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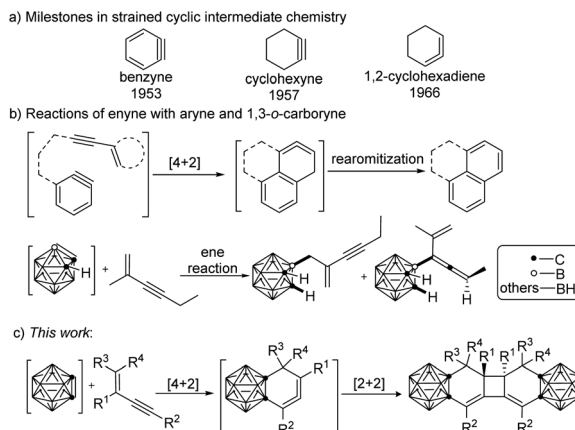
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## Introduction

Strained cyclic organic molecules, such as arynes, cyclic alkynes and cyclic allenes, have intrigued chemists for more than a century with their unusual structures and high chemical reactivity (Scheme 1a).<sup>1</sup> The considerable ring strain (30–50 kilocalories per mole)<sup>2,3</sup> that characterizes these transient intermediates imparts high reactivity in many reactions, including cycloadditions and nucleophilic trappings, often generating structurally complex products.<sup>4</sup> Cyclic allenes are a relatively less studied class of highly strained intermediate as compared with benzynes and cyclic alkynes. The generation and reactivity of 6-membered cyclic allenes such as the parent 1,2-cyclohexadiene have attracted much research interest in recent years.<sup>5</sup> Strained six-membered-ring allenes are also found as common intermediates in the [4 + 2] cycloaddition of conjugated enynes with unsaturated molecules, including alkenes and alkynes.<sup>6,7</sup> Aryne, a very reactive archetypal two-electron component in [4 + 2] cycloadditions, can react with conjugated enynes to generate isoaromatic cyclic allenes, which undergo isoaromatization to afford the polycyclic aromatics (Scheme 1b).<sup>8</sup>

On the other hand, carborane (*o*-C<sub>2</sub>B<sub>10</sub>H<sub>12</sub>), a three-dimensional relative of benzene, is a molecular boron–carbon cluster.<sup>9</sup> Owing to their unique properties, functionalized carboranes are now finding a broad range of applications

encompassing organic synthesis, drug design, polymers, cancer therapy, catalysis, metal–organic frameworks, electronic devices, and more.<sup>9–11</sup> Similarly, 1,2-dehydro-*o*-carborane (*o*-carboryne) can be viewed as a three-dimensional relative of benzene, which has been widely employed as a useful synthon for generating a wide range of functional carboranes over the past two decades.<sup>12</sup> It can undergo cycloadditions,<sup>13–15</sup> the ene reaction<sup>16</sup> and the C–H bond insertion reaction,<sup>17</sup> with a variety of organic molecules to afford a large class of functionalized carboranes. Cycloadditions involving an *o*-carboryne intermediate have been developed to enable the synthesis of various carbocyclic carborane derivatives.<sup>12</sup> In our recently reported work, the ene reaction was observed between 1,3-dehydro-*o*-carboryne and a conjugated enyne due probably to the polarized



**Scheme 1** Reactions involving the enynes, aryne and carboryne: (a) strained cyclic intermediates. (b) Reactions of enyne with aryne and 1,3-*o*-carboryne. (c) This work.

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“C=B” multiple bond (Scheme 1b).<sup>18</sup> Surprisingly, 1,2-dehydro-*o*-carboryne (*o*-carboryne) reacted with conjugated enynes in an unprecedented tandem [4 + 2]/[2 + 2] cycloaddition manner, generating a new class of rigid carborane-fused tricyclo [6.4.0.0<sup>2,7</sup>]dodeca-2,12-dienes (Scheme 1c). Herein, we reported a general method for the construction of such carborane-fused tricyclics.

## Results and discussion

In our initial study, the reaction of *o*-carboryne, generated *in situ* by treatment of 1-OTf-*o*-C<sub>2</sub>B<sub>10</sub>H<sub>11</sub> (**1**) with LiHMDS (lithium bis(trimethylsilyl)amide), with 1.2 equiv. of 2-methyl-1-hexen-3-yne (**2a**) in cyclohexane at room temperature afforded carborane-fused tricyclo[6.4.0.0<sup>2,7</sup>]dodeca-2,12-diene **3a** in 84% GC yield (Table 1, entry 1). Several common bases, such as organic lithium reagents and Grignard reagents, were screened (Table 1, entries 2–5), and the results suggested that LiHMDS was the best choice (Table 1, entry 1).

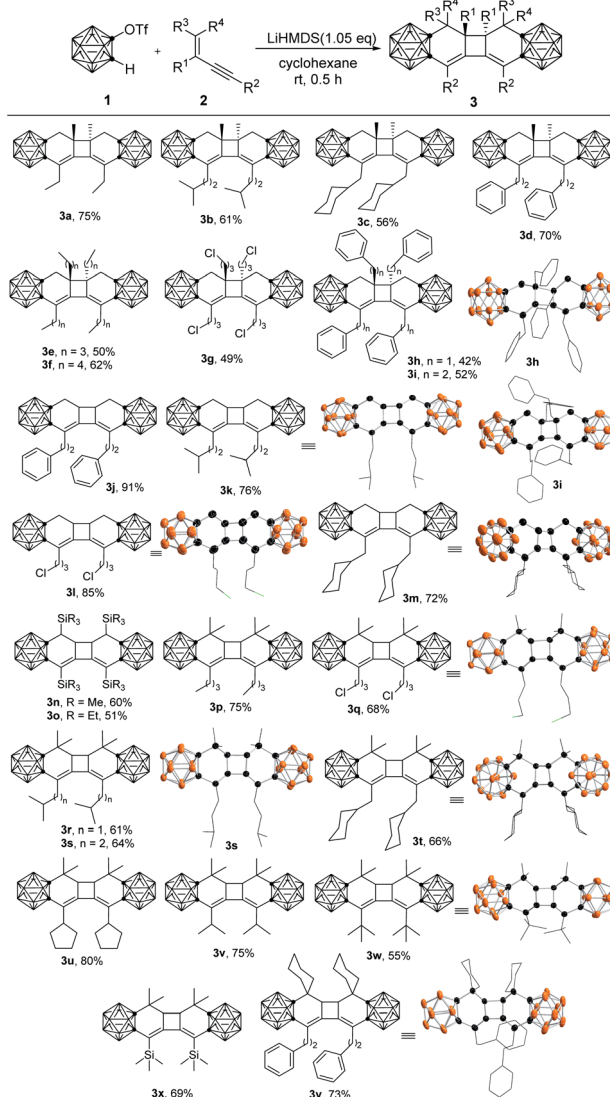
With the optimal reaction conditions in hand, the scope of this tandem [4 + 2]/[2 + 2] cycloaddition of *o*-carboryne with a series of conjugated enynes was examined and the desired carborane-fused tricyclic compounds were obtained in moderate to high yields (Table 2). Various substituents, including linear, branched and cyclic alkyl groups, silyl groups, and distal chloro and phenyl groups, were compatible with this reaction. It was noted that the reaction of the ornamented conjugated enynes directly with the phenyl group led to very complicated results due to the side reactions of the styrene<sup>14*i,l*</sup> or phenylacetylene<sup>15*b*</sup> moiety with *o*-carboryne. It was found that the steric hindrance of the substituents may play a role in the reaction, especially for the ones (*R*<sup>1</sup>) attached to the internal alkenyl carbon atom (**3a**–**3i** vs. **3j**–**3m**). For instance, the reaction of **2j** or **2l** proceeded smoothly to generate cycloadducts **3j** and **3l** in >85% yields, whereas **3d**, **3g** and **3i** were isolated in <70% yields. When the two terminal alkenyl C–H groups were replaced by methyl groups, the yield of the desired products decreased slightly (**3q** vs. **3l**, **3s** vs. **3k**, **3t** vs. **3m**, and **3y** vs. **3j**). Moreover, the substituents (*R*<sup>2</sup>) at the terminal alkynyl carbon atom showed no obvious effect on the yields of **3**.

Table 1 Optimization of reaction conditions<sup>a</sup>

Entry	Base	Yield <sup>b</sup> ( <b>3a</b> , %)
1	LiHMDS	84 (75) <sup>c</sup>
2	<sup>n</sup> BuLi	48
3	<sup>t</sup> BuOLi	33
4	<sup>i</sup> PrMgCl	35
5	CH <sub>3</sub> MgBr	21

<sup>a</sup> Reactions conditions: **1** (0.1 mmol), **2a** (0.12 mmol), base (0.105 mmol) in cyclohexane (2 mL), rt, 0.5 h. <sup>b</sup> Determined by GC. <sup>c</sup> Isolated yield.

Table 2 [4 + 2]/[2 + 2] Cycloadditions of *o*-carboryne with enynes<sup>a,b</sup>

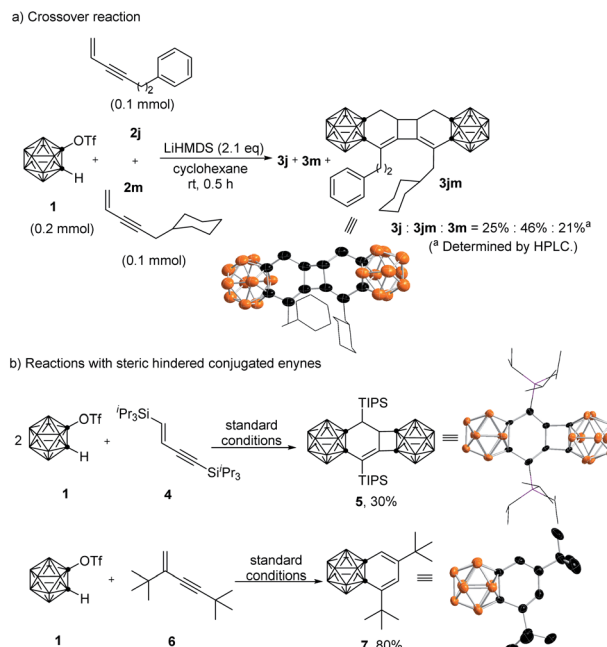


<sup>a</sup> Reactions conditions: **1** (0.1 mmol), **2** (0.12 mmol), LiHMDS (0.105 mmol) in cyclohexane (2 mL), rt, 0.5 h. <sup>b</sup> Isolated yields.

Compounds **3** were fully characterized by <sup>1</sup>H, <sup>13</sup>C, and <sup>11</sup>B NMR spectroscopy as well as by HRMS. The <sup>11</sup>B{<sup>1</sup>H} NMR spectra exhibited a 4 : 2 : 12 : 2 pattern for **3a** and **3d**, a 6 : 14 pattern for **3c**, **3e**–**j**, **3n**, and **3o**, and a 4 : 16 pattern for **3p**–**u** and **3y**, spanning the range  $\delta = -2$  to  $-14$  ppm. The molecular structures of **3h**–**i**, **3k**–**m**, **3q**, **3s**–**t**, **3w** and **3y** were further confirmed by single-crystal X-ray analyses.<sup>19</sup>

To gain some insight into the reaction mechanism, several control experiments were conducted (Scheme 2). Under standard reaction conditions, *o*-carboryne was treated with a mixture (1 : 1 molar ratio) of **2j** and **2m**, yielding three products **3j**, **3jm**, and **3m** in a molar ratio of around 1 : 2 : 1. The isolation of the crossover product **3jm** suggested that this is a stepwise process and the [2 + 2] cycloaddition step is an intermolecular reaction (Scheme 2a). On the other hand, 1,4-





Scheme 2 Control experiments: (a) crossover reaction. (b) Reactions with sterically hindered enynes.

bis(triisopropylsilyl)-1-buten-3-yne **4**, providing two highly sterically demanding silyl groups at the terminals of 1-buten-3-yne, reacted smoothly with *o*-carbonyne to afford **5** in 30% yield (Scheme 2b). In this reaction, the *in situ* generated carborane-fused 1,2-cyclohexadiene intermediate was trapped preferentially by less hindered *o*-carbonyne, which can be ascribed to the fact that the resultant sterically demanding cyclic allene intermediate prevents its dimerization.

Furthermore, the treatment of 2,4-bis(*tert*-butyl)-1-buten-3-yne **6** with *o*-carbonyne gave benzo-*o*-carborane **7** in 80% yield, which might result from the 1,3-H-migration of the extremely sterically demanding carborane-fused 1,3-di(*tert*-butyl)-1,2-cyclohexadiene intermediate (Scheme 2b). These results further supported that this reaction proceeded *via* a carborane-

fused 1,2-cyclohexadiene intermediate. The molecular structures of **3jm**, **5** and **7** were further confirmed by single-crystal X-ray analyses.<sup>19</sup>

Based on the above experimental results, a plausible reaction mechanism is proposed in Scheme 3. At first, *in situ* generated *o*-carbonyne **A** by reaction of **1** with LiHMDS, reacts with conjugated enyne **2** to form highly reactive carborane-fused 1,2-cyclohexadiene **B** *via* [4 + 2] cycloaddition.<sup>20</sup> Two molecules of cyclic allenes **B** undergo a stepwise [2 + 2] cycloaddition *via* a singlet bis-allyl diradical **C** intermediate,<sup>21</sup> affording the desired carborane-fused tricyclo[6.4.0.0<sup>2,7</sup>]dodeca-2,12-diene **3**. This proposed mechanism is supported by DFT calculations (see ESI† for details).

## Conclusions

Using 1-OTf-*o*-C<sub>2</sub>B<sub>10</sub>H<sub>10</sub> (**1**) as an *o*-carbonyne precursor, an unprecedented tandem [4 + 2]/[2 + 2] cycloaddition reaction of *o*-carbonyne with conjugated enynes was developed with a broad substrate scope, affording a series of carborane-fused tricyclo [6.4.0.0<sup>2,7</sup>]dodeca-2,12-dienes in moderate to high yields. In this reaction, a reactive carborane-fused 1,2-cyclohexadiene intermediate was formed, followed by a stepwise [2 + 2] cycloaddition *via* a diallyl diradical to give the final product. This protocol provided a feasible strategy for the synthesis of complex carborane-fused tricyclic compounds in a single process, which is otherwise inaccessible by other means.

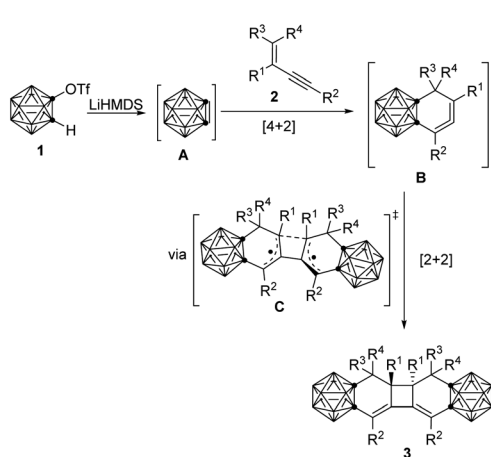
## Conflicts of interest

There are no conflicts to declare.

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Scheme 3 Proposed reaction mechanism.



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