Synthesis by a Cost-Effective Method and Electroluminescence of a Novel Efficient Yellowish-Green Thermally Activated Delayed Fluorescent Molecule

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Supporting Information

ABSTRACT: A new thermally activated delayed fluorescent molecule, TRZ 3(Ph-PTZ), containing three phenothiazines as donor units and a 2,4,6-triphenyl-1,3,5-triazine as the acceptor unit was synthesized using a simple cost-effective method based on a cobalt catalyzed cross-coupling. This compound was tested in organic light-emitting diodes and was found to show superior yellowish-green electroluminescence performance with a maximum external quantum efficiency of 17.4% and a maximum luminance value of 7430 cd/m².

INTRODUCTION

Since the demonstration of the first generation of organic light-emitting diodes (OLEDs) based on conventional fluorescent emitters in 1987 by Tang and VanSlyke,¹ the development of new organic electroluminescent materials as well as OLED device physics and engineering has been the subject of intensive studies. In particular, since the pioneering works of Thompson and Forrest, the development of phosphorescent OLEDs based on organometallic emitters has been proven to be very successful.² These heavy-metal complexes, which exhibit efficient luminescence from the triplet states at room temperature, have been already used in a range of OLEDs exhibiting internal quantum efficiencies of 100%. Recent advances in the field have focused on improving the outcoupling of the devices with a partial horizontal orientation of the light-emitting dipoles, which was shown to be a promising strategy for increasing the external quantum efficiency (EQE) of phosphorescent OLEDs.³ However, the high cost of heavy metals such as iridium, which are needed to improve the spin−orbital interaction and achieve highly efficient radiative transition from the triplet excited state to the ground state, is a major problem for their use in commercial applications. An alternative to highly efficient electroluminescent devices was proposed a few years back, which is based on the use of thermally activated delayed fluorescent (TADF) emitters.⁴ The working principle of this third OLED generation is based on the upconversion of the lowest triplet excited state (T₁) to the lowest singlet excited state (S₁), using thermal energy. For efficient TADF emission, a very small energy gap (ΔE_{ST}) between S₁ and T₁ and a high yield of triplet-to-upconverted-singlet formation are both required.

A large number of efficient TADF materials have been synthesized during the last few years, showing the importance of the molecular design of the emitters to simultaneously achieve large radiative decay rate and small ΔE_{ST}.⁵⁻⁹ This molecular design is generally based on a combination of electron donor and acceptor units, which minimizes the spatial overlap between the lowest unoccupied molecular orbital (LUMO) and the highest occupied molecular orbital (HOMO). In terms of efficiency, a variety of TADF OLEDs has already shown performances equivalent to those achieved in phosphorescent devices. For instance, a green TADF molecule

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with 100% photoluminescence quantum yield (PLQY) and 100% upconversion of the triplet to the singlet excited state was recently used in devices with a maximum EQE of 41.5%. Hirata et al. demonstrated blue TADF OLEDs with 100% internal quantum efficiency and a maximum EQE of 20.6%. According to TADF OLEDs, Zhang et al. reported on the realization of high-performance electroluminescent devices with a maximum EQE of 8.1% for an emission maximum at 624 nm. Such a panel of blue, green, and red (RGB) colors is evidently crucial for different fields of application, such as solid-state lighting and display technologies. In parallel, efficient yellow electroluminescence is also of strong interest because a convenient way to obtain white light emission is to use blue and yellow emitters. In that context, high-performance yellow TADF OLEDs have been already reported with a maximum external quantum efficiency as high as 18.4%. Nevertheless, the design and synthesis of new yellow TADF emitters are still crucial for the development of display and white lighting applications. In addition, further reducing the cost of synthesis of TADF emitters by developing simple and convenient synthetic routes, which do not require the use of expensive catalysts and can be carried out in ambient atmosphere, is also extremely relevant to this research field.

## RESULTS AND DISCUSSION

In this study, we designed, synthesized, and analyzed the electroluminescence properties of a novel, highly efficient, yellowish-green TADF emitter, TRZ 3(Ph-PTZ), containing the 2,4,6-triphenyl-1,3,5-triazine for the central acceptor moiety and peripheral phenothiazine donor units (employed in a wide range of applications). The chemical structure of this molecule is displayed in Figure 1. The design of this molecular structure was motivated by a previous report where a 2,4,6-tri(phenoxazinyl)phenyl-1,3,5-triazine TADF emitter was used in a yellowish-orange OLED with a maximum EQE of about 13%.

To synthesize this compound, a simple and convenient method is proposed, which is based on the cobalt catalyzed cross-coupling of arylzinc with aryl halide. The added value of such reactions is the low-cost reagents employed for the synthesis, leading then to the reduction of cost of OLED. This advantage fits evidently well with the asset of the third OLED generation based on metal-free emitters.

Most of the classical reactions used for the synthesis of OLED molecules based on 1,3,5-triazine are dominated by the use of expensive metal catalysts such as palladium. To avoid the use of such expensive catalysts in the final coupling of the donor part (2,4,6-triphenyl-1,3,5-triazine) with the acceptor one (phenothiazine) in the TRZ 3(Ph-PTZ) molecule, we synthesized a fluorine phenyl-triazine intermediate building block based on a catalyzed cross-coupling reaction between an arylzinc and an aryl halide (Scheme 1). The arylzinc derivative 1 ((4-fluorophenyl)zinc bromide) was first prepared from 4-bromofluorobenzene in the presence of inexpensive reagents such as a cobalt salt and zinc dust in acetonitrile solvent. The resulting organozinc 1 was then engaged in the coupling reaction with the commercially available 2,4,6-trichloro-1,3,5-triazine at room temperature to form the key intermediate building block 2 (2,4,6-tris(4-fluorophenyl)-1,3,5-triazine). To the best of our knowledge, this latter coupling reaction is the first example of one-pot cobalt catalyzed cross-coupling of 3 equiv of (fluorophenyl) zinc bromide with 2,4,6-trichloro-1,3,5-triazine. Finally, the coupling last step of phenothiazine on fluorinated triazine 2 was carried out using the synthetic method reported previously for pyrrole and fluorobenzene derivatives to lead to the TRZ 3(Ph-PTZ) TADF molecule. Chemical characterization of TRZ 3(Ph-PTZ), including NMR and HRMS, is provided in Supporting Information.

To gain insights into the electronic states of TRZ 3(Ph-PTZ), we performed density functional theory (DFT) and time-dependent density functional theory (TDDFT) calculations. The ground-state geometries were optimized with Gaussian09 software at the B3LYP/6-31G* level. Phenothiazine has been described in previous studies to be a nonplanar, central six-membered ring because of the presence of sulfur. In such a moiety, the length of the (C=3) and (C=N) bonds was estimated to be 1.76–1.77 and 1.44–1.45 Å, respectively. The longer length of the (C=S) bond leads to two different molecular conformations because of rotational dynamics of the phenothiazine. In that context, the TRZ 3(Ph-PTZ) molecule, which bears three phenothiazine units, is expected to show different conformations. DFT calculations reveal two equatorial (E-1, E-2) and two axial (A-1, A-2) conformers that show only minor energy differences (see Figure 2 and Tables S1 and S2). Energy difference between E-1 and E-2 (and between A-1 and A-2) is negligible; thus, we will focus only on the first equatorial and the first axial conformers in the following paragraphs for simplicity. Figure 3 shows the HOMOs and LUMOs of the equatorial-TRZ 3(Ph-PTZ) and axial-TRZ 3(Ph-PTZ) obtained. It can be seen that both E-1 and A-1 have degenerate frontier molecular orbitals (Table S3). Such degeneracy is the result of the symmetrical properties of both conformers, implying that all these orbitals can participate in the transition process. In E-1, the HOMOs are mainly distributed on the phenothiazine unit, whereas the LUMOs are localized on the triazine and phenyl units. Only small overlaps between the HOMOs and LUMOs are present at the phenyl ring connecting phenothiazine and triazine units. In the case of A-1, however, the overlaps are quite large.

As shown in Table S4, TDDFT results reveal that S0 → S1 of E-1 is mainly the transitions between its HOMO and LUMO (70.4%). For A-1, HOMO − 1, HOMO − 2, LUMO, and LUMO + 1 played equally important roles in its S0 → S1. It can also be seen that the S0 → T1 transitions for both conformers follow similar trends. According to the orbital distributions shown in Figure 3, one can expect that E-1 should have smaller ΔEST than that of A-1 because ΔEST is proportional to the overlap between involved orbitals. This deduction is

![Figure 1. Molecular structure of TRZ 3(Ph-PTZ).](image-url)
confirmed by our TDDFT results using the optimized S$_0$ structures: the calculated $\Delta E_{ST}$ of E-1 is only 0.0079 eV, whereas $\Delta E_{ST}$ of A-1 is 0.71 eV. Apparently, thermally driven upconversion from T$_1$ to S$_1$ in E-1 could be quite effective.

The UV–visible absorption and steady-state photoluminescence (PL) spectra of TRZ 3(Ph-PTZ) at room temperature in toluene solution with a concentration of 1 × 10$^{-5}$ M are presented in Figure 4a. It should be mentioned that a detailed investigation of the photophysical properties of a TADF emitter containing a phenothiazine electron donor and 2,4,6-triphenyl-1,3,5-triazine electron acceptor unit has been already reported.$^{29}$ In particular, the different conformers of this molecule and their photophysical properties were clearly identified. On the basis of the results published in this previous report, the absorption and PL spectra of TRZ 3(Ph-PTZ) can be easily understood. The absorption spectrum of TRZ 3(Ph-PTZ) shows an intense absorption below 300 nm, which can be assigned to the 1($\pi, \pi^*$) ← S$_0$ transition derived from the triazine acceptor unit. The weak absorption tail between 400 and 500 nm and the broad and structureless absorption peak centered at 366 nm are both related to the intramolecular charge transfer (ICT) character of this molecule. In particular, the absorption tail at longer wavelengths is due to the ICT-S$_0$ transition of the equatorial conformer. As expected, the PL spectrum obtained with an excitation wavelength of 340 nm shows a dual emission with peaks around 431 and 575 nm. Similar to what was observed in a previous study,$^{29}$ the peak at 575 nm shows the highest intensity with a large Stokes shift and a full width at half-maximum of 108 nm. This ICT emission can be straightforwardly assigned to the excited state of the quasi-equatorial conformation (E-1) of TRZ 3(Ph-PTZ). The peak centered at about 431 nm exhibits a much lower intensity with a small Stokes shift, and this emission is attributed to the excited state of the quasi-axial conformer (A-1). It should be emphasized that this quasi-axial conformer shows an overlap between HOMO and LUMO on the triazine acceptor unit. The observation of a smaller Stokes shift for this conformer can be explained by the effect of the $\pi-\pi^*$ character deduced from the non-negligible HOMO–LUMO overlap.$^{29}$

To determine whether the dual emission from TRZ 3(Ph-PTZ) can also be observed in the solid state, an organic thin film based on a blend containing 2 wt % of TRZ 3(Ph-PTZ) in a 3,3′-bis(N-carbazolyl)-1,1′-biphenyl (mCBP) host was prepared by thermal evaporation onto a fused silica substrate. The mCBP host was selected because of the high triplet T$_1$ energy of this material, which prevents back energy transfer from the guest to the host molecules and confines the triplets within the guest emitters. The absorption and steady-state PL spectra of this sample are displayed in Figure 4b. Owing to the small concentration of the guest molecules in the host, the absorption spectrum is completely dominated by the absorption from mCBP. More importantly, the PL spectrum obtained with an excitation wavelength of 340 nm does not show any emission from the mCBP matrix, indicating that an efficient energy transfer takes place from the host to TRZ 3(Ph-PTZ) molecules. In addition, similarly to what was observed in toluene solution, the photoluminescence from the doped film exhibits a dual emission with two peaks at about 427 and 520 nm. Noticeably, whereas the blue emission peak from the quasi-axial conformer presents nearly the same maximum wavelength in solution and thin film, the PL spectrum from the equatorial conformer in the solid state shows a significant blue shift of around 55 nm. Because TADF materials showing ICT generally exhibit a strong solvatochromism effect, a plausible explanation for this observation could be related to the different
polarity of the environments surrounding the TRZ 3(Ph-PTZ) molecules. The transient PL decay of the equatorial conformer in the mCBP blend was then measured using an excitation wavelength of 337 nm. The PL decay curve displayed in Figure 4c was fitted by a sum of two exponential decay functions, the fast and slow components corresponding to the prompt and delayed fluorescence, respectively. The prompt and delayed fluorescence lifetimes are measured to be 6.7 ns and 7.2 μs, respectively. This result suggests that the highly emissive equatorial conformer of TRZ 3(Ph-PTZ) exhibits a TADF activity, which is consistent with the small ΔE_ST value obtained by quantum chemistry calculations.

To investigate the electroluminescence properties of TRZ 3(Ph-PTZ), we fabricated OLEDs with the following architecture: indium tin oxide (ITO) (100 nm)/N,N’-diphenyl-N,N’-bis(1-naphthyl)-1,10-biphenyl-4,4’-diamine (α-NPD) (20 nm)/tris(4-carbazoyl-9-ylphenyl)amine (TCTA) (10 nm)/1,3-bis(N-carbazolyl)benzene (mCP) (10 nm)/5 wt % TRZ 3(Ph-PTZ):mCBP (15 nm)/bis[2-(diphenylphosphino)phenyl] ether oxide (DPEPO) (10 nm)/1,3,5-tris(N-phenylbenzimidazolyl)benzene (TPBi) (55 nm)/lithium fluoride (LiF) (0.8 nm)/aluminum (Al) (900 nm). A schematic representation of this multilayer structure is provided in Figure 5a, and its energy diagram is shown in Figure S4. In this device, α-NPD and mCP are used as hole transport layers, DPEPO as hole blocking layer, and TPBi as electron transport layer. The electroluminescence characteristics of this OLED are displayed in Figure 5b–d. The device shows yellowish-green electroluminescence with color coordinates of CIE (0.23, 0.75) and exhibits a turn-on voltage of 4.2 V, a maximum current efficiency of 58.6 cd/A, a power efficiency of 17.4 lm/W, a maximum luminance up to 7430 cd/m², and a high maximum EQE of 17.4% without any light outcoupling enhancement. As shown in Figure S5, these excellent electroluminescence
performances could be reproduced in several devices. The high EQE value is evidently well beyond the theoretical limit for conventional fluorescent OLEDs and can be attributed to the efficient upconversion of the triplets to singlets through TADF process, which is consistent with the quantum chemistry and PL results presented above. However, it can be seen that the electroluminescence spectra measured in the device for three different current densities do not show any dual emission, in contrast to what was reported in a previous study with a similar emitter. Further photophysical investigations as well as some device optical and electrical device modeling that are both needed to explain this unexpected result and optimize the dual device optical and electrical device modeling that are both subject of more intensive studies in the near future.

CONCLUSIONS

In summary, the compound TRZ 3(Ph-PTZ) shows excellent TADF characteristics and can be successfully used in a high-performance yellowish-green OLED with a maximum external quantum efficiency of 17.4%. Photophysical measurements demonstrate that this molecule exhibits a dual emission because of the introduction of phenothiazine unit in the molecular structure. This moiety is responsible for the presence of conformers showing different photophysical characteristics. Importantly, the synthesis of this compound is carried out using a simple and convenient method based on cobalt catalyzed cross-coupling. This process can be used to reduce substantially the cost of the synthesis of highly efficient TADF molecules, thanks to the low cost of cobalt employed as catalyst for the synthesis. This advantageous method is used for the first time to synthesize TADF emitters and can easily be extended to other synthetic routes.

EXPERIMENTAL SECTION

Materials and Methods. All chemicals are reagent/analytical grade and used without further purification. 1H and 13C NMR spectra were recorded onBruker AC-300 (300 and 75 MHz, respectively) and are internally referenced to residual solvent signals (CDCl3). Data for 1H and 13C NMR are reported as follows: chemical shift (δ, ppm), multiplicity (s = singlet, d = doublet, m = multiplet), coupling constant (J in Hz), integration, and assignment. High-resolution mass spectra were obtained from Ecole Polytechnique Mass Spectral facility. Infrared spectrum was performed with JASCO FT/IR-6100 spectrometer. All of the starting materials were purchased from the usual suppliers (Sigma-Aldrich, Alfa Aesar, and Acros Organics).

Synthesis. 2,4,6-Tris(4-fluorophenyl)-1,3,5-triazine. CoBr2 (0.5 mmol, 109 mg, 0.1 equiv) and Zn dust (16.5 mmol, 1.08 g, 3.3 equiv) were added in MeCN (4 mL). The mixture was stirred at room temperature for 1 h and 4-bromofluorobenzene (5 mmol) was then added. The reaction was stirred to give the (4-fluorophenyl)zinc bromide reagent (80% in GC). 2,4,6-Trichloro-1,3,5-triazine (6.5 mmol, 1.3 equiv) was added at room temperature to the unfiltered solution of (4-fluorophenyl)zinc bromide reagent. In the event of a significant rise in temperature, the reaction mixture was cooled with a water bath. The reaction was then stirred for 1 h. The reaction was quenched by addition of saturated aqueous NH4Cl solution (20 mL), followed by an extraction with DCM (3 × 50 mL). The combined organic layers were dried over MgSO4, and after filtration the solvents were evaporated in vacuum. Purification by flash column chromatography (SiO2, petrol ether/EtOAc = 50:1) afforded the desired product (908 mg) in 50% overall yield. 1H NMR (300 MHz, CDCl3): δ/ppm = 8.77 (dd, J = 8.7, 5.6 Hz, 6H), 7.39–7.16 (m, 6H). 13C NMR (75 MHz, CDCl3): δ/ppm = 170.81, 169.26, 132.30, 131.43, 115.93. 19F NMR (282 MHz, CDCl3): δ/ppm = −106.73. HRMS m/z [M]+ calcd for C21H12F3N3, 363.0983; found, 363.0991.

2,4,6-Tris(4-(10H-phenothiazin-10-yl)phenyl)-1,3,5-triazine (TRZ 3(Ph-PTZ)). An amount of NaH (1.14 mmol, 27 mg,
layers were dried over MgSO₄, and the solvents were tris(4-fluorophenyl)-1,3,5-triazine in 5 mL of dimethylformamide (DMF). After the liberation of H₂ gas had ceased, the orange reaction mixture was stirred for an additional 30 min. Then, 0.28 mmol (101 mg, 1 equiv) of 2,4,6-tris(4-fluorophenyl)-1,3,5-triazine in 5 mL of DMF was added in one batch. The reaction mixture was stirred overnight at 100 °C. It was poured into 100 mL of ice water followed by extraction with DCM (3 × 50 mL). The combined organic layers were dried over MgSO₄, and the solvents were evaporated in vacuum.

**Photophysical Measurements.** For the photophysical measurements, the degassed toluene solution of TRZ 3(Ph-PTZ) was placed in a quartz cuvette. In parallel, the organic TRZ 3(Ph-PTZ) was evaporated under vacuum lower than 3 × 10⁻¹⁰ Torr. The LiF/Al electrodes were deposited through a shadow mask, and the LiF/Al/LiO₂/LiF/Al emitters were evaporated through a mask (100 nm LiF, 100 nm Al) on a glass substrate. The LiO₂/Al/LiF/Al structure was used as an electron-injection layer and a hole-blocking layer, respectively. The LiO₂/Al/LiF/Al/LiF/Al structure was used as an electron-injection layer and a hole-blocking layer, respectively. The LiO₂/Al/LiF/Al/LiF/Al structure was used as an electron-injection layer and a hole-blocking layer, respectively. The LiO₂/Al/LiF/Al/LiF/Al structure was used as an electron-injection layer and a hole-blocking layer, respectively.

**OLED Fabrication and Measurements.** The device was prepared using a pre-cleaned indium tin oxide (ITO) substrate. The different organic layers were sequentially thermally evaporated at vacuum below 3 × 10⁻⁴ Pa. Finally, the LiF/Al electrodes were deposited through a mask shadow by evaporation on the top of the multilayer structure. The device active area was 4 mm². To avoid any degradation and emission quenching due to oxygen and moisture, the OLED was encapsulated in a glovebox filled with nitrogen. The J–V–L characteristics were collected using a source meter (Keithley 2400, Keithley Instruments Inc.) and an absolute external quantum efficiency measurement system (C9920-12, Hamamatsu Photonics). An optical fiber connected to a spectrometer (PMA-12, Hamamatsu Photonics) was used to record the electroluminescence spectrum.


