Showcasing research from the group of Dr Suguru Yoshida and Professor Takamitsu Hosoya at the Institute of Biomaterials and Bioengineering, Tokyo Medical and Dental University (TMDU), Japan.

Rhodium-catalyzed odorless synthesis of diaryl sulfides from borylarenes and S-aryl thiosulfonates

Various diaryl sulfides were efficiently prepared from readily available organoborons and S-aryl thiosulfonates using a rhodium catalyst. This simple and odorless method for sulfide synthesis has been achieved in the middle of Tokyo city.

As featured in:

Rhodium-catalyzed odorless synthesis of diaryl sulfides from borylarenes and S-aryl thiosulfonates†

Kazuya Kanemoto,‡ Yasuyuki Sugimura,†‡ Shigeomi Shimizu,†‡ Suguru Yoshida†‡‡ and Takamitsu Hosoya†‡*‡

Various diaryl sulfides, including heteroaryl- and nitrogen-containing sulfides, have been efficiently prepared by rhodium-catalyzed odorless deborylative arylthiolation of organoborons with S-aryl thiosulfonates. The ready availability of starting materials and further transformation of sulfides have rendered a diverse range of organosulfur compounds easily accessible.

Aromatic sulfides represent a privileged class of compounds that exhibits broad applications in various research fields such as medicinal chemistry1,2 and materials science3 (Fig. 1). This group-guided transformations as well as the ipso-substitution reactions via the C–S bond cleavage of thioarenes have enhanced the utility of aromatic sulfides as synthetic intermediates.4 Although recent developments in the transition-metal catalyzed C–S bond forming reactions have improved the accessibility of aromatic sulfides5,6 the types of sulfides that can be prepared are still limited and a novel method that enables the synthesis of a wide range of sulfides is still required.

In this context, we previously developed a facile synthetic route to aryl and alkenyl sulfides via the copper-catalyzed deborylative arylthiolation of organoborons with thiosulfonates (Fig. 2A).6,8 The use of readily available thiosulfonates7,8 as the source of the thio group rendered the entire transformation free of unpleasant odor and diverse sulfides easily accessible. However, while examining the substrate scope and limitation of this reaction, we found that the reaction between 2-thienylboronic acid (1a) and S-p-tolyl p-toluenethiosulfonate (2a) to prepare 2-thienyl 4-tolyl sulfide (3a) did not proceed under the standard copper-catalyzed conditions (Fig. 2B). This reaction possibly failed because of the deactivation of the catalyst, which is related to the strong coordination of the sulfur atom in 1a to copper. Furthermore, thiolation of 1a was not easy to achieve efficiently by the conventional catalytic thiolation methods using an easily available reagent, including thiold4,e,f,g To address this issue, we focused on a rhodium catalyst because rhodium is analogous to copper in terms of catalytic reactivity,10 such as in 1,4-addition reactions, and several sulfides have been successfully prepared using a rhodium catalyst.11 Based on this idea, we designed a catalytic cycle for the reaction between arylboronic acid and S-aryl thiosulfonate to afford the desired diaryl sulfide (Fig. 2C). This cycle involved the formal substitution reaction of a thiosulfonate with an arylrhodium intermediate II, which is generated from a rhodium catalyst I and arylboronic acid in the presence of a base. Herein, we report that a rhodium catalyst efficiently promoted the deborylthiolation of borylarenes with S-aryl thiosulfonates, including heteroaryl substrates such as 1a, which enabled the synthesis of various diaryl sulfides.

After the catalysts were extensively screened, several rhodium catalysts were found to promote the deborylative p-toluenethiosulfonate 1a with 2a (Table 1, entries 4–8). In particular, the desired sulfide 3a was obtained efficiently when the reaction was conducted in methanol at 50 °C or room temperature using a rhodium catalyst coordinated with cyclooctadiene (5–10 mol% of Rh) in the presence of potassium phosphate (entries 5 and 8–10). Besides potassium phosphate, weaker bases, such as potassium...
carbonate and cesium fluoride, were also suitable (entries 11 and 12). The reactions using a rhodium catalyst coordinated with cyclooctene, 1,5-hexadiene, norbornadiene, or triphenylphosphine gave poor results (entries 2–4 and 13). These results suggested that cyclooctadiene with a suitable coordination ability to rhodium is crucial to efficiently achieve this transformation.

In addition, an iridium catalyst was less effective for this transformation (entry 14). Similar to the copper-catalyzed reaction, the entire experimental process was free from unpleasant organosulfurous odor.

A wide range of arylboronic acids were efficiently deborylthiolated with thiosulfonate 2a under the optimized conditions using 2.5 or 5.0 mol% of [Rh(OH)(cod)]2 with potassium phosphate or potassium carbonate (Fig. 3). Various phenylboronic acids with either an electron-rich or electron-deficient substituent, such as methoxy, hydroxy, dimethylamino, bromo, and methoxycarbonyl groups, were efficiently thiolated to afford diaryl sulfides 3b–f, leaving these groups untouched. The deborylthiolation of a phenylboronic acid bearing a bulky group at the ortho position, such as 2-(pivaloylamino)phenylboronic acid, also occurred, affording 2-thioaniline derivative 3g. Notably, substrates containing a group with an acidic proton, such as hydroxy and amide groups, tolerated the reaction. Moreover, the deborylthiolation of heteroaromatic substrates, such as 2-benzofuranyl- and 3-thienylboronic acids, smoothly proceeded to afford sulfides 3h and 3i in excellent yields. Besides the aromatic substrates, an alkenyl substrate such as β-styrylboronic acid also participated in this reaction to quantitatively afford alkenyl sulfide 3j.

Various S-aryl thiosulfonates were also used to synthesize diaryl sulfides, as demonstrated in the reaction with 3-thienylboronic acid (1i) (Fig. 4). Thiosulfonates bearing an electron-donating or electron-withdrawing group successfully participated in this transformation to afford diaryl sulfides 3k–s. Notably, the reactions with thiosulfonates bearing an unprotected amino or carbamate group, which were unfavorable substrates under copper-catalyzed conditions, also smoothly proceeded to afford diaryl sulfides 3n and 3o in high yields.

To gain insights into the mechanism of the rhodium-catalyzed deborylthiolation, we conducted several control experiments.
was confirmed after the treatment with an aqueous solution of sulfide and an equimolar amount of rhodium salt on deborylthiolation. (D) Plausible reaction mechanism. Yields were determined by HPLC analysis.

(Fig. 5). From a comprehensive survey of the products formed in the reaction between 1a and 2a, the formation of a significant amount of sodium p-toluenesulfinate (4) along with sulfide 3a was confirmed after the treatment with an aqueous solution of sodium bicarbonate (Fig. 5A).12 The addition of 2,2,6,6-tetramethylpiperidine 1-oxyl (TEMPO) did not inhibit the formation of sulfide 3a, suggesting that radical intermediates were not involved in the reaction mechanism (Fig. 5B). The treatment of thiosulfonate 2a with an equimolar amount of rhodium salt at 50 °C for 2.5 h led to the partial consumption (23%) of 2a, and the subsequent addition of arylboronic acid 1a and potassium phosphate to this mixture afforded only a small amount of sulfide 3a (12% yield) (Fig. 5C). This result suggested that arylboronic acid and a base also play a significant role in generating an active catalytic species. Based on these results, the reaction was expected to start with transmetalation between the rhodium catalyst and arylboronic acid in the presence of a base (Fig. 5D). Subsequent σ-bond metathesis of the arylrhodium intermediate I with thiosulfinate (path a) or the oxidative addition of thio sulfinate to the arylrhodium intermediate II, followed by reductive elimination (path b), afforded diaryl sulfide with the regeneration of rhodium sulfinate.

The improvement in the reaction efficiency for the deborylthiolation of heteroatom-containing substrates rendered the synthesis of diverse sulfur-containing molecules easily achievable (Schemes 1–3). For example, the formal synthesis of dithienothio phene (DTT),513 was achieved via the rhodium-catalyzed coupling between 3-thienylboronic acid 1i and 3-thienyl thiosulfonate 2k, affording di(3-thienyl)sulfide 3l, which has been reported as the precursor of DTT (5)11b (Scheme 1). Furthermore, in combination with our aryne chemistry,14 the short synthesis of a highly functionalized triaryl sulfonium salt 8 was accomplished (Scheme 2). The rhodium-catalyzed thiolation of p-anisylboronic acid 1b with thiosulfonate 2f afforded the corresponding diaryl sulfide 3u in high yield. Subsequently, 3u was transformed to sulfilimine 6 based on a reported method.15 The generation of 3-methoxy-5-(3-thienyl)benzene from the precursor 7,14c in the presence of sulfilimine 6 by the treatment of the mixture with cesium carbonate and 18-crown-6 at room temperature unexpectedly afforded multisubstituted triaryl sulfonium salt 8 as the major product. The formation of 2-sulfanylaniline derivative 9, which was assumed to be obtained via the direct thioamination of aryne and subsequent migratory N-arylation,14c was not observed.

The utility of the rhodium-catalyzed deborylthiolation reaction was further demonstrated in the synthesis of diheteroaryl sulfide 14 from simple starting materials, which were easily prepared using the recently emerging C–H functionalization chemistry16,17 (Scheme 3). According to the reported method,16c thiosulfonate 2l was prepared by the cerium trichloride-mediated electrophilic p-toluenesulfonylation of indole 10 using di(p-toluene sulfonyl)sulfide 11 that proceeded in a regioselective manner at the C3-position. Meanwhile, the C5-borylated 2-methylthiophene 13 was prepared regioselectively by the iridium-catalyzed C–H borylation of 2-methylthiophene (12) according to the reported method.17c The rhodium-catalyzed coupling between 2l and 13 successfully proceeded to afford diaryl sulfide 14. In contrast, an attempt of the same transformation under copper-catalyzed
conditions resulted in the formation of only a trace amount of the desired product 14. These results clearly highlighted the advantage of rhodium-catalyzed conditions for synthesizing various di(hetero)aryl sulfides.

In summary, we have developed an odorless synthetic route to diaryl sulfides by the rhodium-catalyzed deborylthiolation of borylarenes using thiosulfonates. Because of the mild reaction conditions, a wide range of substrates could be used, which significantly expanded the scope of the diaryl sulfides that could be synthesized, including heteroaryl- and nitrogen-containing compounds. Further studies of this reaction in terms of the application to the synthesis of bioactive compounds and the detailed mechanism are currently underway.

This work was supported by the Platform Project for Supporting Drug Discovery and Life Science Research funded by AMED; P-CREATE from AMED; JSPS KAKENHI Grant Numbers 15H05118 (B. T. H.), 16H01133 (Middle Molecular Strategy; T. H.), 26350971 (C. S. Y.); Naito Foundation (S. Y.).

Conflicts of interest

There are no conflicts to declare.

Notes and references


3 For selected reviews of sulfur-containing compounds in materials science, see: (a) A. S. Rahate, K. R. Nemade and S. A. Waghule, Rev. Chem. Eng., 2013, 29, 471; (b) S. Dadashi-Silab, C. Aydogan and Y. Yagci, Polym. Chem., 2015, 6, 6595.


12 See the ESF for details.


