A simple molecular design towards the conversion of a MCL backbone to a multifunctional emitter exhibiting polymorphism, AIE, TADF and MCL

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GRAPHICAL ABSTRACT

This work demonstrated a simple molecular design strategy towards the conversion of a MCL backbone to a multifunctional emitter exhibiting polymorphism, AIE, TADF and MCL.

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ABSTRACT

Compared with the large number of single-function materials such as aggregation-induced emission (AIE), mechanochromic luminescence (MCL), or thermally activated delayed fluorescence (TADF) emitters, multifunctional emitting materials offer more opportunities in practical applications. In this report, we provide a simple molecular design strategy towards the conversion of a MCL building block to a multifunctional emitter. Through altering the substituent sites and increasing the number of electron donors and steric hindrance on a normal MCL backbone benzo[d,e]benzo[4,5]imidazo[2,1-a]isoquinolin-7-one, a novel multifunctional material 10,11-bis-(4-diphenylamino-phenyl)-benzo[d,e]benzo[4,5]imidazo[2,1-a]isoquinolin-7-one (10,11-2TPA-BBI) is designed and synthesized. 10,11-2TPA-BBI exhibits simultaneous polymorphism, AIE, TADF and MCL properties. It can form four different aggregate species: yellow solid (YS) and orange solid (OS), orange flake-shaped crystal (OC), and red prism-like crystal (RC). Among them, because of the small energy gaps (ΔE_{ST} < 0.3 eV) between the singlet and triplet excited states, YS show normal fluorescence characteristics with a large ΔE_{ST} of 0.33 eV. OS can be reversibly transformed into YS upon external stimuli, which can be attributed to the emission switch between local excited state and charge transfer state. Crystallographic study indicates that the bulky structure and weak intermolecular interactions account for polymorphism and AIE properties. This work will provide a simple molecular design strategy for multifunctional materials.

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1. Introduction

Luminescent materials based on purely organic compounds have received raising attention in recent years for various applications ranging from chemical sensors, biomedicine, to light-emitting devices etc [1, 2, 3, 4, 5, 6, 7]. Based on the luminescent features, luminescent materials can be classified into aggregation-induced emission (AIE) luminogens, mechanochromic luminescence (MCL) fluorophores, thermally activated delayed fluorescence (TADF) emitters or their combination. AIE luminogens are compounds that show negligible emission in dilute solution while intense luminescence in solid state. In principle, AIE is attributed to the restriction of free intramolecular rotations and vibrations in aggregation state [8, 9, 10]. AIE-active compounds can serve as efficient emitters in light-emitting devices and as photosensitizers in biomedical fields [11, 12, 13]. Interestingly, MCL fluorophores are compounds that exhibit reversible emission color change upon external stimuli (e.g. grinding, pressing, rubbing, heating, organic vapor treatments etc) [14, 15, 16, 17, 18]. Due to their unique features, MCL emitters can be applied in stress-, pressure-, and thermo-indicators in the future [19, 20, 21]. In the past ten years, great efforts have been made to develop TADF emitters, which possess small energy splits (\(\Delta E_{ST}\)) between singlet (\(S_1\)) and triplet (\(T_1\)) excited states to facilitate upconversion from \(T_1\) to \(S_1\) through reverse intersystem crossing (RISC) [22, 23, 24, 25, 26]. TADF emitters have aroused increasing interest for their promising implications in optoelectronic and biomedical fields [27, 28, 29, 30]. Nowadays, compared with single-function (AIE, MCL or TADF) emitting materials, multifunctional emitting materials offer more opportunities in practical applications [31, 32, 33, 34, 35]. The development of multifunctional materials have attracted more and more attention. In 2015, Chi’s group reported the first multifunctional material 4-carbazol-10-phenothiazine diphenyl sulfone that shows AIE, TADF, and MCL properties simultaneously [36]. In 2017, Takeda et al developed a series TADF molecules based on phenothiazine-dibenzo[\(e,d]\]imidazo[2,1-\(a\)]isoquinolin-7-one (BBI) as the electron donor and acceptor, respectively [41, 42]. Due to the change of \(\pi-\pi\) interactions in different aggregation states, these isomers show high-contrast MCL property. However, these compounds have large overlaps between the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO), leading to large \(\Delta E_{ST}\) (>0.5 eV) and no TADF features. In this report, we attempted to provide a simple molecular design strategy towards the conversion of a MCL skeleton to a multifunctional emitter. Through altering the substituent sites and increasing the number of TPA moieties and steric hindrance on the MCL building block BBI, a novel multifunctional material 10,11-bis-(4-diphenylamino-phenyl)-benzo[\(e,d\)]benzo [4,5]imidazo[2,1-\(a\)]isoquinolin-7-one (10, 11-2TPA-BBI, Figure 1) is successfully synthesized. Due to the change of substituent sites and the increase of electron-donating TPA moieties, the overlap between HOMO and LUMO is significantly decreased, inducing a small \(\Delta E_{ST}\) of 0.19 eV. Interestingly, 10,11-2TPA-BBI can form four different aggregate species: yellow solid (YS), orange solid (OS), orange flake-shaped crystal (OC), and red prism-like crystal (RC). Among the four aggregates, OS, OC and RC exhibit TADF properties, while YS show normal fluorescence characteristics. Under external stimuli, OS can be reversibly switched to YS, indicating MCL property. Because of the increased steric hindrance, there are no strong \(\pi-\pi\) interactions in OC or RC. The weak interactions in the crystals are conductive to inhibiting intramolecular rotations, which may help to explain the AIE property of the compound. We believe that the conversion of a normal MCL backbone to a multifunctional emitter would be a significantly powerful strategy for creating multifunctional organic materials.

![Figure 1. Synthesis route of 10,11-2TPA-BBI.](image-url)
2. Experimental

2.1. Chemicals and instruments

The reagents (analytical grade) and solvents (HPLC grade) were purchased from Aladdin Bio-chem Technology Co., LTD (Shanghai, China) and used as received. Mass spectra were performed using a mass spectrometer (Finnigan LTQ, Thermo Electron Corporation). $^1$H and $^{13}$C NMR spectrum of the target compound were measured in CDCl$_3$ on a NMR spectrometer (Bruker AV 400 M) with tetramethylsilane as the internal standard. UV-vis absorption spectrum were performed in the range of 260–600 nm in Agilent 8453 UV-vis spectrophotometer. The photoluminescence spectra (PL), phosphorescence spectra (PH), photoemission spectra, were carried out using Oxford Gemeni S Ultra X-ray diffractometer with Mo-K$_{\alpha}$ radiation at 296 K. The powder X-ray diffraction (XRD) measurements of the samples were measured using a spectrophotometer (Edinburgh FLS 980). The single crystal XRD measurements of the crystals were carried out using Oxford Gemini S Ultra X-ray diffractometer with Mo-K$_{\alpha}$ radiation at 296 K. CCDC 2034918 (RC) and 2034919 (OC) contain the supplementary crystallographic data for this paper [43].

The molecular structure, HOMO and LUMO of 10,11-2TPA-BBI were optimised by density functional theory (DFT) calculations at B3LYP/6-31G(d) level (Gaussian 09 program package) [44]. Based on the optimised geometry of 10,11-2TPA-BBI at ground state, the excitation energies were calculated by time-dependent density functional theory (TD-DFT) method [45].

2.2. Synthesis of the target molecule

2.2.1. Synthesis of 10,11-dibromo-benzo[de]benzo[4,5]imidazo[2,1-a]isoquinolin-7-one (10,11-2Br-BBI)

In a 250 mL three-necked flask, a mixture of 10,11-2Br-BBI (4.0 mmol), 4-diphenylamino)phenyl boronic acid (8.8 mmol) in toluene (60 mL), ethanol (30 mL) and aqueous potassium carbonate solution (1.0 M, 20 mL) was stirred under N$_2$ for 10 min. After the addition of tetrakis(µ-triphenylphosphine) palladium (0.4 mmol), the mixture was refluxed under nitrogen for 18 h. Cooling to room temperature, 80 mL deionized water was added. The mixture was extracted with 100 mL dichloromethane (DCM) for two times. Then the organic phase was combined, washed with saturated aqueous sodium chloride solution (2 × 40 mL), dried over anhydrous sodium sulfate, and concentrated under vacuum. The crude product was purified by column chromatography (silica gel; eluent: 10% ethyl acetate/petroleum ether) to give 10,11-2TPA-BBI as an orange solid with a yield of 70.4%. $^1$H NMR (400 MHz, CDCl$_3$): δ (ppm): 160.66, 149.90, 147.76, 146.36, 143.15, 138.85, 138.64, 136.05, 135.86, 135.37, 132.31, 131.89, 131.73, 131.20, 131.17, 131.06, 129.28, 127.42, 127.25, 126.95, 124.37, 124.27, 123.24, 123.20, 123.05, 122.81, 122.75, 120.94, 120.74, 117.10. MS (MALDI-TOF) [m/z]: Calcd for C$_{54}$H$_{36}$N$_4$O, 756.87; Found: 756.95.

2.2.2. Synthesis of 10,11-2TPA-BBI

In a 250 mL three-necked flask, a mixture of 10,11-2Br-BBI (4.0 mmol), 4-diphenylamino)phenyl boronic acid (8.8 mmol) in toluene (60 mL), ethanol (30 mL) and aqueous potassium carbonate solution (1.0 M, 20 mL) was stirred under N$_2$ for 10 min. After the addition of tetrakis(µ-triphenylphosphine) palladium (0.4 mmol), the mixture was refluxed under nitrogen for 18 h. Cooling to room temperature, 80 mL deionized water was added. The mixture was extracted with 100 mL dichloromethane (DCM) for two times. Then the organic phase was combined, washed with saturated aqueous sodium chloride solution (2 × 40 mL), dried over anhydrous sodium sulfate, and concentrated under vacuum. The crude product was purified by column chromatography (silica gel; eluent: 10% ethyl acetate/petroleum ether) to give 10,11-2TPA-BBI as an orange solid with a yield of 70.4%. $^1$H NMR (400 MHz, CDCl$_3$): δ (ppm): 160.66, 149.90, 147.76, 146.36, 143.15, 138.85, 138.64, 136.05, 135.86, 135.37, 132.31, 131.89, 131.73, 131.20, 131.17, 131.06, 129.28, 127.42, 127.25, 126.95, 124.37, 124.27, 123.24, 123.20, 123.05, 122.81, 122.75, 120.94, 120.74, 117.10. MS (MALDI-TOF) [m/z]: Calcd for C$_{54}$H$_{36}$N$_4$O, 756.87; Found: 756.95.

3. Results and discussion

3.1. Synthesis

Synthetic route of 10,11-2TPA-BBI is illustrated in Figure 1. First, the key intermediate 10,11-2Br-BBI is synthesized by condensation reaction of 1,8-naphthalic anhydride with 4,5-dibromo-1,2-benzenediamine. Then 10,11-2Br-BBI is transformed into the target compound 10,11-2TPA-BBI by Suzuki reaction with 4-(diphenylamino)phenyl boronic acid. The structure of 10,11-2TPA-BBI is confirmed by MS, $^1$H NMR, $^{13}$C NMR, and single-crystal measurements.
3.2. Theoretical calculations

To optimize the geometry and predict the HOMO and LUMO electronic distributions of 10,11-TPA-BBI, DFT calculations were performed. As shown in Figure 2, the HOMO of 10,11-TPA-BBI is mainly localized over the TPA units, while the LUMO is dispersed on the BBI moiety. The HOMO and LUMO energy levels of 10,11-TPA-BBI are estimated to be -4.77 eV and -2.49 eV, respectively. Accordingly, the optical bandgap ($E_g$) is calculated to be 2.28 eV, which is smaller than the reported MCL emitters based on TPA substituted BBI (3-TPA-BBI and 4-TPA-BBI) [41, 42], indicating the increase of molecular conjugation. Compared with 3-TPA-BBI and 4-TPA-BBI, the overlap between HOMO and LUMO of 10,11-TPA-BBI is significantly decreased, which can be attributed to the change of the substituent sites and the increase of the number of TPA moieties. Based on the optimized geometries, TD-DFT calculations were performed to predict the $\Delta E_{ST}$ of 10,11-TPA-BBI. As expected, the small overlap between HOMO and LUMO leads to a small $\Delta E_{ST}$ of 0.19 eV (Table 1), which is much smaller than 3-TPA-BBI and 4-TPA-BBI [41, 42]. The small $\Delta E_{ST}$ indicates that 10,11-TPA-BBI may serve as a typical TADF emitter.

3.3. Spectroscopic properties of 10,11-2TPA-BBI in solvents

UV-vis absorption spectrum of 10,11-2TPA-BBI in toluene is presented in Figure 3a. The absorption profile of 10,11-2TPA-BBI consists three absorption band regions. The two bands at around 300 and 340 nm are attributed to the $\pi-\pi^*$ and $n-\pi^*$ transitions, respectively, while the longer-wavelength absorption at around 450 nm belongs to intramolecular charge transfer (ICT) from TPA moieties to BBI unit. From the onset absorption (520 nm), $E_g$ of 10,11-2TPA-BBI can be calculated to be 2.38 eV, which is in good agreement with the above-mentioned theoretical result (2.28 eV).

PL spectra of 10,11-2TPA-BBI in solution are also investigated. As illustrated in Figure 3b-c, 10,11-2TPA-BBI exhibits faint emission in tetrahydrofuran (THF) solution. With the addition of deionized water fraction ($f_w$), the PL intensity of 10,11-2TPA-BBI first decreases and then enhances gradually. The decrease of PL intensity at low $f_w$ (10%) can be attributed to the slight increase of solvent polarity, which is in favor of molecular motions and non-radiative decay. With $f_w$ increases to 60%, the aggregates may form and restrict intramolecular motions, and thus inhibit non-radiative decay and increase PL intensity. When $f_w$ increases to 90%, the fluorescence intensity reaches to 80-fold of that in pure THF solution, indicating the typical AIE feature of 10,11-2TPA-BBI (Figure 3d) [33]. Note that the reported BBI based compounds 3-TPA-BBI and 4-TPA-BBI are not AIE-active [41, 42]. This verifies that the change of the substituent sites and the increase of the number of TPA moieties are beneficial for restricting intramolecular motions and induce AIE property.

3.4. Polymorphism and TADF properties of 10,11-2TPA-BBI in aggregation states

Through controlling the preparation conditions mentioned above, 10,11-2TPA-BBI formed four different aggregates: YS, OS, OC and RC
Figure 4. (a) Photographs of the four solid states of 10,11-2TPA-BBI under the 365 nm UV excitation (a letter “M” based on YS or OS; scale bar: 1 mm); (b) PL spectra of 10,11-2TPA-BBI in different aggregation states; (c) PL decay curves of 10,11-2TPA-BBI in different aggregation states; (d) The time-resolved PL and PH spectra of 10,11-2TPA-BBI in different aggregation states with 1 ms delay at 77 K: (d) YS; (e) OS; (f) OC; (g) RC.
and finely structured PH, corresponding to charge transfer (CT) excited state and LE state, respectively [47]. In the case of OC and RC, broad PL and PH spectra reflect the CT excited states in crystals. From the PL spectra, the $S_1$ energy values of the four aggregates are calculated to be 2.05 eV (YS), 2.03 eV (OS), 2.23 eV (OC) and 2.24 eV (RC). From the PH spectra at 77 K, $T_1$ energy values are estimated to be 1.72 eV (YS), 1.81 eV (OS) and 2.20 eV (RC), respectively. Correspondingly, $\Delta E_{ST}$ are obtained to be 0.23 eV (YS), 0.22 eV (OS), 0.85 eV (RC) and 0.04 eV (RC). Note that OS, OC and RC have small $\Delta E_{ST}$ (<0.3 eV), which facilitate upconversion from $T_1$ to $S_1$ state through RISC and induce TADF emission. In contrast, YS has a larger $\Delta E_{ST}$ (>0.3 eV) and shows no TADF behavior.

3.5. MCL property of 10,11-2TPA-BBI

Among the four aggregates, OS can be transformed into YS upon heating (170 °C, 5 s), and YS can be converted back to OS form via grinding or fuming with DCM vapor (Figure 4a). Upon external stimuli, the reversible conversion between OS and YS indicates typical MCL property of 10,11-2TPA-BBI. By comparing the PL spectra of YS and OS at 77 K (Figure 4d-e), the MCL behavior of 10,11-2TPA-BBI can be attributed to the emission switch between LE (YS) and CT (OS). The XRD patterns of the four aggregates are shown in Figure 5a, YS and OS show no evident diffraction peaks, indicating that YS and OS are both in amorphous states. In contrast, OC and RC exhibit strong and evident diffraction peaks, suggesting the ordered structures of OC and RC.

Figure 5b shows the maximum PL emission of 10,11-2TPA-BBI upon being treated by heating (or grinding) and fuming with DCM, indicating that the transformation between YS and OS upon external stimulus has good repeatability. Note that the emission color change of 10,11-2TPA-BBI during MCL process is smaller than 3-TPA-BBI and 4-TPA-BBI, in which MCL property may originate from different π−π interactions of the aggregates [41, 42]. In the case of 10,11-2TPA-BBI, it is difficult for the molecules to form π−π stacking in these aggregates because of the bulky structure.

3.6. Single crystallographic analysis of 10,11-2TPA-BBI

To investigate the influence of crystal structures on photophysical properties, the single-crystal analysis of OC and RC were conducted. The details of the structures were summarized in Figure 6a-b and Table 3. OC is triclinic with space group of P-1. In contrast, RC is monoclinic with space group of P2_1/c. The formulas of OC and RC are C_{56}H_{44}N_{4}O_{3} (the solvent chloroform) and 10,11-2TPA-BBI with a ratio of 1:2 and C_{51}H_{44}N_{6}O_{3} (the solvent methanol and 10,11-2TPA-BBI with a ratio of

| Aggregate | PL Value of YS (nm) | PL Value of OS (nm) | PL Value of OC (nm) | PL Value of RC (nm) |
|-----------|---------------------|---------------------|---------------------|---------------------|
| YS        | 583                 | 598                 | 603                 | 613                 |
| OS        | 0.148               | 0.154               | 0.281               | 0.205               |
| OC        | 60.36 (100%)        | 61.42 (13.65%)      | 5.80 (12.27%)       | 10.00 (30.18%)      |
| RC        | /                   | 0.34 (86.35%)       | 0.35 (87.73%)       | 0.15 (69.82%)       |

| Aggregate | $k_f$ (10^7 s^-1) | $k_r$ (10^7 s^-1) | $k_m$ (10^7 s^-1) | $k_m$ (10^7 s^-1) |
|-----------|-------------------|-------------------|-------------------|-------------------|
| YS        | 0.25              | 1.41              | 0.25              | 0.15              |
| OS        | 0.25              | 1.38              | 0.25              | 0.15              |
| OC        | 0.25              | 12.40             | 0.25              | 0.15              |
| RC        | 0.25              | 7.95              | 0.25              | 0.15              |

| Aggregate | $S_1$ (eV) | $T_1$ (eV) | $\Delta E_{ST}$ (eV) |
|-----------|-------------|------------|----------------------|
| YS        | 2.05 (LE)   | 1.72 (LE)  | 0.33                 |
| OS        | 2.03 (CT)   | 1.81 (LE)  | 0.04                 |
| OC        | 2.23 (CT)   | 2.15 (CT)  | 0.14                 |
| RC        | 2.24 (CT)   | 2.20 (CT)  | 0.04                 |

Figure 5. (a) XRD spectra of 10,11-2TPA-BBI in different solid states; (b) maximum PL emission of 10,11-2TPA-BBI upon being treated by heating (or grinding) and fuming with DCM.
As expected, both crystals exhibit twisted molecular structures. In these crystals, the dihedral angles between BBI plane and the neighbouring phenyl planes of TPA units are 59.08°/47.63° (OC) and 53.96°/51.59° (RC). The corresponding bond lengths are 1.459 Å/1.468 Å (OC) and 1.500 Å/1.497 Å (RC), respectively. The bond lengths of C=O are 1.228 Å (OC) and 1.220 Å (RC) (Figure 5a-b). Compared with OC, RC exhibits similarly twisted conformation but longer bond lengths between electron-donating TPA moieties and electron-accepting BBI unit, indicating that the conjugation is increased in RC. The results are consistent with the red-shifted PL spectra of RC comparing to OC (Figure 3b).

As a step further to investigate the different luminous properties of OC and RC, the intermolecular interactions in the crystals were analyzed (Figure 6c-d). Due to the twisted conformations and bulky structures, the 10,11-2TPA-BBI molecules couldn’t form intermolecular aromatic packing, leading to no strong intermolecular interactions such as π...π interactions in OC or RC. In the case of OC, two 10,11-2TPA-BBI molecules form a pair by C–H...C–H and C–H...N intermolecular interactions with distances of 2.856 Å, 2.621 Å and 2.890 Å, respectively. Note that there is no obvious intermolecular interactions between 10,11-2TPA-BBI molecules and the adjacent chloroform molecules. In contrast, the neighbouring 10,11-2TPA-BBI molecules are bound by C–H...π and C–H...O intermolecular interactions distances of 2.843 Å/2.437 Å and 2.597 Å in RC. In addition, there is C–H...N intermolecular interactions with distances of 2.053 Å between methanol molecules and the adjacent 10,11-2TPA-BBI molecules. Compared with OC, the shortened C–H...π, C–H...O and C–H...N distances in RC indicate that intermolecular interactions are enhanced, which also contribute to the red-shifted PL spectra of RC versus OC [48]. Meanwhile, the weak intermolecular interactions in the single crystals are conductive to restraining intramolecular rotations, which may help to explain the AIE property of 10,11-2TPA-BBI.

4. Conclusions

In summary, we successfully converted a MCL-active backbone BBI to a multifunctional emitter 10,11-2TPA-BBI by molecular modification. As expected, the compound exhibits polymorphism, AIE, MCL and TADF properties simultaneously. TADF is achieved by altering the substituent sites and increasing the number of electron-donating TPA moieties on the electron-accepting BBI unit. PL emission switch between LE and CT states is responsible for the MCL behavior. The bulky structure and weak intermolecular interactions contribute to polymorphism and AIE properties. In all, the current work demonstrated a simple molecular design strategy towards the conversion of a MCL backbone to a multifunctional

2:1), respectively. As expected, both crystals exhibit twisted molecular structures. In these crystals, the dihedral angles between BBI plane and the neighbouring phenyl planes of TPA units are 59.08°/47.63° (OC) and 53.96°/51.59° (RC). The corresponding bond lengths are 1.459 Å/1.468 Å (OC) and 1.500 Å/1.497 Å (RC), respectively. The bond lengths of C=O are 1.228 Å (OC) and 1.220 Å (RC) (Figure 5a-b). Compared with OC, RC exhibits similarly twisted conformation but longer bond lengths between electron-donating TPA moieties and electron-accepting BBI unit, indicating that the conjugation is increased in RC. The results are consistent with the red-shifted PL spectra of RC comparing to OC (Figure 3b).

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4. Conclusions

In summary, we successfully converted a MCL-active backbone BBI to a multifunctional emitter 10,11-2TPA-BBI by molecular modification. As expected, the compound exhibits polymorphism, AIE, MCL and TADF properties simultaneously. TADF is achieved by altering the substituent sites and increasing the number of electron-donating TPA moieties on the electron-accepting BBI unit. PL emission switch between LE and CT states is responsible for the MCL behavior. The bulky structure and weak intermolecular interactions contribute to polymorphism and AIE properties. In all, the current work demonstrated a simple molecular design strategy towards the conversion of a MCL backbone to a multifunctional

Table 3. Crystal data and structure refinement for 10,11-2TPA-BBI.

|       | OC                  | RC                  |
|-------|---------------------|---------------------|
| **Formula** | C₁₀₉H₇₃Cl₃N₈O₂ | C₅₆H₄₄N₄O₃        |
| **CCDC No.** | 2034919            | 2034918             |
| **Formula weight** | 1633.10            | 820.95              |
| **Space group** | P-1                | P₂₁/c               |
| **Temperature** | 296                | 296                 |
| **Wavelength/Å** | 0.71073            | 0.71073             |
| **Crystal system** | Triclinic           | Monoclinic          |
| **a/Å** | 13.267(2)           | 12.428(9)           |
| **b/Å** | 16.259(3)           | 14.064(10)          |
| **c/Å** | 22.397(3)           | 25.185(18)          |
| **α/º** | 74.897(6)           | 90.000              |
| **β/º** | 85.589(7)           | 101.955(9)          |
| **γ/º** | 84.927(7)           | 90.000              |
| **Volume/Å³** | 4639.0(13)         | 4307.0(5)           |
| **Z** | 2                   | 4                   |
| **Calculated density/Mg m⁻³** | 1.169              | 1.360               |
| **M/(mm⁻³)** | 0.153              | 0.079               |
| **F(000)** | 1700               | 1728                |
| **Crystal size/mm³** | 0.15 × 0.18 × 0.20 | 0.14 × 0.15 × 0.18 |
| **No. of reflns collected** | 16104             | 7678                |
| **No. of unique reflns** | 4523              | 4430                |
| **Goodness-of-fit on F²** | 0.920             | 1.266               |
| **R₁, wR₂ [I > 2σ(I)]** | 0.1047, 0.2251      | 0.0734, 0.2066      |
| **R₁, wR₂ (all data)** | 0.2762, 0.2671      | 0.1308, 0.2359      |
emitter, which would provide a guideline for the design of multifunctional materials.

**Declarations**

**Author contribution statement**

Bin Huang: Conceived and designed the experiments; Contributed reagents, materials, analysis tools or data; Wrote the paper. Wenbing Yu, Li Yang: Performed the experiments. Yan Li: Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data. Ning Gu: Conceived and designed the experiments; Wrote the paper.

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**Data availability statement**

Data included in article/supplementary material/referenced in article.

**Declaration of interests statement**

The authors declare no conflict of interest.

**Additional information**

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