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Direct introduction of naphthalene-1,8-diamino boryl [B(dan)] group by a Pd-catalysed selective boryl transfer reaction

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A non-symmetrical diboron reagent, B(pin)-B(dan), has been utilised in the Pd-catalysed borylation of aryl bromides and chlorides. Remarkably selective formation of aryl-B(dan) bonds is established. This represents a direct and efficient way to masked boronic acids. The synthetic usefulness of this reaction is demonstrated in the preparation of boron-differentiated di- and polyboron compounds.

Organoboronic acids have gained broad research interests in synthetic chemistry, drug discovery and materials science. As synthetic building blocks, they can react with various functional groups to construct new carbon-carbon or carbon-heteroatom bonds. On the other hand, their high reactivity complicates the chemical selectivity in complex molecule synthesis. Consequently, practical and readily-removable masking groups which render boryl groups temporarily inert have received great development in the last decade, especially in the fields of iterative cross-coupling and functionalised polyboron compounds.

The masked boronic acids are commonly synthesized from the corresponding free boronic acids by condensation with masking molecules such as 1,8-diaminonaphthalene (dan) and Nmethyliminodiacetic acid (MIDA). Although proven successful in many cases, this approach can be troublesome when the starting organoboronic acids are not readily available or unstable. Therefore a direct way to introduce the masked boryl group into simple molecules is more desirable. Suginome and co-workers reported the direct synthesis of alkenyl-B(dan) via C-H borylation⁵ or hydroboration of alkynes^{3e} with 1,8-naphthalenediaminatoborane ((dan)BH). The diboration of alkynes using non-symmetrical diboron reagent, B(pin)-B(dan), was disclosed by the same group in 2010.4d More recently, alkyl and alkenyl 1,8-diaminonaphthalene boronates have been synthesized by hydroboration of alkenes or alkynes using B(pin)-B(dan) with the less Lewis acidic B(dan) moiety being transferred selectively. 6 In contrast, the direct synthesis of aryl B(dan) from commercially available compounds has been rarely studied. The only example of this type was realized via C-H borylation using (dan)BH.⁷ However, the reaction efficiency remained low. Moreover, the regioselectivity of the C-H borylation was governed by steric effects; some regioisomers were not accessible by the method. Consequently, we wanted to explore for a

direct and general approach to the masked aryl boronic acid derivatives.

$$\begin{array}{c} \bigcirc \text{OMe} \\ (\text{pin})B \\ B(\text{dan}) \\ Me \\ \end{array} \begin{array}{c} B(\text{dan}) \\ A \\ \end{array} \begin{array}{c} B(\text{dan}) \\ A \\ \end{array} \begin{array}{c} B(\text{dan}) \\ A \\ \end{array} \begin{array}{c} A \\ B(\text{pin}) - B(\text{dan}) \\ \end{array} \begin{array}{c} B(\text{dan}) \\ A \\ \end{array} \begin{array}{c} B(\text{dan}) \\ A \\ \end{array} \begin{array}{c} A \\ B(\text{pin}) - B(\text{dan}) \\ \end{array} \begin{array}{c} A \\ B(\text{pin}) - B(\text{pin}) - B(\text{pin}) - B(\text{pin}) - B(\text{pin}) \\ \end{array} \begin{array}{c} A \\ B(\text{pin}) - B(\text{pin}) - B(\text{pin}) - B(\text{pin}) - B(\text{pin}) \\ \end{array} \begin{array}{c} A \\ B(\text{pin}) - B(\text{pin}) - B(\text{pin}) - B(\text{pin}) - B(\text{pin}) \\ \end{array} \begin{array}{c} A \\ B(\text{pin}) - B(\text{pin}$$

Scheme 1 Selective boryl transfer of non-symmetrical diboron reagent.

Recently, differentially protected diboron reagents have been applied in chemoselectively transferring one of the two boryl groups to various unsaturated bonds. For example, Santos and co-workers have demonstrated that sp²-hybridized boron of the sp²-sp³ hybridized diboron reagents is selectively transferred in coppercatalyzed β -boration of α , β -unsaturated carbonyl compounds. The sp²-sp² hybridized non-symmetrical diboron reagent B(pin)-B(dan) has recently been applied in the addition reactions of alkenes and alkynes, with the B(dan) moiety transferred selectively (1a and 1b, Scheme 1) as mentioned above. 6 Mechanistically, in both cases, the cause of such selectivity was proposed to be the preferred interaction between the more Lewis acidic B(pin) moiety with basic promoting reagents or catalysts. During our study on preparation of functionalised polyboron reagents, we wondered whether a similar chemoselective borylation might be used for direct introduction of B(dan) onto arenes by Miyaura-type borylation reaction. 9 In classical Miyaura borylation of aryl halides, [LPd(Ar)OAc] is considered as the key intermediate which then undergoes transmetalation process with diboron reagents. We envisioned that under Miyaura-type ChemComm Page 2 of 4

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conditions, B(pin)-B(dan) may undergo a selective transmetalation process with [LPd(Ar)OAc] because of the stronger interaction between the Lewis basic -OAc and the more Lewis acidic B(pin) moiety, generating the [LPd(Ar)B(dan)] intermediate preferentially (1c, Scheme 1). The following reductive elimination would then selectively form the final borylation product ArB(dan). This might provide a straightforward method to aryl-B(dan) from readily available aryl (pseudo)halides.

Table 1. Optimization of reaction parameters.^a

Entry	Ligand	Base	$T(^{o}C)$	Yields (%) ^b
1	PPh ₃	KOAc	80	19
2	tBu-XPhos	KOAc	100	20
3	SPhos	KOAc	100	95
4	XantPhos	KOAc	100	29
5	XPhos	KOAc	100	99
6	XPhos	KO <i>t</i> Bu	100	Trace
7	XPhos	KOAc	r.t.	18
8	XPhos	KOAc	50	76
9	XPhos	KOAc	60	99(98) ^e
10 ^c	XPhos	KOAc	60	64
11 ^d	=	KOAc	60	N.R.

 a Reaction conditions: 4-bromotoluene (0.12mmol), B(pin)-B(dan) (0.10mmol), 1,4-dioxane (0.5 ml) under N₂ atomosphere. b Yields based on 1 H NMR analysis of the crude products with 1,3,5-trimethoxybenzene added as an internal standard. c 4 mol% Pd(OAc)₂ was used as catalyst. d catalyst and ligand were not added. c Isolated yield shown in parenthesis.

With this in mind, we initiated our studies by examining the reaction between B(pin)-B(dan) and 4-bromotoluene under Pdcatalysed conditions. In a preliminary experiment, using a catalytic combination of Pd₂(dba)₃ (2 mol %) and PPh₃ (6 mol %) (entry 1, Table 1), to our delight, the desired arvl-B(dan) product 2a could be obtained in 19% vield based on ¹H NMR analysis of the crude product. Further screening of phosphine ligands afforded more efficient conditions. Thus, when the reaction was conducted at 100 °C for 12 hours using XPhos, a bulky monophosphine ligand, 9c, 10 as the supporting ligand, the desired product could be obtained quantitatively (entry 5). Similar to the classical Miyaura borylation reaction, KOAc was found to be the best base for this system. The reaction temperature could be decreased to 60 °C without loss of yield (entry 9). However, further decreasing the temperature led to dramatically lower yields (entries 7-8). Omitting the Pd catalyst and the ligand, no desired product could be observed (entry 11), therefore a metal-free borylation pathway was not working for this system.

With the optimized reaction conditions in hand, we next explored the substrate scope of this selective boryl transfer reaction. As shown in Scheme 2, various aryl bromides bearing electron-withdrawing and -donating groups at the *para-* or *meta-*positions were borylated in good to excellent yields (for 2a-2l). It should be mentioned that free phenolic hydroxyl group (for 2g) and amino group (for 2h) were well tolerated in the reaction system, while the corresponding free aryl boronic acids bearing these two groups were usually only prepared in moderate yields. Therefore the current method, involving direct formation of C-B(dan) bonds from C-X bonds, is more practical and efficient than the previous condensation route. This

borylation reaction was found sensitive to the steric hindrance; elevated reaction temperature (100 °C) was necessary to give satisfactory yields for *ortho*-substituted aryl bromides (for **2m-2o**). Aryl triflate (for **2q**) could also be converted to aryl-B(dan) directly. In addition, aryl chlorides were also utilized in this reaction, although they generally needed higher reaction temperature (100 °C) than their bromides analogues (for **2a**, **2d**, **2i** and **2s**).

 a Reaction conditions: the aryl halide (0.30 mmol), B(pin)-B(dan) (0.25mmol), 1,4-dioxane (1.0 ml) under N₂ atomosphere. b Isolated yields based on the amount of B(pin)-B(dan) used. c The aryl chlorides and *ortho*-substituted aryl bromides were borylated at 100 o C. Other cases mentioned were all borylated at 60 o C.

Scheme 2 Substrate scope of Pd-catalysed selective boryl transfer.

Following our previous work on synthesis and applications of differentiated polyboron reagents, we applied the present reaction in facile preparation of polyboron compounds containing the B(dan) group. ^{4i, j} Two routes were utilized toward these compounds. The first one was starting from monoboron aryl halides, using the selective naphthalene-1,8-diamino borylation to introduce B(dan) group. Thus, diboron compounds **3a-3c** were conveniently prepared in good yields (Scheme 3A). To the best of our knowledge, there has been no literature precedence on compounds containing both B(dan) and B(MIDA) groups. The second route was using functionalised aryl B(dan) compounds as the starting materials and various known borylation methods to introduce a different type of boronyl group

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(Scheme 3B). For example, Rh-catalyzed borylation of **2d** via C-C bond cleavage under the literature conditions produced **3d** containing a neopentanediolato boronyl group B(neop). A Cucatalysed hydroboration of alkene moiety of also quantitatively furnished diboron reagent **3e** incorporating an alkyl B(pin) group.

Reaction conditions: aryl halides (0.30mmol), B(pin)-B(dan) (0.25mmol), KOAc (0.75 mmol), Pd_2dba_3 (1mol %), Xphos (3 mol %), 4-dioxane (1.0 ml) under N_2 atomosphere. **3a** and **3b** were borylated at 100 °C, **3c** was borylated at 60 °C.

Scheme 3 The applications of Pd-catalysed selective boryl transfer in preparing boron-differentiated diboron compounds

Scheme 4 Chemoselective transformations of functionalised aryl B(dan)

$$\begin{array}{c} \text{B(MIDA)} \\ \text{IIr(OMe)(cod)]}_2 \ (1 \ \text{mol \%}) \\ \text{dtbpy (4 mol\%)} \\ \hline \\ B_2(\text{pin})_2 \ (1.1 \ \text{eq.}) \\ 1.4 \text{-dioxane,} \\ 80 \ ^{\circ}\text{C, 2 h, 95\%} \\ \hline \\ Pd_2(\text{dba})_3 \ (1 \ \text{mol \%}) \\ \text{XPhos (3 mol \%)} \\ \text{XOAc (3.0 \ \text{eq.})} \\ \hline \\ 1,4 \text{-dioxane} \\ 100 \ ^{\circ}\text{C, 12 h, 42\%} \ (\text{dan}) \\ \text{B}(\text{pin}) \\ \end{array}$$

compounds.

Scheme 5 The synthesis of trisboronylbenzene.

Taking use of the relative inertness of B(dan) group, we then assessed the possible utility of the above-prepared functionalised

aryl B(dan) compounds.¹² The *para*-naphthalene-1,8-diamino boryl aniline **2f** was subjected to a chemoselective Pd-catalysed Buchwald-Hartwig coupling ¹³ with bromobenzene and the amination product **4** could be obtained in 75% yield without interference from Suzuki-Miyaura type reaction. Much as anticipated, the reactivities of B(MIDA) and B(dan), two effective protecting groups for boronic acids on **3c**, could also be differentiated in Suzuki-Miyaura coupling under aqueous basic conditions, affording **5** in good yield.

At last, taking advantage of the Ir-catalysed C-H borylation process developed by our group and the methodology reported herein, a 1,3,5-trisboronyl benzene was synthesized in three steps from commercially available 3-chlorophenylboronic acid, with these boronyl groups different from each other. As far as we know, this represents the first example of assembling three different boronyl groups onto the same molecule. This new type of organoboron reagents should be useful building blocks in modular synthesis of multi-functionalised arene compounds.

In conclusion, by employing the non-symmetrical diboron compound B(pin)-B(dan) as the borylating reagent in Pd-catalysed Miyaura-type reaction, we have realized the chemoselective B(dan) moiety transfer to various aryl rings, yielding masked boronic acids directly from abundant and cheap aryl halides. We have also demonstrated the usefulness of this naphthalene-1,8-diamino borylation in facile preparation of boron-differentiated di- and polyboron compounds and other functionalised aryl boron compounds.

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