phenyl and 50% polysiloxane) 10 m × 0.53 mm wide bore column. UV spectra were recorded on either a Hewlett-Packard 8415A single-beam photodiode array or a Perkin-Elmer 452 double-beam spectrophotometer. Mass spectra were recorded on a Kratos MS-30 double-focusing dualbeam spectrometer at 20 eV ionizing voltage.

Materials. 1a and 1b were synthesized according to a previously reported procedure via a Grignard reaction starting with either trans-4pentyl(4-cyanophenyl)cyclohexane (Merck Licristal/ZLI-114) for 1a or trans-4-heptyl(4-cyanophenyl)cyclohexane (Merck Licristal/ZLI-115) for 1b and 1-bromobutane (Aldrich, 99%). Both ketones were more than 99% pure as determined by GLPC after recrystallizations from ethanol.

trans- and cis-3a and 2a were obtained from a solution of 300 mg of 1a in benzene (Baker reagent) that had been purged with nitrogen for 5 min and irradiated to nearly 100% conversion. The photoproducts were separated on a silica (Baker, 60-200 mesh) column with 97/3 (v/v) hexane-ethyl acetate (Aldrich reagents) as mobile phase. trans-3a and 2a were characterized by their ¹H and ¹³C NMR spectra, mass spectra, and co-injection with irradiated samples of 1a whose photoproduct retention times had been previously determined. cis-3a was identified by its proton NMR spectrum. Both diastereomeric cyclobutanols have characteristic cyclobutyl proton resonances in the 1.8-2.8-ppm region.³⁰ The ¹H spectrum of the trans isomer differs from that of the cis in the location of the doublet resonance corresponding to the protons of the methyl group on the cyclobutyl ring. For the cis isomer, the doublet resonated at 0.62 ppm (J = 6.9 Hz) while for the trans isomer it was found at 1.1 ppm (J = 6.9 Hz). The ¹³C spectrum of the trans isomer showed a peak at 78.4 ppm characteristic of tertiary carbons bearing a hydroxyl group and its mass spectrum exhibited an m/e peak at 314, as well as M - 18 and M - 28 peaks characteristic of 1-phenylcyclobutanols.³⁰ On the silica chromatographic column, the cis isomer eluted after the trans

2a has a melting point of 62.0-65.0 °C. NMR spectrum: 7.88 (d, J = 8.25 Hz, 2 H, aromatic), 7.25 (d, 8.25 Hz, 2 H, aromatic), 2.58 (s,

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3 H, methyl α to carbonyl), 2.56 (m, 1 H, cyclohexyl α to phenyl group), 1.89 (d, 4-5 H, cyclohexyl), 1.2-1.1 (m, alkyl and cyclohexyl), 0.9 (t, 3 H, terminal methyl). The mass spectrum of 2a exhibits an m/e peat at 272, its expected molecular ion.

2b was synthesized according to the same procedure used to obtain 1, except methyl iodide (Fisher reagent) instead of 1-bromobutane was employed to form the Grignard reagent. 2b exhibited mp 62.8-64.8 °C and a proton NMR spectrum almost identical to that of 2a. The mass spectrum of 2b displayed an m/e peak at 300, its expected molecular ion.

Treatment of either homologue of 1 with deuterium oxide (MSD Isotopes, 99% d) and Na₂CO₃ in monoglyme³¹ (Aldrich, anhydrous, 99%) yielded ketone with >90% deuterium at the position α to the carbonyl group as determined from comparisons of the ¹H integrals of the α methylene resonances (2.9 ppm) from treated and untreated 1.

Irradiation Procedures. Irradiations were performed with a Hanovia 450 W medium pressure Hg arc lamp. Temperature control was achieved using either a Haake HK-2 or a Forma Scientific circulating water bath. Prior to photolysis, samples were purged with nitrogen for ca. 5 min in their isotropic phases, transferred to $(0.8-1.1) \times 100$ mm capillary tubes, and flame-sealed. Irradiations were performed through water, a Pyrex glass filter, and a Corning 0-51 cutoff (>30% at 380 nm) filter. Irradiations of three samples at each temperature were conducted to less than 10% ketone conversion. The contents of the capillaries were then dissolved in hexane (Aldrich, HPLC grade) and analyzed by gas chromatography. The E/C ratios, uncorrected for detector response, represent an average of triplicate analyses of each of the three samples at each temperature.

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Mixed-Valence, Conjugated Quinone and Imide Anion Radicals. An ESR Investigation

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Abstract: Anion radicals of linear polyacene diquinones and diimides were produced electrochemically and studied by ESR. The odd electron of substituted 1,4,8,11-pentacenetetrone anion radicals is localized (in a naphthoquinoid unit) and hops from one quinone to the other. The hopping rate was measured. The anion radicals of similarly sized anthracenetetracarboxylic acid 2,3:6,7-diimides have a delocalized odd electron, and there is relatively high electron density on the bridge. The results are compared to other unconjugated two-electrophore anion radicals.

Recent studies in our laboratories have been directed toward the synthesis and properties of linear, rigid polyacene quinones and imides.¹⁻⁵ These molecules were designed to have long delocalization lengths and more than one quinone or imide electrophore. These structural features led us to examine the optical and electrical properties of the anion radicals, which displayed unusual properties, indeed, including near-infrared

(near-IR) absorption bands at wavelengths as long as 2000 nm¹⁻³ and electrical conductivities as high as 1 S cm^{-1,4} It was proposed that a central structural issue was the delocalization or localization of the unpaired electron. Based on cyclic voltammetry (CV), near-IR, IR, and ESR spectra, and molecular orbital (MO) calculations, anion radicals containing one aryl ring between the electrophore groups, e.g., 1⁻ and 4⁻, were classified as delocalized.^{1,2} Anion radicals having naphthalene or anthracene bridges connecting quinone electrophores, i.e., 2⁻ and 3a⁻, gave very different optical spectra, and based upon MO calculations, it was suggested that the unpaired electron in these anion radicals was localized on one quinone unit at any instant.³ As such, the anion radicals could be considered mixed-valence species.

 π -Electron localization was unexpected because the two electrophores of 2^- and $3a^-$ are formally part of a planar, conjugated π -system. In the present study ESR was used to examine this

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speculative hypothesis and provide definitive evidence for localization in the quinone anion radicals of the type 3⁻. In contrast, the very similar imide anion radicals of type 5⁻ are shown to be delocalized. The rates of intramolecular electron hopping are reported for 3b- and 3c- and compared to the rates for the unconjugated triptycene derivative 6^- and other anion radicals.



Ar = 4-tert-butylphenyl Ar' = 2,5-di-tertbutylphenyl

Results and Discussion

Synthesis. Anthracene-bridged dielectrophores were chosen for study, in part because they were synthetically accessible. The specific examples were chosen to be more soluble and have better anion-radical stability than previously reported for 3a. Compounds 3b,c were prepared from the precursors 7b,c developed by Hart⁶ and Schlüter,⁷ respectively. The process can be understood in





Compound 3b had only limited solubility in common organic solvents, but the dihexyl-substituted 3c dissolved readily in CH2Cl2,

Table I. Reduction Potentials^a

compd	<i>–E°′</i> (V, SCE)		
1 ^b	0.26	0.74	
2 ^b	0.50	0.75	
3a ^b	0.72	0.84	
3b ^c	(0.79)	(0.94)	
3c ^d	0.83	0.98	
4 ^e	0.71	1.43	
5a"	1.07	1.53	
5b ^e	1.03	1.52	
5c*	1.01	1.45	
6	1.23	1.39	
10 ^b	0.30	0.53	

^a Measured in DMF, 0.1 M nBu_4NBF_4 , scan rate 150 mV/s, glassy carbon working electrode, SCE reference. $E^{\circ \prime} = (E_p^{a} + E_p^{\circ})/2$. The values are reproducible within ±10 mV. ^bReference 3. ^cRecorded in 0.1 M LiClO₄/DMF; values are cathodic peak potentials. ^dRecorded in 0.1 M nBu_4NBF_4/CH_2Cl_2 , scan rate 50 mV/s. ΔE_p (anodic-cathodic) = 80 mV. 'Reference 1.

CHCl₃, or DMF. The orange/red color of 3c in CH₂Cl₂ or CHCl₃ solution dissipated readily in the presence of room light and air to give pale yellow solutions. The product isolated from those solutions was 9, where oxygen is added across the center ring to



9 R = 1-hexyl

give the endo-peroxide. This reaction, similar to that studied for 5c,¹ is a photooxidation, and thus, 3c was handled with minimum exposure to light. 3b was stable in the presence of room light and air.

6 was synthesized by addition of benzyne to the center ring of dimethoxyanthracene diimide 5c.



Cyclic Voltammetry. In general, the voltammograms were recorded in DMF containing 0.1 M Bu₄NBF₄. The working electrode was a glassy carbon disk, and ferrocene was added as an internal standard for the saturated calomel electrode (SCE) reference. The apparent E° was taken as the midpoint between the anodic and cathodic peak potentials. The anodic-cathodic peak separations were 60 ± 5 mV except for 3c which gave 80 mV. The relative anodic/cathodic peak currents were approximately equal, and the peak potentials were independent of scan rate, indicating that the E° values (Table I) represent a thermodynamic equilibrium.

It has been previously reported that 3a reduces in DMF with apparent $E^{\circ} = -0.72$ and -0.84 V.³ Because this compound was so insoluble, it was difficult to get good cyclic voltammograms. Compound 3b was similar. One of the more simple voltammograms was determined using DMF, 0.1 M LiClO₄. It showed two poorly resolved reduction peaks at $E^{\circ} = -0.79$ and -0.94 V. In an attempt to begin with soluble material, bulk electrolysis generated solutions of the anion radical 3b⁻ which was then studied (Figure 1). Complex surface electrochemistry is revealed here and in voltammograms obtained in other solvents and with other electrode materials, which is related to the adsorption of the neutral, but it was not completely unraveled and it will not be discussed.

The voltammogram of 3c is shown in Figure 2. The hexyl groups aid in solubilizing the diquinone and allowed for its examination in CH₂Cl₂, 0.1 M nBu₄NBF₄. Two closely spaced, but distinct, redox couples were present at $E^{\circ} = -0.83$ and -0.98 V.

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E (Volts)

Figure 1. CV of $3b^-$ in DMF, 0.1 M Bu₄NBF₄ (sweep rate 1.0 V s⁻¹, starting potential 0.0 V).



E (Volts) Figure 2. CV of 3c in CH_2Cl_2 , 0.1 M Bu_4NBF_4 (sweep rate 200 mV s⁻¹).

A third, kinetically irreversible peak was present at -1.5 V. No change in the voltammogram was observed during repetitive cycling between -0.3 and -1.2 V.

The reduction potentials of the diquinone series make for an interesting comparison. As expected from the electrochemistry of monoquinones,^{1,2,8} the anthracene diquinone 1 was the easiest to reduce followed by the tetracene diquinone 2^3 and the pentacene diquinones. A similar trend was observed for the second reduction processes. The ΔE° values between the first and second reduction couples were 0.48 V for 1, 0.25 V for 2, and 0.15 V for 3b and 3c. One explanation for the smaller ΔE° for longer molecules is that repulsions in the dianion are diminished as the aryl bridge is extended.

Likewise for the diimides, the first reduction of the pyromellitic diimide derivative 4 (-0.71 V) occurs at a more positive potential than for the anthracene diimide derivatives 5a (-1.07 V), 5b (-1.03 V), and 5c (-1.01 V). However, the ΔE° values for the 5 series were large (0.41-0.49 V) in comparison to the pentacene diquinones. This may imply greater Coulombic repulsions in the diimide dianions than in the diquinone dianions.

CV on the triptycene bis(phthalimide) 6 showed it to be similar to that of the analogous diquinone 10.9 The first reduction of





6 occurred at -1.23 V, midway between the reduction of anthracene diimide **5c** (-1.01 V) and N-phenylphthalmide (-1.36 V). The second reduction of **6** took place at -1.39 V, only 0.16

Table II. ESR Coupling Constants^a

compd	solvent	a (G)
3b-	DMF	1.34 (12 H)
3c⁻	DMF	1.34 (12 H)
3c-	DMF (195 K)	2.89 (6 H)
3c-	CH_2Cl_2	0.20 (4 H), 1.34 (12 H)
3c-	CH ₂ Cl ₂ (185 K)	0.64 (2 H), 2.84 (6 H)
4-	DMF	0.65 (2 H), 1.17 (2 N)
5a-	DMF	4.24 (2 H), 3.07 (4 H)
		0.76 (2 N)
		$4.12(2 H)^{b} 2.31(4 H)^{b}$
5b⁻	DMF	3.19 (4 H), 0.75 (2 N)
5c-	DMF	3.19 (4 H), 0.68 (2 N)
6-	DMF	1.16 (2 N)
6⁻	CH ₂ Cl ₂	1.19 (2 N)

^aSpectra recorded in 0.1 M Bu₄BF₄ solutions at T = 294 K unless otherwise noted. Only coupling constants for fully resolved peaks are listed. ^bCalculated values from the McConnell equation, $a_{\rm H} = Q\rho$, where Q = 23 G. The spin densities, ρ , were determined from a PPP-MO calculation using the b_{2u} SOMO coefficients (symmetry determined with the molecule positioned in the x-z plane).

V more negative of the first reduction. This E° is slightly positive of the second E° for 5c and considerably positive of that for *N*-phenylphthalimide (-2.22 V). It seems clear that the two electrophores of 6 each take up one electron and the Coulombic repulsions are small.

The most revealing comparison is between the polyacene and triptycene analogues. The differences between the first and second E° values for compounds 3 and 10 are quite similar. Since 10 has been shown to be localized,⁹ compounds 3 may be similarly localized. In contrast, comparison of compounds 5 and 6 shows that the first E° values for 5 are more positive and the ΔE° values for compounds 5 are larger than that for 6. This can be understood if 6 forms localized ions, but 5 forms delocalized ions (see below). Delocalization makes it easier to add one electron forming 5⁻, but since the second electron is in a similar space, the electron repulsion in the dianion 5^{2-} is larger.

ESR Spectra. Potentiostatic electrochemical reductions were performed in situ in the spectrometer cavity because some of the anion radicals were of limited stability or had closely spaced couples. It was found that resolution could be improved if the Pt electrode was pulled up out of the cavity after generation of the anion radical. The initial concentration of the electroactive species ranged from 0.1 and 1.0 mM, the solvent was DMF or CH_2Cl_2 , and the electrolyte was 0.05–0.50 M Bu₄NBF₄.

Recall that the odd electron of 1^{-} is delocalized.³ It has been contrastingly demonstrated using variable-temperature ESR studies that the odd electron of the unconjugated 10^{-} is localized.⁹ A main objective of the present investigation was to examine the hypothesis that the odd electron of 3^{-} is also localized.

Electron Localization and Hopping in 3⁻. At room temperature in DMF, 0.1 M Bu₄NBF₄, 3b⁻ and 3c⁻ gave identical 13-line ESR spectra with $a_{\rm H}(12 \text{ H}) = 1.34 \text{ G}$ (Table II). In CH₂Cl₂, 0.1 M Bu₄NBF₄, the 3c⁻ spectrum was better resolved, revealing additional splitting on each of the 13 lines; $a_{\rm H}(4 \text{ H}) = 0.20 \text{ G}$. The couplings are assigned to the 12 methyl hydrogens and the 4 equivalent bridge hydrogens. Thus, at room temperature the odd electron is distributed over both electrophores.

The low-temperature spectra of $3b^-$ and $3c^-$ unequivocally demonstrate that the odd electron is localized on one quinone unit. Thus, as the temperature is lowered, alternant lines broaden, and at the lowest temperatures, spectra with half the number of proton couplings are observed.¹⁰ Using $3c^-$ in DMF, the 13-line 294 K spectrum changed to 7 lines at 210 K. In CH₂Cl₂ the lowtemperature spectrum showed each of the 7 lines split into triplet (Figure 3). The less soluble $3b^-$ could only be studied in DMF. In that solvent, the spectra were not completely isotropic and the

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Figure 3. Experimental and simulated ESR spectra of $3c^{-}$ in CH₂Cl₂, 0.05 M Bu₄NBF₄, at various temperatures.

seven line spectrum was only partially resolved at low temperature (Figure 4). It is noted that at these low temperatures the solvents are rather viscous and anisotropy is quite possible. Also important is the decrease in polarity of the solvent at low temperature. This can affect $a_{\rm H}$ values and can lead to increased ion pairing or aggregation of the sparingly soluble **3b**⁻.

The results lead to a model for 3^{-} that has a double minimum on the potential energy surface so that the electron is localized on one quinone or the other at any instant. The low-temperature $a_{\rm H}(6 \text{ H}) = 2.84$ and $a_{\rm H}(2 \text{ H}) = 0.64 \text{ G}$ show that each of the localized orbitals of 3^{-} is similar in electron distribution to the half-filled orbital of 2,3-dimethyl-1,4-naphthoquinone anion radical, which has $a_{\rm H}(6 \text{ H}) = 2.51$ and $a_{\rm H}(2 \text{ H}) = 0.32 \text{ G}$ for the appropriate hydrogens.¹¹ Both 3^{-} and dimethylnaphthoquinone



Figure 4. ESR spectra of $3b^-$ in DMF, 0.1 M Bu₄NBF₄, at 300 and 220 K.

Table III	Kinetic	Data ^a
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anion radical	conditions	ΔG^{*}_{298} (kcal/mol)	ΔH^* (kcal/mol)	ΔS^* (eu)
3b-	0.1 M Bu ₄ NBF ₄ /DMF	6.6		
3c⁻	$0.1 \text{ M Bu}_4 \text{NBF}_4 / \text{DMF}$ $(230 \rightarrow 200 \text{ K})$	5.2	4.2	-3
3c-	0.05 M Bu ₄ NBF ₄ /CH ₂ Cl ₂	5.4	4.0	-5
3c⁻	$0.50 \text{ M } \text{Bu}_4\text{NBF}_4/\text{CH}_2\text{Cl}_2$ $(273 \rightarrow 210 \text{ K})$	5.3	4.6	-3
6-	$0.1 M Bu_4 NBF_4 / DMF$ $(340 \rightarrow 210 K)$	6.2	4.1	-7
6-	$0.05 \text{ M } \text{Bu}_4\text{NBF}_4/\text{CH}_2\text{Cl}_2$ $(294 \rightarrow 230 \text{ K})$	6.1	4.5	-5
10 ^{- b}	$CH_3CN (298 \rightarrow 248 \text{ K})$	8.3	6.2	-7

^a From variable-temperature ESR measurements. The error in ΔG^* and ΔH^* is estimated to be ±20% and in ΔS^* ±40%. ^bReference 9.

anion radicals have equivalent methyl groups, indicating that on the ESR time scale the electron is delocalized over both carbonyl groups of one quinone. In this regard, consider an alternative structure for 3^- which involves a "top-to-bottom" instead of "side-to-side" exchange. The localized SOMO would encompass not the 1,4-carbonyl groups but the 1,11-carbonyl groups. The electron would then jump to the 4,8-dicarbonyl conjugated system. Since under these conditions neither monosemiquinones nor $1^$ exhibits localization, it seems unlikely that this possibility is realized. Clearly the distance between the quinone groups (and perhaps the polarizability of the long bridge) is important to achieving localization in the absence of strong ion pairing.

The dynamics of the exchange processes were simulated (see Figure 3 for an example), and rate constants for electron hopping were computed at several temperatures. A two-site, intramolecular exchange with no change in structure or ion pairing as the temperature changed was assumed. These data were then used for Eyring plots, leading to the results shown in Table III. The activation parameters for $3c^{-}$ were the same within experimental error in either DMF or CH₂Cl₂ solvent and were unchanged when measured using the electrode in the cavity or out.¹² The simu-

⁽¹¹⁾ Rieke, R. D.; Rich, W. E. J. Am. Chem. Soc. 1970, 92, 7349. (12) Using $3c^{-}$ in CH₂Cl₂, cooling to 185 K, and then reheating did not give the original 294 K spectrum. Cooling to 210 K and reheating did give the 294 K spectrum. We speculate that precipitation occurs at 185 K.

lations for $3b^-$ led to a linear Eyring plot over the range 300-230 K, but because the low-temperature spectra were distorted and could not be simulated, there was considerable experimental uncertainty in the analysis and only ΔG^* is reported.

The absence of an appreciable solvent effect on the rate of electron exchange for $3c^{-}$ made it of special interest to study the possible effects of ion pairing by changing the bulky Bu_4N^+ to Li⁺. Using $3b^-$ in DMF, 0.1 M LiClO₄, a distorted 7-line spectrum was observed at room temperature, which closely resembled the low-temperature spectrum found when Bu_4N^+ was used. The use of HMPA solvent (an effective Li⁺ ligand) led to the 13-line spectrum observed with Bu_4N^+ at high temperature. It is thus clear that Li⁺ ion pairing leads to strong localization at room temperature and that HMPA solvates the cation and increases the rate of electron exchange.

To examine further cation effects, $3c^{-}$ was studied using 0.05 and 0.5 M Bu₄NBF₄ in CH₂Cl₂. The two Bu₄N⁺ concentrations gave the same rate and activation parameters within experimental error. If the anion radical is not ion paired with the bulky Bu₄N⁺ and the electron just jumps from end to end, independence of the rate on cation concentration is expected. If the anion radical is ion paired and there is an *intramolecular* jump of the Bu₄N⁺ along with the electron, the experimental result is also rational. Ruled out is the mechanism in which an ion-paired species, $3c^{-}$, Bu₄N⁺, dissociates to free ions; the electron jumps in the rate-limiting step, and then a new ion pair forms on the other end. This process would be slower at higher Bu₄N⁺ concentration.

Consider the situation more carefully assuming that free ion radicals are present. It is expected that the quinone unit which holds the odd electron in an antibonding orbital would have an expanded molecular framework compared to its counterpart on the other end. Solvent organization should also be different for the ionic and neutral ends of the mixed-valence ion. Electron transfer then involves changes in the nuclear positions of the radical anion and the solvent, and this accounts for the activation energy.

The solvent effect on the reaction rate for $3c^{-}$ can be modeled in a simple way using an equation suggested by Marcus.¹³

$$4\Delta G_{s}^{*} = \lambda_{s} = e\left(\frac{1}{2r_{a}} + \frac{1}{2r_{b}} - \frac{1}{r_{ab}}\right)\left(\frac{1}{D_{op}} - \frac{1}{D}\right)$$

Here, λ_s is the solvent reorganization energy for a thermoneutral reaction, r_a and r_b are the radii of the two reaction centers, r_{ab} is the distance between the centers, D is the dielectric constant, and D_{op} is the refractive index squared. Assuming that $r_a = r_b = 5$ Å and $r_{ab} = 7.5$ Å (center to center) leads to a calculated $\Delta G_s^* = 2.6$ kcal mol⁻¹ in DMF and $\Delta G_s^* = 2.2$ kcal mol⁻¹ in CH₂Cl₂. These values are smaller in magnitude than the experimental ΔG^* (which includes internal reorganizations), and the difference between them is less than the experimental error of the ESR measurements. Therefore, the experimental result is not in conflict with the expectations of theory.

It is of interest that the activation parameters for 3^- are quite similar to those previously reported for the triptycene diquinone anion radical 10^- (Table III)⁹ and measured for the triptycene diimide anion radical 6^- (see below). This can be interpreted to mean that all three rates are determined by the similar changes in nuclear geometry of the ions and the solvent and a similar distance for electron transfer. The small ΔS^+ values are quite appropriate for a reaction with no change in charge.

Anion radicals which are somewhat less similar, but which have short aliphatic bridges between electrophores, are 11^- and 12^- .



In DMF with Bu_4N^+ counterions, 11⁻ has $k \sim 5 \times 10^8 \text{ s}^{-1}$ at 250



Figure 5. ESR spectrum of 5c⁻ in DMF, 0.1 M Bu₄NBF₄, at 298 K.

K.¹⁴ Since the naphthyl and biphenyl groups have about the same E° , 12⁻ also allows an appropriate comparison. Here, $k = 4 \times 10^9 \text{ s}^{-1}$ at room temperature for γ -radiation-generated ions in tetrahydrofuran.¹⁵ Clearly, the exchange rates for anion radicals with short aliphatic bridges are similar to those for 3⁻ (4.2 × 10⁸ s⁻¹ at 273 K). This demonstrates that, for anion radical exchange across short bridges, sp² carbons are similar to sp³ carbons.

Electron Delocalization in 5⁻. Although 3 and 5 differ only in the replacement of the C=C groups with N atoms, the ESR spectra of the 5⁻ species in DMF revealed striking differences with the spectra of 3⁻. The 5a⁻ spectrum was simulated using $a_{\rm H}(4$ H) = 3.07, $a_{\rm H}(2 \text{ H}) = 4.24$, and $a_{\rm N}(2 \text{ N}) = 0.76 \text{ G}$. The simulation and assignment of the 4.24-G coupling to the center pair of hydrogens was confirmed from the spectra of 5b⁻ and 5c⁻ which showed $a_{\rm H}(4 \text{ H}) = 3.19$ and $a_{\rm N}(2 \text{ N}) \sim 0.7 \text{ G}$ (Figure 5; Table II). Clearly, the electron density on the bridge is high and much higher than that found for 3⁻. Indeed, the 5a⁻ $a_{\rm H}$ values are quite comparable to those for the anthracene anion radical, ¹⁶ for which $a_{\rm H} = 5.38 \text{ G}$ for the 9,10-hydrogens and 2.66 G for the 1,4,5,8hydrogens. In a similar light the $a_{\rm N}$ value for 5⁻ is smaller than that for 4⁻.

High electron density on the bridge makes it unlikely that the electron would be localized on the imide moiety, and the low-temperature spectra confirmed this expectation. Only slight broadening of *all* the lines was evidenced at 210 K. Thus, the ESR data demonstrate that *the odd electron of* 5^- *is delocalized*. This is in agreement with previously reported optical data.¹

We have previously discussed the difference in electronic structure of diquinone and diimide anion radicals.¹ Stated concisely, the imide moieties are poorer electron acceptors than quinone units, and therefore, the odd electron will tend to spend more time on the anthracene bridge when it is flanked by imides. This qualitative view is supported by π -MO-PPP calculations on 5. These calculations show that there are a number of low-lying orbitals for 5⁻. Using the calculated SOMO coefficients and the McConnell equation, $a_{\rm H}$ values were calculated for the various hydrogens of 5⁻. As shown in Table II, there was satisfactory agreement between experiment and calculation. Agreement was not achieved for 5⁻ when a Huckel MO calculation was employed, because a different orbital ordering gave a different SOMO.

Electron Localization and Hopping in 6⁻. The ESR spectrum of 6⁻ taken at 294 K in CH₂Cl₂ (Figure 6) displayed a pentet, $a_N = 1.19$ G, as expected for fast exchange between two equivalent nitrogens. At 210 K this spectrum collapsed to give three broad peaks of equal intensity (Figure 6) with $a_N = 2.34$ G indicative of electron localization. The spectra at intermediate temperatures were simulated to estimate rate constants, and an Eyring plot gave the activation parameters shown in Table III. These values were quite similar to those determined for 3⁻ and 10⁻. Thus, the reorganization energies for electron transfer between imides is

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Figure 6. ESR spectra of 6^- in CH_2Cl_2 , 0.1 M Bu_4NBF_4 , at 294 and 210 K.

quite similar to that for electron transfer between quinones. 6^- was also examined in DMF and gave activation parameters similar to those in CH₂Cl₂. At 340 K in DMF, the spectrum is a 1:2:3:2:1 pentet.

Conclusions. The data make it quite clear that the odd electron on 3^- is localized in an orbital which resembles the SOMO of the naphthoquinone anion radical. It is suggested that localization occurs because these large π -systems are quite polarizable.³ If the delocalized SOMO has a high electron density on the end electrophores, small distortions of the nuclear framework can make a localized structure more stable than a delocalized one. This localization process will also be favored by solvent organization and ion pairing of cations with the anionic end of the radical anion. The delocalization of 5^- results because its delocalized SOMO has high electron density on the bridge.

It is revealing to compare the properties of 3^- and 10^- . The small separation between the first and second reduction potentials reveals a similar isolation of the two electrophores. The electron densities on the quinone units are similarly high, and the rate of electron exchange is quite similar. The similar rate arises because the change in internal modes and solvent modes required for electron transfer is essentially the same for the two ions.

Electron localization in conjugated π -systems is an important problem for materials chemistry and especially for conducting polymers. Most of these polymers are composed of partially oxidized or reduced linear π -systems, and electron localization along these linear systems can be an important factor for conductivity. The individual conducting units, polarons or bipolarons, are solid-state analogues of the ion radicals of interest here. Because these polymers are charged and have counterions, and most contain heteroatoms along the π -system, the possibility of localization is quite real. We have shown how easy it is to localize charge even in small molecules and how small structural changes can convert delocalized to localized structures.

Experimental Section

¹H NMR and ¹³C NMR were measured using IBM-NR-300-AF (300 MHz) or IBM-IR-200-AF (200 MHz) FT-NMR spectrometers. Chemical shifts are reported in δ units with respect to the CDCl₃ protonic impurity peak at 7.257 ppm. Infrared spectra were recorded on a Perkin-Elmer 1600 Fourier transform instrument. Fast atom bombardment (FAB) mass spectra were recorded on a high-resolution VG-7070E-HF instrument. Cyclic voltammograms were recorded on a BAS 100 electrochemical analyzer. Bulk electrolyses were performed with a Princeton

Applied Research (Model 173) potentiostat. UV-vis absorption spectra were recorded on a Shimadzu UV-160.

Materials. Anhydrous DMF and CH_2Cl_2 (Aldrich Chemical Co.) were used fresh or after temporary storage over 4-Å sieves. Tetrabutylammonium tetrafluoroborate was recrystallized from H₂O/MeOH, 3:1, and dried in vacuo. 2,3-Dimethyl-1,4-benzoquinone was prepared from 2,3-dimethylphenol.¹⁷ The syntheses of 4, 5a, 5b, and 5c were described previously.

6,13-Dihexyl-2,3,9,10-tetramethyl-1,4,8,11-pentacenetetrone (3c). 2,3-Dimethyl-1,4-benzoquinone (0.59 g, 4.36 mmol) was dissolved in Decalin (4 mL) in a 50-mL round-bottom flask equipped with an addition funnel, reflux condenser, N_2 inlet, and a magnetic stirring bar. The addition funnel was charged with the benzodifuran precursor 9c (0.50 g, 0.436 mmol) suspended in Decalin (10 mL). The suspension of 9c was added to the refluxing quinone solution over a 25-min period. The color changed from yellow to red-brown. After the mixture was refluxed for 30 min, the gray-white precipitate that formed upon cooling was removed by filtration. The filtrate was collected, and the Decalin and unreacted 2,3-dimethyl-1,4-benzoquinone were removed by vacuum distillation. The remaining brown solid was purified by chromatography (silica gel prep plate, CH_2Cl_2 eluent). The material remaining near the origin was collected giving a light brown residue. In the dark, 10 mL of concentrated H₂SO₄ was added to the brown residue. The flask was capped, and stirring was continued for 2 h. The product mixture was poured over crushed ice upon which a dark precipitate formed. The precipitate was collected by filtration, washed with H₂O, and subsequently dried in a vacuum desiccator. A 7-mg portion of the crude solid (121 mg total) was purified by chromatography (silica gel prep plate; CH_2Cl_2 eluent), and the red band $(R_f = 0.27)$ was collected, giving 1.4 mg of the desired product (10% yield). Small amounts of the product were purified in this manner prior to ESR analysis. Initial purification of the crude solid could also be accomplished by sublimation (210 $^{\circ}$ C, 1 torr). ¹H NMR (CDCl₃): δ 9.14 (s, 4 H), 3.76 (m, 4 H), 2.28 (s, 12 H), 1.83 (m, 4 H), 1.63 (m, 4 H), 1.38 (m, 8 H), 0.92 (t, J = 7 Hz, 6 H). ¹³C NMR $(CDCl_3): \delta$ 183.94, 145.93, 142.13, 131.52, 128.20, 127.19, 32.59, 31.66, 29.80, 28.86, 22.66, 14.10, 13.41. IR (KBr, cm⁻¹): 2956, 2924, 2858, 1664, 1648, 1610, 1454, 1369, 1283, 1032, 718. HR-FAB-MS (H₂SO₄ matrix) for $C_{28}H_{43}O_4$ (M + H⁺): calcd 563.3161; found, 563.3138. UV-vis (CH_2Cl_2) (λ , nm (log ϵ): 526 (4.19), 496 (4.09), 457 (3.87), 428 (3.66), 342 (4.80), 2.61 (4.83).

2,3,9,10-Tetramethyl-1,4,8,11-pentacenetetrone (3b). 2,3-Dimethyl-1,4-benzoquinone (0.57 g, 4.19 mmol) and the benzodifuran precursor 9b (0.50 g, 0.511 mmol) were added to Decalin (10 mL) in a roundbottom flask equipped with a reflux condenser, N_2 inlet, and magnetic stirring bar. The reaction mixture was heated to 190 °C and allowed to reflux for 3 h. The heating source was removed, and the product mixture was allowed to stand overnight. The precipitate was collected by filtration and washed with hexane (25 mL), CCl₄ (25 mL), CHCl₃ (15 mL), and subsequently dried in vacuo. The gray solid was transferred to a round-bottom flask containing 15 mL of concentrated H₂SO₄. The flask was securely fastened to a vortex mixer and allowed to shake in the dark for 3 h. The green H_2SO_4 solution was poured over crushed ice at which point a red/brown precipitate formed. The precipitate was collected by filtration and washed with H_2O . The crude product was triturated twice with CHCl₃, giving 3b as a red powder (170 mg, 84% yield). Additional purification of milligram amounts of **3b** was performed by chromatog-raphy (silica gel; CHCl₃ eluent; $R_f = 0.18$). ¹H NMR (C₆D₆/TFA-d (5:1)): δ 8.68 (s, 4 H), 8.35 (s, 2 H), 1.80 (s, 12 H). ¹³C NMR (CDCl₃/TFA-d (5:1)): δ 185.93, 147.84, 134.21, 133.73, 131.93, 128.69, 13.27. IR (KBr, cm⁻¹): 1662, 1601, 1448, 1374, 1283, 1022, 714. HR-FAB-MS (H_2SO_4 matrix) for $C_{26}H_{19}O_4$ (M + H⁺): calcd, 395.1283; found, 395.1262. UV-vis (CHCl₃) (λ, nm): 491, 464, 335, 253.

Triptycene Bis(4-*tert***-butylphenylphthalimide) 6.** The reaction was performed in the dark. N,N'-Bis(4-*tert*-butylphenyl)-9,10-dimethoxy-2,3,6,7-anthracenetetracarboxylic acid 2,3:6,7-diimide¹ (17 mg, 2.6 × 10^{-2} mmol) was suspended in a mixture of CHCl₃ (5 mL) and propylene oxide (1 mL) in a round-bottom flask equipped with a reflux condenser, N₂ inlet, and magnetic stirring bar. α -Diazonium chloride benzoic acid (92 mg, 0.50 mmol) was added to the suspension in one portion, and the reaction mixture was heated to 61 °C and refluxed overnight. The heating source was removed, the solvent was evaporated, and the crude product was collected (some starting material remained). Chromatography (silica gel, CH₂Cl₂ eluent) gave a clear oil (R_f = 0.67) that yielded a white precipitate upon addition of 2 mL of cold Et₂O. The sample was cooled in a refrigerator overnight before the precipitate was collected by filtration (6 mg 8.4 µmol, 31%). ¹H NMR (CDCl₃): δ 8.18 (s, 4 H), 7.72 (m, 2 H), 7.48 (d, J = 8.6 Hz, 4 H), 7.29-7.19 (m, 6 H), 4.45 (s,

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6 H), 1.32 (s, 18 H). IR (KBr, cm⁻¹): 2963, 1777, 1724, 1518, 1376, 1253, 1123, 745. HR-FAB-MS (MNBA matrix) for $C_{46}H_{41}N_2O_6$ (M + H⁺): calcd, 717.2965; found, 717.2978.

endo-Peroxide 9. 3c (0.5 mg) was dissolved in 0.5 mL of CDCl₃ in a standard ¹H NMR tube, giving an orange/red solution. The solution was exposed to room light for 15 h during which time the color changed to pale yellow (λ (nm): 324, 267, 224). The ¹H NMR was examined before and after purification by chromatography (silica gel, CH₂Cl₂ eluent), showing that 9 was formed quantitatively. ¹H NMR (CDCl₃): δ 8.06 (s, 4 H), 2.80 (m, 4 H), 1.60 (m, 4 H), 1.42 (m, 8 H), 0.95 (m, 6 H). IR (KBr, cm⁻¹): 2956, 2924, 2855, 1660, 1611, 1305, 722. HR-FAB-MS (ONPOE matrix) for C₃₈H₄₃O₆ (M + H⁺): calcd, 595.3060; found, 595.3074.

ESR Spectra. ESR spectra were recorded on an IBM-Bruker ESP 300 X-band spectrometer equipped with a variable-temperature control unit. The spectroelectrochemical cell consisted of a 8.5-cm-length quartz

tube that was constricted to 1.2-mm internal diameter at the bottom. A 7-mm Pt wire was encased with insulation tubing except for an exposed end (ca. 1 cm) which was placed in the constricted portion of the cell. A Pt wire counter electrode and a Ag wire reference electrode were positioned away from the working compartment. Following addition of the sample solution, the cell was sealed with a modified septum and degassed with argon. Reductions were performed by setting the potential slightly positive of the corresponding $E^{\circ'}$ values and gradually changing the potential to more negative values. The same ESR tube was used without electrodes present for anion radicals prepared by bulk electrolysis. Temperatures are in Kelvin.

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Uncatalyzed and Chorismate Mutase Catalyzed Claisen Rearrangements of 5,6-Dihydrochorismate and 6-Oxa-5,6-dihydrochorismate

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Abstract: The synthesis of 6-oxa-5,6-dihydrochorismic acid (4) from D-xylose is described. The half-lives for the uncatalyzed Claisen rearrangements of 5,6-dihydrochorismic acid (3) and 4 in D₂O at 30 °C were 49 000 and 1200 h, respectively, compared to a half-life of 15.6 h for chorismic acid (1) under similar conditions. Both 3 and 4 were processed by the mutase activity of chorismate mutase-prephenate dehydrogenase from *Escherichia coli* with $k_{cat}/k_{uncat} = 1 \times 10^6$ and 4×10^5 , respectively, compared to $k_{cat}/k_{uncat} = 2 \times 10^6$ for 1.

The rearrangement of chorismate (1) to prephenate (2) is catalyzed by chorismate mutase. It is the first step in the biosynthesis of phenylalanine and tyrosine from 1, by what is formally a [3,3] sigmatropic rearrangement, and is one of the most intriguing transformations found in nature (Scheme I).¹ The details of the mechanism of the enzymatic process are not understood irrespective of the extensive investigations from numerous laboratories. In a recent publication, we described studies that defined the structural requirements for catalysis by the mutase site of the biofunctional enzyme chorismate mutase-prephenate dehydrogenase from Escherichia coli.² Crucial to the study was the demonstration that 5,6-dihydrochorismic acid (3, Chart I) was a substrate for chorismate mutase. Described herein are the detailed studies of the thermal and enzyme-catalyzed rearrangement of 3 and the related dihydropyran analogue 4, which proved to be another effective substrate for chorismate mutase.

Haslam and co-workers reported that 3 did not display any tendency to rearrange with chorismate mutase, but it was a modest inhibitor.³ The dihydro analogue 3 is, in fact, an excellent substrate for chorismate mutase, but observation of enzymatic catalysis requires special experimental conditions since the uncatalyzed reaction is so slow compared to the uncatalyzed rearrangement of 1. The dihydro analogue 3 and a 1,2-dihydrochorismic acid of undetermined stereochemistry were prepared



Scheme II



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10, X = -CN

by Haslam and co-workers by diimide reduction of (-)-1.³ Selective preparation of the two isomers was accomplished by a clever modification of reaction temperature followed by fractional crystallization. Since we had difficulty with the recrystallization

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