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## Introduction

We recently reported the mild, thermal polycyclization reaction of triynes (*cf.* **1**, Scheme 1) to give complex multi-ring benzenoid products (*cf.* **2**).<sup>1</sup> We call this two-stage process an HDDA cascade. The first step involves generation of a reactive benzyne derivative (*cf.* **3**) by a net 4 + 2 cycloaddition event,<sup>2,3</sup> which we have termed a hexadehydro-Diels–Alder (HDDA) reaction.<sup>1</sup> This benzyne intermediate is then efficiently (and, in the reaction type that is the subject of this paper, intramolecularly) trapped, resulting in the expeditious construction of highly substituted benzene derivatives (*e.g.*, **2**).

Among the hallmarks of this HDDA cascade are: (i) the transformation of triynes like 1 to benzynes like 3 is computed to be exothermic by *ca.* 50 kcal mol<sup>-1</sup> (!)<sup>1</sup> (*cf.*  $\Delta E_{rxn} = -51.4$  kcal mol<sup>-1</sup> for the parent ethyne + 1,3-butadiyne to benzyne<sup>4</sup>); (ii) this thermal mode of benzyne generation is complementary to all previously reported, preparatively practical methods in two important ways—the triyne precursors are not, themselves, benzene derivatives and the aryne is formed in the absence of reagents (*e.g.*, bases or reducing agents) and byproducts (*e.g.*, metal salts, amines, or halide ions) that can affect or, in some

## Tactics for probing aryne reactivity: mechanistic studies of silicon–oxygen bond cleavage during the trapping of (HDDA-generated) benzynes by silyl ethers<sup>†</sup>

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We report mechanistic aspects of the trapping of thermally (HDDA) generated benzyne derivatives by pendant silyl ether groups, which results in net insertion of the pair of benzyne  $C_{sp}$ -hydribized carbon atoms into the silicon-oxygen sigma bond. Cross-over experiments using symmetrical, doubly labeled bis-silyl ether substrates established that the reaction is unimolecular in nature. Competition experiments involving either intramolecular or intermolecular dihydrogen transfer clock reactions (from within a TIPS isopropyl group or cyclooctane, respectively) vs. the silyl ether cyclization were used to gain additional insights. We evaluated effects of the steric bulk of the silyl ether trapping group and of the ring-size of the cyclic ether being formed (furan vs. pyran). These types of competition experiments allow the relative rates of various product-determining steps to be determined. This previously has only rarely been possible because aryne formation is typically rate-limiting, making it challenging to probe the kinetics of subsequent trapping reactions. Solvent effects (polarity of the medium) and computational studies were used to probe the question of stepwise vs. concerted pathways for the Si–O insertion.

instances, interfere with the benzyne trapping reactions; and (iii) a beneficial consequence of point (ii) is that inherent reactivity of benzynes can sometimes be probed more fundamentally and/or that new types of aryne trapping reactions can be uncovered.<sup>5,6</sup> An example of this latter feature is the net addition of a silyl ether (like that in 1) across the pair of strained benzyne sp-hybridized carbons to give an *o*-(trialkylsilyl)aryl



Scheme 1 The hexadehydro-Diels-Alder (HDDA) cascade of triyne 1 gives the highly substituted benzenoid derivative 2 *via* silyl ether trapping of the HDDA-generated benzyne 3.

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ether moiety (like that in 2). Such a process had been previously unknown.<sup>1</sup>

## **Results and discussion**

To account for the trapping reaction that produces 2, we suggested<sup>1</sup> that the benzyne 3 is intercepted by the silyl ether oxygen atom poised five atoms away and that a transient zwitterionic species like 4 is generated. Migrations of silicon from carbon to oxygen are well known in the form of the Brook rearrangement.<sup>7</sup> The opposite process, the reverse (or retro-) Brook rearrangement (i.e., silicon migration from oxygen to carbon), is less common but certainly precedented.8 These reverse Brook reactions are most often encountered within an anionic manifold of intermediates. In a setting like that of 4 to 2, rearrangement would be expected to be particularly favorable since zwitterionic character is quenched as a result of the silyl group migration. A second possibility is that the reaction mechanism involves zwitterion 4 but is not unimolecular. That is, disproportionation of that intermediate with, e.g., another copy of itself or with the silvl ether moiety in substrate 1 could be responsible for formation of 2. A third possibility is that the oxygen-silicon bond adds in concerted (and intramolecular) fashion to the benzyne  $\pi$ -bond in 3, leading directly to 2. This last possibility is of interest because concerted insertion into a sigma bond with accompanying formation of two new sigmabonds (e.g., A-B + X=Y  $\rightarrow$  A-X-Y-B in a single step) is a rare type of transformation.

To probe these mechanistic issues, we prepared the set of three symmetrical tetraynes **6a–c** from the diol precursor **5** (Scheme 2). These differ only in the nature of the alkyl groups in the trialkylsilyl moiety of their silyl ethers. Diol **5** was prepared by Cadiot–Chodkiewicz cross-coupling reaction of the known *N*,*N*-bispropargyl toluenesulfonamide<sup>9</sup> with 4-bromo-3-butyn-1- $ol^{10}$  in the presence of catalytic CuCl in piperidine.<sup>1,11</sup> Bis-silylation of diol **5** gave each of the corresponding bis-TES, -TBS, and -TIPS ethers **6a–c**. As we anticipated, each of these



Scheme 2 Preparation of substrates 6a-c and the efficient HDDA cyclization of each to benzofurans 7a-c.

substrates smoothly underwent a HDDA cascade to provide the corresponding tricyclic benzofuran derivatives **7a–c** when heated alone to the reflux temperature of chloroform. The rate of consumption of each substrate was virtually identical, consistent with the fact that benzyne formation is the rate-limiting step in these HDDA cascade processes. The slightly lower yield of the TIPS-containing product **7c** arises from a competing dihydrogen disproportionation reaction that will be discussed later.

With substrates 6a-c in hand, we designed several experiments to address the question of molecularity of the trapping reaction. First, to distinguish an intra-from an intermolecular process, we carried out a simple crossover/competition study using the pair of substrates 6b and 6c (Scheme 3, panel A). When equimolar amounts of these were heated together in chloroform, the only benzofuran derivatives observed were 7b and 7c. Neither of the analogous products containing one each of the TBS and TIPS groups (*i.e.*, 7d and 7d') was detected  $(\leq 0.5\%)$  by LCMS analysis of the crude product mixture. This result rules out a bimolecular process (*i.e.*, the "second possibility" presented above). A complementary crossover experiment was also performed, this time using the mixed mono-TBS/ mono-TIPS silvlated substrate 6d,12 which gave 7d and 7d' (Scheme 3, panel B). Again, there was no evidence of bimolecularity (crossover); that is, none of 7b or 7c was seen by LCMS analysis.

We then attempted to distinguish between the concerted *vs.* stepwise mechanisms for silyl ether trapping (Scheme 1 bottom, green *vs.* orange) by carrying out the reaction in several solvents of varying polarity. In the stepwise mechanism either of the two



Scheme 3 Crossover experiments establish unimolecularity.

fundamental events could, in principle, be rate-limiting. The rate constant for the first step in that mechanism, formation of the zwitterion **4** from benzyne **3** (*cf.*  $k_{S1}$ , Scheme 1), would be expected to be significantly increased whereas that of the second ( $k_{S2}$ ) decreased as the polarity of the medium is increased.

To assess the impact of solvent polarity on the rate, it was desirable to have a competitive trapping reaction that could be used as a benchmark against which to clock the rate of silvl ether trapping. Fortunately, one was at hand. The reaction of bis-TIPS ether 6c gives a byproduct accompanying 7c—namely, the isopropenyl-containing bicyclic arene compound 9 (Scheme 4, panel A). This arises by an intramolecular net redox process in which two hydrogen atoms have been transferred, intramolecularly, from a TIPS alkyl group to the aryne (cf. red arrows in the benzyne intermediate 8). We have recently reported that this type of unprecedented desaturation reaction is facile for bimolecular H2-transfer reactions to benzynes from certain cyclic hydrocarbons and that it proceeds by a concerted pathway in which all six atoms involved in the bond-breaking and -making events are essentially coplanar.6 Although the geometry of the transition structure (TS) 8 might seem unusual at first glance, notice that the hydrogen atoms being transferred occupy positions 9 and 11 of an array that can be mapped onto a bicyclo[5.3.1]undecane skeleton. Indeed, we were able to locate a computed TS geometry [M06-2X/6-31G(d)] for a variant of this transformation in which the benzyne is minimally substituted-namely, 8-TS (Scheme 4, panel B). Two hydrogen atoms are each partially transferred from the isopropyl to the benzyne sp-hybridized carbon atoms. Assuming that this concerted mechanism is operative in 8, one would expect little dependency on solvent polarity, rendering this competitive transformation a good choice as an internal clock reaction.

Accordingly, we carried out the HDDA cyclization of the bis-TIPS ether **6c** in four solvents (Scheme 4, panel C), each of which we have observed to be inert toward HDDA-generated benzynes. These solvents range in relative permittivity (dielectric constant) from 2.2–37.5 Debyes. As the polarity of the medium was increased, we observed a steady increase in the branching ratio (<sup>1</sup>H NMR analysis of the crude product mixture; *cf.* Scheme 4, panel D) in favor of formation of the silyl ether trapped product **7c** relative to the dihydrogen disproportionation product **9**. We interpret the *direction* of this trend as an argument against the possibility that the rate-limiting event in the silyl ether trapping is the second step in the stepwise mechanism in Scheme 1 (*i.e.*, that associated with the rate constant  $k_{S2}$ ). That event involves conversion of a zwitterion (**4**) to a neutral structure (**2**) and, thus, should be decelerated not accelerated in more polar solvents.

On the other hand, the *magnitude* of the observed rate enhancement in a more polar solvent medium is small. Under the assumption that the rate of formation of the alkene **9** is the same in each solvent, the fastest to slowest  $k_{rel}$ s for silyl ether cleavage differ only by a factor of *ca.* 3. The small size of this effect suggests only a minor degree of polarization in the transition structure for the rate-limiting step in the silyl ether trapping process. This is more consistent with an only modestly polarized and early TS, either for an asynchronous concerted



Scheme 4 Effect of solvent polarity on the silvl ether trapping using an internal clock reaction. The ratio of intramolecular dihydrogen transfer (6c to 9, panel A) to silvl ether trapping (6c to 7c) processes shows a dependence on solvent polarity (results tabulated in panel C). DFT calculations on a truncated analog led to the identification of the concerted TS 8-TS (panel B, non-participating aliphatic hydrogen atoms removed for clarity). <sup>1</sup>H NMR spectrum of the crude product mixture from a typical experiment (panel D), this one for the reaction performed in 1,2-dichloroethane, resulting in a 6 : 1 ratio of products 7c : 9.

process (3 to 2 and  $k_{\rm C}$  in Scheme 1) or for zwitterion formation (3 to 4 and  $k_{\rm S1}$  in Scheme 1).

To gain more information relevant to the possibility of single-step insertion into the Si–O sigma bond, we also examined this benzyne trapping process computationally. We studied each of the model benzyne-containing substrates I and VI (Fig. 1, panels A and B, respectively). Each comprises a simple trimethylsilyl ether and a mono-substituted benzyne, and the two differ only in the length of the methylene chain (two *vs.* three  $CH_2s$ ) linking the aryne and silyl ether moieties. Benzyne I



Fig. 1 Computed energies of minima and transition structures on the potential energy surface for conversion of model benzynes I and VI into silyl ether trapped products V and VIII (panels A and B, respectively). Selected dihedral and internal bond angles are given for TSs II, IV, and VII. [M06-2X/6-31G(d); enthalpic energies (kcal mol<sup>-1</sup>) are given in panels A and B. Energies for each species using three basis sets are given in panel C.

leads to the benzofuran product **V**, the homolog **VI** to the benzopyran **VIII**. We used density functional theory (DFT) to map the reaction potential energy surface (PES) for each substrate/ product pair. The M06-2X functional was used for all calculations and three different basis sets were used (see electronic supplementary information<sup>†</sup>). We were able to locate a zwitterionic species as an intermediate only when we applied an

implicit (or continuum) solvation model (solvation model density, SMD<sup>13</sup>) during geometry optimization and, then, only in the five-membered series (*cf.* the pathway involving **I–V**). The barrier heights leading away from that zwitterion (**III**) are quite low, especially so in the forward direction, suggesting that it is a fleeting intermediate. Computational results (energies for **I–VIII** and geometries for TSs **II**, **IV**, and **VII**) are given in Fig. 1. The trends and conclusions are qualitatively the same regardless of the nature of the basis set that was used.

The overall transformation for either trapping reaction (I to V or **VI** to **VIII**) is highly exothermic (>50 kcal  $mol^{-1}$ ). The geometry of each of the initial (and early) TSs II or VII shows only a slight lengthening of the O-Si bond. The major difference in the two (cf. TS geometries in Fig. 1) is seen in the dihedral bond angle across the pair of benzyne carbon atoms and the silvl ether oxygen and silicon atoms ( $\angle_{1234} = 67^{\circ}$  in **II** vs. 30° in **VII**). TS II was able to settle into the zwitterionic intermediate III, in which this dihedral angle has been significantly reduced to 18°. This then proceeds *via* the second TS IV (dihedral =  $13^{\circ}$ ) to product V (dihedral =  $0^{\circ}$ ) via a second, low barrier and very exothermic elementary step. The TS VII for the six-membered substrate is very similar to III in both geometry and energy. However, it directly evolves into benzopyran **VIII** (dihedral =  $0^{\circ}$ ) without indication of encountering an intervening energy minimum species (i.e., an analogous zwitterionic intermediate). It is also of interest that the benzyne moiety in each of TSs II and VII has undergone an appreciably large, induced distortion of geometry in response to the approach of the oxygen nucleophile. Specifically, the two bond angles within the benzyne ring at atoms C1 ( $\angle_{a12}$ ) vs. C2 ( $\angle_{12b}$ ) reveal a large distortion of the benzyne in each (  $\angle_{12b} - \angle_{a12} = 31^\circ$  for II and 26° for VII). This induced distortion of the transition structure represents an extrapolation of previous explanations of ground-state benzyne distortions that effectively explain the site of greater electrophilic character in unsymmetrical benzynes.14

We have also assessed the dependency of the relative rate of intramolecular silyl ether trapping on the size of the silyl substituent. A different type of dihydrogen transfer competition experiment, this time *vis-à-vis* the external trapping agent cyclooctane,<sup>6</sup> was used. Heating each of **6a–c** to 65 °C in cyclooctane led to a product mixture containing both **7a/b/c** and the reduced benzenoid product **10a/b/c** (Scheme 5). The product ratios reflect the relative ease of trapping by the TES, TBS, and TIPS silyl ethers, respectively, within the benzyne intermediates **11a–c**. As expected, the least hindered TES is the fastest. Specifically, the TES ether traps the benzyne *ca.*  $5 \times$  faster than the TBS and  $20 \times$  faster than the bulkiest, TIPS ether (*cf.*  $k_{rel}$ s in Scheme 5). The magnitude of this effect is consistent with a TS for the product-determining step in which at least some degree of interaction with the silicon center has been established.

Finally, we used the cyclooctane external competition experiment to probe one additional feature—namely, the relative ease of furan *vs.* pyran formation in the silyl ether trapping event. Substrates **6b** and **12** differ in the number of methylenes (2 *vs.* 3, respectively) that tether the TBS ether to the alkyne (Scheme 6).<sup>15</sup> Each was heated in cyclooctane, and the ratio of cyclization to reduction products (*i.e.*, **7b** to **10b** *vs.* **13** to **14**) was



Scheme 5 Effect of steric bulk of the silvl group using an external clock reaction. Competition of intramolecular silvl ether trapping vs. bimolecular dihydrogen transfer from cyclooctane.



Scheme 6 Ring size effects. Competition of intramolecular silyl ether trapping vs. bimolecular dihydrogen transfer from cyclooctane (the external clock reaction).

measured. The ratios turned out to be identical, suggesting a compensating trade-off between entropic (favoring silyl ether trapping enroute to the furan 7b) and enthalpic effects (lower strain in the TS leading to pyran 13) for these two processes. It is notable that the computed activation enthalpies ( $\Delta H^{\ddagger}$ ) for the reactions passing through TSs II *vs.* VII (Fig. 1) are 3.1 *vs.* 1.8 kcal mol<sup>-1</sup>, respectively. The actual rate for each of these reactions would also be, of course,<sup>16</sup> dependent on the mole fraction of each of the reactive conformers I and VI *vis-à-vis* its associated family of all accessible conformations. The additional rotatable bond in the latter substrate reduces the mole fraction of VI (*vs.* that of I), thereby slowing its reaction rate to bring it more in line with that of I than is implied by the computed  $\Delta H^{\ddagger}$  values alone.

### Conclusions

In summary, we have used various competition and crossover experiments to gain mechanistic insights about the siliconoxygen bond cleavage event that attends trapping of HDDAgenerated benzynes containing a well-disposed, pendant silyl ether. The process is unimolecular. Use of dihydrogen transfer reactions, of both an intra- and intermolecular nature, was instrumental to the design of the competition experiments.

These types of competitive reactions allow one to probe the relative rates of various product-determining steps in the reactions of arynes. The rates of trapping reactions are often difficult to unravel because aryne formation is typically ratelimiting, making it challenging to probe the kinetics of the subsequent events. The silvl ether trapping reaction shows a modest rate enhancement in solvents of higher polarity, which we interpret as evidence for only a moderate degree of polarization in the transition structure for the rate-limiting step. Larger alkyl groups on the silyl ether slow the rate of trapping (ca. 20-fold between TES and TIPS). Precursors to benzofuran vs. benzopyran products cyclize at essentially the same rate, suggesting that enthalpic and entropic factors in these two types of tethers are self-compensating. Based on the computational results alone, one would conclude that the silvl ether trapping event in the benzofuran class of precursors occurs in stepwise fashion whereas the benzopyran homologs proceed via a concerted Si-O insertion. However, given the small energetic differences between these two pathways, this conclusion should probably be viewed with a degree of caution. Computed TS geometries suggest that the benzyne distorts in response to the electronic character of the approaching nucleophilic silvl ether oxygen atom. We speculate that this phenomenon may have ramifications extending to many other aryne trapping reactions. Collectively, the studies reported here show how HDDA chemistry can provide access to new fundamental mechanistic insights that are otherwise difficult if not impossible to obtain.

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