_{\Delta}$ ) of Br-NPBI is estimated using Rose Bengal (RB) as a standard in methanol. Singlet oxygen quantum yield for Br-NPBI was found to be 86.0% using Rose Bengal as a standard (Fig. S9, ESI†). nsTA measurements of Br-PBI were carried out in toluene at room temperature. Photoexcitation at 532 nm of nitrogen-purged solution of Br-PBI showed no significant positive absorption, indicating the negligible intersystem crossing in the Br-PBI (Fig. S10, ESI†).

The gated emission measurements of Br-NPBI and NPBI were carried out exciting at 470 nm and 472 nm, respectively, at 77 K in toluene. For Br-NPBI (delay time of 0.05 ms, slit width 1.5 mm) the gated emission band was observed at 599 nm with a lifetime of 11.8  $\mu$ s (Fig. S11, ESI†). Gated measurements of NPBI at 77 K exhibited a band at 597 nm (delay time of 0.05 ms, slit width 1.5 mm) with a lifetime of 7.5  $\mu$ s (Fig. S12, ESI†).

To rationalize the role of bromine and nitrogen substitution on PBI molecule in enhancing the ISC rate, we performed quantum chemical calculations. The computed excitation energies with respect to the  $S_0$  ground state in its equilibrium geometry  $R(S_0)$  using the range-separated CAM-B3LYP functional<sup>67</sup> in combination with aug-cc-pVDZ basis sets<sup>58</sup> are summarized in Table 2. The corresponding data using the smaller def2-SVP basis sets<sup>68,69</sup> are given in Tables S2–S5 (ESI†). The data are also plotted in Fig. S14 and S15 (ESI†) for a better overview.

The absorption spectra of the compounds (Fig. 2) show a three-band vibrational progression so that the most intense band given in Table 1 must be assigned to the transition between the lowest vibrational states of So and the S1 (0-0 transition). The other bands then result from excitations to higher vibrational states of the S<sub>1</sub> (Fig. 4). The small difference between  $\lambda_{abs}$  and  $\lambda_f$  also indicates 0-0 transitions for absorption and emission. For all molecules, they differ only by about 0.05 eV, which most likely results from solvent effects. To model the vibrational progression and the exact energetic location of the 0-0 band, calculations that take into account the influence of nuclear motion are necessary, as we have recently shown in a benchmark.<sup>70</sup> However, this benchmark also proved that adiabatic excitation energies (see Fig. 4 and Fig. S13, ESI†) approximate the 0-0 excitation energy to within about 0.1 eV, while simple vertical energies overestimate 0-0 transition energies by up to 0.3 eV.

Table 2 shows similar effects for the molecules examined in this paper. For PBI, the vertical energy of 2.70 eV calculated with CAMB3LYP/aug-cc-pVDZ (Table 2:  $E(S_1)$  at  $R(S_0)$ ) overestimates the experimentally measured 0-0 transition (Table 1: 527 nm = 2.35 eV) by about 0.35 eV. In contrast, the adiabatic excitation energy of 2.45 eV (Table 2:  $E(S_1)$  at  $R(S_1)$ ) deviates by only 0.1 eV, which is similar to the inaccuracy found in our benchmark for adiabatic energies. If one uses the smaller def2-SVP basis (Tables S2–S5, ESI†), the deviations from the experimental data increase by about 0.1 eV. For Br-PBI, the picture is

Table 2 Energies of the PBI, Br-PBI, NPBI, and Br-NPBI were computed at So, St, Tt, and Tt states using CAM-B3LYP/aug-cc-pVDZ level of theory

	Energy relative to the ground state energy at $R(S_0)$ in eV															
PBI			Br-PBI			NPBI			Br-NPBI							
Geometry	$S_0$	$S_1$	$T_1$	$T_2$	$S_0$	$S_1$	$T_1$	$T_2$	$S_0$	$S_1$	$T_1$	$T_2$	$S_0$	$S_1$	$T_1$	$T_2$
$R(S_0)$	0	2.70	1.64	2.93	0	2.72	1.69	2.92	0	2.82	1.82	2.66	0.0	2.83	1.86	2.64
$R(S_1)$	0.18	2.45	1.37	2.92	0.21	2.49	1.39	2.91	0.15	2.64	1.60	2.65	0.15	2.66	1.63	2.63
$R(T_2)$	0.29	2.61	1.46	2.88	0.29	2.60	1.47	2.84	0.11	2.74	1.71	2.54	0.11	2.74	1.74	2.53

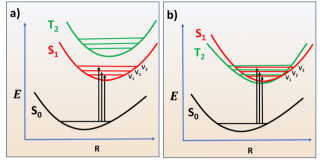


Fig. 4 Schematic representation of potential energy surfaces of S<sub>0</sub>, S<sub>1</sub>, and T<sub>2</sub> of (a) Br-PBI/PBI and (b) Br-NPBI/NPBI demonstrating the possibility of ISC

identical. For NPBI and Br-NPBI, however, the adiabatic energies deviate by about 0.3 eV from the experimental 0-0 transitions listed in Table 1. We attribute this somewhat stronger deviation to a more complex electronic structure, possibly resulting from the influence of the strain resulting from the ring formation.

However, more precise calculations are not necessary because the main difference between PBI/Br-PBI (high fluorescence quantum yield) and NPBI/Br-NPBI (low fluorescence quantum yield) results from the changes in the energetic sequence of S<sub>1</sub> and T<sub>2</sub> (Table 2 and Fig. S14, ESI†), which are clearly indicated by our calculations. For all four molecules, the  $T_1$  is well below the  $S_1$  (>1.0 eV), so an  $S_1$ - $T_1$  singlet-triplet transition as a reason for decreasing (fluorescence quantum yields) can be ruled out. For PBI and Br-PBI, the  $T_2$  for  $R(S_1)$  is about 0.5 eV above the  $S_1$ , so that an  $S_1$ - $T_2$  transition can also be excluded. For NPBI and Br-NPBI, however, the picture changes completely. For both molecules, our calculations predict that the S1 and the T2 are nearly degenerate (Table 2 and Fig. 4) inducing very efficient S<sub>1</sub>-T<sub>2</sub> transitions even for the small spin-orbit couplings calculated for NPBI or Br-NPBI (Table S7, ESI†). The greatly reduced fluorescence yield of Br-NPBI compared to NPBI could result from larger SOC couplings, but this is not supported by the calculations (Table S7, ESI†) which predict a similar size of the couplings. Hence, we attribute this experimental finding to subtle different energetic positions of both states for Br-NPBI and NPBI being outside of the accuracy of the employed theoretical approach. The situation is sketched in Fig. 4. For PBI and Br-PBI, the T<sub>2</sub> lies above the S<sub>1</sub>, so no ISC can happen. For NPBI and Br-NPBI, the T<sub>2</sub> lies

below the  $S_1$  for  $R(S_0)$  but both are nearly energy degenerate so that after the excitation, an efficient S<sub>1</sub>-T<sub>2</sub> ISC can take place.

## Conclusions

In summary, we report the unprecedented triplet population in NPBI and Br-NPBI facilitated by a reduced singlet-triplet energy gap. The moderate and low fluorescence quantum yields were observed for NPBI and Br-NPBI, respectively. The concerted interplay of nitrogen annulation and single bromine substitution fosters efficient intersystem crossing in Br-NPBI with ISC rate,  $k_{\rm ISC}$  = 1.97 × 10<sup>10</sup> s<sup>-1</sup> as observed from the fsTA measurements. The triplet quantum yield is enhanced from  $\Phi_{\rm T}$  = 18.7  $\pm$ 2.3% in NPBI to  $\Phi_{\rm T}$  = 95.2  $\pm$  4.6% for Br-NPBI which could be rationalized as the effect of bromine substitution. The nsTA spectroscopy revealed the long decay times for the triplet states in NPBI and Br-NPBI. Theoretical investigations suggest the significant role of nitrogen annulation in diminishing the singlet-triplet energy gap in NPBI and Br-NPBI. The low  $\Delta E_{S-T}$  value activates the ISC pathway in both NPBI and Br-NPBI, whereas the additional effect of bromine in Br-NPBI enhances the ISC rates to the ultrafast regime. The current work sheds light on the novel design strategies for triplet exciton generation in molecular materials, which have profound implications for developing organic light-emitting diodes and organic solar cells.

## Conflicts of interest

There are no conflicts to declare.

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