

Partial Fluorination Overcomes Herringbone Crystal Packing in Small Polycyclic Aromatics[†]

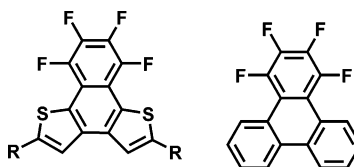
Don M. Cho, Sean R. Parkin, and Mark D. Watson*

Department of Chemistry, University of Kentucky, Lexington, Kentucky 40506-0055

mdwatson@uky.edu

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ABSTRACT



We report the synthesis and characterization of partially fluorinated condensed tetracyclic aromatic compounds. Typical edge-to-face/herringbone packing of nonfluorinated analogues is replaced here by columnar stacks with disk planes orthogonal to the columnar axes. Enhanced π -overlap results with overlaid electron-poor and -rich regions.

Polycyclic aromatic hydrocarbons tend toward edge-to-face (herringbone) solid-state packing motifs.¹ Face-to-face stacking is favored with increasing ratio of π -surface to circumference or with peripheral substituents. On the other hand, the “soft” interaction between arenes (Ar) and highly fluorinated arenes (ArF) leads to alternating face-to-face stacks. This principle appears nearly universally applicable based on the broad range of molecular frameworks with which it has been demonstrated.² Materials with covalently bound Ar and ArF units crystallize likewise with ArF portions stacked face-to-face with Ar portions. In addition to this supramolecular aspect, (partially) fluorinated π -systems show practical promise as active components in organic electronics, with facile electron injection and transport.³

The question addressed here is whether Ar–ArF interactions operate in partially fluorinated, *fused* π -systems, or will the effect be “smeared out” across the π -surface? Based on molecular electrostatic potential⁴ (MEP, Figure 1), one would

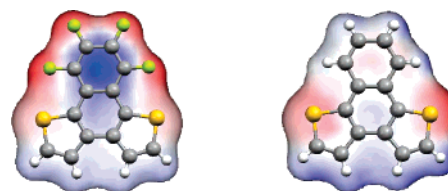


Figure 1. MEP maps of **2a** (left) and **2c** (right).⁴

predict face-to-face stacking in a head-to-tail fashion. As test cases, we chose naphthodithiophenes **2** and triphenylene **3** for which we present here synthesis, preliminary characterization, and single-crystal analysis.

Naphthodithiophenes **2** were prepared in two steps from readily available starting materials via modified published⁵

[†] Dedicated to Prof. Dr. Klaus Müllen.

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(4) Molecular Electrostatic Potential maps generated using ViewerLite.

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