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EDGE ARTICLE

Functionalizing molecular wires: a tunable class of α , ω -diphenyl- μ , ν -dicyano-oligoenes†‡

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We describe the synthesis and characterization of a new class of cyano-functionalized oligoenes and their derivatives. We have made the vinylogous series of α,ω -diphenyl- μ,ν -dicyano-oligoenes (**DPDCn**) comprised of each odd-numbered member from 3 to 13 linear conjugated olefins. Installing cyano groups onto the oligoene backbone lowers HOMO and LUMO energies by up to \sim 0.7 eV, thereby stabilizing the molecule with respect to oxidative decomposition; this exemplifies a new approach to the stabilization of conjugated oligoenes. UV-vis absorption spectra and redox potentials across the **DPDCn** series reveal that the molecular band gap ranges from 2.80 to 1.75 eV. This gap can be further tuned by the facile installation of a variety of aryl end-groups. The choice of end-groups also greatly affects the physical properties such as solubility and the solid-state packing. We also present the longest oligoene crystal structure reported to date. Moreover, we find that the prototypical linear structure makes oligoenes suitable as molecular wires and connectors in the bottom-up construction of nanoscale architectures. As a proof of concept, carboxylic acid terminated oligoenes were used to position 10-nm Fe₃O₄ nanoparticles on a GaAs (100) substrate.

Introduction

In this study we present a new class of tunable, functional and easily derivatized molecular wires. These wires are noteworthy because they are molecularly defined, atomically precise fragments of polyacetylene (PA, Fig. 1a). PA is remarkable because it is a simple hydrocarbon polymer that is highly conductive when

doped.¹ It also has a low band gap and shows large nonlinear optical susceptibilities.² Since the fully conjugated high polymer of –C(H)=C(H)– is electrically conductive, we believe that well-defined oligomers, hereafter referred to as oligoenes, should be useful molecular conductors in nanoscale situations. In addition, naturally occurring terpenoid-based oligoenes, such as carotenoids, have been made into electrical devices and are known to have useful optical properties.³ However, nature provides only a limited number of structures that are often difficult to functionalize further, thus preventing their use in widespread applications.

The synthesis and study of conjugated oligomers has bourgeoned over the past 20 years, however oligoene systems remain understudied. Three major challenges have hampered the development of oligoenes as electronic materials: (1) the most thoroughly studied examples of PA oligomers have been obtained inefficiently through the painstaking isolation of individual oligomers from polymerization reactions terminated at low monomer conversion.4 (2) Previous preparative methods have not supported diverse functionalization to incorporate them into electronic devices. (3) Oligomers of PA are chemically impractical because they are nearly insoluble and oxidatively unstable. Using the new synthetic strategy detailed below for the α,ω -diphenyl- μ,ν -dicyanooligoenes⁵ (DPDC, Fig. 1c) we have overcome these challenges. The key to this synthesis is the use of sterically innocent yet electronically stabilizing cyano groups bound directly to the oligoene chain while protecting the reactive

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 $[\]dagger$ Electronic supplementary information (ESI) available: List of all molecules, detailed synthetic procedures and characterization data of oligoenes, oligoene-aldehydes, Fe $_3O_4$ nanoparticles, and nanoparticle–oligoene SAMs; UV-vis spectroscopic data; cyclic voltammetry; CIF files for DPDCn; DFT theoretical methods and results for DPDCn and DPOn and selected functionalized oligoenes. CCDC reference numbers 847946–847952. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c2sc00770c

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(c) α,ω -diaryl- μ,ν -dicyano-oligoenes (DPDC)

$$Ar = \underbrace{\begin{array}{c} X \\ X = H, DPDCn \\ F, DPDCn-F \\ Br, DPDCn-Br \\ CO_2H, DPDCn-CO_2H \\ CO_2Et, DPDCn-CO_2 \\ NO_2, DPDCn-NO_2 \\ CN, DPDCn-CN \\ C \equiv C - C_6H_{13}, DPDCn-C_8H_{13} \\ SMe, DPDCn-SMe \\ NMe_2, DPDCn-NMe_2 \\ NPh_2, DPDCn-NPh_2 \\ \end{array}}$$

$$X = Ar = \underbrace{\begin{array}{c} X \\ Y = 0, DPDCn-SC_4H_8 \\ 1, DPDCn-SC_5H_{10} \\ X \\ Ar = \underbrace{\begin{array}{c} X \\ Y = 0 \\ Y = 0, DPDCn-C_4H_8 \\ 1, DPDCn-CC_6H_{13} \\ Y = 0 \\ Y = 0$$

Fig. 1 Structure of (a) polyacetylene (PA). (b) Structure of α ,ω-diphenyl-oligoenes (DPO), which we regard as control molecules for comparison with (c) cyano-functionalized α ,ω-diphenyl-oligoenes (DPDC). We identify the oligoene molecules according to the length of their linear conjugated backbone (**n**) and aryl end-group (**X**), **DPDCn-X**. For instance, 1,10-di(4-bromophenyl)-4,7-dicyano-deca-1,3,5,7,9-pentaene is denoted as **DPDC5-Br**. See ESI† for a complete list of molecules.

terminal olefins with bulky phenyl groups. These phenyl endgroups can easily be modified with a diverse range of functionality. We recently reported that alkylthio-terminated members of this family conduct electrically; here we extend that study to reveal the wide variety of compounds that can be produced through our synthetic method. We also highlight the versatility and adaptability of these new electronic materials by using appropriately functionalized oligoenes as electrical conduits that connect magnetic nanoparticles to a semiconductor surface.

Other methods have been used to stabilize oligoenes. Several approaches protect the highly reactive terminal olefins with bulky end-groups such as phenyl (DPO, Fig. 1b)6 or tert-butyl.8,9 Other than such hydrocarbon-end-capped oligoenes, however, there are few other examples of non-terpenoid α,ω-disubstituted oligoenes longer than pentaenes. 10,11 Nature provides its own methods of oligoene stabilization: carotenoids are the most wellstudied oligoenes since they play essential roles in vision (retinal), cell differentiation (retinol) and photosynthesis. 12 In these molecules, methyl groups decorate the oligoene backbone and contribute to both the solubility and stability. However, they also create allylic strain along the poly-olefin backbone, which causes the molecules to bend. These distortions reduce the conjugation and increase the energy difference between the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO), therefore shifting the electronic absorptions to higher energies.

In order to use oligoenes in optical devices, it is desirable to have a family of light-harvesters that cover the entire visible and near-infrared spectral regions. Therefore, we sought small and synthetically available substituents that could confer stability while maintaining the bathochromic absorption shifts expected for longer oligoenes. Cyano groups fulfill these requirements: they are π -electron withdrawing groups that are not bulky enough to disrupt molecular planarity. They have been used to alter the band gaps of aromatic molecules and conjugated polymer materials. Because of their electron-withdrawing ability, they are expected to lower the energies of the frontier orbitals, thus stabilizing the system toward oxidative decomposition. As an ultimate benefit they are easily installed and enable an economical synthesis that allows for diverse end-group functionalization.

Results and discussion

Synthesis

The nexus of our convergent synthesis of the oligoenes is the Knoevenagel condensation¹⁶ between 1,4-dicyano-2-butene and two equivalents of the appropriate aryl-enal (Scheme 1a). We prepare the aryl-enals starting from the analogous benzaldehyde or *trans*-cinnamaldehyde *via* iterative Wittig homologations¹⁷ and subsequent acidic hydrolysis of the intermediate acetal. The conditions for both reactions tolerate a wide variety of functionality and give access to a range of derivatives. Fig. 1 shows those derivatives that we prepared for this study *via* this method. They are made in a small number of steps from commercially available starting materials.

In general the Knoevenagel condensations give moderate yields, which decease with increasing molecular length, producing the **DPDCn** series in 10–60% isolated yields as shown

Scheme 1 Synthesis of DPDCs *via* (a) double-Knoevanegel condensation reaction and (b) Wittig reaction. *Reagents and conditions*: (i) DBU, MeOH, 25 °C, 6–12 h, (60–10%, for **DPDCn** series; from m = 0–5); (ii) LiOMe, THF, 75 °C, 12–24 h; (iii) 10% aq. HCl, 25 °C, 2 h (70–98%). For instance, if m = 1, then the intermediate acetal leads to the product (2*E*,4*E*)-5-phenylpenta-2,4-dienal. A list of all compounds is provided in ESI†.

E/Z mixture

Table 1 Double Knoevenagel condensation yields for the unfunctionalized **DPDCn** series

Oligoene	Oligomeric length (n)	Isolated yield (%)	
DPDC3	3	55	
DPDC5	5	60	
DPDC7	7	54	
DPDC9	9	45	
DPDC11	11	40	
DPDC13	13	10	

in Table 1. Nonetheless, the condensation reaction is operationally simple: it does not require anhydrous solvents and the isolation does not require chromatography. This reaction joins two fragments together and more than doubles the length of the oligomers. The products precipitate from the reaction mixture and are easily isolated by filtration. We purify the crude product by recrystallization from CH₂Cl₂—MeOH. With this approach, the synthesis of **DPDC13** (one of the longest oligoenes prepared to date) is straightforward and involves only five steps. It is also important to mention that the sequence shown in Scheme 1 is scalable; we tested some derivatives on the gram-scale with no loss in their isolated yields.

All of the compounds have exclusively trans-stereochemistry in their double bonds. The stereochemistry of the double bonds in the each aldehyde can be identified by the coupling constants of the olefinic doublet of doublets near 6.5 ppm (this signal belongs to the β-proton of the aryl-enal). This resonance is coupled to the protons of the terminal aldehyde and to a transolefin (-CH=) proton. This resonance resides in an isolated region of the spectra allowing for stereochemical characterization throughout the growth of the aryl-oligoenals. Wittig conditions (Scheme 1b) produce an E/Z-mixture of stereoisomers, however acidolysis of the intermediate acetals exclusively produces the trans-product. Cleavage of the acetal is carried out before separation; however, to further understand our method we tracked the stereoisomeric transformations throughout the extension of trans-cinnamaldehyde to (2E,4E)-5-phenylpenta-2,4-dienal. In this case performing the Wittig reaction on trans-cinnamaldehyde produces the (E)- and (Z)-4-phenylpenta-2,4-dienyl acetals in 16 and 60% yield, respectively. We separated each product by column chromatography; coupling constants from ¹H NMR spectra identified each stereoisomer. Acid hydrolysis of either stereoisomer, E or Z, generates the same product, (2E,4E)-5-phenylpenta-2,4-dienal, in quantitative yields.

In order to verify the practical value of cyano substitution, we have also prepared the unsubstituted α, ω -diphenyl-oligoenes of the corresponding lengths. We synthesized these molecules lacking the cyano groups via the Horner–Wadsworth–Emmons reaction¹⁸ as described by Spangler and co-workers. Dicyano-oligoenes are much less reactive toward ambient oxidation, intermolecular oligomerization or intramolecular decomposition than their unsubstituted relatives and we can conveniently study them under normal aerobic laboratory conditions for extended periods. The thermal stability of these dicyano-oligoenes (**DPDC3–DPDC13**) is similar to that of the $\alpha, \alpha, \omega, \omega$ -tetrakis-tert-butyl-functionalized oligoenes reported by Hopf and

co-workers⁹ and far higher than that of the methyl and unfunctionalized analogs.²⁰ Differential scanning calorimetry shows that under an inert atmosphere all members of the parent series **DPDC3–DPDC13** are thermally stable up to ~250 °C. For a direct comparison between cyano-functionalized and unfunctionalized oligoenes, we photooxidatively decomposed both **DPO5** and **DPDC5** in a side-by-side experiment. We monitored the oligoene starting materials with UV-vis spectroscopy and found that the unsubstituted material decomposes four times faster than the cyano-functionalized material.²¹

Band-gap modulation

In order to appraise the effects of cyano-substitution we first consider the derivatives that have unsubstituted phenyl groups, **DPDCn**. Despite providing stabilization to the oligoene core, the cyano groups do not fundamentally alter the "polyacetylenelike" electronic behavior of oligoenes in the sense that the HOMO-LUMO gap decreases as the length of the oligoene increases, which is typical of other vinylogous series.²² The optical absorptions of the **DPDCn** series for **n** between 3 and 13 span the entire visible spectrum (Fig. 2a). The absorptions are also quite intense with molar extinction coefficients (ε) that exceed 105 M⁻¹ cm⁻¹. We estimate the solution-phase optical band gap (E_{og}) for **DPDC11** and **DPDC13** to be 1.81 and 1.77 eV, respectively. These values approach the band gap observed for PA itself.23 Similarly, we can analyze this series using the quantum mechanical "particle in a box" model, which is often used to describe optical absorptions in one-dimensional systems. This model holds that the excitation energies in a simple system vary with the inverse square of the dimensional length. Fitting the excitation energies in the **DPDCn** series with such an inverse-square expression (see Fig. 2c) leads to an effective mass of 2.15×10^{-45} kg. Extrapolation of the wavelength for the strongest absorption (λ_{max}) and optical band gap (E_{og}) to an infinite "box" length estimates values of 577 nm and 1.59 eV, respectively. This is within the reported range of band gaps (1.4-1.6 eV) for trans-PA.^{1,23} Our extrapolations were carried out as simple linear regression analyses; the Meier correction²⁴ did not provide any benefit in this case because the effective conjugation length (ECL) had not been reached. The ECL of polyenes is higher than other poly-conjugated systems²² and has been estimated to lie between 15 and 20 repeat units. 4,25 We estimate the ECL of a DPDC polymer by selecting the point in our three extrapolations where a change in oligomeric length of one repeat unit corresponds to a negligible energy difference of less than 0.01 eV. In each case, the emergence of size-independent electronic properties (λ_{max} , E_{og} and E_{cv}) is predicted to occur at lengths \geq 19 alkene repeat unites.

Since cyano groups are strongly electron withdrawing, their presence should lower the energies of the frontier orbitals with respect to those of the unfunctionalized α , ω -diphenyl-oligoenes. The redox potentials determined by cyclic voltammetry were used to estimate the energies of the HOMOs and LUMOs of the **DPDCn** series. We found that installation of two cyano groups onto the oligoene core stabilizes the HOMO and LUMO by as much as 0.44 and 0.68 eV with respect to the corresponding unsubstituted **DPOn** molecules (Fig. 2b). This is consistent with UV-vis absorption spectroscopy, in which we observe

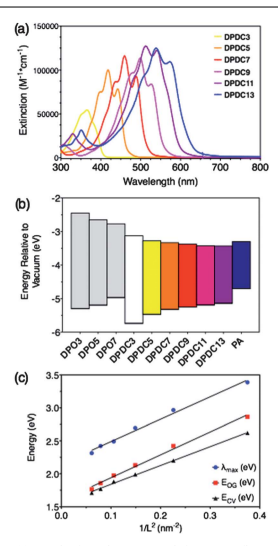


Fig. 2 (a) UV-vis absorption spectra of the parent oligoene series (DPDCn) in CH₂Cl₂. Plot of extinction coefficient vs. absorption wavelength. The wavelength of strongest absorption, λ_{max} (from left to right) at 363, 418, 459, 499, 513 and 536 nm; increasing the length of oligomers tunes the optical absorptions over a range of 350 nm. (b) Effect of the cyano groups on the HOMO and LUMO energies of diphenyl-oligoenes determined by cyclic voltammetry (CV). Redox gap of DPO3, DPO5, DPO7 (grey) and the cyano-functionalized series, DPDCn. CVs were obtained in DMF solution with 0.1 M (n-Bu)₄N+PF₆-. Ferrocene was used as an internal standard for electrochemical measurements. Experimentally determined values for PA (blue) are included for reference. 23 (c) $\lambda_{\rm max}$, the optical band gap $(E_{\rm og})$ and the redox band gap (E_{cv}) for the **DPDCn** series are fit using "particle in a box" energies. Extrapolative analysis estimates the analogous values for an oligomer of infinite length (polymer) of 577 nm, 1.59 eV and 1.55 eV, respectively. Predicted values agree with experimentally determined values for PA.

a bathochromic shift of up to 35 nm between the **DPOn** and **DPDCn** series (see Fig. 3a). The electrochemically determined energy gap, $E_{\rm cv}$, for **DPDCn** molecules can also be extrapolated to predict the band gap of the homologous polymer. In Fig. 2c we again use simple "particle in a box" energies to predict the band gap for the polymer to be 1.55 eV. Both the $E_{\rm cv}$ and the $E_{\rm og}$ converge to similar values.

End-functionalized derivatives

Substitution on the terminal phenyl groups offers another opportunity to tailor the electrical and physical properties of oligoenes. We synthesized a family of derivatives of DPDC5 (Table 2), whose phenyl-substituents varied from strongly electron-withdrawing (NO₂) to electron-donating (NMe₂). We found para-substitution to be most effective, apparently since it is strongly resonance-coupled to the oligoene backbone and at the same time sterically remote (see Fig. S4, ESI†). Spangler and co-workers reported similar results for non-cyano DPOs having para-substituents NO₂, Cl, SMe, OMe and NMe₂. ¹⁰ Examples of the effect of aryl-functionalization on the optical absorption are displayed in Fig. 3a. Strong electron-donating groups give the largest bathochromic shifts and broadened absorption peaks. Functionalization of the **DPDC5** scaffold with p-NMe₂ groups shifts the longest-wavelength absorption from 490 to 640 nm, and this shift is accompanied by the loss of vibronic fine-structure.26 These electronic differences indicate topological changes in the frontier orbitals due to a transformation from an oligoene π - π * excitation to a "push-pull" n- π * transition; giving rise to an intramolecular charge transfer (ICT).27 We performed timedependent density functional theory (TD-DFT)²⁸ calculations at the B3LYP/6-31** level on DPDC5 and DPDC5-NMe2 to investigate their excited states and the topologies of relevant molecular orbitals. In each case we found that the lowest-energy excitation is predominantly promotion of an electron from the HOMO to the LUMO, and in each case this transition is strongly allowed. We show these frontier orbitals for each molecule in Fig. 3c and d, respectively. The LUMOs of DPDC5 and DPDC5-NMe₂ are remarkably similar: each is primarily the expected, lowest-energy π^* -orbital of a conjugated polyene. However, the HOMOs of the two molecules differ. While the HOMO of **DPDC5** is primarily the expected, highest-energy π -orbital of the polyene, in DPDC5-NMe2 there is significant electron density on the terminal aryl groups and the nitrogen lone pairs. A similar trend in the frontier orbitals is predicted even when the cyano groups are removed from the oligoene core (see Fig. S14, ESI†) implying that the cyano groups contribute less to the intramolecular charge transfer (ICT) than do the NMe2 groups. Thus while the HOMO-LUMO excitation in **DPDC5** is largely π - π *, that in DPDC5-NMe₂ has a significant component of $n-\pi^*$ character. This may be considered ICT.29

Since the HOMO–LUMO transition in **DPDC5-NMe₂** is due, at least in part, to $n-\pi^*$ promotion, protonation of the nitrogens should remove the nitrogen lone pair participation in the HOMO and therefore quench any ICT, leaving behind the $\pi-\pi^*$ absorptions of the oligoene. Indeed, addition of trifluoracetic acid to solutions of **DPDC5-NMe₂** and **DPO5-NMe₂** resulted in a hypsochromic shift and resolution of the vibronic fine-structure as seen in Fig. 3b. The effect is reversible; when we neutralize the erstwhile acidic solutions with triethylamine, absorption is again shifted to the red and the vibronic structure is lost.

Solvatochromism may be used to assess the amount of ICT behavior in conjugated push–pull systems, since the polarizability of π -electrons partially depends on the environment provided by the solvent, such as polarizability and solvent cavity size. ²⁹ The absorption behavior of **DPDC5-NMe₂** was surveyed throughout 15 organic solvents. The value of λ_{max} was observed

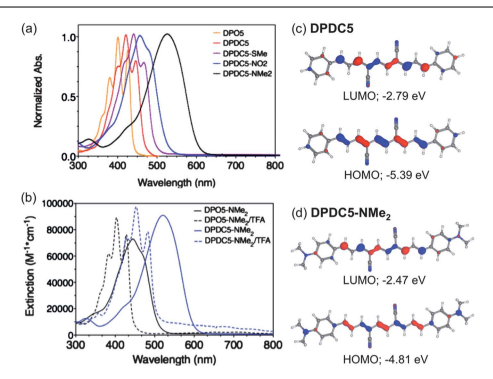


Fig. 3 (a) UV-vis absorption spectra of oligoene derivatives in CH₂Cl₂; **DPO5** (orange), **DPDC5** (red), **DPDC5-SMe** (purple), **DPDC5-NO₂** (blue) and **DPDC5-NMe₂** (black). For easy comparison absorptions have been normalized to 1. (b) Absorption spectra of protonated (solid line) and deprotonated (dashed line) forms of **DPDC5-NMe₂** (black) and **DPO5-NMe₂** (blue) in CH₂Cl₂. TD-DFT calculations compare the LUMO and HOMO of (c) **DPDC5** and (d) **DPDC5-NMe₂**.

to shift 55 nm from solutions of heptane to nitrobenzene, while the longest-wavelength absorption shifted 70 nm from heptane to dimethyl sulfoxide (DMSO) (see Fig. S7, ESI†).

The rigid structure of conjugated oligoenes leaves them sparingly soluble in organic solvents and hinders solution-based processing techniques such as spin-casting or inkjet printing.

Table 2 Comparison of selected DPDC5 derivatives and their UV-vis strongest-wavelength absorptions in CH₂Cl₂ solution

Entry	R	$\lambda_{\max}^{a}/\text{nm (eV)}$	Entry	R	$\lambda_{\text{max}}^{a}/\text{nm} \text{ (eV)}$
1	DPDC5	418 (2.97)	6	$\begin{array}{c} \text{OC}_{6}\text{H}_{13} \\ \text{OC}_{6}\text{H}_{13} \\ \\ \textbf{DPDC5-OC}_{6}\text{H}_{13} \end{array}$	454 (2.73)
2	Br DPDC5-Br	426 (2.91)	7	$OC_{10}H_{21}$ $OC_{10}H_{21}$ $OC_{10}H_{21}$ DPDC5-OC ₁₀ H ₂₁	454 (2.73)
3	Z, CO ₂ H DPDC5-CO ₂ H	434 ^b (2.86)	8	OC ₁₄ H ₂₉ OC ₁₄ H ₂₉ DPDC5-OC ₁₄ H ₂₉	455 (2.73)
4	NO ₂ DPDC5-NO ₂	437 (2.84)	9	SMe DPDC5-SMe	453 (2.74)
5	NMe ₂ DPDC5-NMe ₂	521 (2.38)	10	DPDC5-SC ₅ H ₁₀	463 (2.68)

^a Strongest wavelength absorption is taken at the global λ_{max} and is not indicative of HOMO–LUMO gap energies, E_{og} . ^b Due to the limited solubility absorption spectrum was taken in DMSO.

This is an important obstacle for promising organic electronic materials to overcome for application in film-based technologies. Again, we take advantage of the facile functionalization of phenyl end-groups in order to enhance oligoene solubility. Long alkoxy groups and bulky thioethers (Table 2, entries 6–10) tune the solubility over a wide range while maintaining consonant electronic properties (based on UV-vis absorption, as well as ¹H and ¹³CNMR spectra; see Fig. S5, ESI†). For example, **DPDC5-OC**₁₄H₂₉ and **DPDC5-SC**₅H₁₀ reach molarities up to ~0.10 M in chloroform at room temperature; increasing the solubility from ~10⁻⁴ M for **DPDC5**. These derivatives are also soluble in other common organic solvents such as toluene, tetrahydrofuran and benzene. Using this synthetic approach, longer alkoxy chains or even polymeric attachments are presumed to be possible.

Solid-state structures

We have grown crystals of each member of the **DPDCn** family (from **n** = 3 to 13) and determined the molecular structure of each crystallographically. Each oligoene occurs exclusively in the all-trans form. **DPDC13** (Fig. 4) is the longest oligoene thus far characterized by XRD, longer even than the natural product, rhodopin glucoside, ³⁰ which contains 11 linear conjugated C=C bonds. Although some of the shorter vinylogues have been previously reported in the resin dye patent literature, ³¹ a systematic characterization has been heretofore unavailable. Only a few non-terpenoid structures having more than six conjugated C=C bonds have been previously reported, and none having more than nine. ³²

We observe different packing structures within the parent series of **DPDCn** oligomers. The shortest vinylogues, **DPDC3** and **DPDC5**, are planar and stack in a herringbone pattern, indicating that the cyano groups do not significantly interact with allylic hydrogens (see Fig. S10, ESI†). The longer oligomers, **DPDC7–DPDC13**, co-crystallize with one solvent molecule per unit cell and are not fully planar. The central dicyanobutene moiety is planar and to either side out-of-plane bending of the conjugated backbone is accompanied by a slight twist. DFT calculations at the B3LYP/6-31G** level predict fully planar conformations, however the incorporation of solvent as well as

bending is common in crystal structures of oligoenes having more than six C=C bonds.^{8,9,32}

We were concerned that the cyano groups would disturb the local structure of the oligoene backbone and minimize the degree of conjugation. Despite the observed bending, the central alkene unit (that having doubly allylic cyano groups) remains planar with a dihedral angle of 180.0°. This suggests that deviations from planarity may be attributed to co-crystallization with solvent molecules and not the result of cyano-functionalization.

The molecular organization in the solid state changes when we functionalize the DPDCs with carboxylate groups. Unlike the pentaene of the parent series (DPDC5), crystals of DPDC5-CO₂H contain two molecules of solvent, dimethylformamide (DMF), per oligoene and do not organize in a herringbone structure. In Fig. 5 we show that hydrogen bonding with DMF facilitates the assembly of the carboxylic acids into π -stacked sheets of oligoenes. The lack of solvent between oligoenes allows for the intermolecular distances to decrease to 3.38 Å between π faces of neighboring molecules. These values approach the interplanar distances found for the classic carbon allotrope, graphite; 3.35 Å. 33 DPDC5-CO₂H stacks in an oblique alignment with six close neighbors oriented in a hexagonal pattern when viewed down the length of the molecule.34 All molecules in the crystal are aligned in the same direction and overlap with neighboring molecules through π -stacking interactions. This is an ideal geometry for charge transport through a single crystal.

Self-assembly

As a demonstration of how this synthetic control enables us to incorporate our molecular wires into materials, we used functionalized oligoenes, **DPDC3-CO₂H**, to assemble monolayers of magnetic nanoparticles on GaAs substrates. It has been shown previously that carboxylic acids bind to GaAs³⁵ and Fe₃O₄ through known ligand-exchange procedures.³⁶ First, the native oxide layer of the GaAs substrates were removed through submersion in aqueous ammonia (NH_{3 aq}). After rinsing with ethanol they were submerged in a 1.5 mM solution of oligoene (**DPDC3-CO₂H**) in DMSO. They were then removed from solution, again rinsed with ethanol, and submerged for 1 h in a solution of 10-nm Fe₃O₄ nanoparticles in DMF. Scanning

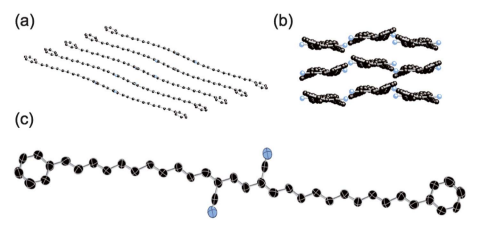


Fig. 4 Packing structure of DPDC13 (a) viewed from the side and (b) down the length of oligoene chain shows deviations from planarity. (c) ORTEP plot of DPDC13. Ellipsoids represent 50% probability levels. Hydrogen atoms have been omitted.

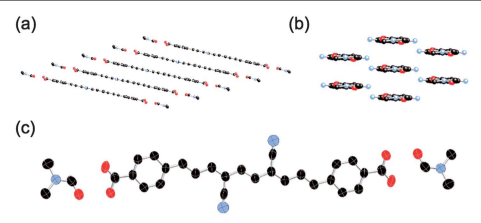


Fig. 5 Packing structure of DPDC5-CO₂H·2DMF (a) viewed from the side and (b) down the length of oligoene chain. (c) Ellipsoids represent 50% probability levels. Hydrogen atoms have been omitted.

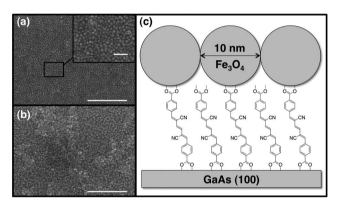


Fig. 6 Scanning electron micrographs (SEM) of 10-nm Fe₃O₄ nanoparticle SAMs made using (a) 1.5 mM and (b) 3.0 mM solution of **DPDC3-CO₂H**, show monolayer coverage at low concentrations and the presence of multilayers at higher concentration. Scale bars are set to 500 nm, while the inset scale bar is set to 50 nm. (c) Schematic of oligoene—Fe₃O₄ composite monolayer.

electron microscopy (SEM) analysis of these films shows clear formation of nanoparticle monolayers as shown in Fig. 6. Nanoparticle coverage was found to be dependent upon the concentration (1.5, 3.0 or 6.0 mM) of the oligoene solution. At higher concentrations (3.0 and 6.0 mM) the appearance of multilayers in addition to monolayers was observed (Fig. 6b; Fig. S12, ESI†). No monolayer formation was observed when only DMSO without oligoene was used or upon the direct submersion of the GaAs substrate into the nanoparticle solution (see Fig. S12, ESI†).

Conclusion

We have developed a new method to synthesize cyanosubstituted oligoenes with up to 13 all-trans conjugated C=C bonds, and we have demonstrated its utility in the synthesis of a wide variety of aryl-functionalized oligoenes. Cyano groups decrease the band gap of oligoenes and stabilize them for study under normal laboratory conditions while preserving the conjugation along the poly-olefin backbone. Derivatization of the terminal phenyl groups allows us to fine tune the electronic structure and physical properties. We can access practically any energy within the visible spectral region by pairing the appropriate oligoene length with the appropriate aryl end-group. Functionalized oligoenes are ideal structures for nanoscale electronic and structural components and can be used to guide oligoene self-assembly and the construction of heterogeneous materials. Furthermore, the utility of these materials in photovoltaics is an obvious application since their band gaps are easily tuned, their absorptions cover the visible spectrum, and they are predicted to be useful in singlet fission.³⁷

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