



## **Strained Molecules**

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## High Reactivity of Strained Seven-Membered-Ring trans-Alkenes

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**Abstract:** trans-Oxasilacycloheptenes are highly reactive strained alkenes. Competition reactions showed that these seven-membered ring trans-alkenes underwent [4+2] cyclo-addition reactions faster than a trans-cyclooctene. They also reacted with quinones and dimethyl acetylenedicarboxylate to form adducts with high diastereoselectivity. Kinetic studies showed that ring strain increases nucleophilicity by approximately  $10^{\circ}$ .

**S**train-promoted reactions have emerged as important transformations in organic synthesis<sup>[1]</sup> and chemical biology.<sup>[2]</sup> Among the strained systems investigated, the eight-membered-ring *trans*-alkenes have found numerous applications.<sup>[3-8]</sup> In contrast, studies of the reactivity of seven-membered-ring *trans*-alkenes are much less common because these compounds are particularly difficult to prepare.<sup>[9-11]</sup> Limited studies of the chemistry of *trans*-cycloheptenes demonstrate that they undergo transformations typical of reactions with acids.<sup>[12-14]</sup> They also participate in reactions observed for *trans*-cyclooctenes, such as [4+2] cycloadditions,<sup>[3,10]</sup> but those reactions were generally low-yielding.

Herein, we demonstrate that highly strained *trans*-oxasilacycloheptenes undergo fast, high-yielding, and, in most cases, stereoselective reactions. These strained alkenes are more reactive in a [4+2] cycloaddition than the most reactive *trans*-cyclooctene.<sup>[15,16]</sup> The tunable synthesis of these sevenmembered-ring *trans*-alkenes allows access to new strainpromoted reactions, including stereoselective reactions with substrates containing electron-deficient  $\pi$ -bonds.

Initial studies revealed that *trans*-alkene **1** was highly reactive in a [4+2] cycloaddition reaction. This sevenmembered-ring *trans*-alkene, which was synthesized from 1,3-pentadiene, benzaldehyde, and a di-*tert*-butyl silylene source,<sup>[14]</sup> reacted with diene **2** within five minutes (Scheme 1). Purification of the product afforded cycloadduct **3** in 98% yield. Although the cycloaddition occurred with retention of the geometry of the *trans*-alkene,<sup>[17]</sup> poor facial selectivity on the diene occurred.<sup>[18]</sup>

A competition experiment established the high reactivity of *trans*-oxasilacycloheptenes. Rates of cycloaddition reactions of the seven-membered-ring system were compared to *trans*-cyclooctene  $\mathbf{6}$ ,<sup>[19]</sup> a protected variant of a *trans*-cyclooctene reported to react faster than other strained alkenes in cycloadditions.<sup>[2,16]</sup> To control for steric effects,<sup>[20]</sup> the *trans*-



Scheme 1. Reaction of trans-oxasilacycloheptene 1 with diene 2.

alkene **4** was used.<sup>[21]</sup> Initial experiments were performed to characterize and isolate cycloadducts **5** and **7** ( $\geq$  97% isolated yield). The relative reactivities of the different strained alkenes were established by a competition experiment (Scheme 2). This experiment revealed that *trans*-alkene **4** reacted seven-times faster with diene **2** than *trans*-cyclooctene **6** did. The enhanced reactivity of alkene **4** is likely due to the higher distortion of the *trans* double bond<sup>[12]</sup> compared to *trans*-cyclooctene **6**, which is itself conformationally distorted.<sup>[16]</sup>

Seven-membered-ring *trans*-alkenes are so reactive that they undergo cycloaddition reactions with the highly sub-



Scheme 2. Competition reactions of diene 2 with trans-alkenes 4 and 6.

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stituted diene 2, even when the alkene is sterically hindered. Cycloaddition of diene 2 with trisubstituted alkene 8 required only 20 minutes to form cycloadduct 9 as a single diastereomer in 96 % yield [Eq. (1)]. This stereoisomer is likely favored because it minimizes developing steric interactions between a phenyl group of the diene and the fused six-membered ring of the alkene.





Scheme 4. Formation of enones 11 a and 11 b.

10a

8 d

25 °C

OTBS

6

Control experiments suggest that strain is responsible for the high reactivity of seven-membered-ring *trans*-alkenes. The less-strained *cis*-alkene isomer of **8** (*cis*-**8**) was prepared by photochemical isomerization of *trans*-**8** (Scheme 3). This alkene was unreactive in cycloaddition reactions even after 100 days.



Scheme 3. Synthesis of cis-8 and attempted reaction with diene 2.

The high reactivity of these compounds permitted the development of new strain-promoted reactions. Trisubstituted alkene 8 reacted with benzoquinone 10a to form enone 11a as a single diastereomer (Scheme 4).<sup>[22,23]</sup> There was even a reaction with the quinone 10b, which is over 1000-times less electrophilic than quinone 10a.<sup>[24]</sup> Although the enones 11a and 11b could arise from concerted ene reactions between the alkene and the quinone, these reactions likely proceed by stepwise mechanisms involving zwitterionic intermediates (intermediate A, Scheme 4).<sup>[24]</sup> As with cyclo-addition reactions, the alkene must be strained: the less strained alkene *cis*-8 did not react with 10a even after several days.

In comparison to *trans*-alkene **8**, *trans*-cyclooctene **6** was much less reactive. Even with the more reactive quinone **10a** [Eq. (2)], reactions were considerably slower, requiring eight



unidentified products

The reaction with quinone **10 a** permits the estimation of how much ring strain increases the nucleophilicity of *trans*alkene **8**. Kinetic studies of the reaction between alkene **8** and quinone **10 a** showed that the rate constant was comparable to that of the reaction between a silyl ketene acetal and quinone **10 a**.<sup>[24]</sup> Considering how much more nucleophilic silyl ketene acetals<sup>[25]</sup> are than trisubstituted alkenes,<sup>[26]</sup> the strain of alkene **8** accounts for a rate acceleration of approximately 10<sup>9</sup>.

The modular assembly of seven-membered-ring *trans*alkenes from dienes and aldehydes by silylene transfer<sup>[14]</sup> enables the synthesis of other electron-rich *trans*-cycloalkenes. Strained silyl enol ether **12**, the first example of a silyl enol ether incorporated into a *trans*-cycloalkene, was even more reactive with quinones.<sup>[27]</sup> Upon exposure to quinone **10 a** or **10 b**, phenols **13 a** and **13 b**, respectively, were formed as single stereoisomers [Eq. (3)]. These reactions, which likely involve addition of silyl enol ether **12** to the quinone followed by cyclization onto the resulting oxocarbenium ion intermediate,<sup>[24]</sup> preserved the geometry of the alkene in the product. The observation that unstrained cyclic silyl enol





ether **14** did not react with quinone **10a** after one week [Eq. (4)] indicates that the reactions of alkene **12** are strain-promoted.<sup>[28]</sup>

Seven-membered-ring *trans*-alkenes were also highly reactive with an electron-deficient alkyne (Scheme 5). *trans*-Enol ether **12** reacted with two equivalents of dimethyl



Scheme 5. Reactions of trans-cycloalkenes 12 and 8 with DMAD.

acetylenedicarboxylate (DMAD)<sup>[29]</sup> in less than ten minutes to form acetal **16** as a single diastereomer.<sup>[30]</sup> An enol ether was not required for this reaction: bicyclic alkene **8** reacted with DMAD in 15 minutes to form acetal **17** as a single stereoisomer (Scheme 5). These reactions likely proceed by conjugate addition to DMAD and ring-closure to form an oxocarbenium ion intermediate. Addition of a second equivalent of DMAD and subsequent cyclization forms the acetal product.<sup>[31]</sup>

The high reactivity of the *trans*-oxasilacycloheptenes with DMAD contrasts with the reactivity of this alkyne with other alkenes. In comparison to the seven-membered-ring system, *trans*-cyclooctene **6** reacted slowly (four days compared to  $\leq 15$  minutes) and unselectively with DMAD, resulting in a mixture of at least four products [Eq. (5)]. Only lactone **18** could be identified from that mixture. The slow and unselective reaction of *trans*-alkene **6** with DMAD is consistent with earlier studies of reactions of strained alkenes with this alkyne.<sup>[30,32]</sup> Unstrained silyl enol ether **14** and alkene *cis*-**8** also did not react with DMAD even after 45 days.



The products of reactions of *trans*-oxasilacycloheptenes can be functionalized to afford products without silicon atoms. Treatment of **13b** with a Lewis acid eliminated Me<sub>3</sub>SiOH [Eq. (6)], and desilylation in the presence of D<sub>2</sub>O provided the deuterium-labeled benzofuran **19**. The benzofuran moiety is found in compounds displaying *anti*-inflammatory and *anti*-tumor activities.<sup>[33]</sup>



In conclusion, seven-membered-ring *trans*-alkenes are highly reactive species in a number of reactions. They are more reactive in Diels–Alder cycloadditions than *trans*-cyclooctenes. Substituted *trans*-alkenes undergo rapid and stereoselective reactions with quinones and DMAD.

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