# **A BN-Benzvalene**

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**ABSTRACT:** The synthesis and crystallographic characterization of BN-benzvalene, the first light-element hetero-benzvalene, is described. BN-benzvalenes are produced via photoexcitation of C5-aryl-substituted 1,2-azaborines under flow conditions. Mechanistic studies support a boron-specific, two-step photoisomerization pathway involving a BN-Dewar benzene intermediate, which is distinct from the photoisomerization pathway proposed in benzene and phospha- and sila-benzenes for the formation of their respective benzvalene analogues.

Benzene has four valence isomers: <sup>1</sup> benzvalene, <sup>2</sup> Dewar benzene, 3 prismane, <sup>4</sup> and bicyclopropenyl. <sup>5</sup> Among them, benzvalene has captured the attention of synthetic chemists due to its unique bonding and reactivity.<sup>2</sup> The development of a practical and facile synthesis of benzvalene by Katz<sup>6</sup> has led to the access to additional strained hydrocarbon scaffolds, including bicyclo $[1.1.1]$ pentane<sup>7,8</sup> and bicyclo<sup>[2.1.1]</sup>hexane<sup>9</sup> derivatives that are of current interest as 3D bioisosteres of benzene in medicinal chemistry.<sup>10</sup>

In comparison to the carbonaceous benzvalene, main group heteroatom-containing benzvalene derivatives have been less developed. Kobayashi pioneered the first example in 1977 by synthesizing the 1,4-diphosphabenzvalene through the photoisomerization of 1,4-diphosphabenzene.<sup>11</sup> Subsequently, Regitz,<sup>12</sup> Ando,<sup>13</sup> Tokitoh, <sup>14</sup> Sekiguchi, <sup>15</sup> Cummins, <sup>16</sup> and Kyushin<sup>17</sup> et al. independently synthesized additional phosphorus- and silicon-containing heteroatom benzvalenes (Schemes 1A and 1B). <sup>18</sup> Conspicuously, second-row element (B, N, O) hetero-benzvalenes have remained elusive to date.<sup>19</sup> Arguably, the involvement of heavier and larger 3<sup>rd</sup>-row main group elements (P and Si) reduces the resonance stabilization (due to weak  $\pi$ -bonding) of the corresponding heteroarenes and attenuates the ring strain, resulting in some thermodynamic driving force for the formation of the heavier hetero-benzvalene structures.<sup>17c, 20</sup>

In this communication, we report the first example of a light-element hetero-benzvalene. Specifically, we describe the synthesis, characterization, and initial reactivity studies of the boron- and nitrogen-containing BN-benzvalene (Scheme 1C). Deuterium labeling analysis is consistent with a distinct mechanism that involves a corresponding Dewar valence isomer species (BN-Dewar benzene) as an intermediate that undergoes a photo-induced 1,2 boron shift to furnish the observed BN-benzvalene.

Our group has had a long-standing interest in the synthetic development and applications of 1,2-azaborines, $^{21}$  which are boron(B)nitrogen(N) containing isosteres of benzene. In collaboration with the Bettinger group, we discovered that 1,2-azaborine undergoes a clean photoisomerization to BN-Dewar benzene upon UV light irradiation ( $hv > 280$  nm).<sup>22</sup> Subsequently, the BN-Dewar benzene species has been investigated as potential molecular solar thermal



**Scheme 1.** Main Group Heteroatom-Containing Benzvalenes

fuels,<sup>23,24</sup> new synthetic building blocks for 1,2-aminoborylated cyclobutane derivatives, $^{25}$  and as a monomer for ring-opening metathesis polymerization. <sup>26</sup> During our investigation of the photochemistry of 1,2-azaborines, we discovered that a C5-aryl-functionalized 1,2-azaborine (with *N*-TBS and *B*-Mes substitution) produced the BN-benzvalene isomer in addition to the BN-Dewar isomer upon photolysis (Scheme 2).



To optimize the formation of BN-benzvalene, we selected C5-*p*-Tol-*N*-TBS-*B*-Mes-1,2-azaborine **1a** as a model substrate using a Vapourtec UV-150 continuous-flow photochemical reactor with adjustable irradiation wavelength (by applying select filters to a mercury lamp). As can be seen from Table 1, we determined that 280 nm UV irradiation with a residence time of 20 minutes in THF (0.10 M) were optimal for the generation of BN-benzvalene.

Under these conditions, we obtained BN-benzvalene **2a** in 88% yield without the formation of BN-Dewar benzene **3a** as a byproduct (Table 1, entry 1). Shortening the residence time (RT) (entry 2) or using a longer wavelength (entry 4) resulted in incomplete conversion, with BN-Dewar benzene **3a** being the major observed product. On the other hand, extending the residence time (entry 3) or using a shorter wavelength (entry 5) resulted in a slightly reduced yield of BN-benzvalene **2a**, likely due to product degradation. Hexane and CH<sub>2</sub>Cl<sub>2</sub> are compatible reaction solvents, however, they produced lower yields compared to THF (entries 6 and 7). The reaction is scalable to at least 0.50 mmol (200 mg) scale without a decrease in yield (entry 8).<sup>27</sup> Notably, both products **2a** and **3a** are stable toward silica gel column chromatography under an inert atmosphere.

**Table 1.** Optimization of the Photoisomerization



<sup>a</sup> Yields were determined by <sup>1</sup>H NMR analysis using 1,3,5-trimethoxybenzene as an internal<br>standard. Scale: 0.10 mmol, unless otherwise stated. Yields of isolated product are reported in parentheses. RT: residence time.

We were able to isolate compounds **2a** and **3a** (Table1, entry 8 and entry 9, respectively), and Figure 1 shows their <sup>1</sup>H NMR spectra. In Dewar isomer **3a**, the two methyl groups on TBS (~0 ppm, orange arrow) are diastereotopic as evidenced by two distinct signals in the 1 H NMR spectrum. Similarly, the benzvalene isomer **2a** also exhibits two signals for the same methyl groups in its <sup>1</sup>H NMR spectrum, albeit with less chemical shift difference. The mesityl *o*methyl groups ( $\sim$ 2.2 ppm, purple arrow) exhibit exchange dynamics (broadened signals) at room temperature for the BN-Dewar benzene **3a** compound, whereas sharp diastereotopic mesityl *o*methyl signals are observed for BN-benzvalene **2a**. These observations are consistent with a more restricted B–Mes bond rotation for **2a** relative to **3a**. The original 1,2-azaborine arene C-H proton signals Hª, H<sup>b</sup>, H<sup>c</sup> are upfield shifted in BN-benzvalene **2a** relative to the signals for BN- Dewar **3a** and are consistent with previously reported similar tricyclic structures.<sup>2a,19a</sup>



**Figure 1.** <sup>1</sup> H NMR spectra in CDCl3 of C5-*p*-Tol-ΒΝ-Dewar benzene **3a** (top trace) and *p*-Tol-BN-benzvalene **2a** (bottom trace) in CDCl<sub>3</sub>.

We next evaluated the scope of C5-aryl functional group under our optimized conditions. A broad range of *p*-substituents are compatible, affording the corresponding products containing alkyl groups (entries **2a–2c**), trifluoromethoxy group (entry **2d**), and halogens (entries **2e–2g**) in good yields. The unsubstituted C5-Ph-1,2 azaborine substrate (entry **2h**), along with C5-3,5-dimethoxy-phenyl- (entry **2i**), and C5-*o*-tolyl- (entry **2j**) derivatives were also successfully converted to the BN-benzvalene products. Moreover, C5-1,2-azaborines containing electronically diverse heterocycles such as thiophene, pyridine, and thiazole (entries **2k–2m**) prove also to be suitable substrates.

### **Table 2.** Reaction Scope at the C5 Position



<sup>a</sup> Yields of isolated products are reported as an average of two trials. Scale: 0.10 mmol, unless<br>otherwise stated. <sup>b</sup> 0.50 mmol scale. <sup>c</sup> Hexane was used as a solvent. <sup>d</sup> Yields were determined by otherwise stated. <sup>b</sup> 0.50 mmol scale. <sup>c</sup> Hexane was used as a solvent. <sup>d</sup> Yields were determined by<br><sup>1</sup>H NMR analysis using 1,3,5-trimethoxybenzene as an internal standard; reported yields are the<br>average of two trials.

We then turned our attention to determine the scope with respect to the nitrogen substituent. The presence of a silicon on nitrogen appears to be crucial for product formation. While silicon-based groups, such as TMS, TBS, and SiPh3 (entries **2a**, **2n**, **2o**) generate the product cleanly, other groups on nitrogen, such as H, Me, Ph, and Ts (entries **2p-2s**) completely shut down the reaction, with unreacted starting material and/or decomposition being observed.

**Table 3.** Reaction Scope at the N Position



 $^a$  Yields were determined by <sup>1</sup>H NMR analysis using 1,3,5-trimethoxybenzene as an internal standard. Scale: 0.10 mmol. Reported yields are the average of two runs. <sup>b</sup> Yields of isolated products are reported as an average of two trials. RT: residence time.

BN-benzvalenes (**2a–2n)** are generally non-crystalline oily substances, with the exception being the *N*-SiPh<sub>3</sub>-substituted **2o**, which is a white solid. We successfully grew crystals of **2o**  (CCDC 2359775) suitable for single crystal X-ray diffraction analysis by slow evaporation from a  $CH_2Cl_2$  solution at -40 ºC. We also obtained the structure of the corresponding 1,2 azaborine starting material **1o** (CCDC 2359773) for comparative bonding analysis. Two structural features of **2o** are noteworthy (Figure 2): 1) The N-B bond distance (1.419(3)Å) in **2o** is shorter than that for **1o** (1.446(3) Å), which is consistent with loss of electron delocalization/aromaticity<sup>28</sup> after the photoisomerization. 2) In **2o**, the C–C bond distances associated with C6 (C6–C5 = 1.497(3) Å, C6–C3 = 1.494(3) Å) are shorter than the C–C bond distances associated with C4 (C4–C3 =  $1.531(3)$ ) Å, C4–C5 = 1.546(3) Å). The bond distance differences are consistent with reported crystallographically characterized aminoborylated cyclopropanes. <sup>29</sup> Notably, density functional theory (DFT) calculations nicely reproduce the experimentally observed structural asymmetry in **2o** whereas the direct carbonaceous benzvalene analogue exhibits symmetrical  $\sigma$ -bonding according to DFT predictions (See Supporting Information for details).



**Figure 2.** Crystal Structure of **2o** and **1o** (hydrogen atoms are omitted for clarity). Thermal ellipsoids are drawn at the 50% probability level. Selected bond lengths in Å for **2o** are: N–B = 1.419(3), N–C6 = 1.479(3), B–C4 = 1.575(3), C4–C3 = 1.531(3), C4–C5 = 1.546(3), C6–C3 = 1.494(3), C6–C5 = 1.497(3); Selected bond lengths in Å for **1o** are: N–B = 1.446(3),  $N-C6 = 1.388(2)$ ,  $B-C3 = 1.519(3)$ ,  $C3-C4 = 1.355(3)$ ,  $C4-C5$  $= 1.421(3)$ , C6–C5  $= 1.355(3)$ .

We also used DFT calculations at the B3LYP/6-311G\*\*// B3LYP/6-31G\*\* level of theory to evaluate the relative energies of **Table 4.** Relative Energies of 1,2-Azaborine's Valence Isomers (calculated at the B3LYP/6-311G\*\*// B3LYP/6-31G\*\* level of theory.)



Dewar benzene and benzvalene isomers vs. their (hetero)arene structures for both the BN and CC series (Table 4). We determined that benzvalenes **2a** and **CC-2a** are slightly less stable than theDewar benzene isomers **3a** and **CC-3a** by 3.2 and 2.8 kcal/mol, respectively). BN-benzvalene **2a** is higher in energy by 49.5 kcal/mol compared to 1,2-azaborine **1a** whereas benzvalene **CC-2a** is higher in energy by 68.1 kcal/mol relative to its benzene isomer **CC-1a**. The difference in the relative stability between the **BN**  and **CC** congeners (**BN**: 49.5 kcal/mol vs. **CC**: 68.1 kcal/mol) can be ascribed to the stronger resonance stabilization energy present in **CC-1a** vs. **1a**. 30

We have also conducted preliminary reactivity studies of BNbenzvalene **2a** and determined that **2a** reacts with thiophenol to cleanly furnish the ring-opened adduct 4a (Scheme 3).<sup>9b,31</sup> Notably, the reaction is sluggish under dark conditions, and irradiation with purple light accelerates the reaction. <sup>32</sup> Compound **4a** (CCDC 2359774), the first example of a BNbicyclo[2.1.1]hexane, is stable towards air and moisture and can be isolated by column chromatography under ambient conditions. **Scheme 3.** Synthesis of BN-Bicyclo<sup>[2.1.1]</sup>hexane



To elucidate the reaction mechanism for the formation of BNbenzvalene, we conducted the experiments as outlined in Scheme 4. The observations during our reaction optimization work (Table 1, entries 2 and 4 vs. entry 1) hinted at BN-Dewar benzene **3a** serving as an intermediate toward the formation of BN-benzvalene **2a**. We were able to isolate pure BN-Dewar benzene **3a** and determine that **3a** cleanly converts to **2a** under the standard reaction conditions (Scheme 4, eq 1). Thus, BN-Dewar benzene **3a** is a chemically and likely kinetically competent intermediate for BNbenzvalene formation. Next, we conducted a deuterium labeling study to identify the bond connectivity over the course of the reaction. We synthesized C6- and C3-deuterated 1,2-azaborine analogues **5a** and **7a**, respectively. These substrates, when subjected to the standard photoisomerization conditions, furnish labeled BN-benzvalene products **6a** and **8a**. Formation of **6a** (Scheme 4, eq 2) is consistent with N-C6 bond remaining connected throughout the reaction. To our surprise, the deuterium in **8a** switched its



location to the bridgehead C-H position, indicating that B-C3 bond is cleaved during the formation of the BN-benzvalene product. By shortening the residence time, we were able to monitor the formation of the deuterium-labeled BN-Dewar benzenes **9a** and **10a**. Consistent with the previously reported mechanism for BN-Dewar benzene formation, <sup>33</sup> no N-C6 and B-C3 bond cleavage was observed (eqs 4 and 5). The findings illustrated in Scheme 4 indicate that the B-C3 bond must be cleaved during the transformation from BN-Dewar benzene to BN-benzvalene.

Scheme 5 illustrates our proposed mechanism that is consistent with our deuterium labeling and reaction intermediate analysis. First, the starting 1,2-azaborine undergoes a photoinduced formal 4*π* disrotatory electrocyclization to generate the BN-Dewar benzene intermediate.<sup>33</sup> Then, the styrene chromophore in the BN-Dewar benzene is excited by UV-light to generate a biradical species. The radical at the C4 position subsequently undergoes a 1,2 boron shift, <sup>34</sup> and producing a C3 radical via a C3-B bond cleavage. Finally, radical-radical recombination of the C3 and C5 radicals creates the central C-C bond of BN-benzvalene. The proposed reaction mechanism is consistent with C5-aryl functional group facilitating the generation of BN-benzvalene by providing extra conjugation in the BN-Dewar intermediate for the photoinitiated formation of the biradical. The proposed 1,2-boron shift for BNbenzvalene formation is distinct from mechanisms proposed in benzene<sup>35, 36</sup> and phospha<sup>12</sup>- and sila<sup>14,15a</sup>-benzene analogues to yield their respective benzvalenes.

**Scheme 5.** Proposed Mechanism for Formation of BN-Benzvalene



In conclusion, we presented the synthesis and crystallographic characterization of the first light-element hetero-benzvalene exemplified as the boron-nitrogen containing BN-benzvalene. Photoexcitation of C5-aryl-substituted 1,2-azaborines under flow conditions furnish BN-benzvalenes with a broad substrate scope, including halogen functional groups and heteroaromatic rings. Mechanistic studies support a distinct stepwise photoisomerization pathway from 1,2-azaborine via a BN-Dewar benzene intermediate involving a radical-induced 1,2-boron shift. BN-benzvalene reacts with thiophenol in the presence of light to generate a BNbicyclo[2.1.1]hexane. Current efforts are directed toward exploring novel modes of reactivity of BN-benzvalene derivatives.

## **ASSOCIATED CONTENT**

The Supporting Information is available free of charge at https://pubs.acs.org. Experimental procedures, compound characterization data, computational and crystallographic information (PDF)

#### Accession Codes

CCDC 2359773, CCDC 2359774, and CDCC 2359775 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data\_request/cif, or by emailing data request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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#### **Notes**

The authors declare no competing financial interests. All authors have given approval to the final version of the manuscript.

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## **DEDICATION**

This work is dedicated to Prof. Lawrence T. Scott on the occasion of his 80<sup>th</sup> birthday.

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