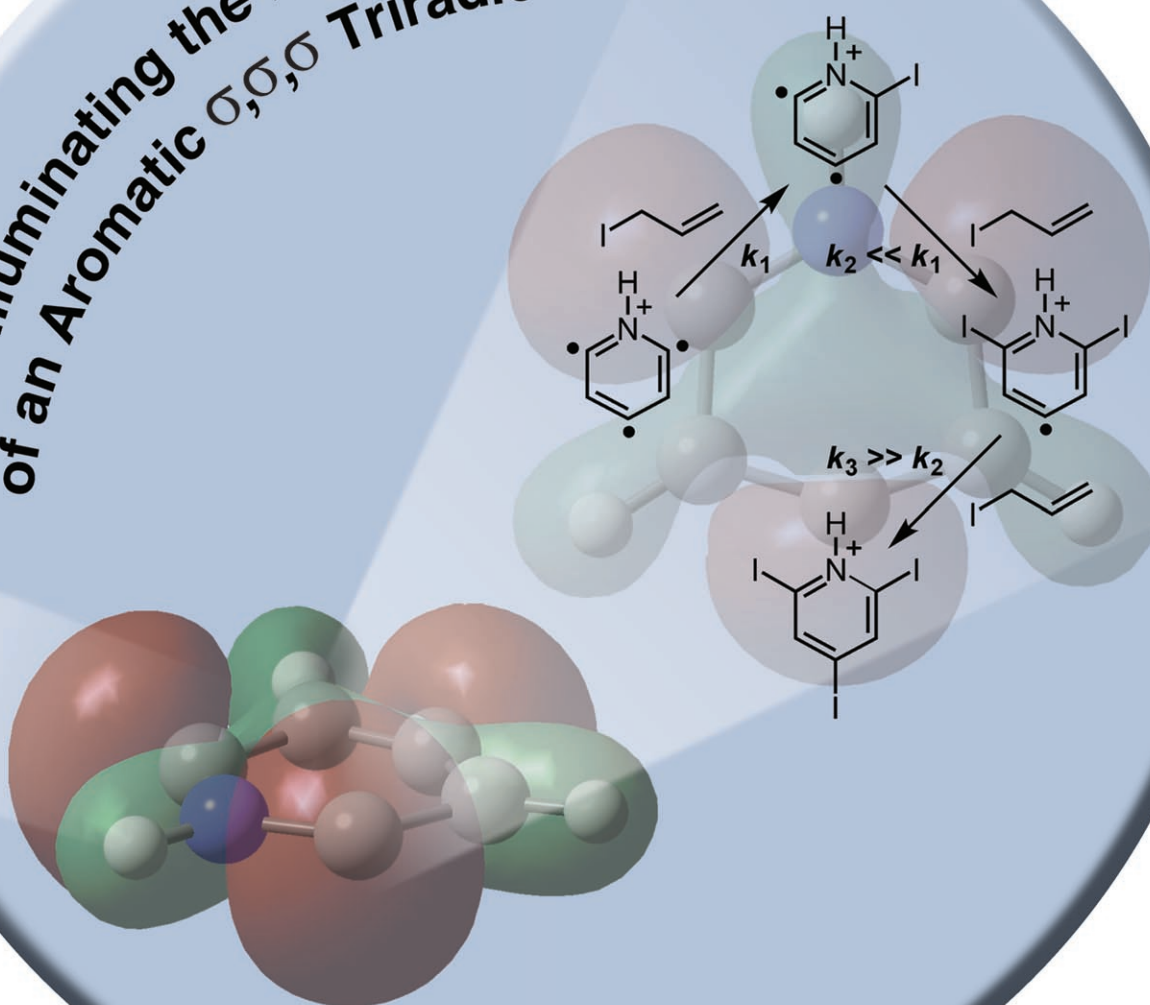


Reactivity of an Aromatic σ,σ,σ -Triradical: The 2,4,6-Tridehydropyridinium Cation**

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Illuminating the Reactivity
of an Aromatic σ,σ,σ Triradical



Certain σ -type, carbon-centered mono- and biradicals, the dehydro- and didehydroarenes, play an important role in organic synthesis, development of new organic materials, and the biological activity of organic compounds.^[1,2] Therefore, numerous investigations have focused on their properties.^[1–5] In contrast, the related σ,σ,σ -triradicals (tridehydroarenes) remain elusive. Almost all studies carried out on tridehydroarenes are computational in nature owing to the difficulty in studying such highly reactive species experimentally.^[6] To the best of our knowledge, the only experimental studies reported for tridehydroarenes are thermochemical measurements on 1,3,5-tridehydrobenzene by Wenthold et al.^[7] and IR detection of 1,2,3-tridehydrobenzene by Sander et al.^[8] The chemical properties of tridehydroarenes appear to be entirely unexplored.

We report here a kinetic reactivity study on a positively charged tridehydroarene, the 2,4,6-tridehydropyridinium cation (**5**). The gas-phase reactions of this triradical are compared to those of two related, previously unreported σ,σ -biradicals, 2,4-didehydropyridinium cation (**3**) and 2,6-didehydropyridinium cation (**4**), as well as two σ -monoradicals, 4-dehydropyridinium cation (**1**) and 2-dehydropyridinium cation (**2**).

Triradical **5** was found to form products similar to those of the mono- and biradicals (Table 1). However, it reacts with most reagents more efficiently than the biradicals and at about equal efficiency as the monoradicals. This behavior can be understood based on a detailed examination of the reactivities of **1–4**.

The monoradicals (**1** and **2**) react rapidly with all the reagents studied. They predominantly abstract a H atom from tetrahydrofuran (THF), HCN and CN groups from *tert*-butyl isocyanide, an I atom from allyl iodide, and a SCH₃ group from dimethyl disulfide, as expected.^[5,9] Monoradical **2** is slightly more reactive than **1** (Table 1), which may be explained by its higher electrophilicity, quantified here by the (calculated) vertical electron affinity (EA) (**1**: 5.84 eV; **2**: 6.59 eV; UBLYP/aug-cc-pVDZ//UBLYP/cc-pVDZ).^[10] For an electrophilic radical, an increase in the EA can lead to a more polar, and hence lower-energy, transition state, thus enhancing its reactivity.^[11]

Addition of a second radical site to monoradical **1** or **2** to generate the singlet biradical **3** results in decreased reactivity. The stabilizing interaction between the two electrons

(revealed by the magnitude of the singlet–triplet (S–T) gap, i.e., the energy difference between the lowest-energy singlet and triplet states) has been used to rationalize this kind of behavior.^[12,13] Part of this interaction is thought to be lost in the transition state of radical reactions owing to a necessity to partially uncouple the biradical electrons.

Based on the above discussion, a larger S–T gap is expected to result in slower radical (but not necessarily nonradical) reactions for singlet biradicals.^[3,5] The S–T gaps of **3** and **4** are calculated to be $-20.0 \text{ kcal mol}^{-1}$ and $-10.0 \text{ kcal mol}^{-1}$, respectively (BD(T)/cc-pVDZ//UBPW91/cc-pVDZ). Based on these values, **4** is expected to be more reactive than **3**. The calculated vertical EAs of **3** and **4** are 6.46 eV and 7.23 eV, respectively (UBLYP/aug-cc-pVDZ//MCSCF/cc-pVDZ), again suggesting that **4** should be more reactive than **3**. Indeed, **4** was found to react faster than **3** with all reagents studied. It is even more reactive than the monoradicals toward three of the four reagents studied. However, the dominating reaction for **4** (but not for **1**, **2**, or **3**) is proton transfer (which is calculated to lead to a *triplet* biradical), a nonradical reaction that does not require uncoupling of the biradical electrons. Hence, the high reactivity of **4** must arise from its relatively high acidity and not from its low S–T gap. Unfortunately, the fast proton-transfer reactions of **4** prevent a comparison of the efficiencies of the radical reactions of **4** with those of **1–3**.

The types of reactions observed for the two biradicals (with the exception of proton transfer) are similar and indicate the presence of two radical sites (e.g., two consecutive I atom abstractions from allyl iodide molecules and two SCH₃ group abstractions from dimethyl disulfide molecules). Most of these reactions were also observed for the monoradicals; however, the reactions with THF are an exception. While the monoradicals predominantly undergo H atom abstraction from THF, the biradicals abstract H₂O, CH₂O, C₂H₄O, and H⁺ faster than they abstract H atoms.

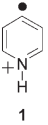
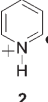
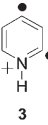
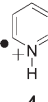
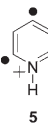
Triradical **5** transfers a proton to the two most basic reagents only, which suggests that it is more acidic than biradical **3** but less acidic than biradical **4**. This observation is supported by the EA calculated for **5** (7.01 eV; UBLYP/aug-cc-pVDZ//UBLYP/cc-pVDZ). Further, its reactivity reveals the presence of three reactive radical sites as it abstracts three SCH₃ groups from dimethyl disulfide molecules, three H atoms from THF molecules, and three I atoms from allyl iodide molecules. However, the three radical sites are not equally reactive. For example, the first and third I atom abstractions occur faster than the second. This finding suggests that the first I atom is abstracted by the radical at the 6-position to generate 2,4-didehydro-6-iodopyridinium cation (Scheme 1). This *meta*-benzyne analogue can be expected to be much less reactive than its 2,6-isomer since **4** is much more reactive than **3**, and addition of an I substituent does not greatly influence the reactivity of this type of a biradical. This expectation was verified by isolating the first I atom abstraction product and allowing it to react with allyl iodide. A remarkably similar reactivity to that of **3** was observed (I abs: 53%, (2°) I abs, (2°) C₃H₅ abs; C₃H₅ abs: 34%, (2°) I abs, (2°) C₃H₅ abs; C₃H₄ abs: 13%; reaction efficiency: 14%). Furthermore, facile proton transfer, the

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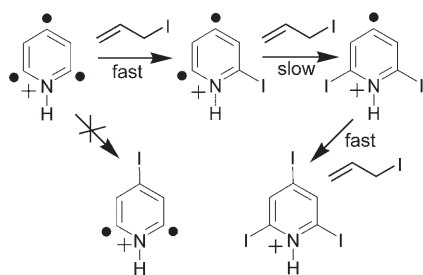
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Table 1: Reaction efficiencies^[a] and branching ratios^[b] for primary products.

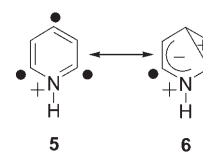
Radical			Reagents 	
 1	H abs 81 % CH ₂ abs 8 % C ₂ H ₃ abs 6 % CHO abs 3 % C ₂ H ₃ O abs 2 % Eff = 28 %	I abs 92 % C ₃ H ₅ abs 8 % Eff = 53 %	SCH ₃ abs 100 % Eff = 75 %	HCN abs 64 % (2°) HCN abs (2°) C ₄ H ₈ abs CN abs 36 % Eff = 90 %
 2	H abs 100 % Eff = 76 %	I abs 84 % C ₃ H ₅ abs 16 % Eff = 69 %	SCH ₃ abs 94 % H abs 5 % SSCH ₃ abs 1 % Eff = 79 %	CN abs 94 % (2°) C ₄ H ₈ abs HCN abs 6 % (2°) C ₄ H ₈ abs Eff = 93 %
 3	H ⁻ abs 36 % CH ₂ O abs 26 % C ₂ H ₄ O abs 20 % H ₂ O abs 14 % 2 × H abs 4 % Eff = 28 %	I abs 49 % (2°) I abs (2°) C ₃ H ₅ abs C ₃ H ₅ abs 42 % (2°) I abs (2°) C ₃ H ₅ abs C ₃ H ₄ abs 9 % Eff = 15 %	SCH ₃ abs 73 % (2°) SCH ₃ abs (2°) SSCH ₃ abs SSCH ₃ abs 23 % (2°) SCH ₃ abs HSCH ₃ abs 4 % Eff = 47 %	HCN abs 54 % (2°) C ₄ H ₈ abs H ⁺ trans and dissociation 46 % Eff = 82 %
 4	H ⁺ trans 78 % (2°) THF add H ₂ O abs 10 % H ⁻ abs 5 % CH ₂ O abs 4 % 2 × H abs 3 % Eff = 85 %	H ⁺ trans 54 % I abs 36 % (2°) I abs C ₃ H ₅ abs 6 % (2°) I abs (2°) C ₃ H ₅ abs C ₃ H ₄ abs 4 % Eff = 34 %	H ⁺ trans 65 % (2°) SCH ₃ abs SCH ₃ abs 17 % (2°) SCH ₃ abs HSCH ₃ abs 8 % e ⁻ abs 8 % SSCH ₃ abs 2 % Eff = 93 %	H ⁺ trans and dissociation 75 % C ₄ H ₈ abs 15 % HCN abs 10 % (2°) C ₄ H ₈ abs Eff = 98 %
 5	H ⁺ trans 39 % 2 × H abs 25 % (2°) H abs (2°) CH ₂ abs (2°) C ₂ H ₃ abs H ⁻ abs 17 % H ₂ O abs 16 % (2°) H abs H abs 3 % Eff = 74 %	I abs 76 % (2°) C ₃ H ₅ abs (2°) C ₃ H ₄ abs (3°) I abs (2°) I abs (3°) C ₃ H ₅ abs (3°) I abs CH ₂ abs 13 % C ₃ H ₅ abs 6 % C ₃ H ₄ abs 5 % Eff = 53 %	SCH ₃ abs 100 % (2°) HSCH ₃ abs (2°) SSCH ₃ abs (2°) SCH ₃ abs (3°) SCH ₃ abs Eff = 72 %	H ⁺ trans and dissociation 98 % HCN abs 2 % Eff = 98 %

[a] Reaction efficiencies^[3,5] (Eff) are reported as $k_{\text{reaction}}/k_{\text{collision}} \times 100\%$. The reaction efficiency is the percentage of collisions leading to a reaction.
 [b] abs = abstraction, trans = transfer, dissociation = dissociation, add = addition; secondary and tertiary products are noted as (2°) and (3°), respectively, and are listed under the primary and secondary products that produce them.


Scheme 1.

reaction characteristic of **4**, was not observed. This finding unambiguously rules out the 2,6-isomer.

A likely ionic resonance structure^[14] of **5**, structure **6**, explains the different rates observed for the three I atom abstractions. This structure indicates a stronger coupling between the radical sites at positions 2 and 4 (or 4 and 6) than at 2 and 6, and suggests that the radical site at position 6 (or 2) should be more reactive than the others. The second I atom abstraction is slow as a result of the strong spin–spin coupling between the



remaining radical sites. However, once the second I atom is abstracted, a phenyl radical (analogous to **1** and **2**) is produced that undergoes a fast I atom abstraction.

The triradical **5** is calculated to have a doublet ground state (D-Q gap: $-26.8 \text{ kcal mol}^{-1}$; BD(T)/cc-pVDZ//UBPW91/cc-pVDZ), which explains why it undergoes radical reactions faster than the analogous biradical **3** whose reactions are hindered by its singlet ground state (based on a comparison of reactions with allyl iodide and dimethyl disulfide where no proton-transfer reactions were observed; note that deprotonation of **5** is calculated to generate a doublet triradical). The triradical reacts with allyl iodide and dimethyl disulfide at about the same efficiency as the monoradicals. Apparently, the large EA of **5** counterbalances most of the radical reactivity hindering effect caused by the coupling of the three unpaired electrons.

Experimental Section

The precursors of monoradicals **1** and **2** were purchased from Sigma Aldrich Co. and used as received. The precursors of the biradicals **3**^[15] and **4**^[16] and the triradical **5**^[16,17] were synthesized using reported methods. The mono-, bi-, and triradicals were generated in a dual-cell Fourier-transform ion cyclotron resonance mass spectrometer by using previously reported methods.^[5,9] Sustained off-resonance irradiated collision-activated dissociation^[18] was used to cleave C–I bonds in the protonated precursors. After isolation, **1–5** were allowed to react with reagents for varying periods of time to determine the second-order reaction rate constants (k_{exp}) and reaction efficiencies ($k_{\text{exp}}/k_{\text{collision}}$) as described previously.^[3,5,19] The absolute values are estimated to be accurate only within $\pm 50\%$, but the relative values are much more accurate ($\pm 10\%$). The structures of all the radical species were confirmed by using structurally diagnostic reactions.^[5,9] All DFT, coupled-cluster, and MCSCF calculations were carried out with the Gaussian 98^[20] and MOLCAS^[21] electronic structure program suites.

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