Protein-like Enwrapped Perylene Bismide Chromophore as a Bright Microcrystalline Emitter Material

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Dedicated to Professor Jürgen Köhler on the occasion of his 60th birthday

Abstract: Strongly emissive solid-state materials are mandatory components for many emerging optoelectronic technologies, but fluorescence is often quenched in the solid state owing to strong intermolecular interactions. The design of new organic pigments, which retain their optical properties despite their high tendency to crystallize, could overcome such limitations. Herein, we show a new material with monomer-like absorption and emission profiles as well as fluorescence quantum yields over 90% in its crystalline solid state. The material was synthesized by attaching two bulky tris(4-tert-butylphenyl)phenoxy substituents at the perylene bisimide bay positions. These substituents direct a packing arrangement with full enwrapping of the chromophore and unidirectional chromophore alignment within the crystal lattice to afford optical properties that resemble those of their natural pigment counterparts, in which chromophores are rigidly embedded in protein environments.

Solid-state fluorescent organic materials are of considerable interest as they enable a broad variety of applications in (opto-)electronics, for example, organic light emitting diodes (OLEDs), waveguiding, solid-state lasers, luminescent sensors, or as fluorescent labels for (bio)medical research as well as security printing technologies. Unfortunately, while pigments are the preferred colorants for these applications compared to dyes owing to their superior thermal, photochemical, and chemical robustness, only very few pigments exhibit a decent fluorescence. In contrast to the bright luminescence with quantum yields of up to 90% in its crystalline solid state, the material was synthesized by attaching two bulky tris(4-tert-butylphenyl)phenoxy substituents at the perylene bisimide bay positions. These substituents direct a packing arrangement with full enwrapping of the chromophore and unidirectional chromophore alignment within the crystal lattice to afford optical properties that resemble those of their natural pigment counterparts, in which chromophores are rigidly embedded in protein environments.

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down by chromophore isolation and rigidification, and high chromophore concentrations are realized without compromising the dyes’ desirable photo-functional properties.

To synthesize a solid-state emitting material with fully enwrapped PBI fluorophores, we introduced bulky 2,4,6-tris(4-tert-butylphenyl)phenoxy substituents at the 1,7-bay positions of regioisomERICALLY pure N,N'-dicyclohexyl-1,7-dibromoperylene-3,4,9,10-tetracarboxylic acid bisamide (Figure 1a). By applying typical reaction conditions for nucleophilic aromatic substitution reactions, PBI 1 could be isolated as a bright fluorescent and thermally highly stable (up to > 350 °C) solid-state material in 78% yield after purification by column chromatography (Supporting Information, Scheme S1). The new solid-state emissive PBI derivative was unambiguously characterized by 1H and 13C NMR spectroscopy, high-resolution mass spectrometry, cyclic voltammetry, and elemental analysis (Supporting Information, Figure S1). Furthermore, solvent-free single crystals suitable for X-ray diffraction (XRD) analysis could be grown by slow evaporation of a dichloromethane/hexane 3:7 solution of PBI 1 at room temperature, providing unique insights into its molecular structure and packing arrangement in the solid state (Figures 1b–d and Supporting Information, Table S1 and Figure S2).

Similar to other 1,7-diphenoxysterminated PBI derivatives,[27,28] PBI 1 is characterized by an almost planar scaffold with dihedral angles of 1.8(4)° and 0.5(3)° for the four bay (ξ(C1-C12b-C12a-C12)) and four inner carbon atoms (ξ(C13a-C12a-C12b-C13b); for numbering see Figure 1a).

Owing to the crystallographic inversion center located in the center of gravity of the molecule (space group P1), the twist angle between both naphthalene mono-imide subunits is 0°. The phenoxy substituents are bent towards opposing faces of the PBI π-scaffold with dihedral angles of 31.1(3)° (ξ(C2-C1-O-Cn)1) and 34.0(5)° (ξ(C8-C7-O-Cn)1)) thereby efficiently shielding the chromophore’s π-scaffold (Figure 1b). Owing to the bulky 4-tert-butylphenyl moieties, the individual chromophores are well-separated in the solid state with an interplanar distance of 8.1 Å and a transversal and longitudinal displacement of 3.7 Å and 7.9 Å, respectively (Supporting Information, Figure S2). The packing arrangement of PBI 1 is characterized by strictly parallel oriented molecules with several C–H···O and C–C···π interactions and a total void volume of 34 Å3. Most importantly, in addition to the jacketing provided by each two tert-butylphenyl units attached at the ortho-position of the 1,7-bay-substituents (dark blue colored, Figures 1b–d), full chromophore enclosure is realized by tert-butylphenyl units of the neighboring chromophores (grey and light blue colored, Figures 1c,d). This desired feature became possible by the molecular design with three tert-butylphenyl units attached to the 2,4,6-positions of the chromophore-bound phenoxy units. The perfect chromophore embedment is corroborated by the absence of co-crystallized solvent molecules and an appreciably high density for an organic crystal of 1.156 g cm−3. All these structural features can likewise be found in the isolated bulk material according to powder XRD analysis (Supporting Information, Figure S3d) and do account for the bright solid-state luminescence of PBI 1 (see below).

The optical properties of PBI 1 were investigated by UV/Vis absorption and fluorescence spectroscopy, both in dichloromethane solutions and in the solid state (Figure 2 and Supporting Information, Figures S3a–c). The solid-state absorption profile of PBI 1 in reflection mode was determined on an ensemble of microcrystals in a BaSO4 trituration, applying the Kubelka–Munk theory (Figure 2).[38] Both absorption spectra reveal sharp S0→S1 transitions located at 559 nm (solution) and 557 nm (solid), respectively, each with a well-resolved vibronic progression at intervals of 1370 cm−1 and a molar extinction coefficient (εmax) of 7.2 × 104 M−1 cm−1.
Table 1: Spectroscopic properties of PBI 1 in dichloromethane solution and in the crystalline solid state at 298 K.

|          | $\lambda_{\text{em}}$ [nm] | FWHM$_{\text{em}}$ [cm$^{-1}$] | $\lambda_{\text{em}}$ [nm] | FWHM$_{\text{em}}$ [cm$^{-1}$] | $\Delta\Phi_{\text{em}}$ [cm$^{-1}$] | $\Phi_0$ [%] | $\tau_1$ [ns] |
|----------|----------------------------|-------------------------------|----------------------------|-------------------------------|-----------------------------------|------------|-------------|
| CH$_2$Cl$_2$ | 559                        | 750                           | 581                        | 1130                          | 680                               | 100        | 5.20        |
| Solid    | 559$^{[p]}$                | 1000                          | 565$^{[p]}$                | 570                           | 260                               | >90$^{[p]}$| 7.70$^{[p]}$|

[a] FWHM was derived as twice the distance between the maximum to the closest edge at half-maximum of the unsymmetrically shaped absorption or emission band to prevent falsification by overlapping transitions. [b] Ensemble property of crystalline material. [c] Single microcrystal investigated with an optical polarization microscope equipped with a fiber-coupled CCD spectrometer.

maximum (FWHM$_{\text{em}}$ = 570 cm$^{-1}$) of the shortest wavelength emission of microcrystalline PBI 1 is approximately half of that of its monomer emission in solution (Table 1). These observations can be rationalized by a rigidification of the chromophore π-scaffold by its enclosing environment in the solid matrix, thereby preventing structural relaxations of the excited molecules (as well as their environment) as it is given in solution. In accordance with this interpretation, an increased fluorescence lifetime of 7.70 ns was observed for the crystalline sample compared to the value of 5.20 ns in dichloromethane solution (Supporting Information, Figure S3b). Most importantly, the bright fluorescence of PBI 1 in solution ($\Phi_0 = 100\%$) is also retained in the solid state with a fluorescence quantum yield $\Phi_F$ of at least 90% after correcting the apparent solid-state emission quantum yield ($\Phi_F^* = 82\%$) for unavoidable reabsorption losses (Figure 2c and Supporting Information, Figure S3c).$^{[30]}

To gain further insights into the optical properties of this unique highly luminescent solid-state material, oriented microplatelets of PBI 1 were grown on quartz substrates$^{[31]}$ from dichloromethane/hexane 4:6 solutions and their anisotropy was investigated by optical polarization (fluorescence) microscopy (Figure 3 and Supporting Information, Figures S5–6). The crystallinity of these samples was demonstrated by selected area electron diffraction (SAED) and out-of-plane XRD experiments performed on microcrystals grown on a carbon-coated copper grid, confirming an identical solid-state packing arrangement as determined by single crystal structure analysis (see above; Supporting Information, Figures S7a–c). Independent on the substrate, the rhomboid crystals exhibit edge lengths of several tens of micrometers and angles of 107° and 73° (Figure 3b), respectively, with thicknesses of several hundred nanometers according to AFM cross-section analysis (Supporting Information, Figure S7d–e).

For an accurate comparison of the fluorescence properties of PBI 1 in solution and in the solid state, the intrinsic steady-state emission spectra have to be acquired without any contamination by reabsorption, which is rather challenging for solid-state samples of chromophores with high tinctorial strengths ($c_{\text{max}}$) and small Stokes shifts ($\Delta\Phi_{\text{em}}$). Whereas highly diluted solutions of PBI 1 ($OD < 0.05$; $c_{\text{max}} < 10^{-6} \text{mol} \cdot \text{L}^{-1}$) are routinely investigated using standard fluorescence spectrometers, the spectroscopic investigations on micrometer-sized single crystals of PBI 1 were performed with an optical polarization microscope equipped with a fiber optic-coupled CCD spectrometer. Thus, single crystals of different sizes and thicknesses were grown from dichloromethane/hexane 3:7 solutions and subsequently investigated (Supporting Information, Figure S4) to obtain nearly reabsorption-free solid-state emission spectra (Figure 2). These data are most accurate and allow for a quantitative analysis of the intrinsic solid-state fluorescence properties of PBI 1.

In contrast to previously reported 1,7-diphenoxy-substituted PBI derivatives,$^{[27,28]}$ PBI 1 displays a solid-state fluorescence spectrum that is an almost perfect mirror image of its absorption profile with distinct vibronic progressions at 565, 608, and 663 nm (each separated by 1360 cm$^{-1}$). It is remarkable that even the emission spectrum of PBI 1 in dichloromethane solution is more broadened, less structured, and more red-shifted by 490 cm$^{-1}$ (Figure 2 and Supporting Information, Figure S3a), which hitherto could only be observed for natural pigments (i.e., protein-encapsulated chromophores)$^{[29]}$ but not for synthetic pigments. Likewise, the Stokes shift ($\Delta\Phi_{\text{Stokes}} = 260 \text{ cm}^{-1}$) and the full width at half

Figure 3. a) Polarization microscopy spectra of a PBI 1 microcrystal on quartz displayed in (c). The gray solid line marks the absorption and the green area the excitation profiles of a band-gap filter cube, respectively. Inset: Respective change of the emission strength at 606 nm. b) Optical microscopy images of a microcrystal for different polarizations (arrow = 15 μm) of the transmitted light along (left) and perpendicular (right) to the long molecular axis including a schematic molecular arrangement. c) Fluorescence microscopy images of a microcrystal upon rotation with respect to parallel aligned polarizer (excitation) and analyzer (detection). The green square marks the area where the spectra were recorded. Scale bar = 50 μm.
f). Accordingly, owing to the high tinctorial strength, the emission spectra of those microplatelets are partially contaminated by reabsorption (see above) and no absorption spectra could be recorded in transmission mode. SAED and XRD experiments unambiguously confirmed that the investigated microcrystals are oriented with their (010) plane parallel to the surface (Figure 3b and Supporting Information, Figures S7a–c). The fast-growing directions of the crystals, which give rise to their rhomboidal shape, are presumably the (100) and (001) lattice planes as evidenced by the simulated Bravais–Friedel–Donnay–Harker (BFDH) morphology (Figure 3b). The long molecular axes of the PBI chromophores and thus their $S_1 - S_0$ transition dipole moments are unidirectionally aligned along the short rhomboid diagonal that intersects the long one at an angle of 86° (Figure 3b). Accordingly, rhomboid crystals of PBI 1 appear deep red in optical polarization microscopy when the transmitting light is polarized along the short rhomboid diagonal but become fully transparent when the polarization is rotated by 90° (Figure 3b and Supporting Information, Figure S6). This behavior becomes even more obvious in polarization-dependent fluorescence microscopy experiments starting with polarizer, analyzer, and short rhomboid diagonal all aligned parallel (Figures 3a,c). By rotating the short diagonal out of this arrangement the bright orange fluorescence gradually decreases and is completely absent at an angle of 90° (Figures 3a,c). Apparently, this process is characterized by a huge optical anisotropy as the emission intensity at 606 nm is reduced by 99% at a perpendicular orientation (Figure 3a). The high quality of the microcrystals and the perfect unidirectional alignment of the chromophores is thereby reflected in an outstanding dichroic ratio, $D_{\text{dichroic}}$ of 82 and an order parameter, $S_{\text{dichroic}}$ of 0.96.[42–44]

The spectroscopic data accordingly fully meet our expectations based on the packing features. Different from conventional pigments or other π-conjugated thin-film and solid-state materials in which electronic couplings from closely packed chromophore units lead to alterations of the spectral features with the commonly observed fluorescence quenching, the outstanding optical properties of PBI 1 are fully conserved and even improved in the solid state. Such features were hitherto exclusive for natural pigments in which evolutionarily optimized protein environments afford chromophore isolation and chromophore rigidification. For this reason, natural light-harvesting pigments can have quite high dye densities to afford the desired high absorption cross sections for the capture of sunlight without undesired “concentration”-quenching of the photo-excited state. For PBI 1-based thin films or crystals, the chromophore concentration is even higher than for any natural pigment because the chromophore part, that is, the perylene bisimide dye, constitutes 27.4% of the total molecular mass of PBI 1 ($c = 0.76 \text{ mol L}^{-1}$)[39] Supporting Information, Figure S9a). For most protein-based colorants this value is considerably smaller (Supporting Information, Figures S8 and S9), for example, 0.8% for the green fluorescent protein,[45] 0.54–0.85% for rhodopsin,[46,47] 9.7% for the Fenna–Matthews–Olson photosynthetic light-harvesting protein complex.[48,49]

In summary, we reported a unique perylene bisimide solid-state emitter, which readily crystallizes from dichloromethane/hexane solutions to form rhomboid microcrystals with absolute solid-state fluorescence quantum yields greater than 90%. The sterically demanding threefold tert-butyl-phenyl-functionalized 1,7-phenoxo substituents efficiently shield the planar chromophore core within the crystal lattice, providing complete wrapping and preventing electronic coupling between adjacent molecules, while maintaining an exceptional high chromophore concentration of more than 27% in the solid state. This well-defined intrinsic dilution of the chromophore core, like it is exclusively illustrated by many natural light-harvesting pigments, accounts for its remarkable solid-state luminescence and gives rise to a unique emission spectrum with a well-resolved vibronic progression. The perfect unidirectional alignment of the chromophores within the single crystalline material enables a huge optical anisotropy with a dichroic ratio of 82, making PBI 1 a highly attractive solid-state material for photonic applications.

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Conflict of interest

The authors declare no conflict of interest.

Keywords: crystal engineering · dyes · fluorescence quantum yield · perylene bisimides · solid-state emitters

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