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Borylation

Acyl-Directed *ortho*-Borylation of Anilines and C7 Borylation of Indoles using just BBr₃

Saqib A. Iqbal, Jessica Cid, Richard J. Procter, Marina Uzelac, Kang Yuan, and Michael J. Ingleson*

Abstract: Indoles are privileged heterocycles found in many biologically active pharmaceuticals and natural products. However, the selective functionalization of the benzenoid moiety in indoles in preference to the more reactive pyrrolic unit is a significant challenge. Herein we report that N-acyl directing groups enable the C7-selective C-H borylation of indoles using just BBr₃. This transformation shows some functional-group tolerance and notably proceeds with C6 substituted indoles. The directing group can be readily removed in situ and the products isolated as the pinacol boronate esters. Acyl-directed electrophilic borylation can be extended to carbazoles and anilines with excellent ortho selectivity. 4-amino-indoles are amenable to this process, with acyl group installation and directed electrophilic C-H borylation enabling selective formation of C5-BPin-indoles.

C-H borylation is a powerful methodology to form synthetically versatile C-B bonds.^[1] Numerous methods have been developed, with iridium-catalysed C-H borylation one of the most notable.^[1] This method functionalises the pharmaceutically important heteroarene indole at the C2-position.^[2] Alternative indole C-H borylation methods include electrophilic borylation (dominated by electronic effects)^[3] and C-H lithiation/borylation (controlled by C-H acidity).^[4] However, these also functionalise the pyrrole unit (at C3 and C2, respectively, Scheme 1 top left). Indole C-H borylation that occurs selectively on the less reactive benzenoid unit is desirable, including for accessing C5 and C7-functionalised indoles which are motifs found in many biologically active natural products and pharmaceuticals (e.g. chloropeptin I, teleocidins, hippadine, tiplaxtinin).^[5] To date the selective C5-H/C7-H borylation of indoles in the presence of C2-H/ C3-H requires prefunctionalised indoles (e.g. halide at C5/ C7) or functionalisation of the more reactive C2-H/C3-H site prior to C5-H/C7-H borylation and then unmasking of the C2-H/C3-H.^[6] To the best of our knowledge, one example of directed iridium-catalysed C-H borylation^[7] provides the only exception to these requirements

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Scheme 1. Select previous work on the borylation of indoles, specifically borylation reactions occurring at the C5 and C7 positions. Bottom inset, this work on acyl-directed electrophilic C–H borylation at C5 and C7 using BBr_{3} .

(Scheme 1, middle left).^[8] This process while notable uses ruthenium and iridium catalysts and substrates containing C6 substituents are not viable (6,7-disubstituted indoles are also bioactive motifs for example, indole isosteres of combrestatins).^[5,6c,9] Therefore a simple, precious metal free route for the C–H borylation of indoles that is selective for: (i) C7 (over C2), including for C6 substituted indoles, and (ii) C5 (over C3), would be highly notable particularly if using a readily removed directing group.

C-H borylation using BX_3 (X = Cl or Br) is an attractive method to form organoboranes,^[3a,b,10,11] and directed borylation using BX₃ has proved to be a powerful route to form B-C bonds for organic materials applications.^[12] Directed electrophilic C-H borylation is dominated by directing R₂N- or Nheterocycle groups with borylation generally forming six membered boracycles preferentially over other ring sizes.^[13] The extension of C-H borylation using BX₃ to the C5/C7 positions of indoles would be highly attractive. However, this requires conditions that disfavour electrophilic C3-H borylation (which is relatively facile) and a directing group that: (i) is compatible with BX₃; (ii) enables selective borylation at the desired position; (iii) is readily deprotected post C-H borylation. Transition metal-catalysed C7-H indole functionalisation often uses bulky phosphinyl directing groups installed at N1 which are challenging to remove (requiring refluxing with $LiAlH_4$,^[5,6a,14] however, in limited cases Nacyl directing groups also have been used^[5,15] and these are

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more readily removed. Herein we demonstrate that *N*-acyl directing groups are compatible with BBr₃ and lead to C7–H borylation of indoles generating useful C7-BPin products on work up (Scheme 1, bottom). Notably, borylation is compatible with C6 substituted indoles in contrast to the iridium-catalysed process. Furthermore, acyl directing groups also enable *ortho* C–H borylation of anilines using BBr₃, including of 4-amino indoles which affords C5-BPin indoles.

To guide our selection of appropriate acyl directing groups initially we probed the thermodynamic outcome from indole borylation at C2 and C7 computationally. Notably, the C7 borylated isomer is calculated to be thermodynamically favoured over the C2 (Scheme 2) isomer in all cases, this is



Scheme 2. Relative energy of C2 and C7 borylated isomers calculated at the M06-2X/6-311G(d,p) level with a polarizable continuum model of DCM.

attributed to (i) the differing degrees of steric clash between R and the C7–H and C2–H hydrogens (as previously noted);^[16] (ii) the differing bond angles in 5 and 6-membered boracycles, with the former leading to compressed O-B-C angles relative to the latter (which approaches the ideal for tetrahedral boron, Scheme 2). C7-borylation is also calculated to be the kinetic outcome (for R = ¹Bu) based on borylation proceeding via acyl→BBr₃ formation, [acyl→BBr₂]⁺ formation and then S_EAr (see SI).

Based on these calculations the borylation of 1-benzoylindole, **1a**, and 1-pivaloyl-indole, **2a**, was targeted. To disfavour borenium cation formation and indole C3 borylation conditions were required avoiding coordinating exogenous base.^[3b,17] For example, using reagents which lead to $[(amine)BX_2]^+$ cations (e.g. BBr₃/ 2,6-lutidine)^[11] led to the borylation of **2a** at C3 selectively (see SI) with no C2 or C7 borylation observed (Scheme 3). Therefore, BCl₃ and BBr₃ in the absence of base were utilised.

While BCl₃ resulted in no borylation of **1a** and **2a**, with BBr₃ C–B bond formation proceeded with both these indoles, forming products with $\delta_{11B} \approx 0$ ppm (distinct to amide-BBr₃ adducts for which δ_{11B} is ca. –10 ppm). Subsequent addition



Scheme 3. Borylation of **2a** under conditions that generate [(amine)- BX_{2}]⁺ borenium cations.

of pinacol/Et₃N led to formation of the pinacol boronate esters **3a–5a** (Scheme 4). The disparity between BCl₃/ BBr₃ also has been observed in N-heterocycle directed borylation and the origin of this has been examined previously.^[18] The



Scheme 4. Borylation of **1a** and **2a** with BBr₃ and subsequent protection with pinacol/Et₃N or ZnPh₂. Right, solid state structure of **7** (hydrogens omitted and ellipsoids at the 50% probability level).^[24]

regioselectivity of borylation using BBr₃ was assessed by NMR spectroscopy in situ and post pinacol protection. This revealed that borylation of 1a led to C7 and C2 borylation products (with 3a and 4a formed in a 4:1 ratio). Borylation of 2a with BBr₃ led to more selective C7 borylation, with compound 5a-BBr₂ the major borylated product observed in situ (in ca. 85-90% conversion, see SI). 5a-BBr₂ and 6a-BBr₂ are more soluble (than benzoyl congeners) enabling in situ reaction monitoring. Notably, while minor amounts of 6a-**BBr**, were observed in situ no **6a** was observed after pinacol protection. To confirm regioselectivity ZnPh₂ was added to the reaction mixture from 2a/BBr₃ to form predominantly 7 (right, Scheme 4) which has a δ_{11B} of 8.6 ppm indicating a four-coordinate boron centre (in contrast 5a has a broad δ_{11B} at 26 ppm consistent with a weaker PinB-O_{pivalov1} interaction). 7 was isolated in 42% yield and subsequently crystallised with X-ray diffraction studies confirming the formulation as the C7-borylated regioisomer. The solid state structure of 7 revealed a B-O distance of 1.610(2) Å and a O-B-C angle of 104.3(1)° that deviates from that calculated for 5a-BBr₂ presumably due to the different steric demand of BPh₂ vs. BBr₂. The complete absence of C3-borylation is consistent with the requirement for boranes more electrophilic than BBr₃ (e.g. borenium salts) to effect intermolecular indole C3 borylation.[3b, 17]

The substrate scope was explored next and notably C6 substituted N-pivaloyl-indoles were amenable to C–H borylation using BBr₃ in moderate to good yields (e.g., 5c and 5d) (Table 1). The 6-methoxy derivative 2e was also a viable substrate, however, it underwent competitive ether cleavage

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Table 1: Substrate scope of pivaloyl-directed C7-borylation.



Conditions A = 1. 2.2 equiv BBr₃ in DCM. 2. + pinacol/Et₃N. Conditions B = 2.2 equiv BBr₃ in DCM, 2. + pinacol/Et₃N 3. + MeOH, 60 °C. Yields are of isolated products post column chromatography. [a] = using 1 equiv BBr₃.

with BBr₃ producing two C7-borylated products (5e and 5f) in varying amounts depending on the amount of BBr3 used. Conditions for one-pot C-H borylation, pinacol protection and pivaloyl deprotection simply required the addition of methanol after BPin formation and heating to 60°C. The removal of the pivaloyl group occurs without any observable C-B cleavage. This enables three steps to be achieved in onepot with no solvent switches with 8a formed in 71% isolated yield. These conditions were applicable to indoles substituted at C2, C3, C4, C5 and C6 (8g-81), and containing electron withdrawing and donating groups. The reaction was performed on a 3 mmol scale to provide 0.82 g of 8g in 86% yield. However, 5-SMe, 5-NO2 and 4-CN substituted indoles did not furnish isolable C-BPin products, while attempts with a bulkier group at C6, 6-(p-tolyl)-N-pivaloyl- indole, led to C2 borylation dominating (35:65 C7:C2). Compounds 8x are useful in Suzuki-Miyaura cross couplings, allylations and halogenations,^[8] and we note that **8a** readily undergoes oxidation with H₂O₂/NaOH to form 7-hydroxy-indole.

During substrate screening minor C2–BBr₂ borylation (forming 6x-BBr₂) often was observed. Attempts to form the C7–BBr₂ products (5x-BBr₂) selectively by heating (in sealed tubes so HBr does not leave the system) failed to change the C2:C7 ratio suggesting that C–H borylation of these indoles is irreversible under these conditions. However, it was observed that the ratio of C2:C7 BBr₂ products was different to that of the C2:C7 BPin products (with C7-BPin increasing). Furthermore, in a number of cases the amount of 5x/8x isolated was greater than that possible based on the observed 5x-BBr₂:6x-BBr₂ ratio (precluding C2-selective protodeborylation during pinacol addition as the only origin of ratio changes). For example, substrate 2k borylates to form a 5k-BBr₂:6k-BBr₂ ratio of ca. 55:45 (by ¹H NMR spectroscopy),

however, post work up **8k** was isolated in 75% yield. This indicates that addition of pinacol enables C2–B protodeborylation and C7–H borylation. As the BBr₂ products are stable to isomerisation in the presence of HBr this suggests that it is a C–B(OR)Br or C–B(OR)₂ species that is undergoing protodeborylation and leading to more selective C7–H borylation.^[19] While the species undergoing C2→C7 isomerisation on pinacol addition is unknown Lewis/Brønsted acid initiated isomerisation of (RO)₂B-Aryl has been previously observed.^[17]

To expand the utility of acyl-directed electrophilic borylation other N-heterocyclic frameworks were explored. However, *N*-pivaloyl-carbazole did not undergo C–H borylation using BBr₃ (even on heating). This is attributed to steric crowding between the two proximal C–H units (at C1 and C8, Scheme 5, top left) and the pivaloyl 'Bu group that presum-



Scheme 5. The directed borylation of *N*-benzoyl carbazole using BBr₃. Inset, the solid state structure of **10** and **11**, ellipsoids at the 50% probability level.^[24]

ably results in large B-O-C-N dihedral angles in the pivaloyl analogue of **10**. Benzoyl contains a smaller R group (phenyl relative to ^tBu), therefore *N*-benzoyl carbazole, **9**, was combined with BBr₃. This did not lead to C–H borylation at room temperature, instead the Lewis adduct, **10**, was formed which was poorly soluble in DCM facilitating isolation and characterisation (including by X-ray diffraction, Scheme 5).

Heating combinations of 9/ BBr₃ led to high yielding C–H borylation at the C1 position. The C–H borylated product, **11**, could be isolated (and structurally characterised by X-ray diffraction studies) or protected at boron in situ to furnish the pinacol boronate ester **12** in excellent yield (96%). For **10** and **11**, the C=O (1.284(3) and 1.296(7) Å) and O–B distances (1.485(3) and 1.504(8) Å) reveal minimal difference, while the O-B-C angle in **11** (109.9(5)°) is comparable to that calculated for **5a-BBr**₂ and is close to ideal for four coordinate boron centres. Notably the B–O distance in **11** is significantly shorter than in **7** indicative of the greater Lewis acidity of the BBr₂ moiety relative to BPh₂.

We next explored the *ortho* borylation of anilines (Scheme 6). In previous work, borenium mediated electrophilic borylation of anilines proceeded at the *para* position.^[17] *Ortho* borylated anilines are accessible e.g., by directed lithiation of carbamate functionalised anilines,^[20] however, this approach has functional group limitations (e.g., C–Br).

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Scheme 6. Directed *ortho* electrophilic borylation of N–H and N–Me anilines. Pivaloyl-directed C–H borylation proceeds at 20 °C (over the course of 3–16 h), whereas as the benzoyl congener requires heating at 60 °C for 16 h.

Both *N*-pivaloyl and *N*-benzoyl anilines were found to undergo selective *ortho* borylation using BBr₃, with no *para*-borylation observed.

This methodology was applicable to *o*-, *m*- and *p*-substituted anilines, forming **13**c-e in good yield, including for a bromo containing derivative (**13**d). Directed borylation with BBr₃ also can be applied to tertiary amides with the *N*-Me derivative, **13**f, formed in good yield (83%). Smith, Chattopadhyay and co-workers have recently developed directed iridium-catalysed *ortho*-borylation of anilines using B₂eg₂ (eg = ethylene glycolate).^[21] This report is notable, but while excellent for N-H systems it is low yielding with *N*-Me substituted anilines (<25%),^[21] in contrast to the high yielding formation of **13f** using just commercially available DCM solutions of BBr₃.

N-Bn-indol-4-yl-2,2-dimethylpropanamide, 14, next was investigated with it hypothesised that borylation would occur at C5 instead of C3 (the preferred site for S_EAr in indoles) due to the preference for the formation of six membered boracycles over seven.[13c] Functionalisation of the C5-H of indoles is important for accessing pharmaceuticals such as C4-amino-C5-functionalised indoles (e.g. Branebrutinib).^[5,22] The thermodynamics of C5 vs. C3 borylation again was probed by DFT calculations which showed the C5 isomers 15A to be more stable than the C3 isomers 15B (inset, Scheme 7) for both halide and pinacol substituents. C5 borylation of 14 was achieved in high selectivity with the pinacol boronate ester 16 formed in moderate yield (77% in situ and 40% post purification). Attempts to monitor the borylation of 14 at the BBr₂ stage were prevented by this intermediate being poorly soluble. Finally, the ability to perform a C5/C7 double C-H borylation using BBr₃ was demonstrated using 17 (made in one step from 4-aminoindole). This formed 18 selectively post pinacol protection.



Scheme 7. Top, relative energy of C3 and C5 borylated isomers at the M06-2X/6-311G(d,p) level, PCM (DCM). Bottom, borylation of **14** and **17**. In situ yields versus an internal standard, isolated yields are provided in parentheses.^[23].

Notably, in situ NMR spectra prior to pinacol addition show that the C3, C7 diborylated compound, **19**, was formed as the major product and this does not isomerise on standing. However, addition of pinacol induces isomerisation of the C3–B moiety to form the thermodynamically favoured C5-BPin unit and yield the desired C5/C7 product in good conversion (72%).

In summary, *N*-pivaloyl is an effective and readily removed directing group enabling C7 borylation of indoles and *ortho* borylation of anilines simply using commercial solutions of BBr₃. The process is complementary to borylation with [(amine)BBr₂]⁺ and to iridium-catalyzed directed borylation as C6-substituted indoles are tolerated using BBr₃, while it has complementary functional group tolerance to directed lithiation methods. Notably, in a number of cases pinacol induced isomerisation of the initial borylated regioisomer is essential to access the desired products containing C5–B and C7–B units. Due to the simplicity of this process and the many heterocycles containing N–H groups we believe acyl-directed borylation with BBr₃ will be applicable to many other systems.

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Conflict of interest

The authors declare no conflict of interest.

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Angew. Chem. Int. Ed. 2019, 58, 1-6



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Communications



Communications



S. A. Iqbal, J. Cid, R. J. Procter, M. Uzelac, K. Yuan, M. J. Ingleson* ____

Acyl-Directed *ortho*-Borylation of Anilines and C7 Borylation of Indoles using just BBr₃



B-directed: Acyl-directed electrophilic C-H borylation provides access to novel C5 and C7 borylated indoles using just BBr₃ as the borylating agent via the formation of six-membered boracycles.

6 www.angewandte.org