Cyclodimers and Cyclotrimers of 2,3-Bisalkynylated Anthracenes, Phenazines and Diazatetracenes

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Abstract: The synthesis of novel (N-)acene-based cyclooligomers is reported. Glaser-Hay coupling of the bisethynylated monomers results in cyclodimers and cyclotrimers that are separable by column and gel-permeation chromatographies. For the diazatetracene, the use of sec-butylsilylethynyl groups is necessary to achieve solubility. Diazatetracene-based cyclodimers and cyclotrimers were used as semiconductors in thin-film transistors. Although their optoelectronic properties are quite similar, their electron mobilities in proof-of-concept thin-film transistors differ by an order of magnitude.

The reaction of diethynylbenzene with Cu(OAc)₂ under oxidative conditions, published by Eglinton and Galbraith, selectively furnished the fairly angle deformed cyclodimer A (Figure 1),[1,2] which polymerizes in the solid state.[2] Later Swager et al.[3] demonstrated that the apparent absence of the expected, less-strained cyclotrimer was due to its insolubility - alkoxysubstituted diethynylbenzenes do give cyclotrimers in good yield upon oxidative homocoupling.[4] Haley et al.[5] published a series of elegant papers in which they connected such cyclotrimers into larger segments of a carbon allotrope called graphdiyne. Yet, only benzene rings[6] were investigated until, in 1998, Komatsu published the dimerization of diethynylnaphtalene derivatives furnishing B,[7] and Anthony prepared structure C as the cyclotrimer of a substituted 2,3-diethynylpentacene.[8]

We prepared the precursor 1,2-diyynes (Scheme 1) by Stille coupling of 1,2-dibromoarenes using TMSCSMe₃ and a bis(benzonitrile)palladium(II) chloride precatalyst[9] in the presence of tBu₃PHBF₄; potassium carbonate liberates the terminal diynes, which were administered by a syringe pump into a solution of copper acetate and pyridine under air. Without a syringe pump, ill-defined polymers formed—we could not isolate or detect cyclooligomers, etc. by mass spectrometry. The purification of the target compounds, formed in 12–13% for the cyclodimers and around 15–25% for the cyclotrimers (Figure 2), was performed by gel permeation chromatography. Surprisingly, dimer 3b₂ could not be isolated, testament to its insolubility - only 3b₃.

Attempts to cyclize diethynyltetraazapentacenes and their dihydro analogues failed, probably due to interfering redox processes. We noticed that the cyclodimers were always considerably less soluble than the cyclotrimers in variance to

Figure 1. Structures of selected known arylene-1,2-bisalkynylene dimers and trimers.

Scheme 1. Synthetic route to (aza)acenylene-alkynylene dimers and trimers.

i) TMSCSMe₃ (5.00 equiv.), (C₆H₅CN)₂PdCl₂ (10 mol%), P(tBu)₃HBF₄ (20 mol%), THF, RT, 10 h; ii) K₂CO₃ (20.0 equiv.), THF/MeOH (1:1), RT, 1 h; iii) Cu(OAc)₂ (22.0 equiv.), pyridine/MeOH (1:1), 60 °C, 24 h.

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the observation made for the parent systems, and that, despite their decoration with R₃Si-ethynyl groups. To assess the thermal stability of the series, TGA/DSC experiments were conducted. All materials are stable up to 300 °C. The least stable compound is strained dimer 3c₂, its decomposition begins at ~300 °C. For the monomeric compound 3c and the trimer 3c₃, decomposition approximately starts at 350 °C (Figures S46–S52 in the Supporting Information).

Figure 3 displays the UV-vis spectra of monomeric 3c and the cycles 3c₂. The cyclooligomerization leads to an increased red shift up to 31 nm for the absorption maximum. This is probably due to the diazatetracene-dialkyne being the major chromophore; the conjugation through the diyne bridges is fairly weak, supported by the lack of ring current over the butadienynes revealed by anisotropy of the induced current density (AICD) calculations[10] (Figure S6). The same trend is visible for 3a and 3a₂, 3b and 3b₃ (red shift in 3b/3b₃, only visible in onset; Figures S31 and S32).

DFT calculations (B3LYP, def2-TZVP) show that the frontier molecular orbitals are evenly distributed over the molecules. Tables 1 and 2 display the experimental and calculated optical and electronic values for the compounds reported herein.
Cyclooligomerization stabilizes the LUMOs by up to 0.29 eV. This effect was not reproduced for the reduction events observed in the cyclovoltammograms, which do not differ significantly.

We notice that the c-series with diazatetraacene as chromophore displays robust electron affinities of approximately −4.0 eV (measured) and −3.5 to −3.7 eV (calculated), thus suggesting that these should be electron transport materials in n-channel transistors.

We obtained a single crystalline specimen of the target compounds (Figure 4 and the Supporting Information). The quasiplanar dimers 3a2 and 3c2 form slightly offset one-dimensional staircases with π−π stacking distances of 351 and 338 pm (averaged through both chromophores and butadiynyl linkers), respectively. In the supramolecular structures, one acenylene subunit is centered above the octadehydro[12]annulene moiety of the neighboring molecule, hinting at less-than-ideal charge transport properties. Unlike the Hay-type dimers, 3a–c, deviate from planarity - the (aza)acenylene subunits are slightly twisted with respect to each other, a feature also observed in Anthony’s pentacene-based dehydroannulene,[11,12] which is most likely a crystal packing effect as gas-phase structures were calculated to be planar. In contrast to the pentacene-based annulene exhibiting three-dimensional π-stacking interactions, only two-dimensional ones are observed in the trimers: In 3a3 (co-crystallizing with chloroform) and 3c3 (solvent-free packing), two of the three “arms” overlap-distances between π-system distances amount to ~3.4 Å. 3b3 forms dimers in which two phenazine chromophores π-stack (~3.1 Å) on top of each other. These then π interact (~4.7 Å with the neighboring dimer pairs forming a one-dimensional slipped dimer stack.

Due to their electrochemical properties and the possibility to obtain solvent-free crystal structures, diazatetracene-based 3c2 were promising candidates for thin-film transistors (TFTs). Note that only Anthony’s trimer was employed in an organic electronic device, albeit in a photovoltaic cell.[10] TFTs were fabricated in bottom gate/top contact architectures (Figure 5 and Supporting Information) with silver as contact electrodes, using a SAM modified dielectric.[156] Thin films of 3c2 and 3c3 were obtained by drop-casting (3c2: toluene, 0.5 mg/mL; 3c3: CHCl3, 0.5 mg/mL). The average electron mobility of the trimer (μave = 1.85x10−2 cm2/Vs) was one order of magnitude higher than that of the dimer (μave = 2.10x10−3 cm2/Vs), probably due to fewer grain boundaries formed upon evaporation of the

![Normalized absorption spectra of diethynyldiazatetracene 3c, its Glaser dimer 3c2, and trimer 3c3 in CHCl3.](image-url)
volatile dichloromethane compared to the microcrystalline film of \(3 \text{c}_2\) from high-boiling toluene solution (see Figure S43 for polarization micrographs), and as a consequence of the higher transfer integrals (38.8 meV for \(3 \text{c}_2\) and 3.55 meV for \(3 \text{c}_3\)) for electron transport (Figure S7). Although TIPS-ethynylated diazatetraconcne,\(^{[17]}\) which was found to pack in a brick-wall motif, exhibited an electron mobility of \(\mu_{\text{brick}} = 5 \times 10^{-2} \text{ cm}^2/\text{V s}\),\(^{[18]}\) the mobilities of trimer \(3 \text{c}_3\) are on the same order of magnitude and that despite bulkier (sBu)Si-ethynyl groups being used to solubilize the material.

In conclusion, we have synthesized and characterized novel acene-based cyclodimers \(3 \text{a}_2\) and \(3 \text{c}_2\) as well as cyclotrimers \(3 \text{a}_3 \cdots \text{c}_3\). Cyclodimer \(3 \text{c}_2\) and cyclotrimer \(3 \text{c}_3\) were used as organic semiconductors in bottom gate/top contact thin-film transistors. The mobility of the trimer is an order of magnitude higher than that of the cyclodimer, although the electronic and optical properties are quite similar - most likely resulting from improved \(\pi-\pi\) interactions between the azaacene subunits. The cyclotrimer \(3 \text{c}_3\) shows mobilities in the same order as that of unsubstituted tetraazatetraconcne in the literature, although sec-

![Figure 4. Single crystal structures and stacking of diazatetraconcnc dimer \(3 \text{c}_2\) (left) and trimer \(3 \text{c}_3\) (right). TIPS-ethynyl substituents were reduced in size for visualization.](image1)

![Figure 5. a) Transfer characteristics of trimer \(3 \text{c}_3\) (left; \(V_{\text{DS}} = 35 \text{ V}\)) and dimer \(3 \text{c}_2\) (right; \(V_{\text{DS}} = 20 \text{ V}\)). b) Schematic bottom-gate/top-contact device architecture. c) Charge transfer mobilities of dimer \(3 \text{c}_2\) and trimer \(3 \text{c}_3\).](image2)
butyl-silyl groups were used. This highlights the importance of the solid-state architecture, which could be influenced by modifying the structure of the single molecule.

**Experimental Section**

Representative procedure towards Glaser–Hay dimers 3c₁ and 3c₂: Under ambient atmosphere, copper(II) acetate (3.04 g, 15.2 mmol, 22.0 equiv.) was dissolved in a mixture of 175 mL pyridine and 175 mL methanol. The mixture was heated to 60°C. Afterwards a solution of 3c (500 mg, 691 μmol, 1.00 equiv.) in 20 mL of pyridine was added dropwise using a syringe pump (rate 0.06 mmol/h). After completion of the addition, the reaction mixture was stirred for additional 4 h. After cooling to room temperature the solvent was removed under reduced pressure. Water and CH₂Cl₂ were added and the mixture was extracted with CH₂Cl₂. The combined organic layers were dried over MgSO₄ and the solvent was removed under reduced pressure. The crude product was purified by column chromatography on silica (PE/CH₂Cl₂ 8:2, 7:3, 1:1) to yield a mixture of 3c₁ and 3c₂ containing some small impurities. Purification of the mixture using a GPC yielded the dimer 3c₂ (60.0 mg, 12 %) as a dark brown solid and the trimer 3c₁ (100 mg, 20 %) as a dark brown solid.

Deposition Numbers 2106563 (for 3a₁), 2106564 (for 3a₂), 2106565 (for 3b₁), 2106566 (for 3c₁), and 2106567 (for 3c₂) contain the supplementary crystallographic data for this paper. These data are provided free of charge by the joint Cambridge Crystallographic Data Centre and Fachinformationszentrum Karlsruhe Access Structures service.

Data related to this article are available through heIDATA, the institutional research data repository of Heidelberg University, under https://doi.org/10.11588/data/24JDZD.

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**Conflict of Interest**

The authors declare no conflict of interest.

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