

Manifestation of Polar Reaction Pathways of 2,3-Dichloro-5,6dicyano-*p*-benzoquinone

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Supporting Information

ABSTRACT: Reactions of 2,3-dichloro-5,6-dicyano-*p*-benzoquinone (DDQ) with silyl enol ethers, silyl ketene acetals, allylsilanes, enamino esters, and diazomethanes have been studied in CH₃CN and CH₂Cl₂ solutions. The second-order rate constants for C attack at DDQ (log $k_{\rm C}$) correlate linearly with the nucleophile-specific parameters N and $s_{\rm N}$ and are 2–5 orders of magnitude larger than expected for SET processes, which strongly supports the polar mechanism for C–C bond



formation. The second-order rate constants for O attack agree well with the calculated rate constants for rate-determining single electron transfer (SET). As a radical clock experiment ruled out outer sphere electron transfer, an inner sphere electron transfer mechanism is suggested for O attack.

INTRODUCTION

2,3-Dichloro-5,6-dicyano-*p*-benzoquinone (DDQ), first synthesized by Thiele and Günther in 1906,¹ has a high redox potential (0.54 V vs SCE in acetonitrile)² and is one of the most important oxidizing reagents in organic chemistry. It usually reacts as a one-electron acceptor to give a radical anion in the first step and has been used for the oxidation of steroid ketones, hydroaromatic compounds, alcohols, phenols, and heterocycles.³ Recently, DDQ was used as an oxidant for several oxidative coupling reactions.⁴

Triggered by the observation of DDQ-substrate adducts during the oxidation of 4-aza-3-ketosteroids by DDQ,⁵ Bhattacharya and co-workers investigated the reactions of cyclic silyl enol ethers with DDQ and found the formation of adducts **3a-C** and **3a-O** (Scheme 1).⁶

The dramatic solvent and temperature effects on the product ratios were considered to be indicative of the operation of two distinctly different pathways for the formation of **3a-C** and **3a-O**. Though both products were suggested to be formed via a radical ion pair, the possibility of a nucleophilic attack of the silyl enol ether on DDQ to form the carbon–carbon adduct **3a-C** was explicitly mentioned as an alternative.⁶ We now report kinetic investigations of the reactions of DDQ with π -nucleophiles, which clearly show that polar mechanisms are rather the rule than the exception for these reactions.

In recent years, we have developed a linear free energy relationship based model for polar organic reactions,⁷ which uses eq 1 to predict rates and selectivities for these reactions:

$$\log k(20^{\circ}\mathrm{C}) = s_{\mathrm{N}}(E+N) \tag{1}$$

In eq 1 the second-order rate constant (log k) is calculated by two nucleophile-specific parameters s_N and N and one electrophile-specific parameter E. A comprehensive nucleophilicity scale covering more than 30 orders of magnitude⁸ has been created by using a series of benzhydrylium ions and structurally related quinone methides as reference electrophiles.⁹ We have now used the linear free energy relationship (1) to elucidate the mechanisms of the reactions of DDQ with π -systems (Table 1).

RESULTS

Product Studies in Acetonitrile. In line with earlier investigations,⁶ the cyclic silyl enol ethers $2\mathbf{a}-\mathbf{c}$ were found to give predominantly $4(\mathbf{a}-\mathbf{c})-\mathbf{C}$, the products of C attack in acetonitrile, and only a trace (~4%) of **4b-O**, the product of O attack, was detected by ¹H NMR spectroscopy (Scheme 2). Compounds $4(\mathbf{a}-\mathbf{c})-\mathbf{C}$ were isolated as mixtures of diastereomers after column chromatography. The major diastereomers of $4(\mathbf{a},\mathbf{c})-\mathbf{C}$ were obtained and characterized as pure compounds by recrystallization from a mixture of dichloromethane and pentane. Additionally, the major diastereoisomer of **4a-C** was characterized by X-ray crystallography (Figure 1).

The reactions of the terminal silyl enol ethers 2d and 2e and silyl ketene acetals 2f and 2g with DDQ yielded the products of C attack (4(d-g)-C) quantitatively (Scheme 3), which were purified by column chromatography.

While the isobutyraldehyde-derived silyl enol ether 2h reacted via O attack to give 68% of 3h-O and 32% of the dehydrogenated product 5h (Scheme 4), according to ¹H NMR analysis of the crude reaction product, the structurally analogous silyl ketene acetal 2i reacted with DDQ preferentially at carbon to give 94% of 3i-C and 6% of 3i-O. At elongated reaction times 3i-C slowly transformed into a mixture of 3i-O and 5i (Scheme 4). The parallel increase of 3i-O and 5i (Figure 2) shows that both products are formed in parallel reactions

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Table 1. Nucleophiles 2a–o and Their Reactivity Parameters N and s_N in CH_2Cl_2

Nucleophile			$N(s_{\rm N})^a$
OSiMe ₃	2a	<i>n</i> = 2	5.21 (1.00)
\bigwedge	2b	n = 1	6.57 (0.93)
└-(ノ) n	2c	<i>n</i> = 3	6.62 (1.00)
	2d	$\mathbf{R} = \mathbf{P}\mathbf{h}$	6.22 (0.96)
OSiMe ₃	2e	$\mathbf{R} = \mathbf{M}\mathbf{e}$	5.41 (0.91)
R	2f	R = OPh	8.23 (0.81)
	2g	R = OBu	10.21 (0.82)
OSiMe₃ I	2h	R = H	3.94 (1.00)
R	2i	R = OMe	9.00 (0.98)
OSiMe ₃			
OSiMe	2ј		$4.61 (1.19)^{b}$
SiMe ₃	2k		4.41 (0.96)
٩́	21		8 52 (0 80)
N_CO ₂ Et	21		8.32 (0.80)
Me ₂ N CO ₂ Et	2m		9.43 (0.80)
н	2n	$\mathbf{R} = \mathbf{P}\mathbf{h}$	9.35 (0.83)
№2=< R	20	R = COOEt	4.91 (0.95)
^a From ref 8. ^b This wor	k.		

from 3i-C and excludes the generation of 5i by elimination from 3i-O. In view of the slow conversion of 3i-C into 3i-O (after 10 min, only 7% of 3i-O are present), the observation of a small amount of 3i-O even after 1 min indicates that 6% of 3i-O is directly formed from DDQ and 2i (kinetic product control). The desilylated products 4h-O and 4i-O were Scheme 2. Reactions of DDQ with Cyclic Silyl Enol Ethers 2a-c



^aIsolated yield. ^bNot observed by ¹H NMR in the crude material.



Figure 1. Crystal structure of 4a-C (major diastereomer, thermal ellipsoids are drawn at the 50% probability level).

obtained by column chromatography on silica gel in 45 and 63% yield, respectively (see the Supporting Information).

The rearrangement of **3i-C** to **3i-O** can be assumed to proceed via a radical pathway, as shown in Scheme 5. When (2,2,6,6-tetramethylpiperidin-1-yl)oxyl (TEMPO) was added to crude **3i-C**, which was formed from **1** and **2i** in CH₃CN within 1 min (Scheme 4), a 45: 55 mixture of **3i-O** and **6** was formed within 72 h at room temperature (¹H NMR). Methyl methylacrylate **5i** probably was lost during evaporation of the solvent. Column chromatography of the mixture provided pure **6** in 49% yield, which was fully characterized.



Scheme 4. Relative Yields (¹H NMR) of the Reactions of DDQ with 2h and 2i in CD_3CN



 $a_{n.d.} = \text{not detected.}$



Figure 2. Time resolved relative ¹H NMR yields of 3i-C, 3i-O, and 5i during the reaction of DDQ with 2i (in CD₃CN, at 20 °C).

The reaction of 1,2-disiloxycyclohexene 2j with DDQ gave exclusively the product of O attack (Scheme 6). The partially desilylated product 4j-O crystallized from CH₂Cl₂/pentane solution and was analyzed by X-ray crystallography (Figure 3).

The reaction of allylsilane **2k** with DDQ initially gave **3k-C**, the product of C attack, that slowly rearranged into the allyl phenyl ether **3k-O** (Scheme 7). After hydrolysis, the desilylated derivative **4k-O** was obtained by crystallization in 81% yield (see the Supporting Information).

Previous investigations have shown that DDQ reacts with the secondary enaminoester 7 to give the spirane 9 (Scheme 8), indicating that 7 attacks at C-2 of DDQ.¹⁰ In this work, we observed the exclusive formation of **3l-O** from **1** and the tertiary enaminoesters **2l** or **2m** after aqueous workup, which can be explained through O attack and formation of **10** as illustrated in Scheme 8. Proton migration, cyclization, hydrolysis, and decarbonylation eventually yield benzofuran

Scheme 5. Trapping of the Intermediate of the Rearrangement of 3i-C to 3i-O



Scheme 6. Reaction of DDQ with 2j





Figure 3. Crystal structure of **4j-O** (thermal ellipsoids are drawn at the 50% probability level).

31-O as the final product. Possibly, **21** and **2m** initially also attack **1** at the carbon, to give a zwitterion analogous to **8**, which cannot cyclize to a spirane because of the absence of an NH proton. Therefore, it may revert to reactants and subsequently react via O attack to give **31-O** as the final product.

While the reaction of phenyldiazomethane with DDQ gave 92% of the bicyclic diketone 3n-C (diastereomer ratio: ca. 55/45), no reaction between ethyl diazoacetate and DDQ was observed at room temperature within 1 h (Scheme 9).

Solvent Effect. Scheme 10 shows that the **4a-O/4a-C** ratio generated by the reaction of DDQ with **2a** depends strongly on the concentration of the reactants and on the solvent. According to entries 1 and 2, the **4a-O/4a-C** ratio increases from 40/60 to 92/8 when reducing the concentration of **2a** from 0.1 to 0.02 mol L^{-1} in CH₂Cl₂. In the dilute solution (0.02

Scheme 7. Reaction of DDQ with 2k



mol L⁻¹), the **4a-O/4a-C** ratio decreases dramatically from 92/ 8 to 7/93 when the solvent is changed from CH_2Cl_2 to $CH_2Cl_2/MeOH$, and CH_3CN (entries 2–7, Scheme 10). In acetonitrile solution, the product ratio is neither affected by the concentration of **2a** nor by the presence of methanol (entries 7–10, Scheme 10). As **3a-C** does not rearrange into the thermodynamically more stable isomer **3a-O** when dissolved in CD_2Cl_2 (Scheme 11), we can rule out isomerization at the product stage.

When the *tert*-butyldimethylsilyl substituted enol ether 2a' was used, more product of O attack was observed than in the corresponding reactions with the trimethylsilyl substituted compound 2a in both CH₂Cl₂ and CH₃CN (entries 2/11 and 7/12, Scheme 10). In the reactions of DDQ with 2a', the 4a-O/4a-C ratio decreases when methanol is added to acetonitrile





"Yield and dr (¹H NMR) correspond to the crude product; only the major isomer was isolated and characterized.

and reaches a constant value of 7/93 for >10% of MeOH (entries 13–17, Scheme 10).

A rationalization for these observations is given in Scheme 12. Attack of 2a at a carbon of DDQ to give 11 is faster than O attack to give 12. As silvl shifts to convert 11 into 3a-C and 12 into 3a-O cannot occur intramolecularly, silyl-carriers are needed. Such a carrier may be the solvent when the reacton is carried out in acetonitrile solution (\rightarrow *N*-trimethylsilyl-nitrilium ions). As the product ratio [4a-O]/[4a-C] obtained from 2a is not affected by addition of the stronger nucleophile methanol in acetonitrile, we conclude that 11 and 12 are formed irreversibly in acetonitrile solution due to the fast subsequent silvl shift, and that the product ratio [4a-O]/[4a-C] = 7/93corresponds to k_0/k_c . The independence of the observed rate constants k_{obs} (for measurements see below) of methanol additives (Figure 4) and NBu₄OTs additives (Tables S1 and S11, see the Supporting Information) are in line with this interpretation.

Because of its lower nucleophilicity, dichloromethane cannot act as a silyl-carrier, and the conversions $11 \rightarrow 3a$ -C and $12 \rightarrow 3a$ -O can only occur in the presence of other nucleophiles, e.g., methanol, or when bimolecular reactions of two molecules of 11 or 12 lead to the formation of 3a-C and 3a-O. In the



^aYield isolated after crystallization from AcOH/CH₃CN

	CN + CN + 2a 2a'	$ \begin{array}{ccc} R & 1. CH_2CI_2 \text{ or } CH_3CN \\ & & 1 \min, 20^{\circ}C \\ \hline & 2. H_2O \text{ or } MeOH \\ \hline & R \\ \hline & Me_3Si \\ {}^{\ell}BuMe_2Si \\ \end{array} $		ci + ci	O H C N O H 4a-C
entry	nucleonhile	solvent	[2a or 2a'] ^a /	relativ	ve yield ^b /%
entry	ndeleophile	Solvent	mol L ⁻¹	4a-0	4a-C
1	2a	CH ₂ Cl ₂	0.1	40	60 (58:42)
2		CH ₂ Cl ₂	0.02	92	8 (50:50)
3		CH ₂ Cl ₂ (1% MeOH)	0.02	39	61 (74:26)
4		CH ₂ Cl ₂ (4% MeOH)	0.02	22	78 (41:59)
5		CH ₂ Cl ₂ (10% MeOH)	0.02	14	86 (36:64)
6		CH ₂ Cl ₂ (20% MeOH)	0.02	9	91 (29:71)
7		CH ₃ CN	0.02	7	93 (25:75)
8		CH ₃ CN	0.1	7	93 (25:75)
9		CH ₃ CN (1% MeOH)	0.1	7	93 (25:75)
10		CH ₃ CN (10% MeOH)	0.1	6	94 (22:78)
11	2a'	CH ₂ Cl ₂	0.02	>95	<4
12		CH ₃ CN	0.02	30	70 (50:50)
13		CH₃CN	0.1	33	67 (51:49)
14		CH ₃ CN (1% MeOH)	0.1	25	75 (51:49)
15		CH ₃ CN (5% MeOH)	0.1	11	89 (45:55)
16		CH ₃ CN (10% MeOH)	0.1	7	93 (41:59)
17		CH ₃ CN (20% MeOH)	0.1	7	93 (34:66)

Scheme 10. Solvent Effect on the Reaction of DDQ with 2a and 2a'

"In all cases, 10 equiv of 2a or 2a' were used, that is, [2a or 2a']/[1] = 10. ^bQuantitative product formation, ratios of diastereoisomers were determined by ¹H NMR of the crude product mixture and are given in parentheses (see the Supporting Information for details).





absence of methanol or in highly dilute solutions in dichloromethane, silyl transfers will be slow, and the formation of 11 that is thermodynamically less stable than 12 will be reversible. As a consequence, in highly dilute dichloromethane solutions (where also the concentrations of 11 and 12 will be small), the ratio [3a-O]/[3a-C] will be large because of the reversible formation of 11. Already small amounts of methanol are sufficient to accelerate the silyl transfer, which results in an increase of the observed rate constant (factor of 2 in a 0.05 mol L^{-1} solution, Figure 4) as well as the yield of 3a-C.

The observations with the $tBuMe_2Si$ substituted enol ether 2a' are in line with these interpretations. As the transfer of the sterically more shielded $tBuMe_2Si$ group is slower than that of the Me₃Si group, in dilute dichloromethane solution, C attack

at DDQ becomes completely reversible, and **4a-O** is the only product (entry 11, Scheme 10). In pure acetonitrile, C attack (which is irreversible with **2a**, see above) becomes partially reversible in the reaction with **2a**', and addition of the more nucleophilic methanol now leads to a decrease of the ratio [**4a-O**]/[**4a-C**] (entries 13–17, Scheme 10).

Charge Transfer (CT) Complexes. The formation of CT complexes between electron-deficient and electron-rich π -systems is well-known.¹¹ As the association constants between the nucleophiles 2 and DDQ (1) are small, at room temperature the corresponding CT complexes could only be observed in few cases as short-lived intermediates when high concentrations of the reactants were employed. Since lowering of the temperature increases the association constants and retards the reactions of 1 with 2, we have been able to observe several CT complexes, however, when mixing the nucleophiles 2 with DDQ in CH₂Cl₂ at -80 °C (Scheme 13). The new absorption bands observed under these conditions can be assigned to charge transfer complexes, because neither DDQ nor the nucleophiles 2 examined absorb beyond 450 nm.

Though the CT complexes are likely to correspond to the encounter complexes preceding the formation of the reaction products from 1 and 2, the kinetic data do not allow us to differentiate whether the CT complexes are intermediates on

Scheme 12. Possible Mechanism of the Reaction of DDQ with 2a





Figure 4. Effect of methanol on the rate of the reaction of DDQ (5.0 \times 10⁻³ mol L⁻¹) with 2a (5.0 \times 10⁻² mol L⁻¹) in CH₂Cl₂ and in CH₃CN (monitoring the CT complex at 570 nm).

Scheme 13. Characteristic Absorption Bands of the CT Complexes of Nucleophiles 2 with DDQ in CH_2Cl_2 at $-80\ ^\circ C$



the way to the products or correspond to dead ends of a nonproductive side channel (Curtin-Hammett principle¹²).

Kinetic Studies. All kinetic investigations of the reactions of DDQ with the nucleophiles **2** were performed in acetonitrile or dichloromethane solution at 20 °C. The reactions were monitored by UV–vis spectroscopy at or close to the

absorption maxima of DDQ (278 nm, Figure 5). For some nucleophiles (2d, 2l, and 2m), the formation of the products



Figure 5. Time-dependent UV–vis absorbance during the reaction of DDQ ($1.0 \times 10^{-4} \text{ mol } L^{-1}$) with 2e ($3.9 \times 10^{-3} \text{ mol } L^{-1}$) in CH₃CN at 20 °C.

was monitored (at 350 nm) because of the intensive absorbance of nucleophiles at lower wavelength (<300 nm). Pseudo first-order rate constants k_{obs} were obtained by leastsquares fitting of the absorbances to the monoexponential function $A_t = A_0 e^{-k_{obs}t} + C$ or $A_t = A_0 (1 - e^{-k_{obs}t}) + C$. Secondorder rate constants k_2 were derived from the linear correlation of k_{obs} (s⁻¹) with the concentrations of the nucleophiles (Figure 6).

In line with the fast reversible formation of the CT complexes, identical rate constants for the reaction of DDQ with **2a** were derived by following the decay of the absorption band of the CT complex (589 nm) and the absorption band of DDQ (286 nm) as well as the increase of the absorption of the product (350 nm) in CH₂Cl₂ at -60 °C (Figure 7).

The observation of second-order kinetics for the reactions of DDQ with nucleophiles **2** in CH₂Cl₂ is surprising in view of the discussion of Schemes 10 and 12. Possibly, the perfect linearity of the k_{obs} vs [**2**] correlations (Tables S15–S21) indicates that



Figure 6. UV-vis spectroscopic monitoring of the reaction of DDQ $(1.0 \times 10^{-4} \text{ mol } \text{L}^{-1})$ with **2e** $(3.9 \times 10^{-3} \text{ mol } \text{L}^{-1})$ at 278 nm in CH₃CN at 20 °C. Insert: Determination of the second-order rate constant $k_2 = 23 \text{ L mol}^{-1} \text{ s}^{-1}$ from the dependence of the first-order rate constant k_{obs} on the concentration of **2e**.



Figure 7. UV-vis spectra of a mixture of DDQ $(1.0 \times 10^{-4} \text{ mol } \text{L}^{-1})$ and 2a $(4.0 \times 10^{-3} \text{ mol } \text{L}^{-1})$ in CH₂Cl₂ at -60 °C and the time dependent UV-vis absorbance at 286, 350, and 589 nm.

also 2a-f,k may act as silyl carriers under the conditions of the kinetic experiments.

Table 2 shows that the ratios of the second-order rate constants for the reactions of DDQ with nucleophiles **2** in CH₂Cl₂ and CH₃CN solutions, $k_2^{CH_2Cl_2}/k_2^{MeCN}$, vary from 0.44 to 3.1 indicating that the kinetics are only slightly affected by solvent polarity.

Ab Initio Calculation of Adiabatic Ionization Potentials. In order to elucidate the feasibility of single electron transfer (SET) processes, we have determined the ionization potentials of the π -nucleophiles 2(a-q) by guantum chemical calculations using the Gaussian 09 program package.¹³ Adiabatic ionization potentials were calculated at the G3(MP2) level of theory, which is a high-level composite ab initio molecular orbital theory method as the sum of the CCSD(T) calculations with a double- ζ basis set and basis set corrections carried out at the MP2 level of theory to approximate CCSD(T) calculations with a large triple- ζ basis. G3(MP2) has been shown to reproduce a large test set of gasphase experimental data within chemical accuracy.¹⁴ The adiabatic ionization potentials listed in Table 3 have been calculated as the difference of the enthalpies of the radical cations and those of the neutral molecules in the gas phase.

Table 3. Calculated Adiabatic Ionization Potentials [G3(MP2)] of Nucleophiles 2 and the Corresponding Electrochemical Oxidation Potentials

nucleophile	IP_{a}/eV	$E_{\rm ox}$ (vs SCE)/V
2a	7.77	1.30 ^{<i>a</i>}
2b	7.83	1.21 ^b
2c	7.71	1.15 ^b
2d	8.10	1.32^{a}
2e	8.24	1.43 ^b
2f	7.89	1.24^{b}
2g	7.65	1.12 ^b
2h	7.77	1.18^{b}
2i	7.22	0.90 ^{<i>a</i>}
2j	7.05	0.80^{b}
2k	8.26	1.44 ^b
21	7.71	1.15 ^b
2m	7.70	1.14^{b}
2n	7.87	1.23 ^b
$2p^{c}$	8.48	1.50 ^{<i>a</i>}
$2q^{c}$	7.18	0.83 ^a

"In CH₃CN from ref 15. ^bCalculated by substitution of calculated ionization potentials into eq 2. ^cSee Figure 8 for the structures of these compounds.

Figure 8 shows that the calculated ionization potentials correlate with the experimental oxidation potentials¹⁵ of **2a**, **2d**,

Tabl	e 2. S	Second	-Order	Rate	Constants	k for	the	Reaction	of DDO	Q wit	h Nucl	leophiles	(2a-n)) at 20)°(
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nucleophile	$k_2^{\rm CH_2Cl_2}/\rm L \ mol^{-1} \ s^{-1}$	$k_2^{\rm MeCN}/{\rm L}~{\rm mol^{-1}}~{\rm s^{-1}}$	$k_2^{\mathrm{CH_2Cl_2}}/k_2^{\mathrm{MeCN}}$	$k_{\rm O}/{\rm L}~{\rm mol}^{-1}~{\rm s}^{-1}~{\rm (in}~{\rm CH}_{3}{\rm CN})$	$k_{\rm C}/{\rm L}~{\rm mol^{-1}}~{\rm s^{-1}}$ (in CH ₃ CN)
2a	1.9×10^{1}	2.5×10^{1}	0.76	1.9	2.3×10^{1}
2b	3.4×10^{2}	5.5×10^{2}	0.62		5.5×10^{2}
2c	9.6×10^{2}	7.1×10^{2}	1.4		7.1×10^{2}
2d	2.0×10^{2}	4.5×10^{2}	0.44		4.5×10^{2}
2e	4.5×10^{1}	2.3×10^{1}	2.0		2.3×10^{1}
2f	1.1×10^{4}	3.5×10^{3}	3.1		3.5×10^{3}
2g		8.5×10^{5}			8.5×10^{5}
2h		1.8×10^{-1}		1.8×10^{-1}	
2i		1.1×10^{7}		6.6×10^{5}	1.0×10^{7}
2j		7.0×10^{4}		7.0×10^{4}	
2k	3.9×10^{-1}	3.9×10^{-1}	1.0		3.9×10^{-1}
21		6.8		6.8	
2m		2.3×10^{1}		2.3×10^{1}	
2n		8.6×10^{3}			8.6×10^{3}



Figure 8. Correlation between calculated adiabatic ionization potentials IP_a and experimental oxidation potentials E_{ox} (vs SCE in CH₃CN).

2i, **2p**, and **2q** as expressed by eq 2, which is similar to the correlation previously reported for aromatic compounds ($IP_a = 1.5E_{ox} + 5.8$).¹⁶ Using correlation (2) we have calculated experimentally not available oxidation potentials from their adiabatic ionization potentials (Table 3).

$$IP_{a} = 1.88E_{ox} + 5.55$$
(2)

DISCUSSION

Correlation Analysis. In order to examine the validity of eq 1 for the reactions of DDQ (C attack) with π -nucleophiles, a plot of $(\log k_{\rm C})/s_{\rm N}$ against the nucleophilicity parameters *N* of nucleophiles is presented in Figure 9. A fair linear correlation ($R^2 = 0.91$) was obtained.



Figure 9. Plot of $(\log k_C)/s_N$ vs *N* for the reactions of DDQ (C attack) with π -nucleophiles in CH₃CN at 20 °C.

According to eq 1, the electrophilicity parameter of DDQ (reactivity at C-2) was determined as E = -3.66 by least-squares minimization of $\Delta^2 = \Sigma (\log k_{\rm C} - s_{\rm N} (N + E))^2$ using the second-order rate constants $k_{\rm C}$ given in Table 2 and the *N* and $s_{\rm N}$ parameters of the nucleophiles **2a-n** from Table 1.

In previous work we have shown that the electrophilicity parameters of benzhydrylium ions and structurally related quinone methides correlate linearly with their reduction potentials (Figure 10).¹⁷ When the data for DDQ are added



Figure 10. Correlation between the electrophilicity parameters *E* and the reduction potentials E_{red} of reference electrophiles¹⁷ and DDQ.

to this correlation, one can see that DDQ reacts much more slowly with π -nucleophiles than expected from its reduction potential. The position of DDQ in Figure 10 implies that the transition states of the reactions of DDQ with $\pi_{\rm CC}$ systems profit much less from the formation of the new CC bond (product stabilizing factor) than the corresponding reactions of benzhydrylium ions and quinone methides. Yu and co-workers have demonstrated recently that a relationship between electrophilicities E and LUMO energies holds only within species that share similar substitution patterns.¹⁸ Thus, arylidene Meldrum's acids and arylidene barbituric acids were found to be 5 orders of magnitude more electrophilic than quinone methides of comparable LUMO energies, indicating that the product stabilizing factor must be more important in the transition states of the former species. The conclusion drawn from Figure 10 that product stabilizing factors are less important in the reactions of DDQ with π -nucleophiles implies that SET processes can be expected to be more likely in reactions of DDQ with nucleophiles than in the corresponding reactions of benzhydrylium ions having similar electrophilicity Ε.

The kinetics of one-electron oxidations of silvlated enol ethers and ketene acetals as well as of allylsilanes with one electron oxidants have previously been studied by Fukuzumi and co-workers.¹⁵ The oxidation potentials of these π -systems as well as the intrinsic barriers for the electron transfer oxidation of these compounds by one-electron oxidants were derived from experimental rate constants using the Rehm-Weller Gibbs relationship (4), which derives the Gibbs energy of activation for the electron transfer $(\Delta G_{et}^{\ddagger})$ from the Gibbs energy of electron transfer (ΔG_{et}^0 given by eq 5) and the intrinsic barrier ($\Delta G_{0,\text{et}}^{\ddagger}$). The latter quantity represents the Gibbs activation free energy for a process where the driving force of electron transfer is zero, i.e. $\Delta G_{et}^{\ddagger} = \Delta G_{0,et}^{\ddagger}$ for $\Delta G_{et}^{0} = 0$. Due to the significant rearrangements of structure accompanying electron transfer, large intrinsic barriers $\Delta G_{0,\text{et}}^{\ddagger}$ (15–20 kJ mol^{-1}) were observed for the one-electron oxidation of a large variety of silvlated π -nucleophiles, including several compounds shown in Figure 8.

By applying the same methodology, we have now used the oxidation potentials in Table 3 to calculate the rate constants for the outer-sphere electron transfer of the reactions of DDQ with the π -nucleophiles **2a**-**n** according to eqs 3–5, assuming an intrinsic barrier of $\Delta G_{0,\text{et}}^{\ddagger} = 18 \text{ kJ mol}^{-1}$, a typical value for such systems according to ref 15.

$$k_{\rm et} = (k_{\rm b}T/h) \exp(-\Delta G_{\rm et}^{\ddagger}/RT)$$
(3)

$$\Delta G_{\rm et}^{\ddagger} = \Delta G_{\rm et}^{0} / 2 + \left[\left(\Delta G_{\rm et}^{0} / 2 \right)^{2} + \left(\Delta G_{0,\rm et}^{\ddagger} \right)^{2} \right]^{1/2} \tag{4}$$

$$\Delta G_{\rm et}^0 = F(E_{\rm ox} - E_{\rm red}) \tag{5}$$

A plot of experimental rate constants log $k_{\rm C}$ (\odot) and log $k_{\rm O}$ (O) for the reactions of DDQ with various π -nucleophiles versus the calculated rates for SET (log $k_{\rm ev}$ from eqs 3–5) is presented in Figure 11. The open circles for systems which



Figure 11. Correlation of experimental rate constants (log k_2 , in CH₃CN) with calculated rate constants for SET (log k_{et}).

react via O attack are close to the diagonal, indicating that these reactions proceed with rates as expected for SET processes, while C attack at DDQ (filled circles) is 2 to 5 orders of magnitude faster than calculated for SET processes. We, therefore, conclude that C attack at DDQ usually occurs by a polar mechanism, which may be reversible if there is no fast subsequent reaction.

It should be noted that all experimental rate constants for the reactions of benzhydrylium ions with π -nucleophiles have been reported to be at least 8 orders of magnitude larger than the calculated rate constants for one-electron transfer processes.¹⁷ Therefore, the reactions of DDQ with π -nucleophiles are generally closer to the borderline between polar reactions and electron transfer than the corresponding reactions of benzhydrylium ions (with π -nucleophiles).

Outer Sphere Electron Transfer (OSET) vs Inner Sphere Electron Transfer (ISET): Radical Clock Experiments. According to Figure 11, the observed rate constant for the reaction of 2j with DDQ corresponds to that expected for a SET process. In order to examine the occurrence of an OSET mechanism, a radical clock¹⁹ substrate 13 was designed based on the structure of 2j (Scheme 14). When 13 was combined with DDQ, only 14, the product of O attack, was observed in the ¹H NMR spectrum of the crude material (>95%), and 15 was isolated in 60% yield after column chromatography, while no cyclization product was observed (Scheme 15). Though cyclization of the radical cation 16 may be slower than analogous cyclizations of radicals, the quantitative formation of





Scheme 15. Exclusion of OSET by a Radical Clock Experiment (CH₃CN, 20 °C)



14 allows us to exclude the operation of an OSET mechanism because Mattay's report on the formation of carbocycles via photoinduced electron transfer oxidative cyclization of silyl enol ethers carrying side chains with olefinic double bonds indicates cyclizations of the intermediate radical cations to be faster than competing intermolecular processes.²¹ Therefore, O attack is suggested to proceed via an ISET mechanism.

CONCLUSION

 π -nucleophiles attack DDQ either at C-2 to give 4hydroxycyclohexadienones or at oxygen to give O-substituted hydroquinones. In several cases initial C attack and subsequent rearrangement to the thermodynamically more stable products of O attack has been observed. Kinetics for the reactions of π systems with DDQ have been determined photometrically, and it was found that the rate constants $k_{\rm C}$ for the attack of the π nucleophiles 2 at C-2 of DDQ can be described by the linear free energy relationship (1), which allowed us to derive the electrophilicity parameter E = -3.66 for the C-2 position of DDQ. It thus possesses an electrophilic reactivity comparable to the flavylium and tropylium ion, and is considerably more reactive than the bis(dimethylamino)-substituted benzhydrylium ion or other highly reactive Michael acceptors (Figure 12).^{8,9a}



Figure 12. Comparison of the electrophilic reactivity of DDQ with that of other electrophiles. 8

The experimental rate constants for the attack of π -systems at C-2 of DDQ are larger than those calculated for SET processes from the electrochemical oxidation potentials of the π -systems and the reduction potential of DDQ, indicating that C attack at DDQ usually occurs via polar processes. However, DDQ reacts about 12 orders of magnitude more slowly than benzhydrylium ions of comparable reduction potential showing that the transition states of the reactions of DDQ with $\pi_{\rm CC}$ systems profit much less from the formation of the new CC-bond (product stabilizing factor) than the corresponding reactions of benzhydrylium ions, which can be explained by the high stabilization of the DDQ radical anion. O-Attack is probably restricted to reactions with sterically shielded nucleophiles or to cases, where the faster C attack is highly reversible because the resulting intermediate cannot undergo fast subsequent reactions. The observed rate constants for O attack are smaller than calculated by eq 1 and thus are similar to those for SET processes (Scheme 16).

Since the radical-clock experiment did not provide evidence for the intermediacy of radical cations of the silylated enol ethers, these reactions are interpreted as inner sphere electron transfer processes.

Scheme 16. C versus O Attack in Reactions of DDQ with π -Nucleophiles



EXPERIMENTAL SECTION

Materials. DDQ was purchased from ABCR and recrystallized from chloroform or dichloromethane before use. Silyl enol ethers $2(a \cdot e, a', h)$ were prepared as reported in the literature.²² Silyl ketene acetals 2(f, g, i) were prepared according to ref 23. The synthesis of disiloxy hexane 2j was achieved by Rühlmann-type acyloin condensation.²⁴ Enamino esters 2l and 2m were prepared from amines and methyl propiolate. Diazomethanes 2n and 2o were prepared according to refs 25 and 26, respectively.

Kinetics. All of the rates of reactions were determined photometrically in dry acetonitrile or in freshly distilled dry CH_2Cl_2 . Fast reactions were determined by using the stopped-flow technique. Slow reactions were measured by conventional photodiode array UV–vis spectrometers. The temperature was kept constant at 20 ± 0.1 °C by using a circulating bath thermostat. In all runs, excess of nucleophiles (at least 8-fold) over DDQ was used to achieve pseudo first-order kinetics.

ASSOCIATED CONTENT

Supporting Information

Details of the kinetic experiments, synthetic procedures, X-ray crystal structure determination, and quantum chemical calculations as well as NMR spectra of all characterized compounds. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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REFERENCES

(1) Thiele, J.; Günther, F. Liebigs Ann. Chem. 1906, 349, 45-66.

(2) Andrieux, C. P.; Merz, A.; Saveant, J.-M.; Tomahogh, R. J. Am. Chem. Soc. 1984, 106, 1957–1962.

(3) (a) Walker, D.; Hiebert, J. D. Chem. Rev. 1967, 67, 153-195. (b) Becker, H.-D. In The Chemistry of the Quinoid Compounds, Part I; Patai, S., Ed.; Wiley: Chichester, U.K., 1974; pp 335-423. (c) Becker, H.-D.; Turner, A. B. In The Chemistry of the Quinoid Compounds Vol. 2; Patai, S., Rappoport, Z., Eds.; Wiley: Chichester, U.K., 1988; pp 1351-1384. (d) Buckle, D. R. In Encyclopedia of Reagents for Organic Synthesis Vol. 3; Paquette, L. A., Ed.; Wiley: Chichester, U.K., 1995; p 1699. (4) (a) Zhang, Y.; Li, C.-J. Angew. Chem., Int. Ed. 2006, 45, 1949-1952. (b) Zhang, Y.; Li, C.-J. J. Am. Chem. Soc. 2006, 128, 4242-4243. (c) Tu, W.; Floreancig, P. E. Angew. Chem., Int. Ed. 2009, 48, 4567-4571. (d) Liu, L.; Floreancig, P. E. Angew. Chem., Int. Ed. 2010, 49, 3069-3072. (e) Guo, C.; Song, J.; Luo, S.-W.; Gong, L.-Z. Angew. Chem., Int. Ed. 2010, 49, 5558-5562. (f) Liu, L.; Floreancig, P. E. Angew. Chem., Int. Ed. 2010, 49, 5894-5897. (g) Hayashi, Y.; Itoh, T.; Ishikawa, H. Angew. Chem., Int. Ed. 2011, 50, 3920-3924. (h) Tsang, A. S.-K.; Jensen, P.; Hook, J. M.; Hashmi, A. S. K.; Todd, M. H. Pure Appl. Chem. 2011, 83, 655-665. (i) Alagiri, K.; Devadig, P.; Prabhu, K. R. Chem.-Eur. J. 2012, 18, 5160-5164. (j) Rohlmann, R.; Garcia Mancheño, O. Synlett 2013, 24, 6-10.

(5) Bhattacharya, A.; DiMichele, L. M.; Dolling, U.-H.; Douglas, A. W.; Grabowski, E. J. J. Am. Chem. Soc. **1988**, *110*, 3318–3319.

Journal of the American Chemical Society

(6) (a) Bhattacharya, A.; DiMichele, L. M.; Dolling, U.-H.; Grabowski, E. J. J.; Grenda, V. J. *J. Org. Chem.* **1989**, *54*, 6118– 6120. (b) For an earlier report on the formation of enones by oxidation of silyl enol ethers with DDQ, see: Ryu, I.; Murai, S.; Hatayama, Y.; Sonoda, N. *Tetrahedron Lett.* **1978**, *19*, 3455–3458.

(7) (a) Mayr, H.; Patz, M. Angew. Chem., Int. Ed. Engl. 1994, 33, 938–957. (b) Mayr, H.; Ofial, A. R. Pure Appl. Chem. 2005, 77, 1807–1821. (c) Mayr, H.; Ofial, A. R. J. Phys. Org. Chem. 2008, 21, 584–595. (d) Mayr, H.; Kempf, B.; Ofial, A. R. Acc. Chem. Res. 2003, 36, 66–77.

(8) For a comprehensive database of nucleophilicity parameters N and s_N as well as electrophilicity parameters E, see: http://www.cup. lmu.de/oc/mayr/DBintro.html.

(9) (a) Mayr, H.; Bug, T.; Gotta, M. F.; Hering, N.; Irrgang, B.; Janker, B.; Kempf, B.; Loos, R.; Ofial, A. R.; Remennikov, G.; Schimmel, H. *J. Am. Chem. Soc.* **2001**, *123*, 9500–9512. (b) Lucius, R.; Loos, R.; Mayr, H. *Angew. Chem., Int. Ed.* **2002**, *41*, 91–95. (c) Richter, D.; Hampel, N.; Singer, T.; Ofial, A. R.; Mayr, H. *Eur. J. Org. Chem.* **2009**, 3203–3211. (d) Ammer, J.; Nolte, C.; Mayr, H. *J. Am. Chem. Soc.* **2012**, *134*, 13902–13911.

(10) Kucklaender, U.; Bollig, R.; Frank, W.; Gratz, A.; Jose, J. Bioorg. Med. Chem. 2011, 19, 2666–2674.

(11) (a) Mulliken, R. S.; Person, W. B. *Molecular Complexes;* Wiley-Interscience: New York, 1969. (b) Mataga, N.; Kubota, T. *Molecular Interactions and Electronic Spectra;* Marcel Dekker: New York, 1970.

(12) (a) Seeman, J. I. Chem. Rev. **1983**, 83, 83–134. (b) Curtin, D. Y. Rec. Chem. Prog. **1954**, 15, 111. (c) Hammett, L. P. Physical Organic Chemistry; McGraw-Hill: New York, 1970.

(13) Frisch, M. J.; Trucks, G. W.; Schlegel, H.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Scalmani, G.; Barone, V.; Mennucci, B.; Petersson, G. A.; Nakatsuji, H.; Caricato, M.; Li, X.; Hratchian, H. P.; Izmaylov, A. F.; Bloino, J.; Zheng, G.; Sonnenberg, J. L.; Hada, M.; Ehara, M.; Toyota, K.; Fukuda, R.; Hasegawa, J.; Ishida, M.; Nakajima, T.; Honda, Y.; Kitao, O.; Nakai, H.; Vreven, T.; Montgomery, J. A.; Peralta, J. E.; Ogliaro, F.; Bearpark, M.; Heyd, J. J.; Brothers, E.; Kudin, K. N.; Staroverov, V. N.; Kobayashi, R.; Normand, J.; Raghavachari, K.; Rendell, A.; Burant, J. C.; Iyengar, S. S.; Tomasi, J.; Cossi, M.; Rega, N.; Millam, J. M.; Klene, M.; Knox, J. E.; Cross, J. B.; Bakken, V.; Adamo, C.; Jaramillo, J.; Gomperts, R.; Stratmann, R. E.; Yazyev, O.; Austin, A. J.; Cammi, R.; Pomelli, C.; Ochterski, J. W.; Martin, R. L.; Morokuma, K.; Zakrzewski, V. G.; Voth, G. A.; Salvador, P.; Dannenberg, J. J.; Dapprich, S.; Daniels, A. D.; Farkas; Foresman, J. B.; Ortiz, J. V.; Cioslowski, J.; Fox, D. J. Gaussian 09, revision A.02; Gaussian, Inc.: Wallingford, CT, 2009.

(14) Henry, D. J.; Sullivan, M. B.; Radom, L. J. Chem. Phys. 2003, 118, 4849-4860.

(15) Fukuzumi, S.; Fujita, M.; Otera, J.; Fujita, Y. J. Am. Chem. Soc. **1992**, 114, 10271–10278.

(16) Pysh, E. S.; Yang, N. C. J. Am. Chem. Soc. 1963, 85, 2124–2130.
(17) Ofial, A. R.; Ohkubo, K.; Fukuzumi, S.; Lucius, R.; Mayr, H. J. Am. Chem. Soc. 2003, 125, 10906–10912.

(18) Zhou, L.-G.; Liao, W.; Yu, Z.-X. Asian J. Org. Chem. 2012, 1, 336–345.

(19) (a) Newcomb, M.; Curran, D. P. Acc. Chem. Res. 1988, 21, 206-

214. (b) Tolbert, L. M.; Sun, X.-J.; Ashby, E. C. J. Am. Chem. Soc. 1995, 117, 2681–2685. (c) Zard, S. Z. Radical Reactions in Organic

Synthesis; Oxford University Press: New York, 2003. (20) Liu, Z.; Peng, L.; Zhang, T.; Li, Y. L. Synth. Commun. 2001, 31, 2549–2555.

(21) (a) Bunte, J. O.; Rinne, S.; Schäfer, C.; Neumann, B.; Stammler, H.-G.; Mattay, J. *Tetrahedron Lett.* **2003**, *44*, 45–48. (b) MacMillan's observation of the intramolecular attack of an enamine radical cation at a 1,2-dialkylated ethylene with formation of a five-membered ring, which has a much lower driving-force confirms this conclusion: Beeson, T. D.; Mastracchio, A.; Hong, J.-B.; Ashton, K.; MacMillan, D. W. C. *Science* **2007**, *316*, 582–585.

(22) (a) Cazeau, P.; Duboudin, F.; Moulines, F.; Babot, O.; Dunogues, J. *Tetrahedron* **1987**, *43*, 2075–2088. (b) Cazeau, P.; Duboudin, F.; Moulines, F.; Babot, O.; Dunogues, J. *Tetrahedron* **1987**, *43*, 2089–2100.

- Article
- (23) Oisaki, K.; Suto, Y.; Kanai, M.; Shibasaki, M. J. Am. Chem. Soc. **2003**, 125, 5644–5645.
- (24) Rühlmann, K. Synthesis 1971, 236-253.
- (25) Wulfman, D. S.; Yousefian, S.; White, J. M. Synth. Commun. 1988, 18, 2349–2352.
- (26) Searle, N. E. Org. Synth. 1956, 36, 25-27.