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Impact of Excited-State Antiaromaticity Relief in a Fundamental Benzene Photoreaction Leading to Substituted Bicyclo[3.1.0]hexenes

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access to complex molecular scaffolds that are difficult to access with reactions in the electronic ground state. While benzene is aromatic in its ground state, it is antiaromatic in its lowest $\pi\pi^*$ excited states. Herein, we clarify to what extent relief of excited-state antiaromaticity (ESAA) triggers a fundamental benzene photoreaction: the photoinitiated nucleophilic addition of solvent to benzene in acidic media leading to substituted bicyclo[3.1.0]hex-2-enes. The reaction scope was probed experimentally, and it was found that silyl-substituted benzenes provide the most rapid access to bicyclo[3.1.0]hexene derivatives, formed as single isomers with three stereogenic centers in yields up to 75% in one step. Two major mechanism hypotheses, both involving ESAA relief, were explored through quantum chemical calculations and experiments. The first mechanism involves protonation of excited-state benzene and subsequent rearrange-



ment to bicyclo[3.1.0] hexenium cation, trapped by a nucleophile, while the second involves photorearrangement of benzene to benzvalene followed by protonation and nucleophilic addition. Our studies reveal that the second mechanism is operative. We also clarify that similar ESAA relief leads to puckering of S₁-state silabenzene and pyridinium ion, where the photorearrangement of the latter is of established synthetic utility. Finally, we identified causes for the limitations of the reaction, information that should be valuable in explorations of similar photoreactions. Taken together, we reveal how the ESAA in benzene and 6π -electron heterocycles trigger photochemical distortions that provide access to complex three-dimensional molecular scaffolds from simple reactants.

INTRODUCTION

Benzene is the archetype aromatic molecule.¹⁻³ Yet, while its reactivity in the electronic ground state (S_0) is well-established, there are many aspects that remain to be explored and exploited related to its photophysics and photochemistry. The inherent stability of benzene in the S₀ state is attributed to its aromaticity, as given by Hückel's 4n+2 rule. In contrast, it displays a marked reactivity in the lowest excited states,⁴⁻¹⁰ and this reactivity change can be attributed to a switch between aromaticity and antiaromaticity whereby benzene becomes antiaromatic in these states.¹¹ The reversal of Hückel's rule was derived qualitatively for the lowest $\pi\pi^*$ triplet state (T₁) by Baird in 1972,^{12,13} and later extended through quantum chemical computations to the lowest $\pi\pi^*$ singlet excited state (S_1) of benzene.^{14,15} The impact of excited-state aromaticity and antiaromaticity for a series of different photophysical and photochemical phenomena has been unraveled in recent years.^{16,17} We now explore to what extent these concepts can be used to rationalize a benzene photoreaction that provides

access to complex three-dimensional molecular structures. We also show how the hypothesis on excited-state antiaromaticity (ESAA) relief applies to the photochemical rearrangements of two 6π -electron heterocycles (silabenzene and pyridinium ion) with S₁ states of $\pi\pi^*$ character.

A connection between the photoreactivity and relief of T₁state antiaromaticity through structural distortions has been revealed as the origin for triplet-state photoreactions of a few different benzene derivatives,^{18–21} and also shown to impact on the bimolecular photoreactivity of benzene and small polycyclic aromatic hydrocarbons (PAHs) in their T₁ states.²² In contrast to the (sensitized) T₁-state reactivity of benzene

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derivatives, herein we probe if structural rearrangements of benzene itself, observed upon direct irradiation to its S_1 state, are also triggered by ESAA alleviation. This aspect of benzene photochemistry has never been analyzed earlier, yet in the analysis one should also consider what other factors impact on the shape of the S_1 -state potential energy surface (PES) of benzene.

When irradiated with 254 nm light, benzene rearranges in the S_1 state to benzvalene and fulvene, and when irradiated by light in the range of 165–200 nm, it also forms Dewar benzene.²³ Irradiation of benzene by 254 nm light in nucleophilic solvents leads to derivatives of bicyclo[3.1.0]-hexene 1 (Scheme 1),²⁴ a reaction that was established for

Scheme 1. Photochemical Rearrangement and Addition of Nucleophilic Solvent to Benzene under Acidic Conditions Leading to Bicyclo[3.1.0]hexenyl Derivatives



1,3,5-tri-tert-butylbenzene,²⁴ benzene,²⁵ and tert-butylbenzene several decades ago.²⁵ However, the low reaction yields, the long reaction times, and the observed complex mixtures of products are severe drawbacks. Hence, the products of the photorearrangement of simple substituted benzenes have never been utilized synthetically in an applied context. A renewed analysis of the reaction, including considerations of excitedstate antiaromaticity relief and the electronic structure of the S₁ state, can provide insights that potentially are useful for synthetic method development. Here, it can be noted that the bicyclo[3.1.0]hexane scaffold is a useful building block for the construction of molecules with rigid three-dimensional shape and unique biological activity, such as glutamic acid analogues,²⁶ β -arabidofuranosyl derivatives,²⁷ nucleosidebased pharmaceuticals,²⁸ and nucleic acid analogues.²⁹ Thus, the photoreaction investigated herein turns a simple flat molecule into a complex three-dimensional scaffold with potential utility in one step.

Formally, the photorearrangement of benzene to 4-methoxybicyclo[3.1.0]hex-2-ene (1, Nu = OMe) in methanol represents an addition of a proton and a methoxide ion onto benzene. The reaction is accelerated in acidic media,³⁰ as compared to the neutral solution, and one can thus argue that it is catalyzed by acid and that protonation could play an important role in the reaction mechanism. Yet, if this is the case, how does protonation play a role?

The photochemical reaction of benzene in nucleophilic media leading to substituted bicyclo[3.1.0]hexenes previously gained considerable attention, and several different mechanisms were proposed.^{4,25,31-33} Our hypothesis is that the ESAA character of benzene impacts its photoreactivity in one of two ways: either through increased basicity (proton affinity) facilitating protonation of benzene in its S₁ and T₁ states followed by structural rearrangements (mechanism A, Scheme 2), or through the unimolecular rearrangement of S₁-state antiaromatic benzene to the prefulvene conical intersection which leads to benzvalene (mechanism B). A third possible mechanism in which benzene is protonated in the S₀ state,

Scheme 2. The Two Mechanisms Explored, A and B, for the Photorearrangement of Benzene in Nucleophilic Media (here MeOH) Leading to 4-Methoxy-bicyclo[3.1.0]hex-2-ene $(1)^a$



^{*a*}Note that benzene in the T_1 state can act as a triplet sensitizer of benzvalene 4, which ring-opens to the S_0 state of benzene.

followed by excitation of the formed benzenium ion, occurs only in an extremely strong superacid, FSO₃H-SbF₅-SO₂CIF.³⁴ In fact, benzene in the S₀ state remains unprotonated even in very strong acids such as triflic acid.³⁵ Two other mechanistic proposals earlier turned out to be invalid—more specifically, mechanisms that involved either prefulvene as a reactive intermediate^{4,31} or a *cis,cis,trans*-cyclohexa-1,3,5-triene with Möbius topology.³⁶

With this foundation we used a combined computational and experimental approach to reinvestigate the photochemical rearrangement of benzenes to bicyclo[3.1.0]hexenes in nucleophilic media under acidic conditions. A complex product with several stereogenic centers and further possibility of functionalization is formed as a result of a single photochemical step from simple and inexpensive starting materials. In this context, we demonstrate the potential of the reaction through preparation of an optically pure single isomer of a bicyclo[3.1.0]hexene derivative in two steps starting from benzene. Although the reaction still has apparent limitations for synthesis, the deepened mechanistic insights could allow for the future identification of conditions and reagents that pave the way to similar photoreactions with widened scopes for synthetic applications. Focus in our investigation is given to the impact of ESAA on the photoreactivity of simple substituted benzenes, and it is revealed that relief of S1-state antiaromaticity is a strongly contributing factor to the observed photochemical deformation of the benzene ring, which ultimately leads to the substituted bicyclo[3.1.0]hexenes.

RESULTS AND DISCUSSION

We begin with an exploration of the reaction scope and the description of the isolation of an enantiomerically pure bicyclo[3.1.0]hexene derivative. The two mechanistic alternatives of Scheme 2 are subsequently investigated. We also

consider two 6π -electron heterocycles, silabenzene and the pyridinium ion, and place the findings on these in relation to what we observe for substituted benzenes. A lower symmetry of the S₁-state electronic structure than that of benzene is key to achieving a high selectivity in the photorearrangements, and such a symmetry lowering is provided by heteroatoms and certain substituents.

Reaction Scope. We first re-examined the synthetic potential of the reaction. The earliest studies reported low reaction yields (few percent) and long irradiation times (weeks).^{24,25} Since then, several substrates (benzene,^{25,33} toluene, *t*Bu-benzene,^{25,37} *o*-xylene, tri-*t*Bu-benzene,²⁴ benzonitriles,^{38–40} and trifluoromethylbenzene and toluenes³⁹) have been reported to undergo the photoinduced rearrangement and addition of methanol,^{24,33,41} acetic acid,^{25,32} water,^{25,32} and 2,2,2-trifluoroethanol^{24,25,32,38–40} under acidic conditions (Figure 1). The quantum yields for several of the substrates were low (~0.05).^{24,30} Also, the published reactions have never been used before for synthetic applications.



Figure 1. Previously reported (*i*) substrates, (*ii*) nucleophiles for photoinduced rearrangement and addition to simple arenes, and (*iii*) observed products.

Irradiation of simple arenes leads not only to a mixture of regio- and stereoisomers of substituted bicyclo[3.1.0]hexenes but also to fulvene and its polymers^{4,31} as well as to phototransposed products.^{39,42} Furthermore, the bicyclic photoproducts 1 are unstable and decompose to fulvenes and polymeric material in acidic as well as in basic media.²⁵ Moreover, different isomers of 1 are known to interconvert between each other, either by acid-catalyzed epimerization³² and racemization,³² by the triplet-benzene-sensitized vinyl cyclopropane rearrangement,³³ or by the bicyclo[2.1.1]hexene rearrangement³⁷ (Scheme 3, pathways a–d). Consequently, the photoproduct composition changes over the course of the reaction as secondary photochemical and thermal processes scramble the isomers, leading to complex mixtures. Still, some isomers are more stable than others, and they accumulate in the system after prolonged irradiation.

We now optimized the reaction conditions for the rearrangement of benzene in acidic water and acetic acid, as well as that of *t*Bu-benzene in acidic methanol (Scheme 4). The reaction yields were optimized by choosing an optimal (i)

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Scheme 3. Rearrangement Mechanisms of Bicyclo[3.1.0]hexenes: (a) Acid-Catalyzed Epimerization,³² (b) Acid-Catalyzed Racemization,³² (c) Sensitized Vinylcyclopropane Rearrangement,³³ and (d) Sensitized Rearrangement through [2.1.1]Hexenyl Derivative 6³⁷







"Reaction conditions: 70 mM substrate, Rayonet reactor, FEP tube reactor, irradiation time 30 min, (a) 1 mM AcOH (aq.), (b) AcOH (conc.), (c) 1.25 mM dichloroacetic acid in MeOH. Isolated reaction yields of the major isomerization product are shown.

concentration of the substrate, (*ii*) amount and strength of the acid used, and (*iii*) irradiation time (for details see Supporting Information, section 2). We also ran the reaction in a FEP (fluorinated ethylene propylene) tubing flow reactor which minimized the irradiation of the reaction mixture and limited the secondary photoprocesses. Finally, we checked the substrate scope of the reaction by testing a series of simple substituted aromatic compounds (Scheme S1 and Table S3). However, for most of the tested substituents the reaction either did not proceed (polar groups, *O*- and *S*-containing arenes) or led to very complicated mixtures and/or to other photochemical transformations (anilines and halobenzenes).

It should be noted that the limits in the substrate selectivity have previously been studied computationally for benzonitriles and anisoles.⁴³ For methylbenzonitrile, prefulvenic biradical

structures were found to be stabilized by the cyano group through π -conjugation, allowing for interconversion between the different isomers. In methylanisoles, on the other hand, the photoisomerization is not possible because biradical prefulvenic structures cannot be formed due to the lack of π conjugation between the C4 atom and the methoxy group, which results in higher barriers in the S1 state. Yet, we found that one substituent, the silvl group, yields arenes with suitable properties for this reaction. Silyl-substituted benzenes react efficiently, and their reactivity increases with higher alkyl substitution at the silicon atom (for the reactivity of a series of methylated phenylsilanes $PhSiH_nMe_{3-n}$ (n = 0, 1, 2, or 3), see Table S4 and Figure S1). Indeed, a test for a linear free energy relationship between the $\log(k/k_0)$ and the Hammett σ constants of the silvl substituents of the four silvl-substituted benzenes reveals an increase in reaction rate with increased electron-donating ability of the substituent, yet the relationship is nonlinear, reflecting an impact of steric bulk on the reaction rate (see Figure S2).

Trimethylsilylbenzene (TMS-benzene) had the optimal reactivity, while dichloroacetic acid (DCA) was found to have the optimal acidity (more acidic media cause decomposition of the formed products; less acidic media slow down the photoreaction). Interestingly, the main silylated photoproduct 1d had a different constitution from the *t*Bu-analogue 1c, which possibly can be rationalized by either (*i*) elimination of the TMS group from the other positions occurring in the photoequilibration or by different substituent effects, or (*ii*) stabilization of a transition-state structure or intermediate by the silyl group (*vide infra*).

While the substrate scope of the photorearrangement is limited and the reaction conditions need to be carefully optimized, the reaction generates highly complex threedimensional structures from simple and easily accessible substrates in one step. Consequently, we attempted to prepare a single enantiomer of the bicyclo[3.1.0]hexenyl photoproduct. Whereas direct methods of enantioselective synthesis turned out to be unsuccessful due to racemization of the photoproduct in the acidic reaction media, we also attempted the chiral resolution of the product (\pm) -1a. Enzymatic kinetic resolution by esterification with Candida antarctica lipase B (CALB) was attempted, yet it could not be optimized to give high enantiomeric excess (ee) (Supporting Information, section 4.4). Instead we managed to prepare diastereomeric esters 7 with Boc-L-Phe (Scheme 5) which could be successfully separated by HPLC. Vibrational circular dichroism spectra⁴⁴ were measured on the separated diastereoisomers in CDCl₃ solution. Comparison of experimental and calculated spectra as well as an X-ray crystal structure determination enabled assignment of the absolute stereochemistry. Further details can be found in the Supporting Information, section 3.

Mechanism Investigations. The two mechanistic pathways shown in Scheme 2 (pathways A and B) both involve excitation of benzene to S_1 as the first step. One may consider intersystem crossing, but it has been shown experimentally through quenching with various triplet-state quenchers that the photorearrangement does not proceed in the T_1 state,⁴ and we can confirm this computationally (*vide infra*). Moreover, when benzene is excited to the S_2 state, Dewar benzene is formed in addition to benzvalene and fulvene.²³ Hence, one can conclude that an S_1 -state process is likely. In this state, benzene either becomes protonated to form a benzenium cation that rearranges to the bicyclo[3.1.0]hexenium cation 3 (pathway

Scheme 5. Synthesis of Diastereomeric Esters 7



A, Scheme 2), or rearranges to benzvalene 4^{45} via the prefulvenic transition state 5 and an S_1/S_0 conical intersection (pathway B).⁴⁶

Both mechanistic pathways involve relief of the S1-state antiaromatic character of benzene, but in different ways. Many pieces of evidence have been acquired for pathway A^{31,32} and pathway B,^{24,33} but no previous study has unambiguously confirmed one mechanism over the other. With regard to ESAA relief, any pathway that allows for disruption of the 6π electron cycle which is antiaromatic in S₁ will provide for such relief. Which pathway is followed depends on (i) the excitedstate lifetime of S₁-state benzene, as this determines the effective concentration of the reactive S_1 -state benzene, (*ii*) the concentration and strength of the acid, and (iii) the activation barriers for the different reaction steps. Thus, ESAA relief combined with other factors (concentrations and excited-state lifetimes) impacts which mechanism is followed. Further factors that can have an impact are the energy penalty for distortion of the σ -framework from the hexagon, which is ideal for sp²-hybridized C atoms, and the ability of substituents to reduce the antiaromaticity of a benzene ring in its S_1 state, similar to what was recently found for the T1-state antiaromaticity.47

Mechanism A. In pathway A, the ESAA is alleviated through protonation, as this breaks the cyclic π -conjugation and transforms the S₁-state benzene into an initially non-aromatic benzenium ion with a linear pentadienyl cationic segment with four π -electrons (Scheme 6). On the basis of such ESAA relief, we postulate that benzene is more basic and has a higher proton affinity in its antiaromatic S₁ state than in its S₀ state. Indeed, benzene has a reported proton affinity which is higher in the S₁ state than in the S₀ state by 26 kcal/mol.⁴⁸ This is in line with an earlier observation on magnitude reversals in proton and hydride affinities of $4n\pi$ - and $(4n+2)\pi$ -electron (benz)annulenyl anions and cations, respectively, caused by switches in aromaticity/antiaromaticity when going from their lowest singlet to their lowest triplet states.⁴⁹

With regard to the benzenium cation in the S_1 state, previous computations revealed that this species adopts a markedly puckered structure, placed close to an S_1/S_0 conical intersection.⁵⁰ This puckered benzenium cation could decay to the S_0 state and collapse to bicyclo[3.1.0]hexenium cation 3

Scheme 6. Formation of the S₁-State Homoaromatic Structure through Relaxation of the Excited Benzenium Cation



(Scheme 6). Subsequent nucleophilic attack on this cation would occur from the less sterically hindered *exo* side. Now, why does the benzenium ion pucker in the S₁ state? Recently, we explored the existence of homoaromaticity in the excited states, primarily in the T₁ state, but a few molecules with S₁-state homoaromaticity were also identified.⁵¹ Thus, a driving force for the puckering of the benzenium ion should be the attainment of excited-state 4π -electron homoaromaticity, which is also in line with the disrotatory path for a photochemically allowed electrocyclic ring-closure of a linear polyenyl chain with four π -electrons.^{52–54}

One may ask if a T_1 -state homoaromatic benzenium ion could be involved, but as noted above, earlier experiments showed that the reaction proceeds in the S_1 state.⁴ Further support for this conclusion is found through computations (Figure 2). Although the benzenium ion has a clear T_1 homoaromatic structure,^{S1} decay from this C_s -symmetric structure to the S_0 state leads back to the C_{2v} -symmetric benzenium ion **2** and not to the bicyclo[3.1.0]hexenium cation **3**. Also, there is no stationary point in the T_1 state that corresponds to a bicyclo[3.1.0]hexenium ion (the vertically excited T_1 -state structure of **3** is a non-stationary point 58.9



Figure 2. Reaction coordinate for benzenium cation (2) leading to bicyclo[3.1.0]hexenium cation (3) in the S₀ and T₁ states calculated at the (U)B3LYP/6-311+G(d,p) level.

kcal/mol above the S_0 state). Thus, if mechanism A is operative, it must involve formation of a benzenium ion in the S_1 state. Now, to what extent does protonation of S_1 -state benzene lead to ESAA relief?

S₁-State Protonation and ESAA Relief. Clearly, excitedstate protonation of benzene disrupts the antiaromaticity, as revealed by changes in the NICS(1)_{zz} values $[\Delta NICS(1)_{zz} =$ NICS(1; benzene)_{zz} – NICS(1; benzenium cation)_{zz}] between benzene and the benzenium ion in their S₁ and T₁ states when optimized with $C_{2\nu}$ symmetry. This reveals large reliefs in ESAA as the benzenium ion in $C_{2\nu}$ symmetry is non-aromatic $(\Delta \text{NICS}(1)_{zz} = -55.6 \text{ and } -28.3 \text{ ppm} \text{ in } S_1 \text{ and } T_1$ respectively, see Table 1; for DFT results, see Table S34). In contrast, protonation of benzene in the S₀ state leads to a loss of aromaticity (Δ NICS(1)_{zz} = 16.0 ppm). This antiaromaticity relief occurs upon protonation of benzene in the S_1 and T_1 states, while aromaticity loss upon protonation in the S₀ state is reflected in the higher calculated proton affinities in the excited states (Table 1), in line with the earlier experimental finding. Furthermore, through comparison of the calculated proton affinities of benzene with those of cyclooctatetraene (COT, Table 1), it is revealed that the proton affinities of benzene in the two excited states are similar to that of COT in the S_0 state. Interestingly, while COT upon protonation in S₀ turns into the aromatic homotropylium ion,⁵⁵ the benzenium ion puckers to C_s symmetry and turns homoaromatic in T_1 and possibly also in S_1 . The same applies for the proton affinities of the planar D_{8h} -symmetric COT in the S₁ and T₁ states when compared to those of benzene in S_0 (Table 1). Opposite effects of aromaticity and antiaromaticity, when compared to benzene, are observed in the S_0 and T_1 states of COT, while its S_1 state displays a different pattern according to NICS, likely due to methodological issues, as it follows the expected trend when other indices are used (see the Supporting Information, section 7.10).

Fluorescence Quenching Attempts. Protonation of benzene in S_1 seems plausible, but in order to examine mechanism A experimentally, we performed a fluorescence quenching study of benzene and TMS-benzene in anhydrous methanol. The fluorescence of benzene is known to be quenched by dissolved oxygen,⁵⁶ a phenomenon which has not yet been fully explained.^{57,58} The quenching was therefore performed with degassed solutions. However, no fluorescence quenching was observed in a concentration range of the acid up to 1.25 M (anhydrous HCl in MeOH) for both TMS-benzene and benzene (Figures 3 and S32, respectively). This finding reveals that there is no interaction of the acid with benzene in its S_1 state. As the photorearrangement occurs in the S_1 state (vide supra),⁴ it can be concluded that excited-state protonation is not operational despite the enhanced proton affinity of S₁-state benzene. On the other hand, in experiments with benzene dissolved in THF/9-BBN mixtures, where 9-BBN is used as a Lewis acidic co-solvent with a potential ability to coordinate to S₁-state benzene, an increased fluorescence quenching is observed with higher concentration of 9-BBN (see Figure S33).

So why is there no fluorescence quenching upon increased concentration of anhydrous HCl? The excited-state lifetime of benzene in the S₁ state is 28 ns,⁵⁹ and with the conditions used in our experiments, the effective concentration of S₁-state benzene (based on its lifetime) can be estimated to merely $\sim 10^{-13}$ M (see Supporting Information, section 5.4). Thus, there is an extremely low concentration of benzene molecules

Table 1. Calculated Gas-Phase Proton Affinities (PA, kcal/mol) and Changes in NICS (Δ NICS(1) _{zz} , ppm) ι	upon Protonation
of Benzene and Cyclooctatetraene (COT) in the S_0 , S_1 , and T_1 States ^{<i>a</i>}	

	S ₀		S1		T_1	
compound	PA	$\Delta \text{NICS}(1)_{zz}$	PA	$\Delta \text{NICS}(1)_{zz}$	PA	$\Delta \text{NICS}(1)_{zz}$
benzene	(183.1) $(181.3)^d$	16.0	214.7 $(C_{2\nu}, i)^b$ 232.1 $(C_s, HA)^b$ $(207.3)^d$	-55.6	217.0 $(C_{2\nu} i)^b$ 218.4 $(C_{s\nu} HA)^b$	$-28.3 (C_{2\nu} i)^{b}$ -48.2 (C _s) ^b
СОТ	220.8 $(C_s, HA)^c$	-6.7^{c}	182.8	-18.7	195.0	15.3

^{*a*}Proton affinities at (U)- and (TD-)B3LYP levels and Δ NICS(1)_{zz} values at CASSCF//B3LYP. NICS(1)_{zz} at CASSCF//B3LYP, B3LYP//B3LYP, and CASSCF//CASSCF levels are given in Table S34. For further computational details and additional discussion, including electronic aromaticity index (MCI and FLU), see Table S36 and section 7.9 in the Supporting Information. Δ NICS(1)_{zz} = NICS(1; non-protonated)_{zz} – NICS(1; protonated)_{zz}. ^{*b*}Benzenium cation in the T₁ and S₁ states has homoaromatic (HA) minimum (C_s), but the values for PA and Δ NICS related to the proton addition to the planar benzene are also given and indicated as "(C_{2w} *i*)" (*i* = imaginary frequency). The NICS(1)_{zz} value of the C_s-symmetric S₁ structure cannot be determined as it is too close to the conical intersection. ^{*c*}The COT and COTH⁺ minima are strongly puckered, and COTH⁺ has homoaromatic (HA) character. NICS(1)_{zz} values for both puckered and planar structures are given in Table S34. ^{*d*}Experimental values from ref 48.



Figure 3. Fluorescence spectra of TMS-benzene (c = 1.0 mM, $\lambda_{\text{exc}} = 254 \text{ nm}$) in anhydrous MeOH, degassed (solid line, purged with argon) and non-degassed (dashed line) in the absence and presence of anhydrous HCl (c = 0-1250 mM). The absorption spectra are shown in the inset.

in their S₁ states available for protonation (note here also that protonation of benzene in the S₀ state occurs only in exceptionally strong superacids).^{34,35} Thus, if mechanism A is followed, it should require an essentially barrierless S₁-state protonation.

Indeed, protonation of benzene in its S1 state is barrierless according to our CASPT2//CASSCF calculations (Figure \$37). Furthermore, the proton migration within MeOH should be rapid, as the free energy of activation for proton transfer in a (MeOH)₈H⁺ cluster, used as protonated methanol clusters $(MeOH)_n H^+$ saturate at n = 8 or 9,⁶⁰ is calculated to be merely 1.3 kcal/mol with an implicit solvation model. This resembles the experimentally determined activation barriers (2-3 kcal/ mol) for proton migration in water and methanol.^{61,62} Additionally, concerted proton transfer in both solvents can involve tunneling.^{63,64} Thus, the computational data speak for mechanism A but the fluorescence quenching does the opposite. Clearly, there are features that make mechanism B more beneficial, the most apparent being the concentration of benzvalene in the S₀ state (included in mechanism B), which is ${\sim}10^{-3}$ M, 45 i.e., 10 orders of magnitude higher than that of S1state benzene ($\sim 10^{-13}$ M). Protonation of benzvalene in the S₀ state, provided it proceeds over a low activation barrier, can therefore be much more efficient than protonation of S₁-state

benzene. Indeed, it has been observed earlier that benzvalene is rapidly attacked by nucleophilic methanol to yield the methoxy adducts 1 in the S_0 state.⁴⁵

Mechanism B. This pathway begins with a unimolecular rearrangement to benzvalene 4 via the prefulvenic transition state, and the puckering of the benzene ring leads to attenuation of the cyclic π -conjugation and relief of ESAA. $NICS(1)_{zz}$ values computed at each point along the intrinsic reaction coordinate (IRC) in the S_1 state show that antiaromaticity is alleviated along this pathway (Figure 4). Clearly, benzene in the S₁ state is antiaromatic, as evidenced by a highly positive $NICS(1)_{zz}$ value of 99.5 ppm at the vertically excited structure and 80.9 ppm at the S₁-state minimum. The prefulvenic transition state, on the other hand, is already moderately influenced by aromaticity, as the structure of highest energy along the S₁-state CASPT2//CASSCF surface has a NICS $(1)_{zz}$ value of -10.1 ppm, while at the CASSCF transition-state NICS(1)_{zz} = -21.2 ppm (for S₀-state benzene NICS(1)_{zz} = -27.4 ppm). Thus, puckering of benzene in S₁ alleviates ESAA, yet is ESAA the factor that triggers the structural deformation?

Origin of the Activation Barrier and ESAA Relief. The S₁state barrier observed computationally for the photochemical transformation of benzene to the prefulvenic S_1/S_0 conical intersection, leading to benzvalene, is well-established from experimental and theoretical studies.⁶⁵⁻⁶⁷ More specifically, the barrier lies \sim 3000 cm⁻¹ (8.6 kcal/mol) above the S₁ planar minimum, and as soon as the excess vibrational energy is enough to overcome it, the non-radiative process known as "channel 3" is opened. $^{68-75}$ Access to "channel 3" leads to the loss of fluorescence and opening of the pathway that leads to the prefulvenic conical intersection and to benzvalene. Now, in order to obtain a reasonable agreement between experiment and computations with regard to the activation barrier, both static and dynamic electron correlations have to be included in the computations-while a CASPT2//CASSCF calculation gives a barrier of 9.4 kcal/mol, the barrier is exaggerated at the CASSCF level (18-20 kcal/mol).^{65,67} Yet, why is there an activation barrier if ESAA is relieved?

Benzene in the S_1 state (${}^{1}B_{2u}$) is D_{6h} symmetric and described to equal extents by two electron configurations coupled in-phase, while the out-of-phase combination constitutes the higher-energy ${}^{1}E_{1u}$ state. It has been concluded that the S_1 state is strongly stabilized by the in-phase



Figure 4. (A) Potential energy surface of benzene in the S₁ state from S₁ planar minimum (D_{6h} symmetry) until the S₁/S₀ conical intersection. The experimentally determined activation energy (~8.6 kcal/mol) given in ref 69 is indicated by a line. (B) The NICS(1)_{zz} at the various structures along the IRC. Geometries were calculated at the SA4-CASSCF(6,6)/ANO-RCC-VTZP level, energies were obtained at the MS4-CASPT2(6,6)/ANO-RCC-VTZP level, and NICS(1)_{zz} values were determined at the CASSCF(6,6)/6-31++G(d,p) level. The plane defined by the C1, C2, C4, and C5 atoms was used to define the central position of the ghost atom located 1 Å below the plane (displayed as a green dot in the inset) for NICS. For computational details and NICS values in the opposite direction, see Figure S47.

coupling.^{76,77} Upon distortion away from the D_{6h} symmetry toward the prefulvenic structure, the S_1 state (former ${}^{1}B_{2u}$) increases in energy while the higher-energy (former ${}^{1}E_{1u}$) state decreases in energy, leading to an avoided crossing and an activation barrier on the S1 PES along the puckering coordinate. Interestingly, it has been revealed that the activation barrier vanishes when going to the less symmetric pyrazine,⁷⁷ and a lack of activation barrier has also been observed for silabenzene.⁷⁸ We now confirm this feature for silabenzene at the CASPT2//CASSCF level (Figure 5), and also for the pyridinium cation, which has a photochemistry $\frac{79-84}{79-84}$ which is of higher synthetic value than that of pyrazine.⁷⁹ Yet, at the lower CASSCF level, which contains only static electron correlation, there are still activation barriers, although they are smaller than for S_1 -state benzene. Thus, the activation barrier observed in the S1 state of benzene is traceable to its high symmetry (for comparisons between the S₁ states of benzene, silabenzene, and pyridinium ion, see Table S19). Here, it should be pointed out that the structural distortion going from the S₁-state minimum to the transition-state structure is much less unfavorable in S₁ than in S₀ because the

energy difference between these two structures in S₀ is 84.3 kcal/mol (see Figure S48), compared to 9.4 kcal/mol in the S₁ state. This large difference in energy loss (74.9 kcal/mol) upon an identical distortion in S₀ vs in S₁ indicates that the puckering goes with aromaticity loss in S₀ and antiaromaticity relief in S₁. Yet, a precise assessment of the various contributions is not possible, as this energy penalty also includes energy contributions to distort the σ -orbital skeleton of the sp²-hybridized C atoms from its ideal hexagon (see Figure S48 and the related discussion in the Supporting Information for an estimate).

Similarly to benzene, drastic decreases in S₁ antiaromaticity, determined by $NICS(1)_{zz}$, are observed for the S₁-state silabenzene and pyridinium ion when going from the CASSCF S_1 -state minima toward the S_1/S_0 conical intersections (Figure 5C,D). Notably, S₁-state silabenzene at the CASSCF level has an optimized nonplanar structure with a considerably reduced antiaromaticity (33.3 ppm) compared to that of S₁-state benzene at its D_{6h} minimum (80.9 ppm). Yet, when regarding the NICS $(1)_{zz}$ of vertically excited silabenzene and the pyridinium ion (99.3 and 91.2 ppm, respectively), it is apparent that the ESAA relief prior to the CASSCF S₁-state minimum of silabenzene is extensive. Clearly, ESAA alleviation seems to be a general process for 6π -electron cycles with S₁ states of $\pi\pi^*$ character. The photochemistry of both silabenzene and pyridinium salts is established (for a summary, see Supporting Information, section 6),^{85,86} and the mechanism and synthetic applications of pyridinium ion photochemistry (Scheme 7) have been reviewed extensively.⁷ Hence, the ESAA relief of 6π -electron cycles is a feature that triggers these molecules to undergo photorearrangements that can be of general synthetic utility. For example, 6azabicyclo[3.1.0]hexene, formed photochemically from pyridinium salts, can efficiently be transformed to polysubstituted cyclopentenes by acid-promoted ring opening (Scheme 7).^{87,8}

Substituent Effects. Now returning to benzene, with mechanism B and the fact that lowered symmetry lowers the activation barrier for puckering, we can explain why the TMSand tBu-substituted benzenes have higher reaction rates than the parent benzene. First, the increased reaction rate of TMSbenzene is not an effect of an increase in S₁-state antiaromaticity when the electron-donating ability increases because the calculated S₁-state antiaromaticity of TMSbenzene is similar to that of benzene $(NICS(1)_{zz} = 78.8 \text{ and})$ 80.9 ppm, respectively). Instead, when benzene, TMSbenzene, and tBu-benzene are calculated at the same computational level, the activation barrier for puckering of benzene in the S₁ state is 9.9 kcal/mol, while the barriers for TMS-benzene and tBu-benzene are 7.6-9.7 and 7.8-9.9 kcal/ mol, respectively, depending on the position of the substituent group (see Table S26). Thus, the lowest activation barrier for puckering of the S1-state TMS-benzene, leading toward the prefulvenic conical intersection, is ~2 kcal/mol lower than for S₁-state benzene. This difference becomes slightly larger when considering ΔG^{\ddagger} , as the puckering toward the prefulvenic conical intersection with the TMS group at the 6-position is 3-4 kcal/mol lower than the free energies of activation along the other paths and \sim 7 kcal/mol lower than that of benzene (Table S26). Additionally, the C_{Ar} -Si(Me₃) bond at this transition-state structure of TMS-benzene is distinctly shorter (1.865 Å, Figure S43) than at the other transition-state structures (1.888-1.895 Å), indicating a hyperconjugative interaction.



Figure 5. Potential energy surfaces at the CASPT2//CASSCF and CASSCF levels from the S_1 minimum to the S_1/S_0 conical intersection for (A) silabenzene and (B) the pyridinium cation. Respective NICS(1)_{zz} for (C) silabenzene and (D) the pyridinium cation. For further computational details, see Supporting Information, section 7.2.

Scheme 7. Photochemistry of Pyridinium Salts and Acid-Promoted Ring Opening of the Bicyclic Product^a



As seen for silabenzene and the pyridinium ion, heteroatoms distort the description of the S1 state so that one of the two electron configurations that are degenerate in the S₁ state of benzene becomes more dominant. This can also be achieved to a smaller extent by certain substituents at the benzene ring, such as the TMS group (Table S24). As a consequence, these substituted benzenes become guided by their electronic structures toward one of the transition-state structures, and at the transition state, as just noted, substituents that are able to interact hyperconjugatively provide for stabilization. Thus, there are benzene-to-prefulvenic S₁-state pathways for the two substituted benzenes that involve slightly lower activation barriers than what is the case for the parent benzene. In summary, substituents that (i) lead to a difference in the weights of the two electron configurations that describe S1state benzene and (ii) are able to interact through (hyper)conjugation with the C6 atom are useful as they will provide access to primarily one of the prefulvenic S1/S0 conical intersections. The TMS group is one such group; others will be reported in forthcoming studies.

Stereochemistry of Mechanism B. Based on mechanism B, how can the observed stereochemistry be rationalized? The preferential formation of exo adducts has previously been used as the main argument for the intermediacy of bicyclo[3.1.0]hexenium ion 3 (mechanism A in Scheme 2),³¹ as the nucleophile should attack from the sterically less hindered exo face. This was explored by Berson et al., who found the main observed photoproduct of irradiation of benzene in acidic D₂O to be exo-8, which was found to be formed by epimerization of the primary photoproduct endo-8 (Scheme 8a).32 He also observed selective endo incorporation of deuterium in position 6 and no addition of deuterium from the exo site leading to 9. We now confirm this through a study of the photoreaction of benzene- d_6 in aqueous solvent under acidic conditions (Scheme 8b). The only observed product was 10, which has the H atom in the 6-endo position and the hydroxyl group in the 4-exo position. This excludes any intermediacy of the benzenium cation (mechanism A), as its puckering would scramble the position of the deuterium/protium label in the 6 position (Scheme 8c). Instead, it is consistent with the concerted attack on the benzvalene tricyclic structure and the subsequent epimerization (Scheme 8d). This can be either concerted or stepwise involving a contact ion-pair. The epimerization goes via protonation of the MeO group, and the exo-8 isomer is lower in energy by 1.4 kcal/mol than the endo-8 when using methanol as implicit solvent (Figure S40). The epimerization of endo-8 to exo-8 is therefore a thermodynamically controlled process and does not scramble the isotope label at the 6-position.

Indeed, the proton affinity of benzvalene in its S_0 state is high: 240 kcal/mol in the gas phase at the B3LYP/6-311+G(d,p) level, i.e., higher than that of S_1 -state benzene and even higher than that of S_1 -state tBu- and TMS-benzene (see Table S38). This supports protonation of benzvalene in the S_0 state, as its concentration in our experiments is 10 Scheme 8. Isotopic Labeling Experiments of Photoaddition of Water to Benzene



orders of magnitude higher than that of S_1 -state benzene (*vide supra*). Moreover, the addition reaction is stepwise according to our computations, progressing with an activation free energy of 7.4 kcal/mol in the first step. The MeOH₂⁺ approaches from

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kcal/mol (exo) and 15.5 kcal/mol (endo), respectively. Cause of Low Quantum Yield. Finally, one can ask why is the quantum yield for the photoreaction not higher than 0.045?³⁰ Experiments reveal that a photostationary state evolves in a solution with both benzene and benzvalene present. Furthermore, earlier experiments give $E(T_1; benzene)$ = 84.1 kcal/mol⁸⁹ and $E(T_1; benzvalene) \approx 65 kcal/mol⁹⁰$ while our (U)B3LYP/6-311+G(d,p) computations give $E(T_1)$ for the two compounds at 83.5 and 60.3 kcal/mol, respectively. Also, the calculated energy needed to reach the vertically excited T₁ state of benzvalene (79.3 kcal/mol) is slightly lower than the triplet energy of benzene (83.5 kcal/mol). As benzene has a non-negligible quantum yield for intersystem crossing $(\Phi_{\rm ISC} = 0.25)^{91}$ it will exist in both its S₁ and T₁ states when excited. Thus, due to its high $E(T_1)$ and the energy match, benzene can transfer its T_1 energy to benzvalene. Now, when benzvalene becomes excited to the T1 state, it will rearrange to T_1 -state benzene through a series of transformations, progressing over low activation barriers according to our calculations (Figure 6). Furthermore, this rearrangement of T_1 state benzvalene to T₁-state benzene, via a T₁-state prefulvene intermediate instead of a transition state as on the T_1 PES (vide supra), is highly exothermic (~60 kcal/mol). Combined, this means that benzene has the ability to catalytically sensitize the back-reaction from benzvalene to benzene, i.e., act as a catalytic sensitizer. Thus, based on the computed S_0 and T_1 PESs, it becomes clear that the low quantum yield (Φ =



Figure 6. Back-reaction from benzvalene to benzene initiated by formation of T_1 -state benzvalene through triplet energy transfer from T_1 -state benzene (inset), and the T_1 and S_0 potential energy curves for its rearrangement to T_1 benzene, calculated at the (U)B3LYP/6-311+G(d,p) level (electronic energies include ZPE corrections; vertically excited energies given in parentheses are purely electronic energies). Energies given in red are relative to S_0 -state benzene, and energies given in blue are relative to S_0 -state benzene.

 $(0.045)^{30}$ could be overcome if T_1 -state benzene (or a derivative) were unable to transfer its triplet energy to benzvalene.

To probe the ability of benzene to act as a catalytic sensitizer, we investigated the effect of the acid and benzene concentrations on the reaction rate and yield. Benzvalene has earlier been shown to give methanol adduct 1 in acidic methanol (Scheme 2),⁴⁵ yet as just noted, benzvalene rearranges to benzene upon T_1 benzene-induced sensitization.⁹² Indeed, high concentrations of TMS-benzene resulted in low yields of the bicyclic photoproducts (for details see Table S1), and the reaction also occurred inefficiently in pure methanol without addition of acid. Thus, benzvalene undergoes a slow reaction in neutral methanol, and this process is enhanced by acid catalysis. A way to increase both the reaction yield and the quantum yield for formation of the bicyclo[3.1.0]hexene would therefore be to identify (hetero)-benzenes which in their T_1 states are unable to transfer their triplet energies to the corresponding benzvalene analogues.

Taken together, the experimental and computational evidence reveals unequivocally that mechanism B is operative, knowledge that could be useful for the rationalization and development of similar photoreactions.

CONCLUSIONS AND OUTLOOK

It is known that benzene becomes antiaromatic in its S₁ and T₁ states, ^{11,14–16} and that relief of this destabilizing antiaromatic character is a driving force for photochemical reactions of various benzene derivatives in their T1 states.^{20,22} Herein we show that S1-state antiaromaticity relief of the benzene molecule itself is strongly involved in the photochemical process that leads up to formation of substituted bicyclo[3.1.0]hex-2-enes in nucleophilic media under acidic conditions. Through combined quantum chemical and experimental studies, we deduce that the first step in the mechanism involves the puckering of S₁-state benzene, which after a conical intersection leads to benzvalene. Importantly, the puckering of S_1 -state benzene to the prefulvenic S_1/S_0 conical intersection is strongly connected to relief of S₁-state antiaromatic character, despite the fact that it passes over an activation barrier of 9.4 kcal/mol. Similar S1-state antiaromaticity relief occurs in silabenzene and the pyridinium ion-and in these species without activation barriers, a feature that results from a lowered symmetry of the electronic structure. Certain substituents at the benzene ring (e.g., a trimethylsilyl group) can, similarly to a heteroatom, lower the activation barrier so that S₁-state TMS-benzene preferentially follows the path to the prefulvenic conical intersection with the TMS group in the 6-position.

When in the S_0 state, nucleophilic attack by MeOH at the benzvalene yields the substituted bicyclo[3.1.0]hex-2-enes, with the highest yield (75%) for TMS-benzene. The synthetic accessibility to the complex bicyclo[3.1.0]hexenyl scaffold offers an interesting possibility for its further utilization in synthetic applications. Generation of three chiral centers in one simple step is often difficult in thermal chemistry. We show this to be a result of a process triggered by ESAA relief of benzene photoexcited to its S_1 state.

We further reveal limiting factors which, if they can be bypassed or removed, indicate how the benzene photochemistry can be further enhanced, as there are obvious drawbacks with the reaction. First, rearrangements of the initially formed products lead to scrambling of substituents. Moreover, the quantum yield for product formation is low ($\Phi = 0.045$),³⁰ as benzene in the T₁ state ($\Phi_{ISC} = 0.25$) transfers its T₁ energy to benzvalene, which subsequently rearranges in the T₁ state and thus regenerates the sensitizer in a unique process of catalytic sensitization. One should identify a 6π -electron (hetero)cycle with lower T₁ energy than the corresponding benzvalene analogue, hindering the re-aromatization.

In conclusion, our combined experimental and computational investigation puts an end to a long debate over the mechanistic photochemistry of benzene, reveals the impact of excited-state antiaromaticity relief in the S_1 state of benzene, and opens up possibilities for new studies within the field of photochemistry of 6π -electron (hetero)cycles.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.9b13769.

Materials and methods, optimization and control experiments, synthetic procedures, reactivity studies, mechanistic experiments (fluorescence quenching, Stern–Volmer analysis, isotope labeling experiments, concentration calculations), computational details, Cartesian coordinates, structures, energies, electronic configurations and the respective weights (for CASPT2//CASSCF calculations), aromaticity indices (NICS(1)_{zz}) MCI, and FLU values), and additional plots for PES and NICS(1)_{zz} for benzene (list of compounds included: benzene and photochemical derivatives, protonated benzene, TMS-benzene, tBu-benzene, silabenzene, pyridinium cation, cyclooctatetraene, and protonated cyclooctatetraene), including Figures S1–S70 and Tables S1–S38 (PDF)

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Notes

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