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Kyoto University
Highlight Review

Ring-Expanding and Ring-Opening Transformations of Benzofurans and Indoles with Introducing Heteroatoms

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Abstract

Skeletal transformations of heteroaromatic compounds with cleaving their endocyclic bonds have been emerging as new synthetic tools in modern organic chemistry. Especially, endocyclic insertion of a heteroatom into a heteroaromatic core is a game-changing approach to exotic classes of heterocyclic compounds. In this Highlight Review is summarized recent progress of ring-expanding and ring-opening reactions of benzofurans and indoles.

Introduction

Heteroaromatic compounds are highly important due to their wide variety and availability, and heteroaromatic cores are often included in functional materials or medicines. Therefore, a huge number of reactions to synthesize and functionalize heteroarenes have been reported. However, transformations of heteroaromatic cores with cleaving endocyclic 

endocyclic carbon–heteroatom bonds are still limited despite great progress of organic synthesis. The infancy of such endocyclic transformations is naturally ascribed to large bond energies of C–O, C–N, and C–S bonds and decisive aromatic stabilization. If one can use heteroaromatic cores as building blocks through endocyclic transformations, novel and useful synthetic approaches would be provided.

In recent years, we have been interested in transformations of heteroaromatic compounds into other skeletons through partial disassembly of their cyclic structures, which we have coined “aromatic metamorphosis”.

Previously our group achieved transformations of dibenzothiophenes, dibenzofurans, benzofurans, and indoles. To establish aromatic metamorphosis as a truly useful strategy in organic synthesis, we need to understand the reactivity of each heteroaromatic rings.

Benzofuran, a singly benzo-fused furan, is distinct from furan and dibenzofuran in that the 2,3-position in benzofuran can be regarded as (Z)-styryl phenyl ether of reasonable reactivity. Therefore, several approaches using the special reactivity of the vinyl ether are conceivable to transform the furan ring in benzofuran. We can categorize the reported cleavages of C2–O bonds of benzofurans via organometallic species into the following four types on the basis of reaction mechanisms: (a) oxidative addition; (b) addition followed by β-elimination; (c) 1,2-metalate migration; and (d) reduction with alkali metal (Scheme 1).

Scheme 1. Ring-opening reactions of benzofuran categorized by reaction mechanism.

(a) Oxidative addition
Ni-catalyzed C–O bond cleavages are generally considered to proceed via oxidative addition due to the high electron density of Ni(0) species. In 1979, Wenkert reported ring-opening arylation and alkylation of benzofuran with the aid of a nickel catalyst and Grignard reagents to afford stilbene derivatives (Scheme 2). This reaction is not only the first example of catalytic ring-opening transformations of benzofuran but also the earliest work of catalytic C–O bond activations.

(b) Addition and β-elimination
Fe and Rh catalysts can promote addition of organometallic reagents to the vinyl ether unit of benzofuran.
and subsequent β-elimination (Scheme 1b). For example, ring-opening arylation of benzofuran with an organoaluminum reagent takes place in the presence of a Rh catalyst (Scheme 3). As well as catalytic systems, Zr complexes react stoichiometrically with benzofuran to give the ring-opening products in a similar manner. Besides alkylations and arylation, Cu-catalyzed cascade reaction composed of reductive ring-opening and the following hydroamination was also reported.

(c) 1,2-Metalate migration

1,2-Migration in metalate complexes of 2-benzofurylmetal with other alkyl or aryl anions results in formation of the corresponding 6-membered 1,2-oxametalacycles (Scheme 1c). For instance, treatment of 2-iodobenzofuran with a manganate complex prepared from MnCl₂ and 3 equivalents of nBuMgBr gives the butylation product with ring-opening of the furan moiety (Scheme 4).

(d) Reduction with alkali metal

As another approach to ring-opening of benzofurans, reductive cleavage of the C2–O bond without transition metal catalysts is known (Scheme 1d). Treatment of lithium metal with benzofuran in the presence of an electron transfer catalyst results in selective cleavage of the C2–O bond to afford α-hydroxy styrene after acidic workup (Scheme 5).

Finally, we revealed that the combination of a nickel catalyst and a diboron reagent can introduce a boron atom into benzofuran with cleaving the C2–O bond (Scheme 7). This reaction is the first example of catalytic introductions of a heteroatom into a heteroaromatic core. Surprisingly, the boron insertion product 2a was stable under air and acidic conditions, and they could be purified on silica gel even with an eluent containing acetic acid.

A plausible reaction mechanism of the boron insertion is described in Scheme 8. The C2–O bond of 1a would undergo oxidative addition to a Ni(0) complex to give oxanickelacycle 3. Transmetalation with B₂(pin): results in formation of borynickel species 4 which affords 2a via reductive
elimination with constructing a C–B bond. Before acidic workup of the reaction mixture, the product 2a’ exists as a precipitate in toluene and 1H and 11B NMR analysis in polar solvents supported the spirocyclic structure of 2a’.

Finally, treatment with aqueous HCl gives 2a.

Scheme 8. A proposed mechanism of boron insertion into benzofuran.

Since the cleavage of the C2–O bond in this reaction proceeds via oxidative addition, Ni catalysts ligated with electron-rich phosphines or NHCs tended to be effective for the boron insertion, and NiCl2(PPh3)2(IPr) complex (IPr = 1,3-bis(2,6-diisopropylphenyl)imidazol-2-ylidene) gave the best result. Although the diol moiety of diboron reagent does not remain in the product, only B2(pin)2 gave 2a in a good yield while other diborons bearing neopentyl glycol and catechol were totally ineffective for this boron insertion.

Notably, the boron insertion reaction had a good compatibility of functional groups (Table 1). For example, methoxy, siloxy, ester, and fluoro groups were compatible with this boron insertion, which would be potentially reactive in the presence of a Ni catalyst (2b–2e). As well as benzofurans, π-extended naphthofurans were also applicable to the boron insertion (2g–2i).

Table 1. Representative reaction scope of boron insertion of benzofurans.

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<tr>
<th>R</th>
<th>% yield</th>
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<tr>
<td>H</td>
<td>75%</td>
</tr>
<tr>
<td>OMe</td>
<td>68%</td>
</tr>
<tr>
<td>OSi(Bu2)Me2</td>
<td>72%</td>
</tr>
<tr>
<td>CO2Me</td>
<td>74%</td>
</tr>
<tr>
<td>2g</td>
<td>62%</td>
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<tr>
<td>2h</td>
<td>81%*</td>
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<tr>
<td>2i</td>
<td>72%*</td>
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*10 mol% catalyst.

To prove the utility of the products as organoboron reagents, we performed several transformations of the resulting oxaborins. For example, Suzuki-Miyaura cross-coupling of borate intermediates 2a’ and 2d’ successfully took place in one pot with high stereoselectivity after etherification (Scheme 9a).

In these coupling reactions, acetonitrile as an additional solvent was important to increase the solubility of borate intermediates.

We also tried applying this boron insertion to the synthesis of a fluorescent molecule. B-aryloxaborins are also known as fluorophores and several dibenzoxaborins were synthesized to explore their photophysical properties. 2a

We conducted introduction of an aryl ring onto the boron atom of naphthoxaborin 2h to yield B-tolynaphthoxaborin 6, which showed blue fluorescence with a quantum yield of 40% (Scheme 9b). DFT calculations of 6 revealed that the oxaborin core and the tolyl group on the boron atom are coplanar at the optimized structure, and the frontier orbitals of 6 are widely delocalized all over the π-system of 6.

Scheme 9. Applications of the boron insertion products.

Cu-Catalyzed Ring-Opening Silylation of Benzofurans

Since catalytic boron insertion into benzofuran was established, we naturally aimed to expand this methodology to insertion of other heteroatoms. Considering the importance of organosilicon compounds as building blocks and functional materials in modern organic chemistry, 22 we chose silicon as the next heteroatom for the insertion.

Initially we applied a disilane reagent instead of B2(pin)2 to some Ni-catalyzed systems expecting formation of 6-membered oxasilacycle, oxasilin 8. 23 However, benzofuran did not react at all and both benzofuran and the disilane were fully recovered (Table 2). Then a Ni/Cu co-catalysis was tested in order to promote transmetalation from disilane via silylcopper species, and a complex mixture was obtained without formation of oxasilin 8. We thought Cu catalyst participated in ring-opening process of benzofuran, and further condition screening with Cu catalysts was conducted. We eventually found that Cu-catalyzed ring-opening silylation of benzofuran with 1,2-di-tert-butoxy-1,1,2,2-tetramethyldisilane (7a) 24 took place instead of silicon insertion to form 8.
The effect of solvent is critical for this silylation of benzofuran. Only THF and pyridine gave the product in moderate yields (Table 3), while other ethereal solvents could not promote the reaction at all and amide solvents such as DMF and DMA resulted in decomposition of the product. Interestingly, a co-solvent system of THF and pyridine increased the yield of 9a. Although the actual reasons of the improvement of reaction efficiency are not clear, addition of pyridine to THF might reduce the aggregation of silylcopper species in the reaction mixture to enhance the reactivity of silylcopper.

Other silicon sources 7b–7e were also tested for the silylation, and a tert-butoxy group on silicon atoms was found to be essential for successful silylation of benzofuran (Table 4). Although diethoxydisilane 7b probably underwent ring-opening silylation, the product was decomposed under the reaction conditions since the steric hindrance around the ethoxysilyl group is insufficient. On the other hand, hexamethyldisilane (7c) did not react at all with 1a. These results indicate that the Lewis acidity of the silicon atoms in 7 is important to activate a disilane with a base. While silylboranes are often used as silyl anion equivalents for many silylation reactions, silylborane 7d gave a complex mixture under the reaction conditions. Unsymmetrical disilane 7e gave a mixture of two silylation products with no selectivity.

Substrate scope of the ring-opening silylation is shown in Table 5. Halogen atoms including chlorine endured reaction conditions (9d and 9e). Benzofurans bearing a monocyclic furan or thiophene underwent silylation only at the benzofuran side (9g and 9h). Attempted double silylation of naphthobifuran gave rise to mono-silylation only at the one side affording 9j–Me after methylation. The exclusive mono-silylation is attributable to the negative charge on the phenoxymoiety emerging after the first ring-opening silylation that would prevent the intermediate from coordination to electron-rich silylcopper species.

A plausible reaction mechanism for this silylation is described in Scheme 10. At the first step, CuO\textsubscript{tBu} should be generated from CuCl and KO\textsubscript{tBu}, and the formation of silylcopper species from the disilane 7a and CuO\textsubscript{tBu} would then take place.\textsuperscript{26} After coordination of 1a to the Cu center, the complex 10 undergoes silylcupration to form 11. The following \(\beta\)-elimination would furnish potassium \(\beta\)-(silylvinyl)phenoxide 9a–K, and regeneration of silylcopper species from CuO\textsubscript{tBu} and 7a would proceed. As a related work, Studer also reported ring-opening silylation of benzofurans and indoles with a silyllithium reagent via silyllithiation.\textsuperscript{17}

Computational studies were also performed to figure out the details of the reaction mechanism of this silylation. DFT calculations\textsuperscript{27} suggested that the rate determining step would be the silylcupration (21.5 kcal mol\textsuperscript{-1}) and the following \(\beta\)-elimination would smoothly proceed (14.1 kcal mol\textsuperscript{-1}). These computational expectations also agreed with the experimental observations that 2-silyldihydrobenzofuran from 11 was not detected after acidic workup. In addition, \(\beta\)-elimination would proceed via syn-configuration to give the silylation product as an (E)-isomer.
Scheme 10. A plausible reaction mechanism of ring-opening silylation of benzofuran.

We conducted transformations of a silylation product 9a. Pd-catalyzed cross-coupling of 9a with iodoarenes smoothly proceeded to furnish the corresponding (E)-stilbene derivatives 12a and 12b in good yields (Scheme 11a). Treatment of 9a with NIS (N-iodosuccinimide) gave (E)-iodostyrene 13 with the hydroxy group protected by a silyl group via intramolecular silicon transfer (Scheme 11b).28

Scheme 11. Transformations of 9a.

Heteroatom Insertion into Benzofurans with Divergency

Considering great interest in a series of oxaheterocycles including oxaborins and demands for efficiently providing them, diversity-oriented approaches from common intermediates would be ideal and robust to rapidly offer a variety of oxaheterocycles. With this in mind, we focused on Mn-mediated 1,2-metate migration which affords a ‘dianion’ of an alkenylphenol from benzofuran (Scheme 4).14b The dianionic species was expected to react with various electrophiles to furnish a series of 6-membered oxaheterocycles. This methodology composed of migratory ring-opening and trapping with a heteroatom electrophile could introduce both an aryl/alkyl group and a heteroatom to benzofuran skeleton in one pot, thereby increasing the diversity of oxaheterocyclic products efficiently.

The reported ring-opening of benzofuran with an alkylmanganese complex employed 2-iodobenzofuran as a substrate, which should be prepared from benzofuran.14b To improve the availability of substrates, we used benzofuran without a substituent at the 2 position and employed a combination of an excess of an organolithium and a catalytic amount of MnCl₂. We expected that an organolithium would participate in deprotonation at the 2 position and a Mn catalyst would be regenerated via transmetalation between the oxamanganacycle and an organolithium (Scheme 12).

Scheme 12. Working hypothesis for divergent heteroatom insertion into benzofuran.

This catalytic system for the ring-opening process indeed worked well, and treatment of benzofuran with 4 equivalents of phenyllithium in the presence of 10 mol% MnCl₂ gave a stilbene derivative 14 in 86% yield (Scheme 13a). Although optimal conditions for ring-opening of benzofuran were in hand, the efficiency of trapping with an electrophile as the following step was unsatisfactory. To solve the problem, we tried a series of additives and finally we found that addition of 1 equivalent of TMEDA (N,N',N',N''-tetramethylethylenediamine) increased the yield of silicon insertion product 15a (Scheme 13b). TMEDA would help to reduce aggregation of dianionic species and thus enhance the reactivity of the intermediate to the electrophile.

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Next, a series of electrophiles were applied to this heteroatom insertion into benzofuran (Table 6). As well as silicon, other elements such as boron, phosphorus,29 germanium, and titanium were successfully introduced to the cyclic structure by employing the corresponding electrophiles (16a–19a). Oxasilin 15a, oxaphosphin 17a, and oxagermin
showed blue fluorescence, and thus these skeletons are considered to be potentially usable for fluorophores. In addition, 18a and 19a were the first examples of syntheses of oxagermin and oxatitin cores, respectively. Although electrophilic trapping with Me₂SnCl₂ or Cp₂ZrCl₂ were likely to give the corresponding cyclic products according to ¹H NMR and MS analysis, they gradually decomposed during purification and could not be isolated in our hand.

Although this reaction proved indoles to undergo transformations with cleaving their C–N bonds, the compatibility of functional groups was insufficient due to the high reactivity of silyllithium.

Table 6. Diverse heteroatom insertion into benzofuran. a) 6 equiv electrophile. b) aq. H₂O₂ was added to the crude mixture after acidic workup.

Table 7. Reaction scope of silicon insertion into benzofurans. a) 20 mol% MnCl₂. b) 2 equiv of TMEDA. c) –78 °C to –20 °C for ring-opening step. d) Et₂O was used instead of THF.

As another advantage of this sequential heteroatom insertion, various substituents could be installed into the products by using other organolithium reagents or other benzofurans (Table 7). Not only (hetero)arylolithiums but also alkylolithiums were also applicable to this heteroatom insertion (15g and 15h). Naphthofurans as well as benzofurans could be used for silicon insertion to afford the corresponding naphthoxasilins (15i–15n).

Lithium-Mediated Ring-Opening and Sequential Boron Insertion into Indoles

Azaborin cores have been attracting great attention for their aromatic system as well as functional materials and bioactive molecules. However, conventional synthetic methods for azaborins are not satisfactory because harsh conditions or complicated substrates are required, and thus skeletons and substituents of reported azaborins are still limited. If direct synthesis of azaborins from indoles, which is one of the most accessible heteroarenes, is achieved on the basis of our methodology of boron insertion, such a reaction should be a powerful method to expand new chemical spaces.

However, as described in the introduction, C–N bond cleavage of indoles is still one of the most challenging reactions in modern organic chemistry. This difficulty is derived from both the high aromaticity of the pyrrole core and the poorer leaving ability of R₂N– anion than that of RO– anion. Recently, Studer reported ring-opening silylation of benzofurans and indoles with the aid of silyllithium, which is one of the most accessible heteroarenes, is achieved on the basis of our methodology of boron insertion, such a reaction should be a powerful method to expand new chemical spaces.

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organoboronic acid esters should be a great advantage over the conventional azaborin syntheses because various organoboronic acid esters are commercially available and easily handled.


A wide range of arylboronates could be used as boron sources for this azaborin synthesis with high compatibility of functional groups (Table 8). Surprisingly, carbonyl groups and halogen atoms endured the reaction conditions although organolithium species existed in the reaction mixture (23ae–23ai). Alkyl-, alkynyl-, and heteroarylboronates were also applicable (23aj–23am). While a variety of boronates could be used, neither electron donating or withdrawing groups at the phenyl ring on the nitrogen atom retarded the reaction (23ca and 23da). This boron insertion of indoles could construct complicated polyarenes bearing multiple azaborin units by employing the corresponding di- and triboronic acid esters as electrophiles (24, 25).

Due to the good compatibility of functional groups on arylboronates with the reaction conditions, this boron insertion into indoles could provide ideal precursors of BN analogues of polyaromatic hydrocarbons. Azaborin 23ah from a bromo-substituted arylboronate successfully underwent intramolecular ring-closure with the aid of a palladium catalyst and a base (Scheme 16). In a similar manner, two-fold ring-closure of 24 successfully proceeded to form 27.

**Conclusion**

This Highlight Review briefly summarizes skeletal transformations of benzofurans and indoles with breaking their aromatic cores. As shown above, those two heteroarenes have obtained new roles of synthetic building blocks for other heterocycles, and the concept of “aromatic metamorphosis” has been growing as a reliable synthetic methodology to provide novel and valuable molecules. Further progress in this field will disclose new chemical space for both synthetic and material chemistry.

Table 8. Reaction scope of boron insertion into indoles. ‘2 equiv of PhB(pin) was used. ‘0.5 equiv of the corresponding arylboronate was used. ‘0.2 equiv of the corresponding arylboronate was used.

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References and Notes


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