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## A mild carbon-boron bond formation from diaryliodonium salts†

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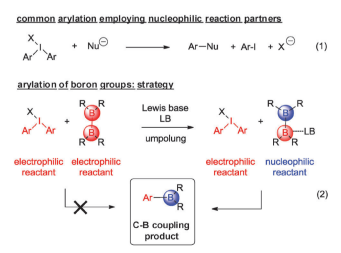
The direct metal-free borylation of diaryliodonium salts with diboron reagents is now demonstrated to be a feasible process toward formation of aryl boronic esters without any additive or catalysts, and it can be extended to a two-step C-C coupling of both aryl groups of the initial diaryliodonium reagent.

Due to their ready availability and high stability, diaryliodonium salts 1 constitute an attractive class of compounds that have been recognized as particularly versatile aryl transfer reagents.<sup>1</sup> Consequently, they have been involved in a series of arylation reactions, which have been mostly aided by the presence of transition metal catalysts. While initially C-C bond forming reactions had been explored to a larger extent, carbon-heteroatom bond forming events based on diaryliodonium salts have been investigated in greater detail in recent years (Scheme 1, eqn (1)).<sup>2–10</sup> Among all these accomplishments, carbon-boron bond formation toward aryl boronic acid derivatives is notably absent.

Arylboronic acids and esters are key components for modern cross-coupling reactions and advanced synthetic transformations. 11 Their high versatility has made C-C coupling events based on them one of the most versatile themes in the field, 12 which was recognized by the Nobel prize for Suzuki in 2010. 13 Common approaches to this type of reagents employ the original Miyaura-type borylation reaction between aryl halides and bis(pinacolato)diboron (pinB-Bpin, 2a) in the presence of catalytic amounts of palladium complexes. The key intermediate [LPd(Ar)(OAc)] undergoes a transmetallation process with 2a.<sup>14</sup> Other transition metal complexes have catalyzed the aryl halide borylation<sup>15</sup> complementing the C-H activation of arenes by borane reagents. 16 We here report the unprecedented direct

To accomplish the targeted carbon-boron bond formation, we envisioned that a Lewis base could be employed for the activation of the diboron reactant. Such an interaction should be favorable as it provides an umpolung of the native electrophilic boron atom. The resulting nucleophilic character<sup>17</sup> at one of the boron centers then ensures for effective C-B bond formation with the electrophilic iodine(III) reagent. This approach is reminiscent of previous alkoxide base mediated reactions developed by some of us18,19 and others. 20,21 A subsequent screening between various diphenyliodonium salts 1a-d and bis(pinacolato)diboron (2a) was undertaken and for the acetate derivative 1d confirmed the

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Scheme 1 Conceptual approaches to carbon-heteroatom bond formation using diaryliodonium salts.

formation of aryl boronic esters, in particular aryl pinacolboranes, through an effective coupling between diaryliodonium salts and bis(pinacolato)diboron 2a under mild conditions and without any requirement for a metal promoter. In principle, the synthetic approach to such a type of direct C-B bond formation appears unconventional at first sight, as it involves the combination of two reactants that are of exceedingly electrophilic nature (Scheme 1, eqn (2)).

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Scheme 2 Borylation of diphenyliodonium salts: optimization with  $B_2pin_2$ . Reaction conditions:  $(C_6H_5)_2I^+X^-$  (0.2 mmol),  $B_2pin_2$  (0.3 mmol), 50 °C, 1.25 mL solvent, 24 h; average yield from two independent runs calculated by GC spectroscopy with mesitylene as the internal standard; the value in parenthesis refers to an isolated yield based on (C<sub>6</sub>H<sub>5</sub>)<sub>2</sub>IOAc.

correctness of the initial hypothesis (Scheme 2). In agreement with previous experience, 18,19 methanol was identified as the best solvent.<sup>22</sup> For the reaction between acetate derivative 1d and 2a, the expected product 3a was formed as an isolated high yield of 77%. This successful transformation leads to two important conclusions: first, methanol as the solvent may play a crucial role in activating the diboron reagent through a state such as A. Secondly, the pronounced dependence of the reaction on the counterion of the diphenyliodonium reagent suggests a crucial participation of this anion as well. This may include a direct interaction between the solvated acetate with the diboron reagent (state B) or a participation of its negative charge throughout the hydrogen-bonding network of the protic solvent (state C). For the latter scenario, the activation is reminiscent of the more common activation with methoxide base. Obviously, the anions from compounds 1a-c do not display sufficient basicity to induce activation of the diboron reagent. The importance of the polar protic solvent is evident from a comparison of a reaction in polar unprotic THF, which led to a significantly lower yield of 39% of 3a.<sup>22</sup>

With the conceptual verification of the C-B bond formation in hand, the reaction was further extended to various diboron compounds for aryl-boron coupling (Scheme 3). For example, products 3b and 3c are conveniently generated by treatment between diphenyliodonium acetate and bis(neopentylglycolato) diboron (2b) and bis(hexyleneglycolato)diboron (2c), respectively, as in the case of the parent transformation with bis(pinacolato)diboron (2a). The same protocol was also applicable for the related bis(catecholato)diboron (2d) to furnish the coupling product 3d. Due to the notorious instability of the catecholboryl derivative, the boronic ester was transformed into phenol upon oxidative work-up.<sup>22</sup> Finally, an excellent result was obtained using the mixed diboron reagent Bpin-Bdan (2e) (dan = 1,8-diaminonaphthalene),<sup>23</sup> which underwent selective activation at the more electrophilic Bpin center. 19 This promotes the transfer of the Bdan entity to selectively generate C-B coupling product 3e in 87% conversion

Scheme 3 Borylation of diphenyliodonium salts: influence of the diboron reagent X = O, NH. Reaction conditions:  $(C_6H_5)_2IOAc$  (0.2 mmol), diboron reagent (0.3 mmol), 50 °C, 1.25 mL methanol, 24 h; yield calculated by GC spectroscopy with mesitylene as the internal standard as an average of two reactions; isolated yields given in brackets calculated from (C<sub>6</sub>H<sub>5</sub>)<sub>2</sub>IOAc; [a] isolated yield of the corresponding alcohol after oxidative work up.<sup>22</sup>

and 52% isolated yield. There is only one precedence of borylation of aryl halides with Bpin-Bdan (2e), which required the presence of a palladium-XPhos catalyst under basic conditions.<sup>24</sup>

The scope of the reaction has been evaluated for a number of symmetric diaryliodonium salts, which includes ortho-, meta and parasubstitution patterns as well as higher substituted aromatic entities (Scheme 4). As to a general trend, electron releasing groups on the 4-substituted aryls of Ar<sub>2</sub>IOAc contribute to a higher percentage of borylated product formation (8a-10a) in comparison with the electron-withdrawing functional groups (products 4a-7a). In contrast, 3-substituted aryl groups in Ar<sub>2</sub>IOAc bearing electron-withdrawing substituents favor the borylation with the activated B<sub>2</sub>pin<sub>2</sub> (products 11a-13a), while the corresponding diaryliodonium salt with 3-tolyl groups is transformed into coupling product 14a in moderate yield. As regularly encountered in arylboronates, yields can diminish during the purification step, being comparable to the most recent achievement in Zn catalyzed borylation of aryl halides.<sup>25</sup>

Interestingly, the highly substituted aryl(pinacolboronate) esters 15a-18a are quantitatively formed demonstrating that the present method provides great tolerance to formation of sterically encumbered aryl boronates.

With the aim to explore a selective mixed borylation we conducted two parallel strategies: (i) the reactivity of the novel nonsymmetric diaryliodonium salts 19-23 with B2pin2 and (ii) the reactivity of electronically and sterically mixed diaryliodonium salts 19 and 20 with the mixed diboron reagent Bpin-Bdan (2e). Scheme 5 shows that sterically encumbered aryl boronates exert a higher degree of selective borylation when reacted with both symmetrical and mixed diboron compounds. Therefore, the diaryliodonium salt 19 reacts with 2a to give 7a and 19a in a 1:1 ratio, which is comparable to the reaction with 2e providing 7e and 19e in a ratio of 63:37. However, the higher the steric hindrance on one of the aryl groups, the more pronounced chemoselectivity favoring 3a is observed, regardless the electronic nature of the substituents on the neighboring aryl group. Interestingly, the

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AcO 
$$\bigoplus$$
 + B<sub>2</sub>pin<sub>2</sub>  $\bigoplus$  + B<sub>2</sub>pin<sub>2</sub>

Scheme 4 Borylation of diphenyliodonium salts: reaction scope with  $B_2$ pin<sub>2</sub>. Reaction conditions:  $Ar_2$ IOAc (0.2 mmol),  $B_2$ pin<sub>2</sub> (0.3 mmol), 50 °C, 1.25 mL methanol, 24 h; average yield from two independent runs calculated by GC spectroscopy with mesitylene as the internal standard; values in brackets refer to isolated yields based on  $Ar_2$ IOAc.

$$F_{3}C \bigcirc \bigoplus_{\text{19}} \begin{array}{c} B_{2}\text{pin}_{2} \\ \text{(1.5 equiv)} \\ \text{MeOH, 50 °C} \end{array} \qquad F_{3}C \bigcirc \bigoplus_{\text{80\%, 7a:19a}} \begin{array}{c} 19a \\ \text{19a} \\ \text{80\%, 7a:19a} = 1:1 \end{array}$$

Scheme 5 Borylation of non-symmetric diphenyliodonium salts.

reaction of 23 with 2e is conducted towards the formation of 3e and 3a in 9:1 ratio, showing that steric factors are predominant.

Scheme 6 One pot cross coupling from diaryliodonium salts via  $B_2pin_2$ . Reaction conditions for activation:  $Ar_2IOAc$  (0.2 mmol),  $B_2pin_2$  (0.3 mmol), 50 °C, 1.25 mL MeOH, 24 h, reaction conditions for cross-coupling:  $Pd(PPh_3)_4$  (10 mol%),  $Cs_2CO_3$  (2 eq.), 80 °C, 2 mL toluene, 16 h. Yields refer to isolated yields based on  $Ar_2IOAc$ .

Finally, the successful development of the C-B bond forming reaction was extended to an in situ cross coupling reaction. Since the borylation reaction of the diaryliodonium salts 1 generates an equimolar amount of free aryl iodides, subjecting the crude reaction mixture to Suzuki-Miyaura cross-coupling should result in an overall diaryl synthesis. Such a process would make economic use of both of the aryl groups of the diaryliodonium precursor. Transformations of this kind are rare. Within this context, Beletskaya reported an exhaustive Suzuki-Miyaura coupling between diaryliodonium salts and sodium tetraphenylborate,26 Nachtsheim developed two sequential C-N bond forming events of cyclic diaryliodonium salts with anilines to yield N-arylated carbazoles227 and Greaney recently reported on the use of both aryl groups of diaryliodonium salts in 1,3-difunctionalization of indoles.<sup>28</sup> Indeed, the anticipated reaction sequence comprising the C-B bond formation as a transient path to C-C coupling could be realized. After the borylation reaction, the solvent was changed from MeOH to toluene followed by addition of the palladium catalyst Pd(PPh<sub>3</sub>)<sub>4</sub> and carbonate base (Scheme 6). By this, the mentioned product mixtures from C-B bond formation with symmetric diaryliodonium salts engage in the desired direct Suzuki-Miyaura coupling to provide the corresponding diaryl compounds 25-28 as the only C-C coupling products.

In summary, we have developed a new protocol for the metal-free formation of aryl-boron bonds. It employs readily available diaryliodonium acetates and commercially available diboron reagents, which in a methanol solution engage in direct aryl-boron bond formation through a formal umpolung of the electrophilic boron center. The reaction is selective, proceeds under mild conditions and does not require any metal reagent or other promoter. It opens a new methodological venue for the use of hypervalent diaryliodonium reagents in carbon-heteroatom bond formation. By simply adding a palladium source, the reaction can be directly expanded to biaryl synthesis through the cross coupling of symmetric diaryliodonium salts.

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