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Regioselective *ortho* halogenation of *N*-aryl amides and ureas *via* oxidative halodeboronation: harnessing boron reactivity for efficient C–halogen bond installation†

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The installation of the C–halogen bond at the *ortho* position of *N*-aryl amides and ureas represents a tool to prepare motifs that are ubiquitous in biologically active compounds. To construct such prevalent bonds, most methods require the use of precious metals and a multistep process. Here we report a novel protocol for the long-standing challenge of regioselective *ortho* halogenation of *N*-aryl amides and ureas using an oxidative halodeboronation. By harnessing the reactivity of boron over nitrogen, we merge carbonyl-directed borylation with consecutive halodeboronation, enabling the precise introduction of the C–X bond at the desired *ortho* position of *N*-aryl amides and ureas. This method offers an efficient, practical, and scalable solution for synthesizing halogenated *N*-heteroarenes under mild conditions, highlighting the superiority of boron reactivity in directing the regioselectivity of the reaction.

Introduction

The amide functional group is an important structural motif found in a variety of natural products and biologically active molecules. For example, 25% of known drugs exhibit an amide motif in their complex structure and several of these substances contain the *N*-aryl amide functional group, for example, atorvastatin, aubagio, and prilocaine.¹ Thus, transformations that are designed to cause structural modifications in the proximity of the amide functional group are of utmost importance as structural modifications, often drastically change the physiological and biological properties of pharmaceutically active molecules.² In this regard, regioselective *ortho*-halogenation of *N*-aryl amides is of high interest as the halogen serves as one of the most versatile synthetic handles for functional group interconversions.³

However, traditional electrophilic aromatic halogenation of *N*-aryl amides typically results in mixtures of *para* and *ortho*/

para substitution (Fig. 1Aa).^{4b,d,e} A higher selectivity can be obtained with transition metal-catalyzed C–H functionalization (Fig. 1Ab).⁴ However, these halogenations are limited by the need for tailored activating groups and inert atmosphere conditions, which can impede synthetic efficiency.⁵ In addition, these transformations are generally restricted to *meta* and *para*-substituted aryl amides due to the formation of di-substituted halogenated side products.^{4b,f}

Despite the maturity of the field *ortho* functionalization remain elusive. To address the selectivity issue, an alternative strategy would be to install boron on the aromatic ring to alter the aromatic reactivity (Fig. 1Ac) and achieve *ipso*-substitution *via* an oxidative halodeboronation at the C–B aromatic carbon instead of electrophilic aromatic substitution (EAS), providing boron control over nitrogen control. An attractive approach to installing the boron is to use the anilide amide as a handle to direct the aromatic borylation, as first developed by Murakami in 2010 (Fig. 1B). Murakami disclosed a BBr_3 -mediated synthesis of pyridine-based boracycles, which has proven to be an exceptionally attractive approach in functionalizing *N*-heteroarenes under metal-free conditions.⁶ Recently, BBr_3 -mediated borylative functionalization *via* stable boracycles has developed into a reliable synthetic route for installing hydroxyl and Bpin functionality in the *ortho*-position to a variety of directing groups (Fig. 1B).⁷ More recently, we showed that the chemistry is not limited to only hydroxyl and Bpin, but it can also be used for installing halogens to phenyl pyridines and aryl aldehydes (Fig. 1B).⁸ One of the key findings in this study is that

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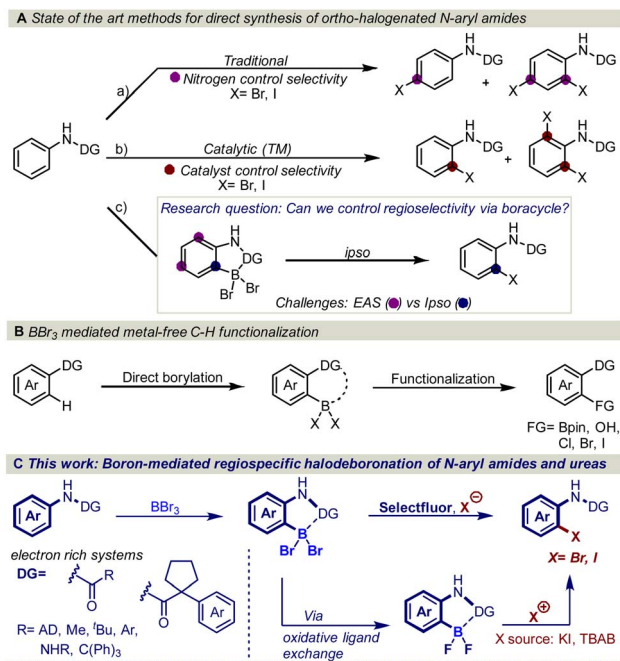


Fig. 1 (A) Literature reports on *ortho* halogenation of *N*-aryl amides and our research question. (B) Recent reports on BBr_3 -mediated C-H borylation and functionalization. (C) This work: boron-mediated regioselective halodeboronation of *N*-aryl amides and ureas.

the reaction proceeds through a difluoroboracycle, which plays a critical role in controlling the reaction's regioselectivity and preventing side reactions. However, the reaction has a limitation in that it requires an electronically deficient system. This is because electronically rich systems can lead to challenges in achieving regioselective halogenations, as competing electrophilic aromatic substitution (EAS) reactions may occur. The reactivity of electronically rich systems is highly dependent on the directing groups, making it difficult to control regioselectivity, especially with electron-rich directing groups. Inspired by the synthetic opportunities that would arise from an oxidative halodeboronation on an electronically rich system, we set out to test the capabilities of a new difluoroborane system on electronically rich *N*-aryl amides (Fig. 1C). Herein we report a convenient protocol for the unexplored regioselective *ortho*-halogenation of *N*-aryl amides and ureas. Our developed protocol shows that by using a boron handle, it is possible to control the regioselectivity of the halogenation through *ipso*-addition at the C-B carbon.

Results and discussion

Our study was initiated by examining the sequential addition of BBr_3 and Selectfluor to *N*-phenylpivalamide (**1a**) in the expectation of forming BBr_2 -complex **2a**, which would undergo oxidative ligand exchange with Selectfluor to yield BF_2 -complex **4a** and cationic bromine. Ultimately, this reaction pathway was expected to generate *ortho*-brominated anilide **5a** via *ipso*-substitution. However, a mixture of regioisomers (**5a** and **5b**) was obtained, which suggests that the electronically rich anilide

substrate was challenging to react selectively and led to side reactions involving the radical and cationic forms of bromine promoted by Selectfluor. It is known that the combination of bromide and Selectfluor generates the cationic form of bromine, and we hypothesize that over time, the cationic form of bromine will generate bromide radicals,⁹ bromine, or HBr, which could explain the formation of the side product **5b**. A potential solution to address this issue would be to avoid adding Selectfluor directly to boracycle **2a** and instead prepare the electrophilic halogenating reagent by mixing Selectfluor with an anionic halogen source prior to its addition to **2a**. This would create a mixture of a cationic halogen source and fluoride, both of which are necessary for the formation of aryl- BF_2 species (**4a**). Our experiments showed that by premixing 1 equivalent of Selectfluor with 1.2 equivalents of potassium iodide (KI) and subsequently adding this mixture to a solution of **2a** in an acetonitrile-water system at 60 °C, we were able to obtain *ortho*-iodinated product **3a** exclusively with a yield of 41% (Table 1, entry 2). We attempted to further enhance the yield by increasing the loading of Selectfluor and KI, which led to the successful generation of **3a** in 93% yield (Table 1, entry 4 vs. entries 2–3). Further, we investigated other inorganic and organic iodine sources and the result showed that KI delivered the best result (ESI, optimization Table S1,† entries 5–8). To be noted, without BBr_3 no product formation was observed, which shows the importance of the six-member ring boron handle in this transformation (Table 1, entry 5). Next, subjecting boracycle **2a** with direct electrophilic iodinating reagents provided **3a** in a poor yield (Table 1, entries 6–8) and no reaction, respectively (Table 1, entry 9).

With our optimized reaction conditions in hand, the reaction scope was investigated. Several *para*-substituted *N*-aryl amides including donating groups, halogenated, and withdrawing groups, were subjected to the protocol, yielding *ortho*-iodinated products **3a–3h** in moderate to excellent yields (42–93%, Scheme 1). Similarly, substitution at *meta*-position including methyl, strongly donating methoxy, halogens and withdrawing trifluoromethyl ($-\text{CF}_3$) were also well tolerated providing exclusively the desired iodinated product **3i–3m** in good to excellent yields (51–96%). Suggesting that the boracycle formation on the less hindered carbon is much faster than the hindered one. A sluggish reaction was observed in the case of substitution at *ortho*-position for substrates **1n** and **1o**. This could be explained by the steric hindrance at the *ortho*-position which potentially hinders the necessary planar conformation of the amide in the boracycle intermediate. However, by modifying the first step time, this issue could be addressed and **3n** could be obtained in 65% and **3o** 48%, respectively. Disubstitution on the *N*-aryl ring provided **3p–3r** in good to excellent yields (57–92%). Furthermore, extended aromatics **3s–3u** and substrates with a higher degree of complexity **3v–3w** are also compatible with the reaction, provided the desired products in moderate to excellent yields without any side reactions on the phenyl rings. Notably, under similar conditions diiodination was performed on oxydibenzene substrate **1x** giving desired product **3x** in 62% yield. The reaction is scalable and can be performed on gram-scale (5.64 mmol) without much loss of efficiency and



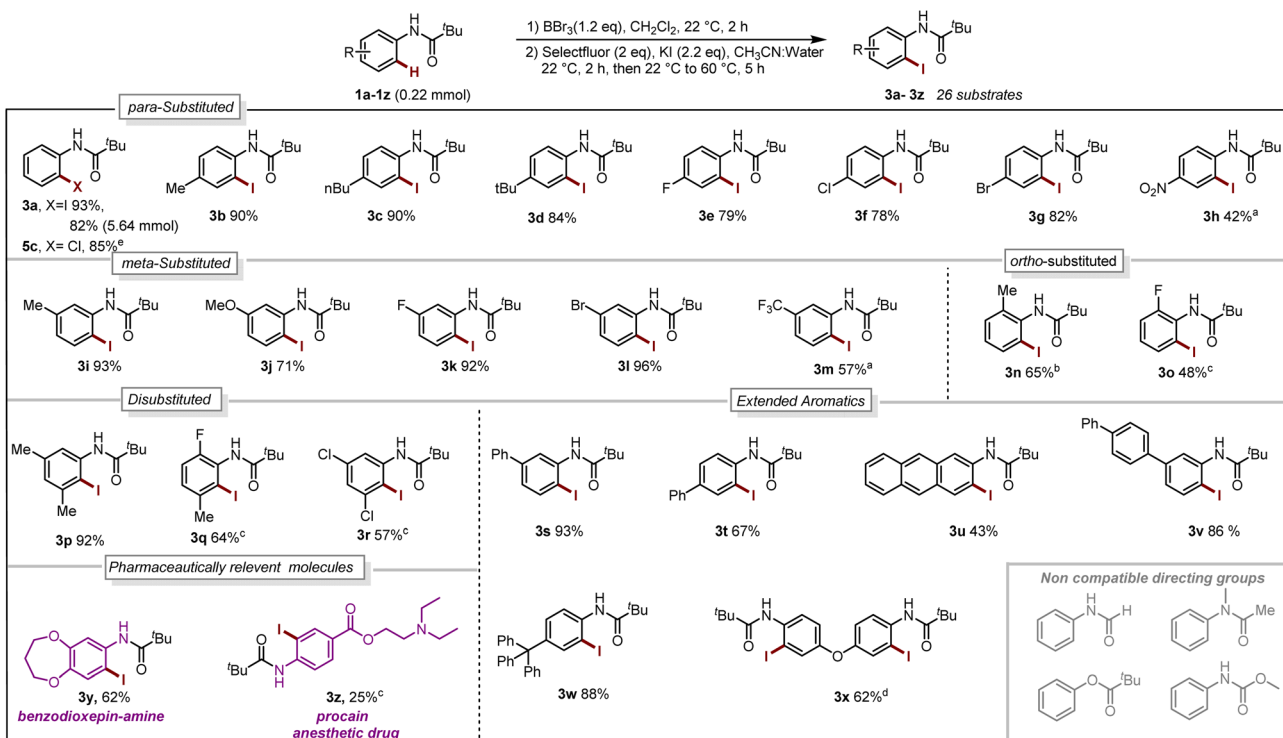
Table 1 Selected reaction optimization (0.22 mmol Scale)^a

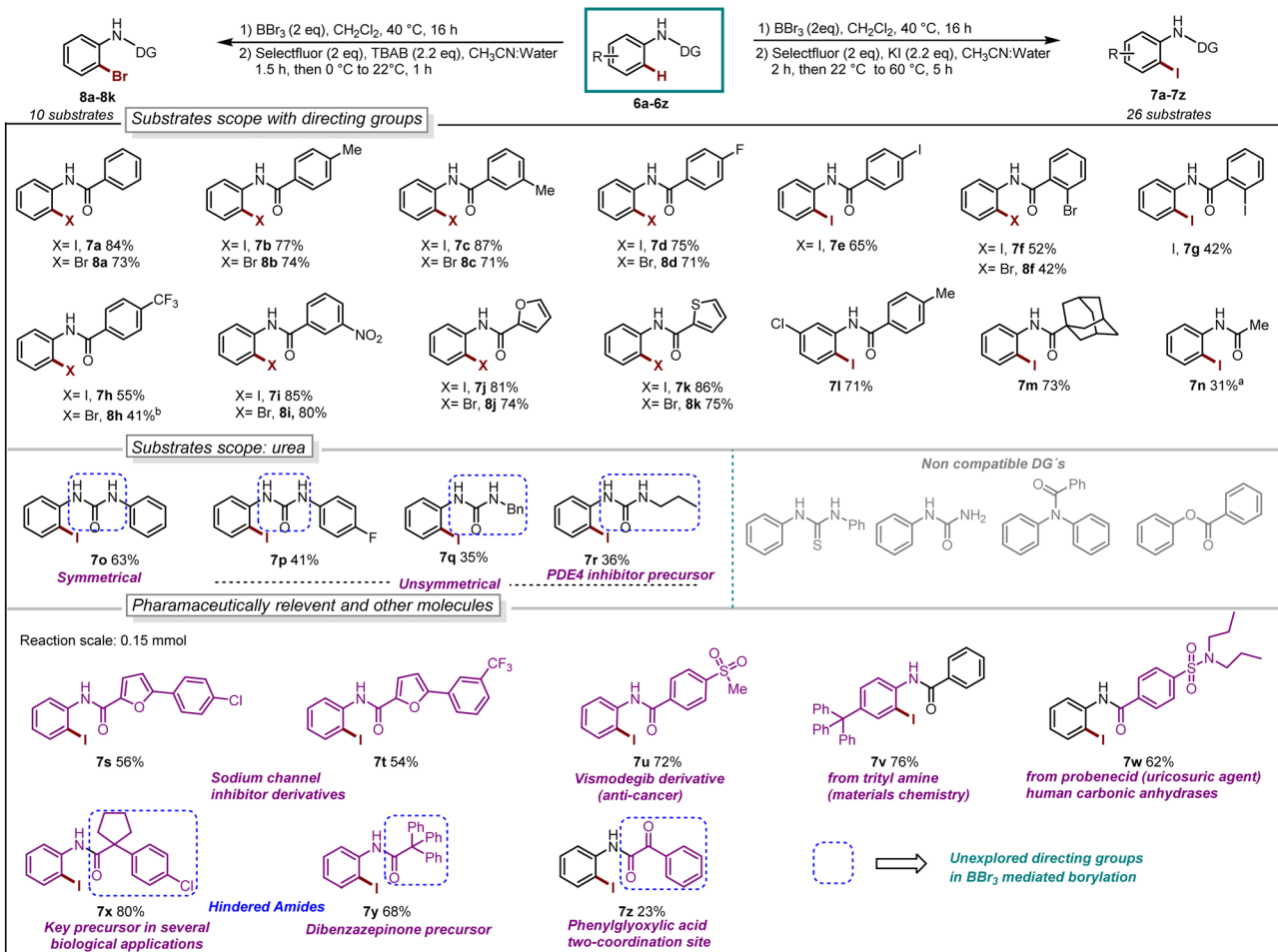
1a	1) BBr_3 (1.2 equiv.), CH_2Cl_2 , 22 °C, 2 h 2) Selectfluor (2 equiv.), halogen source (2.2 equiv.), CH_3CN :Water, 60 °C, 5 h	3a	5a	5b	2a	4a
Entry	Halogen source (equiv.)	Oxidant (equiv.)	Solvent	Time (h)	Yield 3a ^c	NMR ratio 5a:5b
1	—	Selectfluor (2)	CH_3CN :Water	2	—	0.26 : 1
2	KI (1.2)	Selectfluor (1)	CH_3CN :Water	5	41%	—
3	KI (1.7)	Selectfluor (1.5)	CH_3CN :Water	5	66%	—
4 ^a	KI (2.2)	Selectfluor (2)	CH_3CN :Water	5	93%	—
5 ^b	KI (2.2)	Selectfluor (2)	CH_3CN :Water	5	0%	—
6	NIS (1)	—	CH_3CN :Water	5	14%	—
7	ICl (1)	—	CH_3CN :Water	5	12%	—
8	Barluenga's reagent (1)	—	CH_3CN :Water	5	17%	—
9	I2 (2)	—	CH_3CN :Water	5	0%	—

^a Reaction conditions iodination: step (i) 1a (0.22 mmol), BBr_3 (0.26 mmol), in 0.5 mL anhydrous CH_2Cl_2 at 22 °C, 2 h; step (ii) Selectfluor (SF) (0.44 mmol), potassium iodide (KI) (0.48 mmol) in 1.5 mL CH_3CN and 1 mL water at 22 °C, 2 h then 60 °C for 5 h. ^b Without BBr_3 . ^c Isolated yields. Barluenga's reagent = bis(pyridine)iodonium(I) tetrafluoroborate.

compound 3a can be obtained in 82% yield (Scheme 1). Compound 1y exhibiting an unsymmetrical *N*-aryl ring delivered the single isomer exclusively in 62% yield (3y). The biologically active procain derivative 1z with an electron-deficient system also provided desired product albeit in lower yield (3z). Next, we investigated the generality of the protocol with respect to the directing group. It was found that with slight alterations

in the reaction conditions, including heating at 40 °C, the reaction was compatible with a range of directing groups, and the *ortho*-iodination exhibited excellent regioselectivity, yielding products in good to excellent yields (Scheme 2). For instance, electron-donating groups at both *para* and *meta* positions (7a–7c, Scheme 2) were well tolerated, providing the desired iodinated products in yields ranging from 77–87%. Halogen-bearing





Scheme 2 Reaction scope, iodination: ^a BBr_3 (0.66 mmol), 60 °C, 24 h. Bromination: ^bstep-2 at 40 °C, 1.5 h.

substrates on *para* and *ortho* positions also delivered iodinated products in moderate to good yields (7d–7g, 42–75%, Scheme 2). It is noteworthy to mention that these di-halogenated substrates possess high importance due to their facile cyclization for the synthesis of biologically active compounds.¹² Electron withdrawing groups on the directing group, such as $-CF_3$, $-NO_2$, do not impede the reaction (7h 55%, 7i 85%, Scheme 2). Electron-rich heteroaromatics worked exceptionally well without any side reaction (7j 81%, 7k 86%, Scheme 2). Substitution on the aniline part was also well tolerated under similar conditions (7l 71%). Notably, alkyl-based directing groups such as adamantane-1-carboxamide and acetamide also provided desired products (7m 73%, 7n 31%).

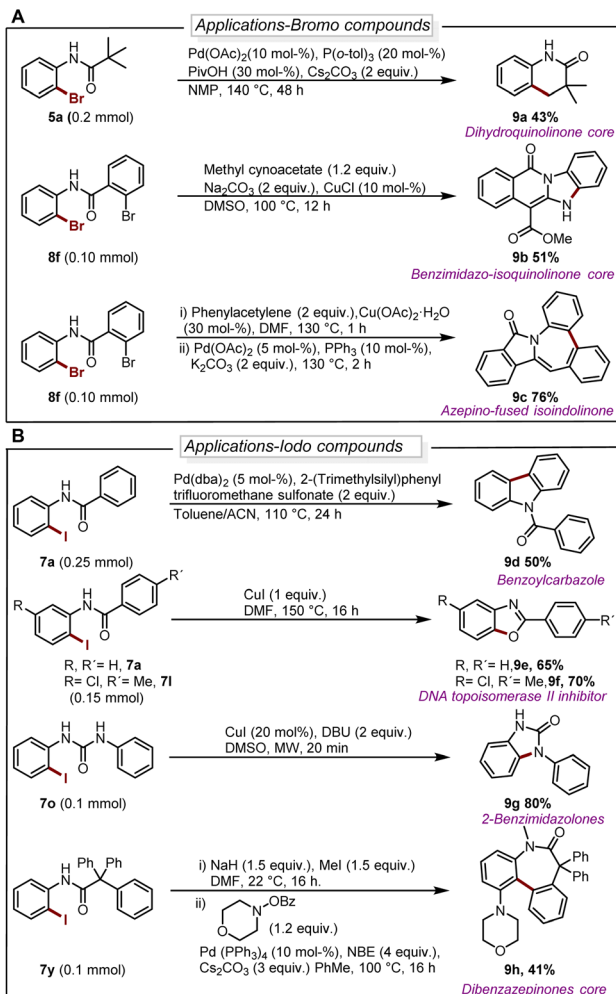
It is also possible to install bromine to the *ortho*-position of anilides by using TBAB as the bromine source. The bromination worked exceptionally well for a series of substrates such as donating groups (8a–8c, 71–74%), halogens (8d 71% and 8f 42%), withdrawing groups (8h 41% and 8i 80%), heteroaromatics (8j 74% and 8k 75%).

Next, we explored urea substrates for our deborylative iodination. Urea represents a more complex substrate class in BBr_3 chemistry due to its polarity, which gives rise to difficulties in purification. The reaction was found to proceed efficiently

with a range of ureas, affording low to moderate yields of the corresponding iodinated ureas with tolerance for phenyl (7o 63%), *p*-fluorophenyl- (7p 41%), benzyl- (7q 35%) and propyl- (7r 36%) substituents. It is noteworthy to mention that this is the first approach for the halogenation as well as the borylation of *N*-aryl urea using BBr_3 . Next, we examined late-stage skeletal editing on a series of bioactive molecules and other applicative substrates.

The deborylative iodination worked exceptionally well for sodium channel inhibitors^{10a} with tolerance for furan heterocycle and $-CF_3$ functionality (7s 56%, 7t 54%, Scheme 2), and vismodegib derivative precursor (7u 72%), hindered *p*-trityl anilide^{10b} (7v 76%), probenecid derivative^{10c} (7w 62%) respectively. Notably, hindered amide substrates such as 6x^{10d,e} and 6y undergo smooth transformation under our condition providing valuable and challenging substrates in good to excellent yields (7x 80%, 7y 68%). Substrate with two coordinating sites such as 6z also provided desired iodination product albeit in lower yield (7z 23%). To explore the utility of deborylative halogenation we carried out a few transformations on halogenated products (Scheme 3A and B). The biologically active 3,4-dihydroquinolinone derivative 9a was synthesized from substrate 5a using a known palladium-catalyzed approach in 43% yield.¹¹



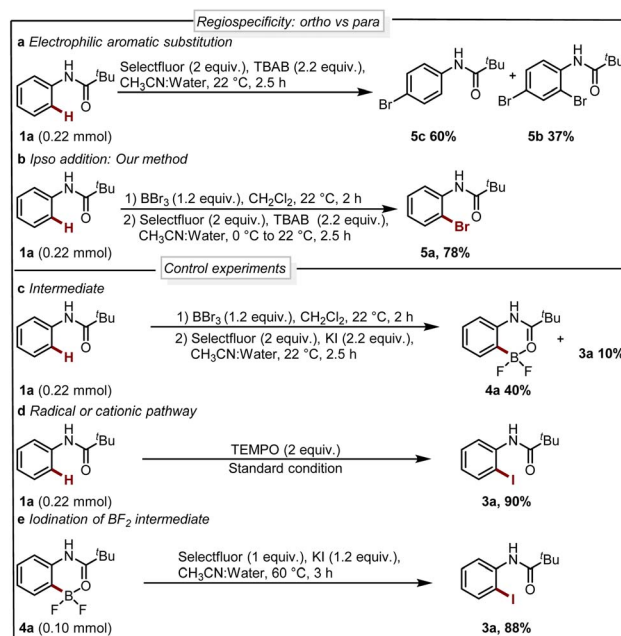


Scheme 3 (A) Diversification of brominated substrates. For reaction details see ESI.† (B) Diversification of iodinated substrates. For reaction details see ESI.†

Next, the benzimidazo-isoquinolinone^{12a} **9b** and azepino-fused isoindolinone core^{12b} **9c** were synthesized from **8f** in 51% and 76% yield respectively (Scheme 3A). Further, the biologically active benzoyl carbazole^{13a} **9d** and benzoxazole core^{13b} **9e–9f** were synthesized using known palladium-aryne and copper chemistry from substrates **7a** and **7f** in 50%, 65% and 70% yield, respectively (Scheme 3B). In a copper-catalyzed reaction, iodo urea **7o** was converted into 2-benzimidazolones¹⁴ **9g** in 80% yield (Scheme 3B).

Finally, we demonstrated the utility of this strategy in the synthesis of the dibenzazepinones core¹⁵ in a reaction where **9h** can be obtained from **7y** in a yield of 41% (Scheme 3B).

Next, we analyzed the selectivity of our protocol, we conducted experiments with the oxidative halogenation conditions and the results are collected in Scheme 4a and b. In the presence of Selectfluor and bromine source tetra butyl ammonium bromide (TBAB), we observed electrophilic aromatic substitution (EAS) of **1a** providing a mixture of brominated products **5b** and **5c** with no selectivity for the *ortho*-bromination (Scheme 4a). Further, with similar conditions in the presence of a boron



Scheme 4 (a) Nitrogen control reaction (EAS). (b) Boron control reaction (*ipso*). (c–e) Control experiments.

handle, we observed exclusively the *ortho* bromination and **5a** could be isolated in 78% yield (Scheme 4b). This result reveals that, using the boron complex the reactivity completely shifted to *ipso* rather than EAS. This experiment supports the fact that our protocol is regioselective where the reactivity of the system is governed by boron rather than nitrogen.

Seeking to gain mechanistic insight into the bromide-to-fluoride ion exchange behavior and C–B bond cleavage, we performed a series of experimental and computational studies. We identified two crucial steps to be investigated, *i.e.* (i) the generation of the difluoroborane species and (ii) the occurrence of ionic *ipso*-addition. To confirm the difluoroborane species **4a** we quenched the reaction prematurely and isolated the **4a** in 40% yield along with 10% of **3a** (Scheme 4c). This result shows the formation of intermediate **4a** in the reaction. The compound **4a** is also confirmed by a single crystal X-ray structure (Fig. 2, ESI Section 9†). In order to get insights into the radical or ionic *ipso* addition mechanism, we conducted the reaction in the presence of radical scavenger TEMPO (Scheme 4d). However, the excess TEMPO did not affect the reaction outcome providing desired iodinated product in 90% yield. Furthermore, to reveal the reactivity and support the mechanism we subjected BF_2 intermediate **4a** to the reaction conditions of the oxidative halideboronation (Scheme 4e) and obtained 88% yield of desired product **3a**.

We performed a computational investigation intended to model the *ipso* addition using trimethyl amine-iodo as a model source for iodination (Scheme 5). Thus, we adopted DFT at the $\omega\text{B97xD/LACVP}^*$ level with a PBF continuum model for water solvation, to evaluate the effect of dibromo-boracycle *vs.* difluoro-boracycle on *ipso* addition. The water model may slightly overestimate the stabilization provided by solvent

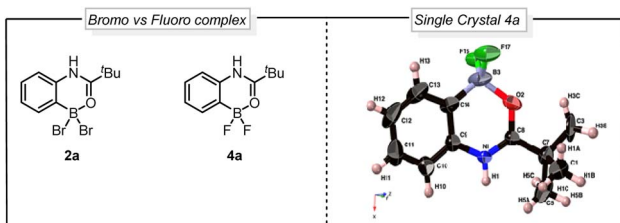


Fig. 2 Comparison between bromo vs. fluoro complex and single crystal X-ray structure of **4a**.

compared to the actual solvent mix but is expected to give results superior to any other type of solvent model available to us.

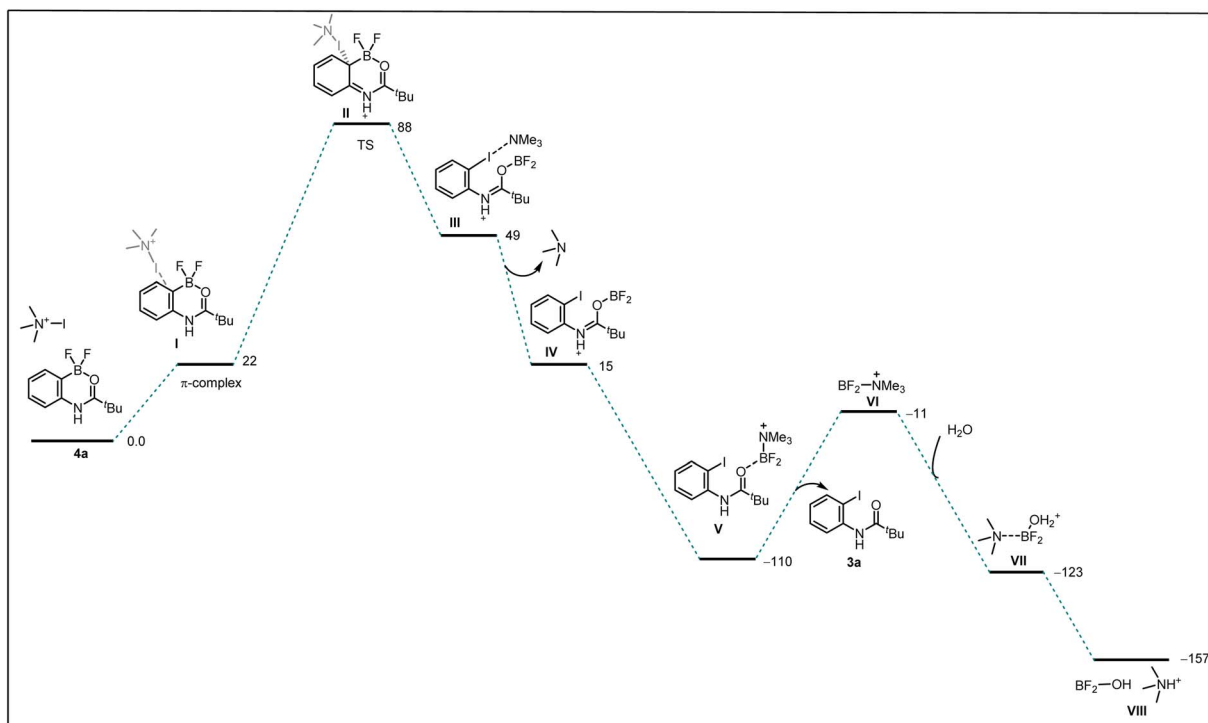
We started by considering tetrahedral difluoro-boracycle **4a** reacting with iodo reagent to π -complex (I) which is endergonic by 22 kJ mol^{-1} . This reacts through TS(II) to complex (III), with a barrier of 88 kJ mol^{-1} compared to separated starting materials. In complex (III) boron is trigonal and has a weak interaction with π electrons of the aromatic ring. The iodine carrier, here modelled by NMe_3 , is weakly associated to the iodine by a halogen bond.

Dissociation of NMe_3 is monotonous on the potential energy surface, but exergonic. Reassociation to boron is very strongly exergonic and drives the overall reaction. Dissociation of final product **3a** is slightly endergonic, but boron can in turn associate to water, and through dissociation and final proton transfer arrive at the endpoint of the computational investigation, (VIII). Overall, formation of **3a** and byproducts from **4a** is

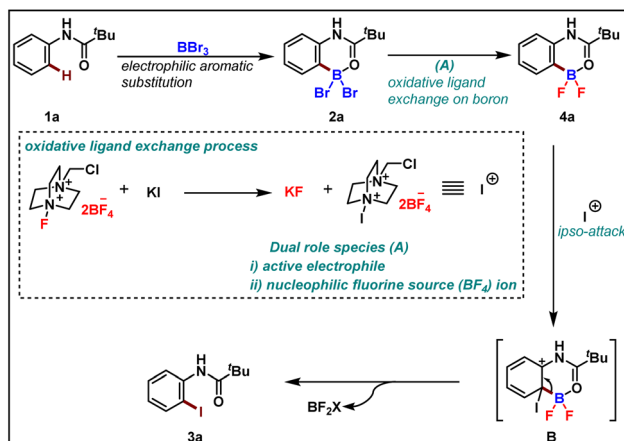
exergonic by 157 kJ mol^{-1} , with a barrier of 88 kJ mol^{-1} . The limiting TS, (II), was located by coordinate driving of the iodine-carbon distance to shorter values. At a C-I distance of 2.2 \AA , the structure is similar to the final structure of the TS, with similar energy, and could be used as a starting point for the subsequent TS search. Similar coordinate drives were performed for all positions of the aromatic ring, as well as for the *ipso*-position of the bromo-boracycle substrate **2a**. In all cases, the energy rose to $>150 \text{ kJ mol}^{-1}$ without snapping into a product-like geometry, clearly showing that iodination is very strongly preferred at the *ipso* position of **4a** compared to **2a**.

To understand the reactivity difference between bromo boracycle **2a** and fluoro-boracycle **4a** (Fig. 2, single crystal X-ray structure), we analyzed the structures of both compounds, as well as the immediate product (III) from *ipso*-substitution of both complexes, where the C-B bond has been broken. In the bromo complex, the boron is electron deficient as compared to the fluoro complex. Despite the higher electronegativity of fluorine, the shorter B-F bonds, and the potential for π -donation from fluorine to the formally empty orbital of boron in (III) stabilize the boron. The BBr_2 complex **2a** has significantly shorter bonds to oxygen and carbon.

In a tetrahedral environment, the boron can compensate for the low electron density by stronger bonds to the other atoms, but in the trigonal planar complex (III), the BBr_2 moiety lacks the stabilization which can be found from *p*-donation in the BF_2 moiety. In other words, RBBR_2 is a stronger Lewis acid than RBF_2 . In the tetrahedral complex, there is not much π -donation from fluorine to boron; the boron *p*-orbital is not empty as it is



Scheme 5 Free energy surface of *ipso*-substitution. We use a simplified model for the base, which is hypothesized to act as a carrier of electrophilic I^+ , to drive the final dissociation of boron, and to deprotonate the water adding to the boron product.



Scheme 6 Proposed mechanism.

occupied with lone pair of carbonyl. Thus, the empty p-orbital can only be stabilized by π -donation from the three attached atoms, oxygen and two halides. The oxygen contribution is the same in both complexes, but the halides are very different. For instance, Br basically cannot do π -stabilization, and the orbitals of Br lone pairs are of the wrong size and distance to give efficient π overlap. On the other hand, F has a perfectly matched, filled p-orbital, which gives a very good π -stabilization.

To rationalize the observed reactivity, based on control experiments and DFT studies we propose a mechanism (Scheme 6) that starts with the formation of boracycle 2a *via* an electrophilic aromatic substitution in the presence of boron tribromide. Next, the bromo-boracycle 2a converts into fluoro boracycle 4a in the presence of an oxidative system (A). In the classical oxidative process,⁹ Selectfluor oxidizes halogen anion X^- to halogen cation X^+ . However, in our case, the oxidative condition plays a dual role which is the generation of electrophile and nucleophilic fluorines for the ligand exchange on boron. In the presence of this oxidative condition, the fluoro boracycle undergoes *ipso* addition on cationic iodine to form intermediate B which upon deborylation forms desired iodinated product 3a.

Conclusion

In summary, we have demonstrated a regioselective, efficient, and practical *ortho* halogenation of *N*-aryl amides and ureas promoted by a boron handle. The power of this method is showcased by halogenation on a set of different carbonyls as directing groups. Our strategy leverages the dual role of Selectfluor as a fluoride source and oxidant, enabling regioselective deborylative *ortho*-halogenation of *N*-aryl amides and ureas. The proposed mechanism involves oxidative ligand exchange on boron followed by *ipso*-addition with concomitant release of the boron species. This newfound reactivity of the C–B bond and B–X bond holds promise as an alternative for the functionalization of a broad range of *N*-heterocycles in the coming years. Overall, our protocol demonstrates its applicability in the synthesis of urea substrates, pharmaceutically

active molecules, and hindered amides, showcasing its strength in material science, medicinal chemistry, and synthetic organic chemistry.

Data availability

All detailed procedures, characterization data, and spectra are available in the ESI.†

Author contributions

H. S. supervised the overall project. G. H. S. conceived the idea and designed the study with G. S. G. F. M. A. N. and L. Ö. contributed to the crystal structure. P. O. N. contributed to the computational study. G. S. G., F. M. A. N., L. Ö. and P. O. N. provided valuable suggestions on the manuscript. H. S. and G. H. S. co-wrote the manuscript.

Conflicts of interest

The authors declare no conflict of interest.

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Notes and references

- (a) N. A. McGrath, M. Brichacek and J. T. Njardarson, *J. Chem. Educ.*, 2010, **87**, 1348–1349; (b) X. D. Yang, X. H. Zeng, Y. H. Zhao, X. Q. Wang, Z. Q. Pan, L. Li and H. Bin Zhang, *J. Comb. Chem.*, 2010, **12**, 307–310; (c) E. Vitaku, D. T. Smith and J. T. Njardarson, *J. Med. Chem.*, 2014, **57**, 10257–10274; (d) M. M. Heravi and V. Zadsirjan, *RSC Adv.*, 2020, **10**, 44247–44311.
- S. Kumari, A. V. Carmona, A. K. Tiwari and P. C. Trippier, *J. Med. Chem.*, 2020, **63**, 12290–12358.
- (a) U. Ladziata, A. Y. Koposov, K. Y. Lo, J. Willging, V. N. Nemykin and V. V. Zhdankin, *Angew. Chemie-Int. Ed.*, 2005, **44**, 7127–7131; *Angew. Chem.*, 2005, **117**, 7289–7293; (b) N. Zheng, K. W. Anderson, X. Huang, H. N. Nguyen and S. L. Buchwald, *Angew. Chemie-Int. Ed.*, 2007, **46**, 7509–7512; *Angew. Chem.*, 2007, **119**, 7653; (c) E. A. Merritt and B. Olofsson, *Angew. Chemie-Int. Ed.*, 2009, **48**, 9052–9070; *Angew. Chem.*, 2009, **121**, 9214–9234; (d) J.-X. Yan, H. Li, X.-W. Liu, J.-L. Shi, X. Wang and Z.-J. Shi, *Angew. Chemie-Int. Ed.*, 2014, **53**, 4945–4949; *Angew. Chem.*, 2014, **126**, 5045–5049; (e) S. Z. Tasker and T. F. Jamison, *J. Am. Chem. Soc.*, 2015, **137**, 9531–9534; (f) Shaifali, P. Mehara, A. Kumar and P. Das, *ChemCatChem*, 2021, **13**, 2459–2464.
- (a) J. M. Fourquez, A. Godard, F. Marsais and G. B. Quéguiner, *J. Heterocycl. Chem.*, 1995, **32**, 1165–1170; (b) R. B. Bedford, M. F. Haddow, C. J. Mitchell and



R. L. Webster, *Angew. Chemie-Int. Ed.*, 2011, **50**, 5524–5527; *Angew. Chem.*, 2011, **123**, 5638; (c) B. Urones, Á. M. Martínez, N. Rodríguez, R. Gómez Arrayás and J. C. Carretero, *Chem. Commun.*, 2013, **49**, 11044–11046; (d) J. Chen, X. Xiong, Z. Chen and J. Huang, *Synlett*, 2015, **26**, 2831–2834; (e) D. Liang, X. Li, C. Wang, Q. Dong, B. Wang and H. Wang, *Tetrahedron Lett.*, 2016, **57**, 5390–5394; (f) S. Kathiravan and I. A. Nicholls, *Chem. - Eur. J.*, 2017, **23**, 7031–7036; (g) R. Das and M. Kapur, *J. Org. Chem.*, 2017, **82**, 1114–1126; (h) Z. lin Li, K. kang Sun and C. Cai, *Org. Biomol. Chem.*, 2018, **16**, 5433–5440; (i) Y. Du, Z. Xi, L. Guo, H. Lu, L. Feng and H. Gao, *Tetrahedron Lett.*, 2021, **72**, 153074; (j) E. Kianmehr and H. Afaridoun, *Synthesis*, 2021, **53**, 1513–1523.

5 (a) D. A. Petrone, J. Ye and M. Lautens, *Chem. Rev.*, 2016, **116**, 8003–8104; (b) R. Das and M. Kapur, *Asian J. Org. Chem.*, 2018, **7**, 1524–1541.

6 N. Ishida, T. Moriya, T. Goya and M. Murakami, *J. Org. Chem.*, 2010, **75**, 8709–8712.

7 (a) L. Niu, H. Yang, R. Wang and H. Fu, *Org. Lett.*, 2012, **14**, 2618–2621; (b) S. A. Iqbal, J. Cid, R. J. Procter, M. Uzelac, K. Yuan and M. J. Ingleson, *Angew. Chem., Int. Ed.*, 2019, **58**, 15381–15385; *Angew. Chem.*, 2019, **131**, 15525–15529; (c) J. Lv, X. Chen, X. S. Xue, B. Zhao, Y. Liang, M. Wang, L. Jin, Y. Yuan, Y. Han, Y. Zhao, Y. Lu, J. Zhao, W. Y. Sun, K. N. Houk and Z. Shi, *Nature*, 2019, **575**, 336–340; (d) J. Lv, B. Zhao, Y. Yuan, Y. Han and Z. Shi, *Nat. Commun.*, 2020, **11**, 1316; (e) G. Wu, X. Fu, Y. Wang, K. Deng, L. Zhang, T. Ma and Y. Ji, *Org. Lett.*, 2020, **22**, 7003–7007; (f) G. Wu, B. Pang, Y. Wang, L. Yan, L. Chen, T. Ma and Y. Ji, *J. Org. Chem.*, 2021, **86**, 5933–5942; (g) S. Li, C. Hu, X. Cui, J. Zhang, L. L. Liu and L. Wu, *Angew. Chem., Int. Ed.*, 2021, **60**, 26238–26245; *Angew. Chem.*, 2021, **133**, 26442–26449; (h) Z. J. Wang, X. Chen, L. Wu, J. J. Wong, Y. Liang, Y. Zhao, K. N. Houk and Z. Shi, *Angew. Chem., Int. Ed.*, 2021, **60**, 8500–8504; *Angew. Chem.*, 2021, **133**, 8581–8585; (i) S. Rej and N. Chatani, *J. Am. Chem. Soc.*, 2021, **143**, 2920–2929; (j) S. Rej, A. Das and N. Chatani, *Chem. Sci.*, 2021, **12**, 11447–11454; (k) J. Lv, X.-J. Zhang, M. Wang, Y. Zhao and Z. Shi, *Chem. - Eur. J.*, 2022, **28**, e202104100; (l) O. Sadek, A. L. Gac, N. Hidalgo, S. Mallet-Ladeira, K. Miqueu, G. Bouhadir and D. Bourissou, *Angew. Chem., Int. Ed.*, 2022, **61**, e202110102; *Angew. Chem.*, 2022, **134**, e202110102; (m) G. Wu, Z. Yang, X. Xu, L. Hao, L. Chen, Y. Wang and Y. Ji, *Org. Lett.*, 2022, **24**, 3570–3575; (n) X. Xu, G. Wu, Z. Yang, X. Liu, L. Hao, Y. Wang, Z. Ma and Y. Ji, *Org. Lett.*, 2022, **24**, 7163–7167; (o) Review: S. Rej and N. Chatani, *Angew. Chem., Int. Ed.*, 2022, **61**, e202209539; *Angew. Chem.*, 2022, **134**, e202209539; (p) G. Berionni, *Angew. Chem., Int. Ed.*, 2022, **61**, e202210284; *Angew. Chem.*, 2022, **134**, e202210284; (q) S. A. Iqbal, M. Uzelac, I. Nawaz, Z. Wang, H. Jones, G. S. Nichol, K. Yuan, C. Millet, G. A. Chotana and M. J. Ingleson, *Chem. Sci.*, 2023, **1**, 3865–3872; (r) D. Zhao and G. Yang, *Synfacts*, 2023, **19**, 0554; (s) Z. Yang, L. Hao, X. Xu, Y. Wang, G. Wu and Y. Ji, *Org. Lett.*, 2023, **25**, 5875–5879; (t) P. Kumar Someswara Ashwathappa, T. Higashi, V. Desrosiers, A. A. Omaña and F. G. Fontaine, *Angew. Chemie-Int. Ed.*, 2023, **62**, 1–7; (u) T. Wang, Z.-J. Wang, M. Wang, L. Wu, X. Fang, Y. Liang, J. Lv and Z. Shi, *Angew. Chem., Int. Ed.*, 2023, e202313205. 8 G. H. Shinde and H. Sundén, *Chem. - Eur. J.*, 2023, **29**, e202203505.

9 (a) R. G. Syvret, K. M. Butt, T. P. Nguyen, V. L. Bulleck and R. D. Rieth, *J. Org. Chem.*, 2002, **67**, 4487–4493; (b) C. Ye and J. M. Shreeve, *J. Org. Chem.*, 2004, **69**, 8561–8563; (c) Z. Dağalan, R. Koçak, A. Daştan and B. Nişancı, *Org. Lett.*, 2022, **24**, 8261–8264.

10 (a) M. E. Kort, I. Drizin, R. J. Gregg, M. J. C. Scanio, L. Shi, M. F. Gross, R. N. Atkinson, M. S. Johnson, G. J. Pacofsky, J. B. Thomas, W. A. Carroll, M. J. Krambis, D. Liu, C. C. Shieh, X. F. Zhang, G. Hernandez, J. P. Mikusa, C. Zhong, S. Joshi, P. Honore, R. Roeloffs, K. C. Marsh, B. P. Murray, J. Liu, S. Werness, C. R. Faltynek, D. S. Krafte, M. F. Jarvis, M. L. Chapman and B. E. Marron, *J. Med. Chem.*, 2008, **51**, 407–416; (b) F. Dumur and F. Goubard, *New J. Chem.*, 2014, **38**, 2204–2224; (c) M. D'Ascenzio, S. Carradori, D. Secci, D. Vullo, M. Ceruso, A. Akdemir and C. T. Supuran, *Bioorg. Med. Chem.*, 2014, **22**, 3982–3988; (d) G. Tang, Z. Qiu, X. Lin, W. Li, L. Zhu, S. Li, H. Li, L. Wang, L. Chen, J. Z. Wu and W. Yang, *Bioorg. Med. Chem. Lett.*, 2010, **20**, 3507–3510; (e) J. Tng, J. Lim, K. C. Wu, A. J. Lucke, W. Xu, R. C. Reid and D. P. Fairlie, *J. Med. Chem.*, 2020, **63**, 5956–5971.

11 J. Yan, H. Li, X. Liu, J. Shi, X. Wang and Z. Shi, *Angew. Chemie - Int. Ed.*, 2014, **53**, 4945–4949; *Angew. Chem.*, 2014, **126**, 5045–5049.

12 (a) J. Lu, X. Gong, H. Yang and H. Fu, *Chem. Commun.*, 2010, **46**, 4172–4174; (b) H. K. Saini, N. K. Nandwana, S. Dhiman, K. Rangan and A. Kumar, *Eur. J. Org. Chem.*, 2017, **2017**, 7277–7282.

13 (a) R. W. Bowman, H. Heaney and P. H. G. Smith, *Tetrahedron Lett.*, 1982, **23**, 5093–5096; (b) C. Lu, N. A. Markina and R. C. Larock, *J. Org. Chem.*, 2012, **77**, 11153–11160.

14 Z. Li, H. Sun, H. Jiang and H. Liu, *Org. Lett.*, 2008, **10**, 3263–3266.

15 X. Abel-Snape, A. Whyte and M. Lautens, *Org. Lett.*, 2020, **22**, 7920–7925.

