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Synthesis of arrays containing porphyrin, chlorin, and perylene-imide constituents for panchromatic light-harvesting and charge separation†

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Achieving solar light harvesting followed by efficient charge separation and transport is an essential objective of molecular-based artificial photosynthesis. Architectures that afford strong absorption across the near-UV to near-infrared region, namely panchromatic absorptivity, are critically important given the broad spectral distribution of sunlight. A tetrapyrrole-perylene pentad array was synthesized and investigated as a means to integrate panchromatic light harvesting and intramolecular charge separation. The pentad consists of three moieties: (1) a panchromatically absorbing triad, in which a porphyrin is strongly coupled to two perylene-monoimides via ethyne linkages; (2) a perylene-diimide electron acceptor; and (3) a chlorin hole-trapping unit. Integrating the three components with diphenylethyne linkers generates moderate electronic coupling for intramolecular energy and hole/electron transfer. The construction of the array relies on a stepwise strategy for incorporating modular pigment building blocks. The key building blocks include a trans-A₂BC porphyrin, a chlorin, a perylene-monoimide, and a perylene-diimide, each bearing appropriate (halo, ethynyl) synthetic handles for Pd-catalyzed Sonogashira coupling reactions. One target pentad, three tetrads, four triads, and four monomeric benchmark compounds were synthesized from six building blocks (three new, three reported) and 10 new synthetic intermediates. Four of the tetrapyrrole-containing arrays are zinc chelated, and four others are in the free base form. Absorption and fluorescence spectra and fluorescence quantum yields were also measured. Collectively, investigations of the arrays reveal insights into principles for the design of novel reaction centers integrated with a panchromatic antenna for artificial photosynthetic studies

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Introduction

A chief objective of artificial photosynthesis is to create model systems that mimic native photosynthetic processes and through systematic modifications can be used to probe relevant physicochemical features. The processes encompass diverse phenomena including light capture, excited-state energy transfer, excited-state electron transfer, dark electron/hole transfer, proton pumping, carbon fixation, and water splitting. While early model systems were first reported decades ago, 1-12 the ensuing half-century of research has not resolved

fundamental synthetic limitations to the creation of molecular architectures that rival those in native photosynthesis. Such architectures require large numbers of pigments and redox cofactors organized in exacting 3-dimensional spatial arrangements over mesoscale dimensions. Given the challenges of creating such architectures, an alternative tack entails development of more compact structures with designs complementary to those in native photosynthesis. 13-27

As one case in point, we recently developed a design motif that engenders panchromatic absorption yet affords a lowest singlet excited-state of discrete energy. The motif relies on a perylene-monoimide attached to a porphyrin via an ethyne linker, where the linker joins the perylene 9-position and the porphyrin meso-position, both of which are sites of high electron density in the highest occupied molecular orbital (HOMO) of each constituent. Such a triad (PMI_{2T}P) composed of two perylene-monoimides tightly coupled electronically (via ethyne linkers) to one porphyrin is shown in Fig. 1. The triad absorbs strongly across the 400–700 nm region (average peak $\varepsilon \sim 80~000~\text{M}^{-1}~\text{cm}^{-1}$), yet exhibits excited-state features similar to those of the tetrapyrrole Indeed, the fluorescence emission

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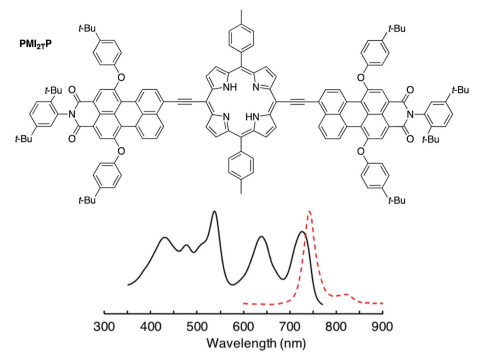


Fig. 1 The absorption (black solid) and fluorescence (red dashed) spectra of a prior panchromatic triad PMI_{2T}P

spectrum generally resembles that of a free base tetraarylporphyrin, although shifted bathochromically, and the fluorescence quantum yield (Φ_f) is considerably enhanced versus that of the porphyrin alone. This latter effect arises from a dramatic shift in radiative probability from the normally intense porphyrin near-UV Soret band(s) to the normally weak porphyrin visible bands including the $S_0 \rightarrow S_1$ absorption (and $S_1 \rightarrow S_0$ fluorescence) transition.³⁰ Because the triad exhibits tetrapyrrole-like lowest singlet excited-state properties we considered that the triad could be used as an antenna element in conjunction with an integrated charge-separation unit. The resulting integrated pentad array (C-T-PDI) contains a perylenemonoimide-porphyrin-perylene-monoimide triad unit (T), a perylene-diimide (PDI), and a chlorin (C) (Chart 1).

The processes envisaged for light harvesting and charge separation in the pentad C-T-PDI are illustrated in Fig. 2. The central triad acts as a supermolecule with panchromatic absorption (see e.g. Fig. 1) as a result of the substantial electronic interactions between the porphyrin and perylenemonoimides afforded by the direct ethyne linkers. Excitation in any of the absorption bands of the triad is followed by rapid, quantitative energy flow to the tetrapyrrole-like lowest energy singlet excited state (S₁) of this array, analogous to internal conversion from upper to lower energy excited states of a simple chromophore. The core porphyrin is of a trans-A₂BC design. The perylene-diimide and chlorin are each attached to the core porphyrin via a diphenylethyne linker, which affords electronic interactions that while weaker than a direct ethyne linkage, enable electronic coupling sufficient to support rapid energy and hole/electron transfer with only slight perturbation of energy levels and electronic spectra. The perylene-diimide serves as an acceptor upon excited-state electron transfer from

the excited triad (T*), whereas the chlorin functions as a hole trap via a ground-state process involving the oxidized triad (T⁺). The overall light-driven process is thus expected to afford the chlorin cation radical and perylene-diimide anion radical upon charge separation across the pentad (forming C⁺-T-PDI⁻). While the chief functions of the perylene-diimide and the chlorin are for charge separation, both also absorb light and are expected to afford additional light-harvesting capacity to the pentad by funneling harvested excitation energy to the S1 excited state of the central triad.

Herein, we report the synthesis of the pentad array as well as all companion arrays for full photophysical characterization. The synthesis relies on a building block approach³¹⁻³⁴ wherein constituent chromophores are synthesized separately and then joined together through Pd-mediated coupling reactions. The companion arrays include three tetrads (Chart 2), four triads (Chart 3), and four monomeric compounds (Chart 4). The 12 targets (one pentad and 11 benchmark compounds) were prepared from six building blocks (three new, three reported) and 10 new synthetic intermediates. Absorption and fluorescence studies are also reported as a first step toward in-depth analysis of integrated light-harvesting and charge-separation processes in the arrays.

Results and discussion

Molecular design

The rationale for the selection of components of the pentad was based on optical and redox properties. In particular, a chlorin with sparse substitution35 was chosen given that such chlorins exhibit a wavelength absorption band at shorter wavelength, and thus an S1 excited-state at higher energy, than that of the

Chart 1 Target pentad array.

central bis(perylene-monoimide)porphyrin triad. The lower S_1 energy of the central panchromatic porphyrin-based triad unit *versus* that of a standard chlorin is one of the functionally important features of the C-T-PDI pentad. Redox potentials of reported benchmark compounds with similar structures of the chlorin, panchromatic bis(perylene-monoimide)porphyrin triad, and the perylene-diimide constituents^{28,36–38} were considered in the design of an array to achieve photoinduced charge separation. The envisaged processes based on the choices are illustrated in Fig. 2: the central panchromatic triad (in both the ground and excited state) is more readily oxidized compared to the perylene-diimide and more readily reduced relative to the chlorin moiety, whereupon a charge-separated state is expected with the electron localized at the perylene-diimide and the hole trapped at the chlorin.

One slight structural modification in the arrays reported here compared with the previously made triad $PMI_{2T}P$ is the N-substituents of the imido groups in the perylenes.^{29,39} The prior

2,5-di-*tert*-butylphenyl substituents are replaced with 2,6-diiso-propylphenyl groups, because the former substituents can give rise to atropisomers,⁴⁰⁻⁴² thereby complicating NMR analysis. The use of 2,6-diisopropyl *versus* 2,5-di-*tert*-butyl groups for solubilization has insignificant effects on perylene photophysics and energetics,⁴³ in large part due to the presence of a node at the perylene-imide nitrogen atom in both the HOMO and the lowest unoccupied molecular orbital (LUMO).⁴¹

Synthesis

Eleven target compounds (Charts 2–4) were prepared for comparison with the pentad, including two free base tetrads (T-PDI and C-T, omitting either the chlorin or the perylene-diimide moiety in the pentad) and a zinc chelate (ZnT-PDI); two free base triads (T-Ph and C-P-PDI) and their zinc chelates (ZnT-Ph and ZnC-ZnP-PDI); a monomeric perylene-diimide (PDI-Ph); a monomeric chlorin (C-Ph); and a monomeric porphyrin (P-TMS/TIPS) bearing ethynes with trimethylsilyl and

excited-state

energy transfer

Paper

Light
Harvesting

excited-state
energy transfer

Charge
Separation

ground-state
hole transfer

Perylene
Monoimide

N HN

Perylene
Monoimide

Linker

Fig. 2 Illustration of the expected light-harvesting and charge-separation processes in pentad C-T-PDI.

Perylene

Diimide

excited-state

electron transfer

triisopropylsilyl protecting groups, and related zinc chelate (**ZnP-H/TIPS**). Zinc chelates were prepared to tune the electrochemical properties (because zinc chelates are more easily oxidized than the free base forms), providing more insights to the intramolecular electron–transfer processes in the arrays.

The pentad **C-T-PDI** contains four functional entities attached to the core porphyrin *via* ethynyl and ethynylphenyl linkers. The building blocks include a *trans*-A₂BC-porphyrin,⁴⁴ which constitutes the core of the pentad and establishes a starshaped architecture; an iodophenyl–chlorin,³⁵ an ethynyl-perylene-monoimide,⁴⁵ and an iodophenyl-perylene-diimide.^{37,42} The preparation of tetrapyrrole arrays with direct ethynyl linkages was first reported by Arnold,⁴⁶ next developed by Anderson⁴⁷ and Therien,⁴⁸ and has since found widespread use.^{18,26,49–53} The chief reaction for joining building blocks to construct either direct ethynyl linkers or longer diphenylethyne linkers entails Pd-mediated Sonogashira coupling of halo-(iodo or bromo) and ethynyl-substituted reactants. Such reactions with free base tetrapyrroles must be carried out in the absence of copper to avoid unwanted metalation.^{54,55}

Porphyrin building blocks. Two *trans*-A₂BC-porphyrins, 2-Br₂/TMS and 4-Br₂/TMS/TIPS, provided building blocks for the benchmark tetrads (C-T and T-PDI) and the target pentad C-T-PDI, respectively. The "2 + 2" condensation⁵⁶ of the diimino derivative of 1⁵⁷ and dipyrromethane 1-TMS^{57,58} generated the corresponding *trans*-AB zinc porphyrin Zn2-TMS in 40% yield. Subsequent demetalation⁴³ (58%) and bromination⁵⁹ with *N*-bromosuccinimide (NBS) of the two open *meso* positions (74%) afforded the desired free base *meso*-dibromoporphyrin 2-Br₂/

TMS bearing a phenyl group and a TMS-protected ethynylphenyl group (Scheme 1).

Porphyrin building block 4-Br₂/TMS/TIPS (Scheme 2) bears two distinct ethynylphenyl units (with TMS and TIPS protecting groups for successive coupling reactions) and was synthesized in a strategy similar to that for porphyrin 2-Br₂/TMS. A dibutyltin-chelated diformyldipyrromethane 3,56 which was more readily purified than the free base counterpart, was synthesized according to the literature. The zinc trans-ABporphyrin Zn4-I/TMS, which bears one iodophenyl group and one TMS-protected ethynylphenyl group, was prepared therefrom with 1-TMS in 15% yield. Sonogashira coupling of Zn4-I/ TMS and TIPS-acetylene in the presence of catalytic CuI (1 mol% of the porphyrin) produced the zinc porphyrin Zn4-TMS/TIPS in 80% yield. Subsequent bromination⁵⁹ and demetalation⁴³ generated the free base dibromoporphyrin 4-Br₂/TMS/TIPS in a yield of 27% for two steps. The zinc-chelated benchmark compound ZnP-H/TIPS was also synthesized from the trans-ABporphyrin Zn4-TMS/TIPS in 44% yield, through three successive steps of bromination, coupling with phenylacetylene, and cleavage of the TMS group.

Chlorin building block and benchmark. The known iodophenyl-chlorin 5⁶⁰ was employed as a building block for integration into the pentad. Pd-mediated coupling of the zinc chlorin counterpart Zn5⁶⁰ with phenylacetylene in tetrahydrofuran (THF) containing triethylamine (TEA) followed by demetalation with trifluoroacetic acid afforded the chlorin benchmark C-Ph in 84% yield (Scheme 3).

Perylene building blocks. The iodophenyl-perylene-diimide 7 was prepared by statistical imidation of the perylene-bis(anhydride) 6 with two anilines—2,6-diisopropylaniline (3.0 equiv.) and 4-iodoaniline (2.0 equiv.)—in 30% yield (Scheme 4), following a reported method.³⁷ The difference in the loading equivalents of two anilines is due to their unequal reactivity. Surprisingly, the reported synthesis of the unsymmetric perylene-diimide by partial hydrolysis of a symmetric perylene-diimide precursor^{37,42} failed in our hands to generate the desired product 7. The perylene-diimide benchmark PDI-Ph was prepared thereafter by coupling with phenylacetylene in 52% yield. Synthesis of the perylene-monoimide building block 8 was previously reported.³⁴

Construction of the benchmark triads and tetrads. With the five building blocks in hand (porphyrin 2-Br2/TMS and 4-Br2/ TMS/TIPS, chlorin 5, perylene-diimide 7, and perylenemonoimide 8), the target arrays and benchmark compounds (Charts 2 and 3) were prepared through a series of Pd-mediated coupling reactions. Syntheses of triads T-Ph and ZnT-Ph, tetrads T-PDI, ZnT-PDI, and C-T all started from the shared porphyrin building block 2-Br₂/TMS. Stepwise coupling with perylenemonoimide 8 followed by chlorin 5 (for C-T) or perylenediimide 7 (for **T-PDI**) afforded the free base arrays (Scheme 5). Pd-mediated Sonogashira coupling of porphyrin 2-Br₂/TMS and perylene-diimide 8 produced the triad T-Ph in 86% yield. Metalation of T-Ph with zinc acetate gave the zinc-chelated triad ZnT-Ph in 45% yield. The TMS-protecting group in T-Ph was removed by treatment with K2CO3 in a mixture of toluene/ CH₃OH, affording the ethynylphenyl triad T-Ph-H

Chart 2 Constituent tetrad benchmarks.

quantitatively. Thereupon, **T-Ph-H** was coupled in the presence of $Pd_2(dba)_3/P(o\text{-tol})_3$ either with iodophenyl-chlorin 5 or iodophenyl-perylene-diimide 7 to generate the tetrad **C-T** or **T-PDI** in 24% or 40% yield, respectively. Metalation of **T-PDI** with zinc acetate afforded the zinc tetrad **ZnT-PDI**.

Syntheses of the target pentad and chlorin-porphyrin-perylene-diimide triads. The target pentad C-T-PDI and the chlorin-porphyrin-perylene-diimide triads (C-P-PDI and ZnC-ZnP-PDI) were also prepared through stepwise coupling reactions with the same building block porphyrin 4-Br₂/TMS/TIPS. The distinct protection of the ethyne units with trimethylsilyl

Chart 3 Constituent triad benchmarks.

and triisopropylsilyl groups afforded the opportunity for selective reactions.

Phenylethynyl groups were installed at the meso-positions of 4-Br₂/TMS/TIPS through a Sonogashira reaction to afford the

porphyrin benchmark **P-TMS/TIPS** in 78% yield (Scheme 6). The TMS group in **P-TMS/TIPS** was thereafter removed with K₂CO₃ at room temperature, whereupon coupling of the resulting porphyrin and perylene-diimide 7 afforded the porphyrin–perylene-diimide

Chart 4 Constituent monomer benchmarks.

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Scheme 1 "2 + 2" condensation to prepare porphyrin building block 2-Br₂/TMS

dyad 9 (49% yield). Dyad 9 has limited solubility in many organic solvents (toluene, CH₂Cl₂, chloroform, THF, acetonitrile, methanol, or DMSO), but the solubility could be improved upon zinc chelation (not shown). Despite the meager solubility, dyad 9 in a saturated solution (\sim 1 mM) was deprotected with tetrabutylammonium fluoride (TBAF) and then coupled with the chlorin building block 5 in the presence of Pd₂(dba)₃/P(o-tol)₃ to produce the chlorin-porphyrin-perylene-diimide triad C-P-PDI. Subsequent zinc chelation generated the triad ZnC-ZnP-PDI, wherein both the porphyrin and the chlorin macrocycles are zinc chelated.

The pentad C-T-PDI was synthesized similarly through the stepwise incorporation of various building blocks (Scheme 7). Perylene-monoimide 8, instead of phenylethynyl groups in the case of C-P-PDI, was installed on the meso-positions of the porphyrin 4-Br₂/TMS/TIPS, generating the panchromatic triad T-TMS/TIPS. A succession of reactions – deprotecting with K₂CO₃, coupling with perylene-diimide 7, and again deprotecting with TBAF - afforded the tetrad T-H/PDI in 12% overall yield for the three steps. The final coupling entailed reaction of the tetrad with

Scheme 2 Preparation of the porphyrin building block 4-Br₂/TMS/ TIPS and benchmark ZnP-H/TIPS.

iodophenyl-chlorin 5 to complete the synthesis of the target pentad C-T-PDI in 27% yield. The pentad-forming reaction was monitored with analytical size exclusion chromatography (SEC), as has been done previously with multiporphyrin arrays (see ESI†).31,32 The pentad array and all benchmarks were also characterized by analytical SEC and MALDI-MS prior to any photophysical studies. On the basis of the analytical SEC traces, the purity of each array was estimated to be >99%, which is suitable for further photophysical characterization.

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Scheme 3 The chlorin building block 5 and the preparation of benchmark compound C-Ph.

Photophysical studies

Absorption spectra. Steady-state absorption spectra of the pentad C-T-PDI and all benchmark arrays were measured (in toluene at room temperature) as a first step toward understanding intramolecular photophysical processes. Spectra for the pentad and constituent tetrads and triads containing panchromatic bis(perylene-monoimide)porphyrin (denoted T), along with relevant monomers, are shown in Fig. 3 (solid). Spectra for related reference arrays in which the bis(perylene-monoimide)porphyrin core is replaced by a simple porphyrin, along with associated monomers, are shown in Fig. 4 (solid). The absorption spectral properties are summarized in Table 1. Consistent with the previously reported panchromatic triads, 29,39 the triad T-Ph shows a broad distribution of absorption from the near-UV to near-IR region. Compared to the spectrum of a typical porphyrin (see that for P-TMS/TIPS in Fig. 4C), the spectral distribution for **T-Ph** (average $\varepsilon \sim$ 80 000 M⁻¹ cm⁻¹) reflects a substantial shift of absorption strength from the typically intense porphyrin near-UV Soret feature(s) (typical peak ε of 350 000–400 000 M^{-1} cm⁻¹) into the typically weak visible transitions (typical peak ε of 10 000-20 000 M⁻¹ cm⁻¹), accompanied by a bathochromic shift of features such as the $S_0 \rightarrow S_1$ band along with the possible generation of new bands (e.g. at ~650 nm). Corresponding differences can be seen for the corresponding zinc chelatespanchromatic triad ZnT-Ph (Fig. 3F) compared to porphyrin ZnP-H/TIPS (Fig. 4D). The substantial shift of intensity from the porphyrin near-UV Soret region into and across the visibleregion features upon attachment of perylene-monoimides via

Scheme 4 Synthesis of the perylene-diimide building block 7, and the benchmark PDI-Ph.

direct ethyne linkers arises at least in part from a significant diminution in mixing of the key porphyrin one-electron excited-state configurations within the four-orbital model. Such a diminution in configuration interaction reduces the constructive configurational interference that affords the normal intense near-UV bands and the destructive interference that gives the typically weak visible bands within the Gouterman four-orbital model. A decrease in configuration interaction is further accentuated by the shifts in energies of the frontier molecular-orbital, which also contributes to the bathochromic shift in the $S_0 \rightarrow S_1$ transition.

As expected, the absorption spectra of the pentad (C-T-PDI) and two tetrads (C-T and T-PDI) show panchromatic features similar to those of triad T-Ph (Fig. 3, panels A-C and E). In analogy, the spectrum of the tetrad containing a zinc porphyrin (ZnT-PDI) has similar panchromaticity as reference triad ZnT-Ph (Fig. 3, panels D and F). The spectra of the tetrads generally reflect the contributions of the corresponding triad (T-Ph or ZnT-Ph) and monomer (C-Ph or PDI-Ph), with some modest changes in relative intensities and peak positions. In turn, the spectrum of pentad C-T-PDI has features that correspond to those in the spectra of tetrad C-T and monomer PDI-Ph, or tetrad T-PDI and monomer C-Ph, again with some shifts in intensities and wavelengths. For example a slight bathochromic shift in the Soret band of the chlorin moiety (at 418 nm) is

Scheme 5 Preparation of triads (T-Ph, ZnT-Ph) and tetrads (C-T, T-PDI, ZnT-PDI).

observed in the pentad compared to the tetrad C-T (2 nm) and chlorin benchmark C-Ph (4 nm). Broadening of this band is also observed for the pentad [full width at half maximum (FWHM) = 65 nm] compared to the tetrad C-T (40 nm) and the monomer C-Ph (33 nm).

The band at \sim 530 nm due to perylene-diimide absorption is essentially superimposable on that of the tetrad **T-PDI** and the monomeric PDI-Ph, except for a 6 nm hypsochromic shift for the former relative to the latter. For pentad C-T-PDI, the $S_0 \rightarrow S_1$ feature at ~741 nm due to the absorption of the panchromatic bis(perylene-monoimide)porphyrin moiety shows a bathochromic shift and broadening (FWHM = 54 nm) compared with that of tetrad C-T (727 nm, FWHM = 46 nm), tetrad T-PDI (728 nm, FWHM = 71 nm), and triad T-Ph (725 nm, FWHM = 46)nm). This effect on the $S_0 \rightarrow S_1$ energy and also in the relative intensity of the absorption band is likely due in part to substituent effects on the electronic properties of the central triad. In particular, T-Ph contains one phenyl ring and one ethynylphenyl group at the two meso-positions apart from the meso-positions that bear ethynyl linkers to the perylenemonoimides (Chart 3). In each tetrad the ethynylphenyl group

is converted to a diphenylethyne linker to the chlorin or perylene-diimide (Chart 2). The pentad has two such diphenylethyne groups, terminated with the chlorin or perylene-diimide (Chart 1). Thus, in progressing from the triad to pentad, the change in terminal moieties on the ethynylphenyl and phenyl groups propagates to the porphyrin and sufficiently alters the porphyrin molecular orbital energies and configuration interaction to affect the characteristics of the $S_0 \rightarrow S_1$ transition.

ZnT-PDI

Despite the above-noted differences in certain spectral characteristics, the generally similar spectral features of the pentad and the corresponding constituents indicate that ground-state electronic interaction between the panchromatic core and either the perylene-diimide or the chlorin (*via* a diphenylethyne linker) are relatively weak, but not zero. Thus, addition of the chlorin and perylene-diimide moieties does not impart a substantial electronic perturbation to the central triad. In contrast, replacing the perylene-monoimide-ethynyl units attached to the central porphyrin with phenylethynyl groups restores the strong Soret bands and effectively diminishes panchromaticity, affording the spectrum of the triad **C-P-PDI** as a linear combination of those from the three monomeric

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Scheme 6 Synthesis of triads C-P-PDI and ZnC-ZnP-PDI from porphyrin 4-Br₂/TMS/TIPS.

components (Fig. 4, panels A, C, E and F). The same is true for the triad containing a central zinc porphyrin (ZnC-ZnP-PDI) relative to its three components (Fig. 4, panels B, D-F).

Fluorescence spectra. The fluorescence emission spectra of the pentad and all benchmark arrays measured in toluene at room temperature are also shown in Fig. 3 and 4 (dashed lines). The emission spectral properties including the (Stokes) shift between the $S_0 \rightarrow S_1$ absorption and $S_1 \rightarrow S_0$ emission maxima are listed in Table 1. The emission pattern of the panchromatic

arrays (tetrad C-T and T-PDI, and the triad T-Ph) is porphyrinlike, albeit at much lower energy than a typical monomeric porphyrin, or even P-TMS/TIPS for which the meso-substituents (two phenylethynyl groups, an ethynylphenyl and a phenyl ring) cause a significant bathochromic shift compared to a porphyrin lacking ethyne-containing substituents. As reported previously, 28,30 arrays that contain strongly electronically coupled porphyrin-perylene-monoimide units exhibit a chromophoreinduced reduction in configuration interaction involving the four frontier molecular orbitals of the central porphyrin, which results in a redistribution of intensity among the bands in the absorption spectra. This effect accounts at least in part for the non-mirror-image relationship observed for the absorption and the fluorescence of the porphyrin-bis(perylene-monoimide)containing architectures. A change in nuclear coordinates of the array and the solvent may also contribute to diminished mirror symmetry, and also to a somewhat larger Stokes shift for such architectures compared to a typical porphyrin.³⁰

There is a resemblance of the fluorescence spectra of the two tetrads (C-T and T-PDI) and that of the triad T-Ph, as well as a similar resemblance between the emission of the zinc-chelate tetrad **ZnT-PDI** and reference monomer (Fig. 3, panels D and F). Fluorescence emission of the pentad and tetrads are predomithe central panchromatic bis(perylenefrom monoimide)-porphyrin unit, regardless of excitation wavelength. This is shown in Fig. 4A for pentad C-T-PDI, for excitation wavelengths that should primarily excite the chlorin or perylenes. This finding is consistent with efficient energy transfer from either the chlorin (in C-T) or the perylene-diimide (in **T-PDI**) to the central bis(perylene-monoimide)porphyrin.

As noted above, prior work on arrays that contain one (or more) perylene-monoimide unit(s) coupled via an ethyne to a porphyrin are best treated as a supermolecule: excitation within any absorption band, including features in the green region that one would normally associate with the pervlene, results in rapid internal conversion within the array to produce the tetrapyrrole-like S₁ excited state, from which fluorescence occurs. Thus, it is expected that rapid energy transfer from the excited chlorin or excited perylene-diimide to the bis(perylenemonoimide)porphyrin unit in the pentad and tetrads will give rise to tetrapyrrole-like emission from the S₁ excited state of that triad unit. This is observed for the pentad and tetrads, even if the (Förster) spectral overlap of the emission from the excited perylene-diimide unit (e.g., in C-T-PDI and T-PDI) would be the greatest with the green-region absorption of nominal perylenemonoimides of the central panchromatic bis(perylenemonoimide)porphyrin triad. Thus, observations here support the view that the arrays containing a bis(perylene-monoimide) porphyrin unit are efficient in absorbing light spanning the near-UV and into the near-IR and producing a discrete tetrapyrrole-like S1 state from which charge separation can ensue.

In comparison with the aforementioned arrays, the fluorescence of the non-panchromatic triad C-P-PDI is at a higher energy, because no strongly coupled pervlene-monoimide pigments are integrated at the central porphyrin. However, the presence of the two meso-phenylethynyl groups on the

ZnC-ZnP-PDI

C-T-PDI

Scheme 7 Synthesis of the target pentad C-T-PDI.

central porphyrin causes sufficient bathochromic shifts in the $S_0 \rightarrow S_1$ absorption and corresponding $S_1 \rightarrow S_0$ fluorescence bands that the S_1 excited state remains at a lower energy than the S_1 of the chlorin and the perylene-diimide pigments (Fig. 4). The same is true for triad **ZnC-ZnP-PDI**, in which the central bis(phenylethynyl)porphyrin is a zinc chelate, which tends to hypsochromically shift the $S_0 \leftrightarrow S_1$ transitions compared to

those of the free base analogue (Fig. 4F). Thus, the observation again of porphyrin fluorescence independent of excitation wavelength indicates that these triads afford efficient energy transfer from the two terminal chromophores to the central porphyrin, from which charge separation can occur.

Fluorescence quantum yields. The fluorescence quantum yields (Φ_f) for all the arrays prepared herein along with those for

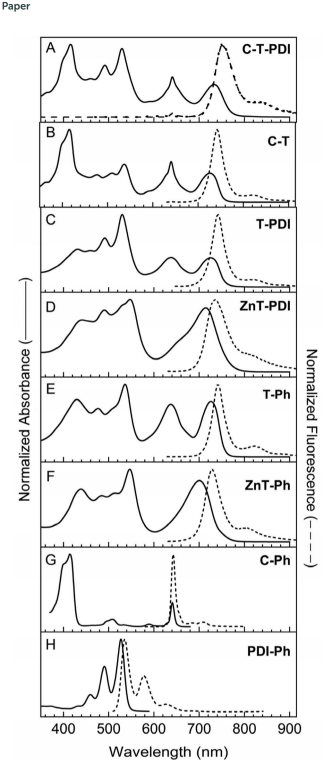


Fig. 3 The absorption (solid) and fluorescence (dashed) spectra of pentad (A) containing a central panchromatic bis(perylene-monoimide)porphyrin triad (denoted T) and benchmark tetrads (B–D), triads (E and F) and monomers (G and H). In (A), fluorescence spectra for the pentad were obtained using primarily excitation of a nominal absorption band of the chlorin or perylenes, with essentially the same result.

relevant benchmarks are given in the last column of Table 1. Starting with the panchromatic arrays, pentad C-T-PDI and tetrads C-T, T-PDI, and ZnT-PDI show no apparent dependence

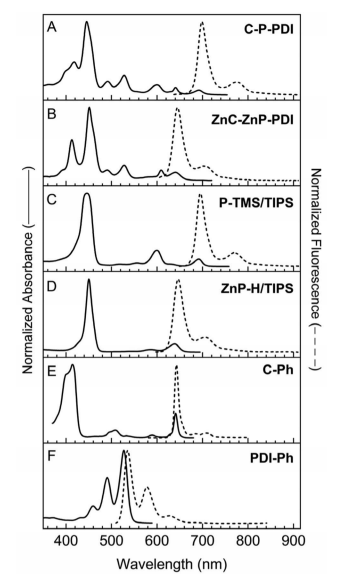


Fig. 4 The absorption (solid) and fluorescence (dashed) spectra of triads (A and B) containing a central porphyrin (denoted P) and benchmark monomers (C–F).

of the $\Phi_{\rm f}$ value on excitation wavelength. The same is true of benchmark triads **T-Ph** and **ZnT-Ph**. As can be inferred from the fluorescence spectra described in the prior section, these $\Phi_{\rm f}$ values reflect emission from the excited bis(perylene-monoimide)porphyrin triad unit (T*) in these arrays.

The $\Phi_{\rm f}$ of C-T (0.26) and T-PDI (0.23) are reduced from that for the benchmark panchromatic triad (0.30) by 13% and 23%, respectively, indicating modest yields of excited-state electron transfer. On the basis of the redox properties, the most likely product of hole/electron transfer would be C⁺-T⁻ and T⁺-PDI⁻ in the two tetrads, respectively. The $\Phi_{\rm f}$ for pentad C-T-PDI (0.16) is reduced by 47% from the panchromatic triad benchmark, suggesting that the yield of hole/electron-transfer initiated in the excited triad (T*) in this case is roughly the sum of the yields in the two tetrads. These measurements do not identify the product or product ratio, but the results tend to suggest that both C⁺-T⁻-PDI and C-T⁺-PDI⁻ form in the pentad. A subsequent ground-

Table 1 Spectral properties of perylene-porphyrin arrays and benchmarks"

Compound	Abs $\lambda 1 \text{ (nm)}$	Abs λ1 (nm) Abs λ2 (nm) Abs λ3 (nm)	Abs $\lambda 3 \text{ (nm)}$	Abs $\lambda 4 \text{ (nm)}$	Abs λ5 (nm)	Abs $\lambda 6 \text{ (nm)}$	Abs $\lambda 4$ (nm) Abs $\lambda 5$ (nm) Abs $\lambda 6$ (nm) Abs $\lambda 6$ (nm) Flu $\lambda 1$ (nm)	Flu $\lambda 1~(\mathrm{nm})$	Flu $\lambda 2~(nm)$	Flu $\lambda 2 \text{ (nm)}$ Abs–Flu $(\text{cm}^{-1})^b$	$\phi_{ m f}$
Panchromatic arrays	arrays										
C-T-PDI	398^{c} (0.78)	416 (1.0)	492(0.71)	530(0.95)		641(0.55)	734 (0.45)	753(1.0)	841 (0.23)	342	0.16
C-T	403^{c} (0.84)	415 (1.0)	476 (0.38)	537 (0.52)		640 (0.56)	726 (0.40)	742 (1.0)	824 (0.10)	293	0.26
I-PDI		432(0.52)	491 (0.77)	530(1.0)		638(0.41)	725 (0.41)	742 (1.0)	829 (0.10)	309	0.23
ZnT-PDI		441(0.72)	491 (0.85)	547 (1.0)		663^{c} (0.46)	714 (0.88)	735(1.0)	816 (0.23)	400	0.13
T-Ph		431(0.79)	476(0.67)	536 (1.0)		637 (0.73)	726 (0.77)	741 (1.0)	824 (0.15)	275	0.30
ZnT-Ph		439 (0.71)	485 (0.64)	547 (1.0)		$662^{c} 0.48$	700 (0.85)	728 (1.0)	807 (0.20)	547	0.22
Non-panchromatic arrays	natic arrays										
S-P-PDI	$418\ (0.45)$	445 (1.0)	492 (0.19)	528(0.27)	600 (0.14)	641 (0.10)	692(0.66)	698 (1.0)	775 (0.18)	118	0.17
ZnC-ZnP-PDI	413 (0.56)	451 (1.0)	490(0.15)	528 (0.21)		609 (0.15)	641 (0.12)	646 (1.0)	706 (0.20)	130	0.04
Benchmark monomers	onomers										
C-Ph	401(0.87)	414 (1.0)	496 (0.075)	508 (0.10)	589 (0.04)	640 (0.29)		641 (1.0)	706 (0.09)	30	0.32
PDI-Ph			491(0.61)	521 (1.0)				536 (1.0)	578 (0.57)	301	0.99
P-TMS/TIPS		445 (1.0)			600(0.22)		692 (0.10)	695(1.0)	772 (0.19)	77	0.16
ZnP-H/TIPS		450 (1.0)			587 (0.03)		639 (0.11)	647 (1.0)	707 (0.20)	201	0.10

All data were acquired in toluene at 298 K. The value in parenthesis following the absorption or fluorescence wavelength is the intensity relative to the maximum in the spectrum. b Shift between $s_0 \rightarrow s_1$ absorption and $s_1 \rightarrow s_0$ fluorescence maxima. c Shoulder. the S_0

state hole/electron step would be needed for each intermediate to produce the full charge-separated state C^+ -T-PDI $^-$.

Tetrad **ZnT-PDI** and benchmark panchromatic triad **ZnT-Ph** were prepared to test whether the more facile oxidization of a zinc chelate *versus* the free base form would enhance excited-state electron transfer in these types of systems. The Φ_f of **ZnT-PDI** (0.13) is reduced by 41% compared to benchmark panchromatic triad **ZnT-Ph** (0.22), indicating that the electron-transfer yield (to produce $\text{ZnT}^+\text{-PDI}^-$) is roughly doubled relative to the yield in the analogous tetrad **T-PDI** that contains the free base porphyrin in the panchromatic triad unit. The simplest interpretation is that stabilization of the electron-transfer product ($\text{ZnT}^+\text{-PDI}^-$ *versus* $\text{T}^+\text{-PDI}^-$), together with the higher S_1 excited-state energy of ZnT^* *versus* T^* (Fig. 3 and 4) places the process in the zincated array on a more favorable location on the Marcus rate *versus* free energy curve.

The analysis now turns to the triads C-P-PDI and ZnC-ZnP-PDI in which the central panchromatic triad of the abovementioned pentad (and tetrad benchmarks) is replaced by a porphyrin, albeit one with two phenylethynyl and one ethynylphenyl substituents that renders the porphyrin lowest singlet excited state the S₁ excited state of the entire array (vide supra). As noted above, the redox properties of analogous arrays suggest that the central porphyrin in these triads will be harder to oxidize than the panchromatic triad unit in C-T-PDI, T-PDI or **ZnT-PDI**. The Φ_f in triad **C-P-PDI** (0.17) is essentially the same as that in porphyrin benchmark P-TMS/TIPS (0.16) consistent with no excited-state electron transfer. On the other hand, the $\Phi_{\rm f}$ in triad ZnC-ZnP-PDI (0.04) is reduced by 60% versus porphyrin benchmark ZnP-H/TIPS (0.10) indicating a substantial yield of excited-state electron transfer. Whether this reflects formation of ZnC⁺-P⁻-PDI or ZnC-P⁺-PDI⁻ or some combination in the first step and whether this is followed by the ground-state process(es) that would produce the ZnC+-P-PDI fully chargeseparated array requires in-depth time-resolved absorption studies, which are beyond the scope of this current work. Regardless, the present studies show that making both the chlorin hole trap and the porphyrin excited-state hole/electron donor simultaneously more easily oxidized compared to the free base porphyrin analogue has a dramatic effect on the photodynamics of the arrays.

Conclusions and outlook

One target pentad and 11 benchmark compounds were prepared at the 10 mg scale. The two *trans*-A₂BC dibromoporphyrin building blocks anchor the star-shaped architecture and were readily accessible through an established "2 + 2" condensation of dipyrromethanes followed by bromination. A modular strategy was employed for the construction of the arrays, in which several pigment building blocks (porphyrin, chlorin, perylene-diimide, and perylene-monoimide) were readily synthesized and then integrated into arrays *via* copper-free Pd-mediated Sonogashira coupling reactions. This building block approach opens new possibilities for next-generation design and synthesis of molecular systems for studies in the broad field of artificial photosynthesis. The availability of the *trans*-A₂BC

porphyrin building blocks in conjunction with successive Pd-

mediated coupling reactions provides access to star-shaped architectures that are compositionally richer than those we have prepared previously for light-harvesting and charge separation.⁶⁴ The inclusion of a panchromatic supermolecule constitutes a molecular entity that differs considerably from chromophores found in natural systems.

The absorption spectrum of the pentad array shows panchromatic absorption from below 400 nm to above 700 nm due to the central bis(perylene-monoimide)porphyrin unit, complemented by absorption features due to the perylenediimide and chlorin. The latter two constituents can efficiently transfer energy to the central unit and also serve as electron acceptor and hole trap, respectively, for charge separation that is initiated in the core triad. Two chlorin-porphyrinperylene-diimide triads that are capable of charge separation but lack the panchromatic-absorbing core were also prepared as functional benchmarks. The results described herein provide the foundation for in-depth photophysical investigations of all the arrays.

Experimental section

General methods

Paper

All chemicals obtained commercially were used as received unless otherwise noted. Reagent-grade solvents (CH₂Cl₂, CHCl₃, methanol, toluene, ethyl acetate) and HPLC-grade solvents (toluene, CH2Cl2, hexanes) were used as received. ACS-grade hexanes was distilled (with a rotary evaporator) and dried (over 4 Å molecular sieves) immediately before use. THF was freshly distilled from sodium/benzophenone ketyl and used immediately. Matrix-assisted laser-desorption ionization mass spectrometry (MALDI-MS) was performed with the matrix 1,4bis(5-phenyl-2-oxaxol-2-yl)benzene (POPOP)65 unless noted otherwise. ESI-MS data are reported for the molecular ion or cationized molecular ion. Noncommercial compounds 1,57 3,56 5,60 Zn560 and 834 were prepared following literature procedures, whereas the known dipyrromethane 1-TMS58 was synthesized following a more recent method.⁵⁷ All other compounds were used as received from commercial sources.

Chromatography

A sequence of three-chromatography procedures³¹ was employed for purification of arrays following Pd-mediated coupling reactions. First, adsorption column chromatography (flash silica, Silicycle) was applied to remove catalysts and other materials. Second, preparative-scale SEC was applied. A glass column (o.d. 4.3 cm) was packed using BioRad Bio-Beads S-X1 (MWCO = 12 000 Da) in HPLC-grade toluene with a length of 53 cm. The chromatography was performed with gravity flow (~1 drop per s). Eluate was collected in 3 mL portions and analyzed with MALDI-MS and absorption spectrometry. The portion with the desired product was combined and concentrated. Third, adsorption column chromatography (flash silica, Silicycle) was performed with HPLC-grade CH₂Cl₂ and n-hexane unless noted otherwise.

Analytical-scale SEC (styrene-divinylbenzene copolymer columns) was performed to characterize the purity of the resulting samples. Analytical SEC was performed with 100 Å, 500 Å, and 1000 Å columns in series eluting with HPLC-grade toluene (flow rate = 1.0 mL min⁻¹, 25 °C). Sample detection was achieved by absorption spectroscopy using a diode-array detector ($\lambda_{\text{det}} = 430, 476, 537, 638 \text{ nm}$).

General procedure for Pd-catalyzed coupling reactions

Toluene (or THF) and TEA were deaerated by bubbling with argon for 1 h. The solid starting materials, including Pd₂(dba)₃ and P(o-tol)3, were charged in a Schlenk flask. The flask was evacuated under high vacuum for 5 min, then purged with argon. The evacuation-purge procedure was repeated three times. The deaerated solvents were then added to the Schlenk flask via a syringe. The resulting mixture was again deaerated by three freeze-pump-thaw cycles. Then liquid reagents, if any, were added to the flask. The mixture was then heated to initiate reaction. The copper-free method has been described in detail.54,55

5-Phenyl-15-[4-(2-trimethylsilylethynyl)phenyl] porphyrinatozinc(II) (Zn2-TMS)

Following a reported method,56 a solution of diacyldipyrromethane 1 (42 mg, 0.15 mmol) in THF (0.50 mL) was treated with propylamine (0.18 g, 3.0 mmol) and stirred at room temperature for 1 h. The solution was concentrated to dryness. The resulting solid was dissolved in ethanol (15 mL) and treated with dipyrromethane 1-TMS (48 mg, 0.15 mmol) and Zn(OAc)₂ (0.28 g, 1.5 mmol). The mixture was then refluxed open to the air for 18 h. Upon cooling to room temperature, the mixture was concentrated and chromatographed [silica gel, hexanes/CH2Cl2 (2:1)] to afford a purple solid (37 mg, 40%): ¹H NMR (THF- d_8 , 400 MHz) δ 10.28 (s, 2H), 9.40 (d, J = 4.4 Hz, 4H), 9.02 (d, J =4.4 Hz, 4H), 8.21-8.23 (m, 4H), 7.78-7.88 (m, 5H), 0.38 (s, 9H); 13 C NMR (THF- d_8 , 100 MHz) δ 150.9, 150.6, 150.5, 144.9, 144.3, 135.6, 132.7, 132.3, 130.8, 128.1, 127.2, 123.2, 120.5, 119.2, 106.63, 106.56, 106.4, 95.2, 0.127; MALDI-MS obsd 620.7 [M⁺], ESI-MS obsd 621.1428 $[(M + H)^{+}]$, calcd 620.1375 $(M = C_{37}H_{28}-C_{37}H_{2$ N_4SiZn); λ_{abs} (CH₂Cl₂) 409, 536, 572 nm.

5-Phenyl-15-[4-(2-trimethylsilylethynyl)phenyl]porphyrin (2-TMS)

Following a reported method43 with slight modifications, a solution of zinc porphyrin Zn2-TMS (33 mg, 0.053 mmol) in CHCl₃ (50 mL) was treated dropwise with trifluoroacetic acid (0.20 mL, 1.2 mmol). The solution was stirred at room temperature for 30 min. Then the reaction was quenched by the addition of saturated aqueous NaHCO3 solution. The mixture was extracted with CH2Cl2. The organic extract was washed with brine, dried with Na₂SO₄, concentrated and chromatographed [silica gel, hexanes/CH₂Cl₂ (2:1)] to afford a purple solid (17 mg, 58%): 1 H NMR (THF- d_{8} , 300 MHz) δ 10.38 (s, 2H), 9.44 (m, 4H), 9.04 (m, 4H), 8.22-8.26 (m, 4H), 7.82-7.94 (m, 5H), 0.38 (s, 9H), -3.12 (br, 2H); 13 C NMR (THF- d_8 , 100 MHz) δ 150.8, 150.6, 150.5, 148.1, 147.8, 146.3, 142.9, 142.4, 135.7, 132.8,

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132.7, 132.6, 132.5, 131.6, 131.3, 131.2, 128.6, 127.8, 123.7, 120.0, 118.7, 106.2, 106.1, 95.6, 0.127; MALDI-MS obsd 558.4 [M⁺], ESI-MS obsd 559.2326 [(M + H)⁺], calcd 558.2240 (M = $C_{37}H_{30}N_4Si$); λ_{abs} (CH₂Cl₂) 406, 503, 537, 575, 630 nm.

10,20-Dibromo-5-phenyl-15-[4-(2-trimethylsilylethynyl)phenyl] porphyrin (2-Br₂/TMS)

Following a standard bromination method,59 a solution of porphyrin 2-TMS (12 mg, 0.021 mmol) in CHCl₃ (6.8 mL) containing pyridine (9.0 µL) was cooled to 0 °C and treated with NBS (9.2 mg, 0.052 mmol). The mixture was stirred at 0 °C for 1 h. The reaction was quenched by the addition of saturated aqueous NaHCO3 solution. The mixture was extracted with CH₂Cl₂. The organic extract was washed with brine, dried with Na₂SO₄, concentrated, and chromatographed [silica gel, hexanes/CH₂Cl₂ (4:1)] to afford a purple-red solid (11 mg, 74%): ¹H NMR (THF- d_8 , 300 MHz) δ 9.62 (d, J = 4.5 Hz, 4H), 8.82 (d, J = 4.5 Hz, 4H), 8.12-8.16 (m, 4H), 7.78-7.89 (m, 5H), 0.38 (s,)9H), -2.74 (s, 2H); 13 C NMR (THF- d_8 , 100 MHz) δ 142.6, 142.3, 135.4, 131.2, 129.0, 127.8, 124.2, 122.7, 121.5, 105.9, 104.2, 95.9, 0.089; MALDI-MS obsd 717.1 $[(M + H)^{+}]$, ESI-MS 715.0520 $[(M + H)^{+}]$ H)⁺], calcd 714.0405 (M = $C_{37}H_{28}Br_2N_4Si$); λ_{abs} (CH₂Cl₂) 422, 485, 520, 557, 661 nm.

5-(4-Iodophenyl)-15-[4-(2-trimethylsilylethynyl)phenyl] porphyrinatozinc(II) (Zn4-I/TMS)

Following a reported method,⁵⁶ a solution of diformyldipyrromethane–tin complex 3 (0.40 g, 0.63 mmol) in propylamine (1.0 mL) was stirred at room temperature for 2 h and then concentrated to dryness. A solution of the resulting solid and Zn(OAc)₂ (1.20 g, 6.3 mmol) in ethanol (63 mL) was refluxed open to the air for 17 h. The cooled mixture was passed through a pad of silica gel (CH₂Cl₂), concentrated, and chromatographed [silica gel, hexanes/ethyl acetate (4 : 1)] to afford a purple-red solid (73 mg, 15%): ¹H NMR (THF- d_8 , 300 MHz) δ 10.29 (s, 2H), 9.42 (d, J = 4.4 Hz, 4H), 9.02 (dd, J = 5.6, 4.4 Hz, 4H), 8.22 (d, J = 8.0 Hz, 2H), 8.16 (d, J = 8.0 Hz, 2H), 8.02 (d, J = 8.0 Hz, 2H), 7.88 (d, J = 8.0 Hz, 2H), 0.38 (s, 9H); MALDI-MS obsd 748.0 [(M + H)⁺], ESI-MS obsd 747.0392 [(M + H)⁺], calcd 747.0414 (M = $C_{37}H_{27}IN_4SiZn$); λ_{abs} (CH₂Cl₂) 411, 538, 572 nm.

5-[4-(2-Triisopropylsilylethynyl)phenyl]-15-[4-(2-trimethylsilylethynyl)phenyl]porphyrinatozinc(II) (Zn4-TMS/TIPS)

Following the general procedure for Pd-catalyzed coupling reactions described above, a mixture of **Zn4-I/TMS** (67 mg, 0.090 mmol), PdCl₂(PPh₃)₂ (3.2 mg, 4.5 µmol), and CuI (1.7 mg, 1.0 µmol) in THF/TEA (9:1, 9.0 mL) was treated with TIPS-acetylene (0.100 mL, 0.45 mmol) under argon and then stirred at 60 °C for 4 h. The reaction was quenched by the addition of water. The mixture was extracted with CH₂Cl₂. The organic extract was washed with brine, dried, concentrated, and chromatographed [silica gel, hexanes/ethyl acetate (9:1)] to afford a red solid (58 mg, 80%): 1 H NMR (THF- d_{8} , 300 MHz) δ 10.29 (s, 2H), 9.42 (d, J = 4.5 Hz, 4H), 9.02 (dd, J = 7.2, 4.8 Hz, 4H), 8.22 (dd, J = 7.8, 1.8 Hz, 4H), 7.89 (dd, J = 9.9, 7.8 Hz, 4H), 1.50 (m,

3H), 1.10 (m, 18H), 0.37 (s, 9H); MALDI-MS obsd 802.4 [(M + H) $^{+}$], ESI-MS obsd 801.2775 [(M + H) $^{+}$], calcd 801.2782 (M = $C_{48}H_{48}N_4Si_2Zn$); λ_{abs} (CH $_2Cl_2$) 411, 538, 576 nm.

5,15-Dibromo-10-[4-(2-triisopropylsilylethynyl)phenyl]-20-[4-(2-trimethylsilylethynyl)phenyl]porphyrin (4-Br₂/TMS/TIPS)

A solution of Zn4-TMS/TIPS (54 mg, 67 μmol) in CH₂Cl₂ (22 mL) containing pyridine (0.030 mL, 0.33 mmol) was treated with NBS (0.030 g, 0.17 mmol) at 0 °C for 1 h. The solution was allowed to warm to room temperature and then treated with saturated aqueous NH₄Cl solution. The mixture was extracted with CH₂Cl₂. The organic extract was washed with brine, dried, and concentrated to dryness. A solution of the resulting solid in CH₂Cl₂ (40 mL) was treated dropwise with trifluoroacetic acid (0.26 mL, 3.4 mmol) and stirred at room temperature for 30 min. The reaction was quenched by the addition of saturated aqueous NaHCO3 solution. The mixture was extracted with CH₂Cl₂. The organic extract was washed with brine, dried, concentrated and chromatographed [silica gel, hexanes/CH2Cl2 (3:1)] to afford a red solid (16 mg, 27%): ¹H NMR (THF- d_8 , 300 MHz) δ 9.64 (d, I = 5.1 Hz, 4H), 8.84–8.87 (m, 4H), 8.14 (d, I =7.8 Hz, 4H), 7.90 (m, 4H), 1.49 (m, 3H), 1.29, (s, 18H), 0.37 (s, 9H), -2.74 (s, 2H); MALDI-MS obsd 896.6 [(M + H)⁺], ESI-MS obsd 895.1836 $[(M + H)^{+}]$, calcd 895.1857 $(M = C_{48}H_{48}Br_{2}N_{4})$ Si₂); λ_{abs} (CH₂Cl₂) 429, 524, 562, 605, 659 nm.

5-(4-Ethynylphenyl)-10,20-bis(2-phenylethynyl)-15-[4-(2-triisopropylsilylethynyl)phenyl]porphyrinatozinc(II) (ZnP-H/TIPS)

A solution of Zn4-TMS/TIPS (118 mg, 0.146 mmol, 1.0 equiv.) in CH₂Cl₂ (49 mL, [Zn4-TMS/TIPS] ~3 mM) containing pyridine (65 μL, 0.81 mmol, 5.5 equiv.) was cooled to 0 °C under argon, and treated with NBS (65 mg, 0.37 mmol, 2.5 equiv.) at 0 °C for 1 h. The reaction was then quenched by the addition of saturated aqueous NaHCO3 solution. The mixture was extracted with CH2Cl2. The organic extract was dried, concentrated, and chromatographed (silica gel, hexanes/ $CH_2Cl_2 = 1:1$) to afford a blue solid (140 mg): TLC (silica, hexanes/ $CH_2Cl_2 = 1:1$) $R_f =$ 0.28; MALDI-MS obsd 957.0 $[(M + H)^{+}]$. Following the general procedure for Pd-catalyzed coupling reactions described above, a mixture of the resulting solid (140 mg, 0.146 mmol, 1.0 equiv.), phenylacetylene (35 μL, 0.32 mmol, 2.2 equiv.), Pd₂(dba)₃ (40 mg, 0.044 mmol, 0.30 equiv.), P(o-tol)₃ (107 mg, 0.350 mmol, 2.4 equiv.), and CuI (1.4 mg, 7.3 μmol, 0.050 equiv.) in deaerated THF/TEA (5:1, 14.6 mL) was stirred at 60 °C for 17 h. The mixture was allowed to cool to room temperature, dried, and chromatographed (silica gel, hexanes/ $CH_2Cl_2 = 2:1$ to 3:2) to afford a blue solid (72.6 mg): TLC (silica, hexanes/ $CH_2Cl_2 = 3:2$) $R_f = 0.30$; MALDI-MS obsd 1001.9 [(M + H)⁺]. Then, a mixture of the resulting solid (72.6 mg) and K₂CO₃ (1.00 g, 7.24 mmol, 100 equiv.) in THF/CH₃OH (1:1, 15 mL) was stirred at room temperature for 3 h. MALDI-MS analysis showed the completion of the reaction [obsd 929.8, $[(M + H)^{+}]$]. The reaction was then quenched by the addition of water. The mixture was extracted with CH2Cl2. The organic extract was dried, concentrated, and chromatographed (silica gel, hexanes/

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CH₂Cl₂ = 3 : 2 to 1 : 1) to afford a blue solid (60.2 mg, 44% for three steps): ¹H NMR (THF- d_8 , 400 MHz) δ 9.76 (d, J = 4.3 Hz, 4H), 8.86 (d, J = 4.7 Hz, 4H), 8.20–7.47 (m, 18H), 3.28 (s, 1H), 1.30–1.22 (m, 21H); MALDI-MS obsd 929.8 [(M + H)⁺], ESI-MS obsd 929.3038 [(M + H)⁺], calcd 929.3012 (M = C₆₁H₄₈N₄SiZn).

10-Mesityl-18,18-dimethyl-5-(4-(2-phenylethynyl)phenyl) chlorin (C-Ph)

Following the general procedure for Pd-catalyzed coupling reactions described above, a solution of **Zn5** (5.0 mg, 6.9 μmol) in THF/TEA (9:1, 0.70 mL) was treated with phenylacetylene (3.8 μ L, 0.034 mmol) in the presence of PdCl₂(PPh₃)₂ (0.70 mg, 1.0 μ mol) and CuI (0.20 mg, 1.0 μ mol) and stirred at 60 °C for 24 h. The reaction was quenched by the addition of water. The mixture was extracted with CH2Cl2. The organic extract was washed with brine, dried, and concentrated. A solution of the resulting sample in CH₂Cl₂ (4.1 mL) was treated dropwise with trifluoroacetic acid (0.027 mL) and stirred at room temperature for 30 min. The reaction was quenched by the addition of saturated aqueous NaHCO3 solution. The mixture was extracted with CH₂Cl₂. The organic extract was washed with brine, dried, concentrated, and chromatographed [silica gel, hexanes/CH₂Cl₂ (3:1)] to afford a green solid (3.7 mg, 84%): ¹H NMR (CDCl₃, 300 MHz) δ 8.93 (s, 1H), 8.86 (s, 1H), 8.83 (d, J = 4.5 Hz, 1H), 8.74-8.76 (m, 2H), 8.60 (d, J = 4.5 Hz, 1H), 8.43 (d, J = 4.5 Hz, 1H), 8.46 (d, J = 4.5 Hz, 1H), 8.08 (d, J = 8.1 Hz, 2H), 7.69 (d, J =8.1 Hz, 2H), 7.55-7.58 (m, 2H), 7.32-7.34 (m, 3H), 7.23 (s, 2H), 4.60 (s, 2H), 2.59 (s, 3H), 2.06 (s, 6H), 1.84 (s, 6H), -1.83 (s, 2H); MALDI-MS obsd 634.8 $[M^{+}]$, ESI-MS obsd 635.3169 $[(M + H)^{+}]$, calcd 634.3096 (M = $C_{45}H_{38}N_4$); λ_{abs} (toluene) 414, 508, 589, 641 nm.

N-(4-Iodophenyl)-*N*'-(2,6-diisopropylphenyl)perylene-3,4,9,10-bis(dicarboximide) (7)

Following a reported procedure,³⁷ a mixture of 6 (3.00 g, 7.64 mmol), 2,6-diisopropylaniline (4.3 mL, 23 mmol), 4-iodoaniline (3.35 g, 15 mmol), Zn(OAc)₂ (1.05 g), and imidazole (22.5 g) was heated to 190 °C under argon. After 20 h, the mixture was allowed to cool to room temperature and then was treated with 60% ethanol (180 mL) for another 2 h. The suspension was then filtered. The filter cake was washed with 60% ethanol. The dry sample was suspended in 10% aqueous KOH solution and refluxed for 30 min. Upon cooling to room temperature, the mixture was filtered. The filter cake was washed with water and dried in an oven. The resulting solid was soaked in CH2Cl2 (100 mL), sonicated for 5 min, and filtered to remove insoluble perylene-diimide byproducts. The CH₂Cl₂ filtrate was concentrated and chromatographed [silica gel, hexanes/ethyl acetate (7:1 to 7:3)] to afford a red solid (1.52 g, 30%): ¹H NMR $(CDCl_3, 300 \text{ MHz}) \delta 8.66-8.80 \text{ (m, 8H)}, 7.91 \text{ (dd, } J = 6.6, 1.8 \text{ Hz},$ 2H), 7.51 (t, J = 7.5 Hz, 1H), 7.36 (d, J = 7.5 Hz, 2H), 7.12 (dd, J6.6, 1.8 Hz, 2H), 2.76 (quintet, J = 6.6 Hz, 2H), 1.18 (d, J = 6.6 Hz, 12H); MALDI-MS obsd 754.0 $[(M + H)^{+}]$, ESI-MS obsd 753.1235 $[(M + H)^{+}]$, calcd 752.1172 $(M = C_{49}H_{29}IN_{2}O_{4})$; $\lambda_{abs}(CH_{2}Cl_{2})$ 434, 459, 490, 527 nm.

N-[4-(2-Phenylethynyl)phenyl]-*N*-(2,6-diisopropylphenyl) perylene-3,4,9,10-bis(dicarboximide) (PDI-Ph)

Following the general procedure for Pd-catalyzed coupling reactions described above, a solution of 7 (3.0 mg, 4.0 µmol) in toluene/TEA (5:1, 1.8 mL) was treated with phenylacetylene (2.2 µL, 0.020 mmol) in the presence of PdCl₂(PPh₃)₂ (0.50 mg, 0.60 µmol) and CuI (0.10 mg, 0.60 µmol) and stirred at 60 °C for 3 h. Then the reaction was quenched by the addition of water. The mixture was extracted with CH₂Cl₂. The organic extract was washed with brine, dried, concentrated, and chromatographed [silica gel, hexanes/ethyl acetate (7:3)] to afford a red solid (1.5 mg, 52%): ¹H NMR (CDCl₃, 300 MHz) δ 8.69–8.81 (m, 8H), 7.73 (d, J = 8.7 Hz, 2H), 7.49–7.59 (m, 3H), 7.35–7.40 (m, 7H), 2.76 (pentad, J = 6.9 Hz, 2H), 1.19 (d, J = 6.9 Hz, 12H); MALDI-MS obsd 726.1 [M⁺], ESI-MS 727.2553 [(M + H⁺)], calcd 726.2519 (M = C₅₀H₃₄N₂O₄); λ _{abs} (toluene) 459, 491, 527 nm.

5,15-Bis[2-(3,4-(N-(2,6-diisopropylphenyl)iminodicarbonyl)-1,6-bis(4-tert-butylphenoxy)perylen-9-yl)ethynyl]-10-phenyl-20-(4-(2-trimethylsilylethynyl)phenyl)porphyrin (T-Ph)

Following the general procedure for Pd-catalyzed coupling reactions described above, a mixture of porphyrin 2-Br₂/TMS (5.4 mg, 7.5 μmol), 8 (12 mg, 0.015 mmol), Pd₂(dba)₃ (2.1 mg, 2.2 μ mol), and P(o-tol)₃ (5.5 mg, 18 μ mol) in toluene/TEA (5 : 1, 3.4 mL) was stirred at 60 °C for 4 h. The mixture was allowed to cool to room temperature and then was purified according to the three-chromatography method. The product was concentrated to afford a black solid (14 mg, 86%): ¹H NMR (CDCl₃, 300 MHz): δ 9.73 (d, J = 5.1 Hz, 4H); 9.27 (d, J = 7.2 Hz, 2H), 9.21 (d, J= 7.8 Hz, 2H, 9.02 (d, J = 8.1 Hz, 2H), 8.86 (d, J = 5.1 Hz, 2H),8.80 (d, J = 5.1 Hz, 2H), 8.33 (s, 2H), 8.28 (s, 2H), 8.20–8.10 (m, 4H), 7.90 (d, J = 7.8 Hz, 2H), 7.79 (m, 3H), 7.67 (t, J = 7.8 Hz, 2H), 7.39–7.49 (m, 12H), 7.32 (d, J = 7.8 Hz, 4H), 7.08 (d, J = 9.0 Hz, 4H), 7.05 (d, J = 9.0 Hz, 4H), 2.77 (quintet, J = 6.6 Hz, 4H), 1.34 (s, 36H), 1.18 (d, J = 6.6 Hz, 24H), 0.089 (s, 9H), -1.90 (s, 2H); MALDI-MS obsd 2158.0 $[(M + H)^{+}]$, calcd 2156.9563 (M = $C_{149}H_{128}N_6O_8Si$); λ_{abs} (toluene) 429, 476, 537, 638, 726 nm.

5,15-Bis[2-(3,4-(N-(2,6-diisopropylphenyl)iminodicarbonyl)-1,6-bis(4-tert-butylphenoxy)perylen-9-yl)ethynyl]-10-phenyl-20-(4-(2-trimethylsilylethynyl)phenyl)porphyrinatozinc(π) (ZnT-Ph)

A solution of **T-Ph** (22 mg, 10 μmol) was treated with $Zn(OAc)_2 \cdot 2H_2O$ (44 mg, 0.20 mmol) in CH_2Cl_2/CH_3OH (2:1, 3.0 mL) for 18 h. The resulting mixture was washed with water. The organic layer was dried and purified according to the three-chromatography method to afford a black solid (10 mg, 45%): 1H NMR (CDCl₃, 300 MHz): δ 9.77 (m, 4H); 9.30–9.00 (m, 6H), 8.90–8.82 (m, 4H), 8.35 (s, 2H), 8.27 (s, 2H), 8.22–8.10 (m, 4H), 7.90–7.65 (m, 7H), 7.39–7.49 (m, 12H), 7.34 (d, J = 8.1 Hz, 4H), 7.08 (d, J = 8.7 Hz, 4H), 7.07 (d, J = 8.6 Hz, 4H), 2.77 (quintet, J = 6.5 Hz, 4H), 1.34 (s, 36H), 1.18 (d, J = 6.6 Hz, 24H), 0.090 (s, 9H); MALDI-MS obsd 2220.0 [(M + H)⁺], calcd 2219.8771 (M = $C_{149}H_{126}N_6O_8SiZn$).

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5,15-Bis[2-(3,4-(N-(2,6-diisopropylphenyl)iminodicarbonyl)-1,6-bis(4-tert-butylphenoxy)perylen-9-yl)ethynyl]-10-(4-ethynylphenyl)-20-phenylporphyrin (T-Ph-H)

A solution of **T-Ph** (7.0 mg, 3.2 μmol) in toluene (3.0 mL) and methanol (3.0 mL) was treated with $\rm K_2CO_3$ (44 mg, 0.32 mmol) and stirred at room temperature for 3 h. The reaction was quenched by the addition of water. The mixture was extracted with $\rm CH_2Cl_2$. The organic extract was washed with brine, dried, and concentrated to afford a black solid (6.7 mg, quant.): $^1\rm H$ NMR (CDCl₃, 300 MHz): $^5\rm 0.73$ (d, J=5.1 Hz, 4H); 9.27 (d, J=7.2 Hz, 2H), 9.21 (d, J=7.8 Hz, 2H), 9.02 (d, J=8.1 Hz, 2H), 8.84 (d, J=5.1 Hz, 2H), 8.80 (d, J=5.1 Hz, 2H), 8.34 (s, 2H), 8.28 (s, 2H), 8.10–8.20 (m, 4H), 7.91 (d, J=7.8 Hz, 2H), 7.79 (m, 3H), 7.67 (t, J=7.8 Hz, 2H), 7.39–7.49 (m, 12H), 7.32–7.00 (m, 12H), 3.36 (s, 1H), 2.77 (quintet, J=6.6 Hz, 4H), 1.34 (s, 36H), 1.18 (d, J=6.6 Hz, 24H), -1.88 (s, 2H); MALDI-MS obsd 2085.7 [(M+H) $^+$], calcd 2084.9168 (M = $\rm C_{146}\rm H_{120}\rm N_6\rm O_8$); $\lambda_{\rm abs}$ (toluene) 429, 476, 537, 638, 726 nm.

5,15-Bis[2-(3,4-(*N*-(2,6-diisopropylphenyl)iminodicarbonyl)-1,6-bis(4-*tert*-butylphenoxy)perylen-9-yl)ethynyl]-10-[4-(2-(4-(9,10-(*N*-(2,6-diisopropylphenyl)iminodicarbonyl)perylene-3,4-dicarboximido)phenyl)ethynyl)phenyl]-20-phenylporphyrin (T-PDI)

Following the general procedure for Pd-catalyzed coupling reactions described above, a mixture of the triad T-Ph-H (34 mg, 16 μmol), iodophenyl-perylene-diimide 7 (12 mg, 16 μmol), Pd₂(dba)₃ (2.2 mg, 2.4 μmol), and P(o-tol)₃ (5.4 mg, 18 μmol) in toluene/TEA (5:1, 4.0 mL) was stirred at 60 °C for 4 h. The mixture was allowed to cool to room temperature and then was chromatographed according to the three-chromatography method to afford a black solid (17 mg, 40%): ¹H NMR (CDCl₃, 400 MHz) δ 9.78 (d, J = 5.1 Hz, 4H); 9.30 (d, J = 7.2 Hz, 2H), 9.20 (d, J = 7.8 Hz, 2H), 9.01 (d, J = 8.1 Hz, 2H), 8.88 (d, J = 5.1 Hz, 2H)2H), 8.80-8.60 (m, 10H), 8.34 (s, 2H), 8.28 (s, 2H), 8.10-8.20 (m, 4H), 7.94–7.90 (m, 4H), 7.79 (m, 3H), 7.67 (t, J = 7.8 Hz, 2H), 7.30-7.52 (m, 19H), 7.15-7.05 (m, 10H), 2.80-2.77 (m, 6H), 1.34 (s, 36H), 1.20-1.18 (m, 36H), -1.95 (s, 2H); MALDI-MS obsd 2711.0 [(M + H)⁺], calcd 2709.1217 (M = $C_{188}H_{148}N_8O_{12}$); λ_{abs} (toluene) 430, 463, 492, 530, 639, 727 nm.

5,15-Bis[2-(3,4-(*N*-(2,6-diisopropylphenyl)iminodicarbonyl)-1,6-bis(4-*tert*-butylphenoxy)perylen-9-yl)ethynyl]-10-[4-(2-(4-(9,10-(*N*-(2,6-diisopropylphenyl)iminodicarbonyl)perylene-3,4-dicarboximido)phenyl)ethynyl)phenyl]-20-phenylporphyrinatozinc(II) (ZnT-PDI)

A solution of **T-PDI** (0.020 g, 7.4 μmol, 1.0 equiv.) in CH₂Cl₂/CH₃OH (2:1, 2.2 mL) was treated with Zn(OAc)₂·2H₂O (32 mg, 0.15 mmol, 20 equiv.) at room temperature for 22 h, then washed with water. The organic layer was dried, concentrated, and chromatographed by SEC (HPLC-grade toluene) followed by silica gel (CH₂Cl₂/CH₃OH = 1:1 to 9:1) to afford a black solid (0.020 g, quant.): ¹H NMR (CDCl₃, 400 MHz) δ 9.78 (d, J = 5.1 Hz, 4H); 9.30 (d, J = 7.2 Hz, 2H), 9.20 (d, J = 7.8 Hz, 2H), 9.01 (d, J = 8.1 Hz, 2H), 8.88 (d, J = 5.1 Hz, 2H), 8.80-8.60 (m, 10H), 8.34 (s, 2H), 8.28 (s,

2H), 8.10–8.20 (m, 4H), 7.94–7.90 (m, 4H), 7.79 (m, 3H), 7.67 (t, J=7.8 Hz, 2H), 7.30–7.52 (m, 19H), 7.15–7.05 (m, 10H), 2.80–2.77 (m, 6H), 1.34 (s, 36H), 1.20–1.18 (m, 36H); MALDI-MS obsd 2772.9 [(M + H)⁺], calcd 2771.0352 (M = $C_{188}H_{146}N_8O_{12}Zn$).

5,15-Bis[2-(3,4-(*N*-(2,6-diisopropylphenyl)iminodicarbonyl)-1,6-bis(4-*tert*-butylphenoxy)perylen-9-yl)ethynyl]-10-[4-(2-(4-(10-mesityl-18,18-dimethylchlorin-5-yl)phenyl)ethynyl) phenyl]-20-phenylporphyrin (C-T)

Following the general procedure for Pd-catalyzed coupling reactions described above, a mixture of the triad T-Ph-H (33 mg, 16 μmol), iodophenyl-chlorin 5 (11 mg, 16 μmol), Pd₂(dba)₃ (2.2 mg, 2.4 μmol), and P(o-tol)₃ (5.4 mg, 18 μmol) in toluene/ TEA (5:1, 4.0 mL) was stirred at 60 °C for 3 h. The mixture was allowed to cool to room temperature and then was chromatographed according to the three-chromatography method to afford a black solid (10 mg, 24%): ¹H NMR (CDCl₃, 400 MHz) δ 9.79–9.77 (m, 2H), 9.53–9.44 (m, 4H), 9.10 (d, J = 7.6 Hz, 2H), 8.98 (s, 1H), 8.92–8.85 (m, 4H), 8.78 (d, J = 4.0 Hz, 1H), 8.63 (d, J = 4.0 Hz, 1 = 4.8 Hz, 1H, -1.89 (s, 2H), 8.54 (d, J = 4.4 Hz, 1H), 8.50 (m,1H), 8.42 (d, J = 4.4 Hz, 1H), 8.36 (s, 1H), 8.35 (s, 1H), 8.32 (d, J =8.0 Hz, 2H, 8.19 - 8.21 (m, 3H), 8.08 (d, J = 8.4 Hz, 1H), 8.01 (d, J= 8.4 Hz, 2H, 7.87-7.77 (m, 5H), 7.52-7.42 (m, 11H), 7.31 (d, J = 0.0000 m)8.0 Hz, 6H), 7.13-7.10 (m, 8H), 4.65 (s, 2H), 2.74 (quintet, J =6.8 Hz, 6H), 2.62 (s, 3H), 2.10 (s, 6H), 1.89 (s, 6H), 1.34 (s, 36H), 1.16 (d, J = 6.8 Hz, 36H), -1.81 (s, 2H), -1.90 (s, 2H); MALDI-MSobsd 2618.6 $[(M + H)^{+}]$, calcd 2617.1795 $(M = C_{183}H_{152}N_{10}O_{8})$; λ_{abs} (toluene) 416, 477, 511, 537, 641, 728 nm.

5,15-Bis(2-phenylethynyl)-10-[4-(2-trimethylsilylethynyl) phenyl]-20-[4-(2-triisopropylsilylethynyl)phenyl]porphyrin (P-TMS/TIPS)

Following the general procedure for Pd-catalyzed coupling reactions described above, a mixture of porphyrin **4-Br**₂/**TMS**/**TIPS** (47 mg, 52 µmol, 1.0 equiv.), phenylacetylene (13 mg, 0.13 mmol, 2.5 equiv.), Pd₂(dba)₃·CHCl₃ (8.1 mg, 7.9 µmol, 0.15 equiv.), and P(o-tol)₃ (19 mg, 62 µmol, 1.2 equiv.) in toluene/TEA (5:1, 10.2 mL) was stirred at 60 °C for 2 h. The mixture was then washed with water. The organic layer was dried, concentrated, and chromatographed (silica gel, hexanes/CH₂Cl₂, 2:1) to afford a blue solid (38 mg, 78%): TLC (silica, hexanes/CH₂Cl₂, 3:2) R_f = 0.54; ¹H NMR (CDCl₃, 300 MHz) δ 9.65 (d, J = 4.7 Hz, 4H), 8.78 (dd, J = 9.3, 4.8 Hz, 4H), 8.12 (d, J = 8.0 Hz, 4H), 8.02 (d, J = 6.6 Hz, 4H), 8.00–7.88 (m, 4H), 7.60–7.50 (m, 6H), 1.27–1.25 (m, 21H), 0.40 (s, 9H), -2.09 (s, 2H); ESI-MS obsd 939.4105 [(M + H)⁺], calcd 938.4200 (M = C₆₄H₅₈N₄Si₂).

5-[4-(2-(4-(9,10-(*N*-(2,6-Diisopropylphenyl)iminodicarbonyl) perylene-3,4-dicarboximido)phenyl)ethynyl)phenyl]-10,20-bis(2-phenylethynyl)-15-[4-(2-triisopropylsilylethynyl)phenyl] porphyrin (9)

A mixture of **P-TMS/TIPS** (10 mg, 0.011 mmol, 1.0 equiv.) and K_2CO_3 (0.15 g, 1.1 mmol, 100 equiv.) in THF/CH₃OH (1:1, 10 mL) was stirred at room temperature for 30 min. The reaction was then quenched by the addition of 2 M HCl solution (10 mL). The mixture was extracted with CH_2Cl_2 . The organic extract was

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dried and concentrated to dryness. Following the general procedure for Pd-catalyzed coupling reactions described above, a mixture of the resulting sample, iodophenyl-perylene-diimide 7 (8.3 mg, 0.011 mmol, 1.0 equiv.), $Pd_2(dba)_3 \cdot CHCl_3$ (1.7 mg, 1.6 µmol, 0.15 equiv.), and $P(o\text{-tol})_3$ (4.0 mg, 0.013 mmol, 1.2 equiv.) in toluene/TEA (5 : 1, 2.4 mL) was stirred at 60 °C for 2 h. The mixture was then allowed to cool to room temperature and passed through a pad of silica gel (CH_2Cl_2). The eluate was concentrated and chromatographed (silica gel, CH_2Cl_2) to afford a green solid (7.9 mg, 49%): TLC (silica, CH_2Cl_2) to afford a green solid (7.9 mg, 49%): TLC (silica, CH_2Cl_2) $R_f = 0.22$; Analytical SEC (toluene, 1.0 mL min⁻¹) $t_R = 21.12$ min; MALDI-MS obsd 1491 (M^+), calcd 1490.5854 ($M = C_{103}H_{78}N_6O_4Si$); λ_{abs} (toluene) 445, 491, 528, 600, 692 nm. The compound was not characterized by NMR spectroscopy due to limited solubility in organic solvents.

5-[4-(2-(4-(9,10-(*N*-(2,6-Diisopropylphenyl)iminodicarbonyl) perylene-3,4-dicarboximido)phenyl)ethynyl)phenyl]-15-[4-(2-(4-(10-mesityl-18,18-dimethylchlorin-5-yl)phenyl)phenyl]-10,20-bis(2-phenylethynyl)porphyrin (C-P-PDI)

A solution of 9 (6.5 mg, 0.0044 mmol, 1.0 equiv.) in THF (2.2 mL) was treated with 1 M TBAF solution in THF (8.7 µL, 0.0087 mmol, 2.0 equiv.) at room temperature for 2 h, then treated with water. The mixture was extracted with CH₂Cl₂. The organic extract was dried and concentrated to dryness. Following the general procedure for Pd-catalyzed coupling reactions described above, a mixture of the resulting sample, iodophenyl-chlorin 5 (2.9 mg, 0.044 mmol, 1.0 equiv.), Pd₂(dba)₃·CHCl₃ (1.4 mg, 0.00132 mmol, 0.30 equiv.), and P(o-tol)₃ (1.6 mg, 0.0053 mmol, 1.2 equiv.) in toluene/TEA (5:1, 1.2 mL) was stirred at 60 °C for 2 h. The mixture was then allowed to cool to room temperature. Purification according to the threechromatography method afforded a green solid (1.4 mg, 17%): TLC (silica, $CH_2Cl_2/CH_3OH = 99:1$) $R_f = 0.52$; ¹H NMR (THF d_8 , 400 MHz) δ 9.68 (dd, J = 11.8, 4.3 Hz, 4H), 8.87–8.86 (m, 4H), 8.68-8.64 (m, 2H), 8.58 (s, 2H), 8.54 (d, J = 4.5 Hz, 1H), 8.40-8.32(m, 4H), 8.29-8.27 (m, 5H), 8.21-7.98 (m, 15H), 7.71-7.51 (m, 12H), 7.24 (s, 2H), 4.56 (s, 2H), 3.10-3.02 (m, 2H), 2.05 (s, 9H), 1.89 (s, 6H), 1.30-1.29 (m, 16H), -1.82 (s, 2H), -1.88 (s, 2H); MALDI-MS obsd 1867.6 $[(M + H)^{+}]$, calcd 1866.7143 (M = $C_{131}H_{90}N_{10}O_4$; λ_{abs} (toluene) 418, 446, 528, 600, 642, 692 nm.

5-[4-(2-(4-(9,10-(*N*-(2,6-Diisopropylphenyl)iminodicarbonyl) perylene-3,4-dicarboximido)phenyl)ethynyl)phenyl]-15-[4-(2-(4-(10-mesityl-18,18-dimethylchlorinatozinc(п)-5-yl)phenyl) ethynyl)phenyl]-10,20-bis(2-phenylethynyl) porphyrinatozinc(п) (ZnC-ZnP-PDI)

A solution of C-P-PDI (1.0 mg, 0.54 μmol, 1.0 equiv.) in THF (1.0 mL) was treated with $Zn(OAc)_2$ (2.3 mg, 0.011 mmol, 20 equiv.) at room temperature for 18 h. The sample was treated with water and extracted with CH_2Cl_2 . The organic extract was dried and concentrated to afford a green solid (1.0 mg, quant.): TLC (silica, $CH_2Cl_2/CH_3OH = 99:1$) $R_f = 0.44$; ¹H NMR (THF- d_8 , 400 MHz) δ 9.65 (dd, J = 11.8, 4.3 Hz, 4H), 8.87–8.86 (m, 4H), 8.68–8.64 (m, 2H), 8.58 (s, 2H), 8.54 (d, J = 4.5 Hz, 1H), 8.40–8.36 (m, 4H), 8.29–8.27 (m, 5H), 8.21–7.98 (m, 15H), 7.71–7.41 (m, 12H),

7.24 (s, 2H), 4.53 (s, 2H), 3.06–3.02 (m, 2H), 2.05 (s, 9H), 1.89 (s, 6H), 1.30–1.29 (m, 16H); MALDI-MS obsd 1991.5 [(M + H)⁺], calcd 1990.5 (M = $C_{131}H_{86}N_{10}O_4Zn_2$); λ_{abs} (toluene) 418, 446, 528, 600, 642, 692 nm.

5,15-Bis[2-(3,4-(*N*-(2,6-diisopropylphenyl)iminodicarbonyl)-1,6-bis(4-*tert*-butylphenoxy)perylen-9-yl)ethynyl]-10-[4-(2-trimethylsilylethynyl)phenyl]-20-[4-(2-triisopropylsilylethynyl)phenyl]porphyrin (T-TMS/TIPS)

Following the general procedure for Pd-catalyzed coupling reactions described above, a mixture of dibromoporphyrin 4-Br₂/TMS/TIPS (14 mg, 16 μmol), ethynylperylene-monoimide 8 (26 mg, 32 μmol), Pd₂(dba)₃ (4.4 mg, 4.8 μmol), and P(o-tol)₃ (12 mg, 38 µmol) in toluene/TEA (5:1, 6.4 mL) was stirred at 60 °C for 4 h. The mixture was allowed to cool to room temperature and then chromatographed according to the threechromatography method to afford a black solid (28 mg, 76%): ¹H NMR (CDCl₃, 300 MHz), δ 9.78 (d, J = 4.8 Hz, 4H), 9.48 (dd, J= 7.2, 3.9 Hz, 2H), 9.11 (d, J = 7.8 Hz, 2H), 8.86 (dd, J = 7.8, 2H)4.5 Hz, 4H), 8.37 (s, 2H), 8.36 (s, 2H), 8.34 (d, J = 8.4 Hz, 2H), 8.15 (d, J = 7.8 Hz, 4H), 7.92 (dd, J = 8.4, 3.0 Hz, 4H), 7.86 (d, J =8.1 Hz, 2H), 7.73 (t, J = 5.7 Hz, 2H), 7.40–7.43 (m, 10H), 7.32 (s, 2H), 7.30 (s, 2H), 7.12 (dd, J = 9.0, 2.1 Hz, 8H), 2.75 (hept, J =6.9 Hz, 4H), 2.04 (hept, J = 6.9 Hz, 3H), 1.36 (d, J = 1.8 Hz, 36H), 1.17 (d, J = 6.6 Hz, 24H), 0.99 (d, J = 6.9 Hz, 18H), 0.40 (s, 9H),-1.81 (s, 2H); MALDI-MS obsd 2337.2 [M⁺], calcd 2337.0897 (M = $C_{160}H_{148}N_6O_8Si_2$); λ_{abs} (CH₂Cl₂) 423, 471, 531, 636, 725 nm.

5,15-Bis[2-(3,4-(*N*-(2,6-diisopropylphenyl)iminodicarbonyl)-1,6-bis(4-*tert*-butylphenoxy)perylen-9-yl)ethynyl]-10-[4-(2-(4-(9,10-(*N*-(2,6-diisopropylphenyl)iminodicarbonyl)perylene-3,4-dicarboximido)phenyl)ethynyl)phenyl]-20-[4-(2-triisopropylsilylethynyl)phenyl]porphyrin (T-H/PDI)

A solution of T-TMS/TIPS (28 mg, 12 μ mol) in THF/MeOH (3:1, 12 mL) was treated with K₂CO₃ (11 mg, 80 μmol) at room temperature for 1 h. The reaction was quenched by the addition of water. The mixture was extracted with CH2Cl2. The organic extract was washed with brine, dried, and concentrated to dryness. Following the general procedure for Pd-catalyzed coupling reactions described above, a mixture of the resulting solid, iodophenyl-perylene-diimide 7 (11 mg, 14 μmol), $Pd_2(dba)_3$ (1.6 mg, 1.8 µmol), and $P(o-tol)_3$ (4.3 mg, 14 µmol) in toluene/TEA (5:1, 4.8 mL) was stirred at 60 °C for 4 h. The mixture was washed with water and extracted with CH₂Cl₂. The organic extract was washed with brine, dried, and concentrated to dryness. A solution of the resulting solid in THF (2.0 mL) was treated dropwise with 1.0 M TBAF solution in THF (7.8 µL, 7.8 μmol) and stirred for 1 h at room temperature. The reaction was quenched by the addition of saturated aqueous NaHCO3 solution. The mixture was extracted with ethyl acetate. The organic extract was washed with brine, dried, concentrated, and chromatographed according to the three-chromatography method to afford a black solid (11 mg, 40%): MALDI-MS obsd 2889.7 (M^+) , calcd 2889.2551 $(M = C_{199}H_{168}N_8O_{12}Si)$; λ_{abs} (toluene) 431, 459, 491, 530, 641, 730 nm. The purity of the sample was **RSC Advances**

estimated to be >95% according to analytical SEC. The sample was used directly in the next step without further characterization.

5,15-Bis[2-(3,4-(N-(2,6-diisopropylphenyl)iminodicarbonyl)-1,6-bis(4-tert-butylphenoxy)perylen-9-yl)ethynyl]-10-[4-(2-(4-(9,10-(N-(2,6-diisopropylphenyl)iminodicarbonyl)perylene-3,4-dicarboximido)phenyl)ethynyl)phenyl]-20-[4-(2-(4-(10mesityl-18,18-dimethylchlorin-5-yl)phenyl)ethynyl)phenyl] porphyrin (C-T-PDI)

Following the general procedure for Pd-catalyzed coupling reactions described above, a solution of T-H/PDI (11 mg, 4.0 μmol), iodophenyl-chlorin 5 (4.0 mg, 6.0 μmol), Pd₂(dba)₃ (1.0 mg, 1.1 µmol), and P(o-tol)₃ (1.5 mg, 4.8 µmol) in toluene/ TEA (5:1, 1.6 mL) was stirred at 60 °C for 18 h. The mixture was chromatographed according to the three-chromatography method to afford a black solid (3.5 mg, 27%): ¹H NMR $(CDCl_3, 700 \text{ MHz}) \delta 9.77-9.73 \text{ (m, 4H)}, 9.37-9.11 \text{ (m, 4H)}, 8.93-$ 8.64 (m, 12H), 8.55-8.44 (m, 6H), 8.42-8.32 (m, 4H), 8.25-7.91 (m, 12H), 7.82 (d, J = 6.2 Hz, 4H), 7.52 (d, J = 7.7 Hz, 3H), 7.43-7.49 (m, 16H), 7.35–7.43 (m, 6H), 7.32 (d, J = 8.2 Hz, 4H), -1.91 (br, 1H), 7.04-7.13 (m, 6H), 4.66 (s, 2H), 2.76 (pent, J = 7.1 Hz, 6H), 2.64 (s, 3H), 2.01-2.09 (m, 6H), 1.36-1.38 (m, 36H), 1.18 (d, J = 5.8 Hz, 42H, -1.73 (br, 1H), -1.79 (br, 1H), -1.84 (br, 2H);MALDI-MS obsd 3265.8 [M⁺], calcd 3265.3844 (M $C_{227}H_{180}N_{12}O_{12}$); λ_{abs} (toluene) 418, 494, 530, 641, 742 nm.

Photophysical properties

Static absorption (Shimadzu UV-1800) and emission (Horiba Nanolog) and other measurements were performed on dilute (µM), Ar-purged solutions in toluene at room temperature. Fluorescence quantum yields (Φ_f) utilized samples with $A \leq 0.1$ at the excitation wavelength and were obtained by absolute measurements (Horiba QuantiPhi). Values for several arrays were also obtained by relative $\Phi_{\rm f}$ measurements using mesotetraphenylporphyrin in non-degassed toluene ($\Phi_{\rm f}=0.070$)³⁰ as a standard and gave the same values as the absolute measurements within experimental uncertainty ($\pm 5\%$ of the reported value).

Conflicts of interest

The authors declare no competing financial interests.

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