

## C–H Activation of Ethers and Alkanes by Gernylene–Aryl Halide Complexes

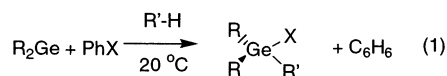
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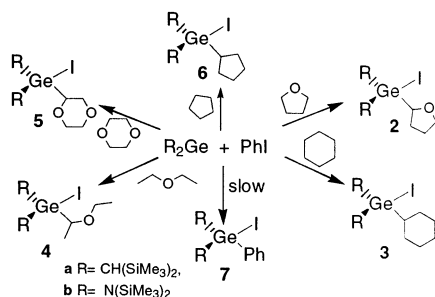
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Selective intermolecular C–H activations and insertions have been the focus of much research both in academia and industry.<sup>1–3</sup> Main group radical-based C–H activations of alkanes have focused on halogenation, sometimes under phase-transfer conditions.<sup>4,5</sup> Although these reactions are selective for tertiary and secondary C–H bonds, most of the tertiary activation products are unstable to the reaction conditions and are observed only in trace amounts.

We have recently discovered a novel C–H activation of ethers and alkanes using Ge[CH(SiMe<sub>3</sub>)<sub>2</sub>]<sub>2</sub> (**1a**) or Ge[N(SiMe<sub>3</sub>)<sub>2</sub>]<sub>2</sub> (**1b**) and PhX (X = I, Br, Cl) (eq 1).



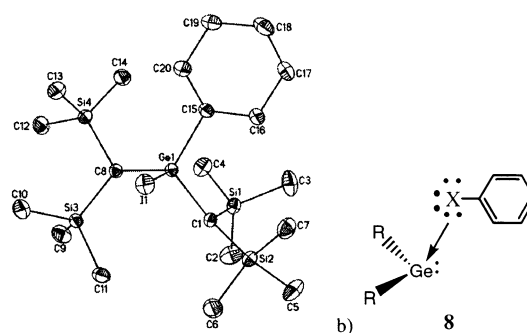
For ethers such as tetrahydrofuran (THF) ( $\alpha$ -C–H 92 (kcal/mol),  $\beta$ -C–H 97.6),<sup>7,8</sup> and Et<sub>2</sub>O ( $\alpha$ -C–H 89,  $\beta$ -C–H 112)<sup>9</sup> the weakest C–H bonds are selectively activated. Activation of the C–H bonds in 1,4-dioxane (96),<sup>7</sup> cyclohexane (97),<sup>7</sup> cyclopentane (93),<sup>10</sup> and butane (97, 98)<sup>11</sup> have also been observed (Scheme 1). Benzene,

**Scheme 1.** Activation of Alkanes and Ethers by GeR<sub>2</sub> and PhI

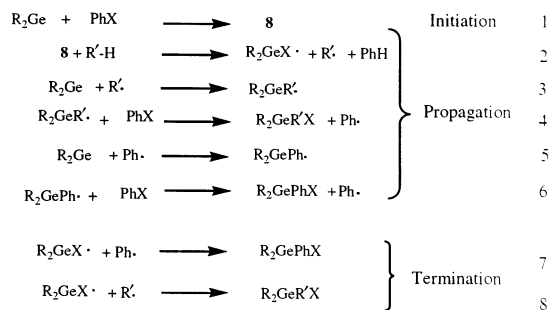
derived from the phenyl halide, is produced along with the C–H activation product. Labeling studies with deuterated solvents and PhI or 2-bromotoluene, show monodeuterio arene by GC/MS. No trace of radical coupling products was observed by <sup>1</sup>H NMR spectroscopy or GC/MS for the reactions shown in Scheme 1.

The CH-activation products are synthesized by slow, room-temperature addition of **1a** or **1b** (0.02 M) dissolved in the alkane or ether of interest to an equimolar solution of PhI dissolved in the same alkane or ether. For **1a**, the optimal rate of addition can be judged by noting that the yellow-orange gernylene color quickly fades to give a colorless solution before the next drop of gernylene is added. The products in Scheme 1 have been characterized by <sup>1</sup>H and <sup>13</sup>C NMR spectroscopies, elemental analysis, and in the case of **3a**, X-ray crystallography (Figure 1a).

Oxidative addition (OA) of aryl halide to gernylene is a concentration-dependent side reaction. At high concentration of



**Figure 1.** (a) ORTEP of **3a**. Selected bond lengths (Å) and angles (deg): Ge–I, 2.5946(4); Ge–C15, 1.9866(13); C1–Ge–C8, 107.10(5); C8–Ge–C15, 117.51(5); C8–Ge–I, 106.52(4); C15–Ge–I, 102.40(4). (b) Proposed intermediate.

**Scheme 2.** A Proposed Mechanism for C–H Activation Involving Free Phenyl Radical

reactants (0.2 M), the OA product, **7a**, is formed in up to 40% yield, and **7b**, in 82% yield. At low concentration of PhI (0.02 M) and **1a** (<0.03 mM) the C–H activation product is produced almost exclusively, with less than 1% of **7a** formed (as determined by <sup>1</sup>H NMR spectroscopy). Related oxidative addition products have been reported for Sn and Si analogues. Lappert has investigated the reaction chemistry of the analogous stannylene Sn[N(SiMe<sub>3</sub>)<sub>2</sub>]<sub>2</sub> toward alkyl and aryl halides and reported that the reaction of Sn[N(SiMe<sub>3</sub>)<sub>2</sub>]<sub>2</sub> with PhBr in THF gives a 9:1 ratio of [(Me<sub>3</sub>Si)<sub>2</sub>-N]<sub>2</sub>Sn(Ph)Br:[(Me<sub>3</sub>Si)<sub>2</sub>N]<sub>2</sub>SnBr<sub>2</sub>.<sup>12</sup> Similarly, West reported that reaction of a stable silylene with PhBr in hexanes results in oxidative addition and disilane formation.<sup>13</sup> In neither case were the analogous CH-activation products reported.

A proposed mechanism is presented in Scheme 2. Abstraction of a hydrogen radical by free phenyl radical is not supported by key experiments reported herein. Instead, we propose that the hydrogen radical abstraction occurs at the *ipso*-carbon of species **8** as the Ge–X bond and an incipient phenyl radical form and the C–X bond breaks. This is consistent with a recent theoretical analysis of the related West silylene chemistry.<sup>14</sup>

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The regioselectivity of this reaction has been studied using **1a** and PhI. Reaction with butane at  $-30\text{ }^{\circ}\text{C}$  results in a 1°:2° ratio of 0.08: 1 and reaction with methylcyclopentane at  $20\text{ }^{\circ}\text{C}$  gives a 2°:3° ratio of 1: 7.1. These ratios indicate a radical-like selectivity related to that observed for free phenyl radical generated by phenylazotriphenylmethane (PAT) at  $60\text{ }^{\circ}\text{C}$  (0.11–0.13:1.01; and 1:4.8 respectively).<sup>15</sup>

A difference in reactivity is observed when the halide is varied. Virtually all **1a** is consumed instantly upon addition to a PhI/THF solution. When PhBr is used, the reaction takes 3 days to go to completion. In the case of PhCl, the reaction is 33% complete after 12 days at room temperature, or complete after 2 days at reflux. This suggests that cleavage of the C–X bond is involved in the rate-limiting step.

Differing, detailed reactivity is observed for germynes **1a** and **1b**, and is attributed to their different Lewis acidities. Although both activate alkanes and ethers, **1b** reacts more slowly than **1a**. Parallel activations of methylcyclopentane run under identical conditions show that all of **1a** has been consumed, while up to 35% of **1b** is still observed via  $^1\text{H}$  NMR spectroscopy. Reactions run with **1b** produce more **7**. For example, when the synthesis of **2b** is undertaken employing the optimized conditions for **2a**, **7b** is produced in 13% yield, whereas the yield for **7a** is <1%. When aromatic substrates, such as toluene, ethyl benzene, or cumene are used, the products from benzylic C–H activation are observed when **1a** is employed. However, reaction of aromatic substrates with **1b** results only in slow formation of **7b**.

Competition experiments were performed at  $60\text{ }^{\circ}\text{C}$  using a 50: 50 mol:mol solution of cumene:THF- $d_8$ . When **1a** and PhI are added to the solution, a  $4.7 \pm 0.4$  ratio of C–H activation products is observed. When PAT is used to generate phenyl radical, the ratio of cumene/THF activation is  $3.4 \pm 0.3$ , suggesting that a species other than free phenyl radical is involved in the C–H activation initiated by **1a** and PhI. If this competition is performed using **1b** and PhI, only **7b** and the C–H activation product derived from THF- $d_8$  are observed; no trace cumene activation is observed. Free phenyl radical, generated by PAT, readily abstracts H radical from aromatic substrates such as cumene and toluene.<sup>15</sup> Hence, these results in which C–H activation of THF by **1b** and PhI occurs in the presence of cumene are inconsistent with formation of free phenyl radical. Had free phenyl radical formed, one would have expected to observe a  $\sim 3.4:1$  ratio of cumene/THF- $d_8$  C–H activation.

Competition experiments at  $60\text{ }^{\circ}\text{C}$  using a 1:1 ratio of THF: THF- $d_8$  with germylene **1a** and PhI gives a  $k_{\text{H}}/k_{\text{D}}$  of  $5.0 \pm 0.2$  as determined by GC/MS; for **1b** the ratio is  $4.1 \pm 0.2$ ; for phenyl radical generated by PAT the ratio is  $4.2 \pm 0.2$ . The different  $k_{\text{H}}/k_{\text{D}}$  values for **1a** and **1b** argue against a common intermediate, including free phenyl radical, being formed in reactions employing **1a** or **1b**. The different  $k_{\text{H}}/k_{\text{D}}$  values for **1a** and phenyl radical generated from PAT also argue against free phenyl radical being generated in the C–H activation. Although the  $k_{\text{H}}/k_{\text{D}}$  values for **1b** and PAT are not statistically different, the cumene/THF competition experiment above argues against free phenyl radical playing a role in the case of **1b**.

The competition experiments and the  $k_{\text{H}}/k_{\text{D}}$  ratio observed for **1a** are inconsistent with the formation of free phenyl radical. The experimental evidence is consistent with hydrogen abstraction occurring via an incipient radical present on the *ipso*-carbon of the phenyl ring in **8** (Figure 1b). The germynyl-X radical may then combine with the solvent radical to yield the observed products (step 8). Fast recombination within the solvent cage is consistent with the absence of R'–R' coupling products. Note that high

dilution conditions minimize the likelihood of reaction between R'• and GeR<sub>2</sub>, thus avoiding steps 3–7 and the manifold of reactions required to form OA product **7**. Species **8** is consistent with the different isotope effects observed for **1a** and **1b**, the differing isotope effect observed for **1a** and PAT, and the differing substrate reactivity observed. Steps 3–7 are similar to the mechanism proposed by Lappert for the reaction of stannylene chemistry and aryl halides.<sup>11</sup>

Single-electron transfer (SET) is not uncommon in germanium chemistry<sup>16,17</sup> and plays an important role in main group C–H activations as well. To test whether the reactions reported herein proceed by SET from PhI to Ge, forming a radical germynyl anion, the radical anion of **1a** was generated using sodium metal in THF according to the method of Gaspar et al.<sup>18,19</sup> No trace of a C–H activation product was observed by  $^1\text{H}$  NMR spectroscopy. Additionally, during the reaction of **1a** with PhI, UV–vis spectroscopy showed only loss of the germylene peak at 420 nm as the reaction progressed; the characteristic green color of the germynyl radical anion ( $\lambda_{\text{max}} = 666\text{ nm}$ ) was not observed. Thus, the reactions described in this communication likely do not involve radical anion species.

In summary, a new reaction for regioselective activation of C–H bonds using a germylene/aryl halide reagent has been discovered. The reaction works for both straight-chain and cyclic alkanes and a variety of ethers. In contrast to previously reported main group C–H activations that result in a carbon–halogen bond, these C–H activated products have a C–Ge bond. High yields of C–H activation products can be obtained through the use of high-dilution techniques.

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**Supporting Information Available:** Experimental details (PDF). An X-ray crystallographic file in CIF format for compound **3a**. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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