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Tuning the redox profile of the 6,6'-biazulenic platform through functionalization along its molecular axis†

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The $E_{1/2}$ potential associated with reduction of the linearly-functionalized 6,6'-biazulenic scaffold is accurately correlated to the combined σ_p Hammett parameters of the substituents over >600 mV range. X-ray crystallographic analysis of the 2,2'dichloro-substituted derivative revealed unexpectedly short C-Cl bond distances, along with other metric changes, suggesting a nontrivial cycloheptafulvalene-like structural contribution.

Originally isolated from plant essential oils in the nineteenth century, azulenic compounds continue to fascinate scientists to this day. The correct structure of the polar azulenic scaffold, $C_{10}H_8$, ($\mu \approx 1.1 \text{ Debye}^1$) that features fused five- and sevenmembered sp²-carbon rings was first recognized by Pfau and Plattner in 1936.² The X-ray crystallographic confirmation of azulene's molecular structure was reported 20 years later.3,4 Azulenic building blocks are attractive for developing π -functional molecules and materials.⁵⁻⁷ Among the three possible linear biazulenes, 6,6'-biazulene has recently been emerging as a particularly effective oligoazulenic π -linker in the design of molecular electron reservoirs,8 on-chip microsupercapacitors,9 quasi-molecular rectifiers,10 organic and organometallic selfassembled monolayer (SAM) films, 8,11 and organic field-effect transistors. 12 Since the synthesis of 6,6'-biazulene in 1980, 13 access to its derivatives, especially those functionalized along its molecular axis, has been quite limited. 11,14,15

Given that azulene's highest occupied and lowest unoccupied molecular orbitals (HOMO/LUMO) have complementary orbital density distributions, their energies can be varied independently, to a first approximation. 16-18 This is accomplished by considering the position of attachment of a substituent to the azulenic core, as well as its electron withdrawing/ donating ability. 19 In this communication, we envisioned a similar approach for tailoring the energetics of the frontier molecular orbitals of the linear 6,6'-biazulenic framework. Indeed, Fig. 1 illustrates the orbital density complementarity between the HOMO and LUMO of 6,6'-biazulene.

Herein, we demonstrate that the redox potential $(E_{1/2})$ of the 6,6'-biazulenic scaffold is accurately predictable based on the Hammett σ_p parameters²⁰ of the substituents X and X' incorporated along its molecular axis (Fig. 2). To the best of our knowledge, this is the first study unveiling such quantitative redox correlations in the context of azulenic π -systems. In addition, we discuss how dichloro substitution exerts unexpected structural permutations within compound 3a (X = X' =Cl) in the solid state.

To facilitate synthetic accessibility of 2,2'-functionalized 6,6'-biazulenes, we selected the 1,1',3,3'-tetraethoxycarbonyl-6,6'-biazulene (1) as the "parent" platform. Rather than employing a toxic organotin reagent to assemble 1 via Stille cross-coupling,²¹ our synthesis of this compound involved simple deamination of the biazulenic derivative 2a (Scheme 1). On the other hand, subjecting 2a to Sandmeyer chlorination afforded dark purple, nearly black, crystalline

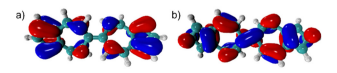


Fig. 1 Highest occupied (a) and lowest unoccupied (b) molecular orbitals (HOMO and LUMO) of 6,6'-biazulene (B3LYP functional and cc-pVDZ basis set).

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Fig. 2 2,2'-Fuctionalized 6,6'-biazulenes considered in this work.

Scheme 1 Syntheses of 1,1',3,3'-tetraethoxycarbonyl-6,6'-biazulene (1) and 2,2'-dichloro-1,1',3,3'-tetraethoxycarbonyl-6,6'-biazulene (3a).

2,2'-dichloro-1,1',3,3'-tetraethoxycarbonyl-6,6'-biazulene (3a) in an 85% yield (Scheme 1). Thus, 3a can be prepared from 2-amino-6-bromo-1,3-diethoxycarbonylazulene²² in two steps with an overall yield of *ca.* 80% (Scheme S1, ESI†).¹⁴ In addition to 3a, we synthesized its unsymmetric congener, 2-chloro-1,1',3,3'-tetraethoxycarbonyl-6,6'-biazulene (3b), *via* Sandmeyer chlorination of our recently reported 2-amino-1,1',3,3'-tetraethoxycarbonyl-6,6'-biazulene (2b) (Scheme 2).⁸ Both 3a and 3b constitute versatile precursors to a variety of 2-, and 2,2'-functionalized 6,6'-biazulenes.

Given that unsymmetrically functionalized **3b** is a structural hybrid of centrosymmetric **1** and **3a**, it is reasonable that its ¹H NMR spectrum (Fig. S7, ESI†) appears as a superposition of the ¹H patterns observed for the latter compounds (Fig. S1 and S2, ESI†).

We assigned all resonances in the ¹³C NMR spectrum of **3a** *via* synergistic consideration of its HSQC, HMBC, and 1,1-ADEQUATE²³⁻²⁶ 2-D maps (Fig. S3–S6, ESI†). Notably, this is the only unambiguously assigned ¹³C NMR profile of any 6,6′-biazulene reported in the literature to date.

Our TD-DFT calculations (Table S4, ESI†) corroborate that the lowest energy broad band centered around 500 nm in the electronic absorption spectrum of 3a (Fig. 3a) corresponds to the HOMO \rightarrow LUMO and HOMO-1 \rightarrow LUMO transitions. The broad nature of this band can be attributed, in part, to the range of accessible microstates pertaining to the

Scheme 2 Synthesis of 2-chloro-1,1',3,3'-tetraethoxycarbonyl-6,6'-biazulene (**3b**).

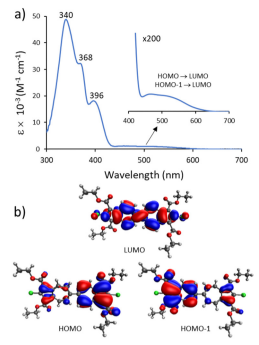


Fig. 3 (a) Electronic absorbance spectrum of ${\bf 3a}$ in CH $_2$ Cl $_2$ at 22 °C. (b) TD-DFT calculated molecular orbitals of ${\bf 3a}$ involved in the S $_0 \to S_1$ transition.

azulene–azulene interplanar angle within 3a in CH_2Cl_2 solution. While the LUMO of 3a involves the entire biazulenic core, 3a's nearly degenerate HOMO and HOMO-1 are each primarily localized on one of the azulenic units (Fig. 3b).

We have recently shown that 2,2'-functionalized 6,6'-biazulenes undergo reversible one-step, 2e⁻ reduction in CH₂Cl₂/ⁿ-Bu₄N⁺PF₆⁻ solutions under the potential inversion regime²⁷ regardless of whether they feature symmetric or asymmetric substitution along their molecular axis.8 After recognizing a qualitative relationship between the $E_{1/2}$ values associated with the 2e⁻ reduction of the 6,6'-biazulenic core and the electron donating/withdrawing characteristics of the 2,2'-substituents, we set out to identify a quantitative approach for tuning the 6,6'-biazulenic redox profile (Fig. S13–S16, ESI†). Remarkably, plotting the half-wave redox potential $(E_{1/2})$ against the combined σ_p Hammett parameters²⁰ of the substituents X and X' for the seven 6,6'-biazulenic derivatives listed in Fig. 2 revealed a nearly perfect linear correlation over >600 mV range (Fig. 4 and Table S5, ESI†). While attempts to correlate molecular redox potentials with the Hammett parameters of functional groups have been reported in the past for benzenoid organic/organometallic and ferrocene-based compounds, 28-31 this is the first example of invoking such a relationship for a cohort of azulenic derivatives. Notably, a similar trend $(E_{\rm p,c} \ \nu s. \ \sigma_{\rm p})$ holds for the family of 2-substituted 1,3-diethoxycarbonyl azulenes shown in Fig. 5 and Table S6 (ESI†). However, the 1e⁻ reduction of the latter compounds is invariably irreversible (Fig. S17-S20, ESI†).11 Coupling any pair of 2-substituted azulenes to form the corresponding 6,6'-biazulene leads to full reversibility of the reduction, with the biazulenic $E_{p,c}$

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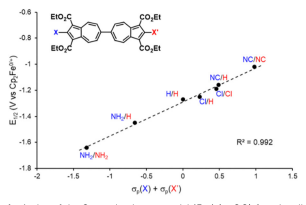


Fig. 4 A plot of the $2e^-$ reduction potential ($E_{1/2}$) for 2,2'-functionalized 6,6'-biazulenes vs. the combined σ_p Hammett parameters of the substituents X and X'.

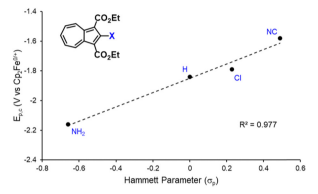


Fig. 5 A plot of the reduction potential $(E_{\rm p,c})$ for 2-functionalized 1,3-diethoxycarbonylazulenes vs. the $\sigma_{\rm p}$ Hammett parameter of the substituent X.

being 0.51 ± 0.04 V more positive compared to the average $E_{\rm p,c}$ of the monoazulenes (Tables S5 and S6, ESI†). This is a consequence of the substantially greater resonance stabilization energy of the 6,6'-biazulenic framework.

The molecular structure of **3a** is illustrated in Fig. 6. The two chemically identical, but crystallographically unique C-Cl bonds, C2-Cl1 and C2'-Cl2, within **3a** have the lengths of 1.674(2) Å and 1.682(2) Å, respectively. The longer C-Cl bond

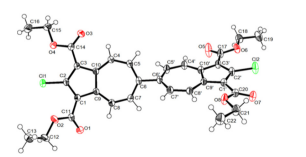


Fig. 6 Molecular structure of 3a (50% thermal ellipsoids). Selected interatomic distances (Å) and dihedral angle (deg): C2-Cl1 1.674(2), C2'-Cl2 1.682(2), C6-C6' 1.478(2), C9-C10 1.437(2), C9'-C10' 1.440(3), O2···Cl1 2.918(2), O4···Cl1 2.834(1), O6···Cl2 2.816(2), O7···Cl2 2.876(2), C5-C6-C6'-C5' 48.1(2).

Table 1 Comparison of C(2)-C(1) and C(9)-C(10) bond distances in X-ray structurally characterized 2-chloroazulenic derivatives

Compound	$d(C^2$ –Cl) Å	$d(C^9-C^{10})$ Å	CCDC identifier
EtO ₂ C 10 CO ₂ Et CI 2 CI EtO ₂ C CI ETO ₂	1.674(2) 1.682(2)	1.437(2) 1.440(3)	This work
EtO ₂ C Cl ² EtO ₂ C	1.712(2)	1.471(2)	HOMRAE
$\begin{array}{c} \text{EtO}_2\text{C} \\ \text{Cl} \xrightarrow{2} \\ \text{EtO}_2\text{C} \end{array}$	1.714(4)	1.470(6)	XIGPOU
EtO ₂ C Cl ² 9 EtO ₂ C	1.709(2)	1.470(3)	AYIDIX

Fig. 7 Minor zwitterionic resonance forms of 3a

belongs to the half of the molecule featuring the shortest Cl···O contact (Cl2···O6) of 2.816 Å. Both C-Cl bonds in **3a** are significantly shorter than any C-Cl bond in all structurally characterized chloroazulenes featuring Cl-substitution at the five-membered ring of the azulenic scaffold (Fig. S22 and Table S14, ESI†). In particular, the C-Cl bonds in **3a** are statistically shorter, per the 3- σ criterion, 32 than those in the X-ray structures of the other three 2-chloroazulenes known to date (Table 1). $^{33-35}$

The C–Cl bond contractions in **3a** may be rationalized by invoking the minor heptafulvalene-like zwitterionic resonance forms depicted in Fig. 7, which show C—Cl double-bond character. This argument mirrors the crystallographic analysis of the chlorodiphenylmethylium cation by Laub *et al.*³⁶ The short C–Cl bond distance of **1**.668(8) Å within this carbocation, which is statistically indistinguishable from those in **3a**, was attributed to chlorine backdonation exerting partial double-bond C—Cl⁺ character (Fig. S23, ESI†).³⁶ Unusually short C–Cl bonds, including various C–Cl multiple bonding scenarios, continue to attract both experimental and theoretical interest.^{37–39}

The C6–C6′ bond distance of 1.478(2) Å within 3a is markedly shorter than those in all other crystallographically characterized 6,6′-biazulenes (Table 2).^{8,11,12,40} In addition, as summarized in Table 2, the C–C bonds at the fusion of the five- and seven-membered rings in 3a are substantially contracted as well. Both of these observations are consistent with a nontrivial heptafulvalene-like contribution to the structure of the biazulenic scaffold of 3a.⁴¹ Moreover, the azulene–azulene interplanar angles in dichloro 3a and diisocyano 4a, which share the same 1,1′,3′3′-tetraethoxycarbonyl-6,6′-biazulenic core, are 48.1° and 66.9°, ¹¹ respectively, although such comparison should be viewed *cum grano salis* as this torsion parameter is undoubtedly sensitive to crystal packing.

In summary, we demonstrated that the redox profile of the 6,6'-biazulenic scaffold functionalized along its molecular axis is quantitatively tuneable within a wide window of potentials.

Table 2 Comparison of C(6)-C(6') and C(9)-C(10) bond distances in Xray structurally characterized 6,6'-biazulenes

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Compound	$d(C^6-C^6)$, Å	$d(C^9-C^{10})$ Å	CCDC identifier
$\begin{array}{c} EtO_2C \\ Cl \\ EtO_2C \\ \end{array} \xrightarrow{\begin{array}{c} G \\ G \\ \end{smallmatrix}} \xrightarrow{\begin{array}{c} G \\ G \\ \end{smallmatrix}} \xrightarrow{\begin{array}{c} CO_2Et \\ G \\ \end{smallmatrix}} \\ CO_2Et \\ \end{array}$	1.478(2)	1.437(2) 1.440(3)	This work
10 6: 10 9·	1.498(2)	1.485(2) 1.488(2)	NAZROZ
EtO ₂ C 10 6 CO ₂ Et C=N 9 6 CO ₂ Et EtO ₂ C CO ₂ Et	1.512(4)	1.475(3)	OSIGED
$(OC)_{5}Cr-C\equiv N \xrightarrow{g} \begin{matrix} CO_{2}Et \\ g & g \end{matrix}$	1.497(3) 1.499(3)	1.465(3)-1.476(3)	$HITRIP^a$

^a Two crystallographically independent molecules in the asymmetric

This was accomplished by considering, for the first time in the context of azulenic systems, the well documented σ_n Hammett parameters²⁰ reflecting the net electronic influence of the substituents. We anticipate that the facile access to the crystallographically unusual dichloro derivative 3a offers not only new opportunities to engage the 6,6' architecture as an attractive molecular template in the realm of organic charge transfer and/ or conductive materials, but also to explore unorthodox approaches for pursuing C-Cl multiple bonding.

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Conflicts of interest

There are no conflicts to declare.

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