

UvA-DARE (Digital Academic Repository)

Photoinduced Halogen-Atom Transfer by N-Heterocyclic Carbene-Ligated Boryl Radicals for C(sp3)-C(sp3) Bond Formation

Wan, T.; Capaldo, L.; Ravelli, D.; Vitullo, W.; de Zwart, F.J.; de Bruin, B.; Noël, T. **DOI**

10.1021/jacs.2c10444

Publication date 2023 Document Version Final published version

Published in Journal of the American Chemical Society

License CC BY

Link to publication

Citation for published version (APA):

Wan, T., Capaldo, L., Ravelli, D., Vítullo, W., de Zwart, F. J., de Bruin, B., & Noël, T. (2023). Photoinduced Halogen-Atom Transfer by N-Heterocyclic Carbene-Ligated Boryl Radicals for C(sp³)-C(sp³) Bond Formation. *Journal of the American Chemical Society*, *145*(2), 991-999. https://doi.org/10.1021/jacs.2c10444

General rights

It is not permitted to download or to forward/distribute the text or part of it without the consent of the author(s) and/or copyright holder(s), other than for strictly personal, individual use, unless the work is under an open content license (like Creative Commons).

Disclaimer/Complaints regulations

If you believe that digital publication of certain material infringes any of your rights or (privacy) interests, please let the Library know, stating your reasons. In case of a legitimate complaint, the Library will make the material inaccessible and/or remove it from the website. Please Ask the Library: https://uba.uva.nl/en/contact, or a letter to: Library of the University of Amsterdam, Secretariat, Singel 425, 1012 WP Amsterdam, The Netherlands. You will be contacted as soon as possible.

UvA-DARE is a service provided by the library of the University of Amsterdam (https://dare.uva.nl)

Photoinduced Halogen-Atom Transfer by *N*-Heterocyclic Carbene-Ligated Boryl Radicals for C(sp³)–C(sp³) Bond Formation

Ting Wan,[∥] Luca Capaldo,[∥] Davide Ravelli, Walter Vitullo, Felix J. de Zwart, Bas de Bruin, and Timothy Noël*



XAT step, delivering the desired nucleophilic alkyl radicals. The present XAT strategy is very mild and accommodates a broad scope of alkyl halides, including medicinally relevant compounds and biologically active molecules. The key role of NHC-ligated boryl radicals in the operative reaction mechanism has been elucidated through a combination of experimental, spectroscopic, and computational studies. This methodology stands as a significant advancement in the chemistry of NHC-ligated boryl radicals, which had long been restricted to radical reductions, enabling C-C bond formation under visible-light photoredox conditions.

INTRODUCTION

Downloaded via UNIV AMSTERDAM on February 6, 2023 at 10:18:04 (UTC). See https://pubs.acs.org/sharingguidelines for options on how to legitimately share published articles.

The possibility to exploit photonic energy in organic synthetic endeavors has dramatically impacted the way chemists assemble molecules. In particular, photocatalysis has enabled a convenient entry to open-shell intermediates,¹ spurring the development of efficient manifolds for the generation of C-,² N-,³ and O-⁴ centered radicals, as well as halogen radicals,⁵ which can be subsequently used to forge new chemical bonds.⁶ In contrast, boron-based congeners have long remained in obscurity,⁷ mainly due to the intrinsic difficulties associated with the handling of these highly electron-deficient and unstable intermediates. However, ligated boryl radicals (LBRs),⁸ i.e. boron-centered radicals where the boron atom is coordinated with a Lewis base, are more stable and provide a suitable entry for use in radical chemistry (Figure 1A).^{9,10}

In particular, *N*-heterocyclic carbene-based (NHC) boranes are emerging as a convenient source of LBRs: NHC boranes are stable crystalline compounds, and a diverse array of NHCs can be ligated to boranes allowing the LBR properties to be fine-tuned.¹¹ Notably, these boranes can be uniquely paired with photocatalysis to generate the targeted boron-centered radicals.¹² Herein, a photocatalyst absorbs visible light and engages with the ligated borane in a single-electron transfer step which, upon deprotonation, generates the corresponding boryl radical. Although these boron-centered radicals have attracted mainly interest from the synthetic community as nucleophilic radicals,^{13,14} they have also been used in the role of halogen-atom transfer (XAT) agents.

In the latter scenario, the halogen-affinity of the LBR is exploited for the homolytic activation of a C-X bond to yield

carbon-centered radicals. However, this manifold has been so far mainly used to reduce C-X bonds into the corresponding C-H bonds via a radical chain mechanism.^{11,12,15} Surprisingly, boryl radicals have been largely overlooked for the construction of C-C bonds (Figure 1B). In an early example, the radical silvldifluoromethylation of electron-deficient alkenes was reported.¹⁶ Herein, a very specific interaction, based on halogen-bonding between the substrate and an NHC borane, was needed to trigger the desired C-X bond photolysis and subsequently initiate the radical chain mechanism sustained by the LBR (Figure 1C). Inspired by this report, we questioned whether it would be possible to realize a more general strategy to generate the pivotal ligated boryl radical. Such a pathway might allow the engagement of a broader array of substrates, ultimately leading to a general approach for C-C bond formation. Moreover, succeeding in this challenge would provide a cheap, tunable, and sustainable alternative for other XAT-based approaches¹⁷ using silyl¹⁸ and α -aminoalkyl radicals.¹⁹ To this end, we disclose our results regarding the development of a mild and broadly applicable protocol for $C(sp^3)-C(sp^3)$ bond formation using photoinduced XAT by NHC-ligated boryl radicals under blue-light irradiation (Figure 1D). The key role of the NHC-ligated boryl

Received: October 1, 2022 Published: December 30, 2022



A Boryl Radicals (BRs) vs Ligated Boryl Radicals (LBRs)



C SilyIdifluoromethylation of electron-deficient alkenes



Figure 1. (A) Boryl radicals (BRs) versus ligated boryl radicals (LBRs). (B) NHC boranes as a bench-stable source of LBRs. (C) Silyldifluoromethylation of electron-deficient alkenes. (D) This work. L_B: Lewis base.

radicals in the operative reaction mechanism has been uncovered through a combination of experimental, spectroscopic, and computational studies.

RESULTS AND DISCUSSION

At the outset of our investigations, we recognized the importance of finding the ideal combination of ligated borane and photocatalyst to establish an efficient and competent system to promote the desired reactivity. In detail, the photocatalyst would absorb the visible light and subsequently generate the LBR via a SET followed by deprotonation.¹² The resulting boryl radical would be ultimately entrusted with the XAT step.

We immediately realized that the success of our plan hinges on (i) the redox potentials of the borane and the photocatalyst and (ii) the halogen-affinity of the resulting LBR. Based on literature and our experimental data (see Section 11 in the Supporting Information), we identified NHC-ligated borane B1 as an ideal candidate for our purposes: in fact, its oxidation potential $(E_{pa}(B1^{\bullet+}/B1))$ is +0.89 V vs SCE,²⁰ which suggests that this approach should be feasible in combination with routinely used photoredox catalysts. Other ligated boranes tested (see Section 11 in the Supporting Information) were found to have an exceedingly high redox potential, thus preventing the formation of the desired ligated boryl radical.

pubs.acs.org/JACS

Next, we started our investigation by screening different photoredox catalysts that possess a higher excited state reduction potential $(E(PC^*/PC_{red}))$ than that of B1 (Table 1). Our experiments revealed that, when a degassed CH₃CN

Table 1. Optimization of Conditions^a



^aGC yields are given using biphenyl as external standard.

solution of 2a (0.1 M), 1a (2 equiv), and B1 (1 equiv) was irradiated with blue light (λ = 456 nm, 12 h, rt) in the presence of 2 mol % of the organic photocatalysts PC1 or PC2, product 3 could be obtained in 43-44% GC yield (Table 1, entries 1 and 2). In contrast, $Ru(bpy)_3(PF_6)_2$ (PC3) gave worse results (Table 1, entry 3).

Next, we screened the effect of the solvent on the transformation and noticed that protic reaction mixtures boosted the reaction yield (Table 1, entry 4: up to 68% in CH₃CN/H₂O 9:1). Fine-tuning the ratio of the reagents, the photocatalyst loading, and reaction time allowed an excellent 79% yield to be obtained for the targeted hydroalkylation (Table 1, entries 5-6). Several control experiments revealed that excluding light, PC1, or B1 did not lead to any product formation (Table 1, entries 7-8). Moreover, 3 was not produced at elevated temperatures either (Table 1, entry 9). The exclusion of molecular oxygen appeared to be crucial as air-equilibrated conditions led to a significantly reduced yield (Table 1, entry 10: 37%), while O_2 -saturation shut down reactivity (Table 1, entry 11). It is important to mention that,

pubs.acs.org/JACS



Figure 2. Substrate scope for LBRs-mediated XAT under visible-light irradiation. For secondary and tertiary organic halides: **2** (0.5 mmol), **1** (2 equiv), **B1** (1.2 equiv) in CH₃CN/H₂O 9:1 (5 mL) in the presence of **PC1** (5 mol %), 3 h. For primary organic halides: **2** (2 equiv), **1** (0.5 mmol), **B1** (1.2 equiv) in CH₃CN/H₂O 9:1 (5 mL) in the presence of **PC1** (5 mol %), 12 h. Solutions were bubbled with N₂ (5 min) prior to irradiation ($\lambda = 456$ nm). ^a Reaction time: 18 h. ^b Solvent: ethyl acetate. ^c See Supporting Information for further details. ^d NaI (2 equiv) was added to the reaction mixture. brsm: based on remaining starting material.

when direct UV-A light irradiation (Kessil lamp, $\lambda = 390$ nm, full intensity) was used, product **3** was formed in a 75% assay yield, without any photocatalyst added.

Using the optimized set of conditions (Table 1, entry 6), we next evaluated the scope of the visible-light induced hydro-alkylation protocol (Figure 2).

Hereto, dimethyl maleate was combined with several alkyl iodides, and we found that the expected products were obtained in all cases (4-23). Notably, the acid-sensitive acetal function is well tolerated under our optimized conditions (8, 80%). Also iodides of medicinally relevant nitrogen-containing scaffolds, including Boc-protected azetidine, pyrrolidine, and piperidine, were competent reaction partners, allowing isolation of the corresponding adducts in good yields (9-12, 57-73%). In a similar vein, oxygen- and sulfur-containing alkyl iodides could be engaged in the reaction protocol (13-15, 40-77%). Next, we employed 1-iodoadamantane as a model for tertiary alkyl iodides, and we found that the hydroalkylated product was obtained in very good yield (16, 79%). The presence of a free carboxylic acid slightly reduced the reaction efficiency; however, the targeted compound could still be accessed in synthetically useful quantities (17, 56%).

Finally, we focused on primary alkyl iodides, which are interesting yet more challenging to engage in the reaction protocol. A slight modification of the reaction conditions (see GP4 in the Supporting Information, including an inverted organic halide/olefin ratio and extended light exposure) resulted in complete conversion and yielded the compounds **18–23** in satisfactory yields (40–57%). Of note are the unprotected aliphatic alcohols (**20**) and easily oxidizable phenols (**23**).

With respect to the SOMOphile scope, we found that different olefins could successfully take part in the transformation (3, 24-30). The product of our benchmark reaction (3) was obtained in 73% yield. Notably, when using phenyl vinyl sulfone (2c), we were able to trap stabilized benzyl radicals, while the reaction with dimethyl maleate did not afford the expected product. Also, norbornenone, N-phenyl acrylamide, methyl acrylate, and diethyl vinyl phosphonate could be engaged as SOMOphiles in the reaction protocol, showing its tolerance toward a wide variety of functional groups, such as ketones, amides, esters, and phosphonates (27-30, 40-59% yield). The applicability of our visible-light photocatalytic hydroalkylation process was also demonstrated by the fact that several derivatives of biologically active molecules could be readily modified; these include densely functionalized compounds, such as derivatives of sugars (i.e., α -D-glucofuranose and α -L-sorbofuranose), caffeine, cyclouridine, and even a dipeptide (31-36, 43-76%).

While alkyl bromides were not reactive under our original reaction conditions, we found that simple addition of NaI (2 equiv) could obviate this issue by generating the corresponding alkyl iodide *in situ* via $S_N 2$. With this operationally facile approach, we could subject various primary alkyl bromides to the hydroalkylation strategy and obtain the targeted compounds in decent yields (37–40, 44–50% yield).

Finally, we also successfully translated our batch protocol to a fast and scalable continuous-flow process, which should enable a fast transition between medicinal and process chemistry (see GP6 described in the Supporting Information).²¹ In flow, we were able to prepare compounds **3**, **35**, and **36** in good yields requiring only 30 min of light exposure. We also exploited the continuous-flow technology to scale our benchmark reaction up to 5 mmol (78% yield of the isolated product, Figure 3), corresponding to a productivity of 21 g d⁻¹ of compound **3**.

To gain insight into the reaction mechanism, we performed a series of experimental and computational studies. In particular, we identified two crucial steps to be investigated, i.e. (i) the generation of the ligated boryl radical²² and (ii) the occurrence of a radical chain process. To reveal the presence of the ligated boryl radical, we recorded an EPR spectrum of a deoxygenated benzene solution of **PC1** (0.05 M) and **B1** (0.05 M) containing phenyl *N-tert*-butylnitrone (PBN, 0.0125 M) as a spin trap. Prior to irradiation no signal was observed; however, after continuous irradiation for 15 min ($\lambda = 460$ nm)



pubs.acs.org/JACS

Figure 3. Translation to continuous-flow conditions (see GP6 in the Supporting Information) and scale-up.

we clearly saw the appearance of two distinct features: one can be attributed to the reduced photocatalyst (acridine radical), while the second feature is derived from the trapping of the ligated boryl radical with PBN (Figure 4a; for further details, see Section 6.1 in the Supporting Information).^{15a} The assignment is further supported by the observation of the adduct of the LBR with PBN (PBN*NHCBH2) in HRMS (Figure S5). These experiments unequivocally show the formation of the desired LBR under photoredox conditions. Direct observation of the ligated boryl radical via EPR was unsuccessful due to its short lifetime.²⁴ As further evidence, we also tracked the interaction between PC1 and B1 via UV-vis spectroscopy. In particular, when a deoxygenated CH₃CN solution (10^{-5} M) of PC1 and B1 was irradiated at 456 nm, we observed that the former was directly converted into a species absorbing in the 450-700 nm range (two isosbestic points were discerned; see Figure 4b). We propose this species to be the acridine radical generated upon single-electron reduction by B1, based on a comparison with the literature for a similar compound.25

In order to get some insights into the radical chain mechanism, we conducted experiments with deuterium-labeled substrates, and the results are collected in Figure 4c. The deuterium incorporation was calculated via ¹H NMR on purified products. Overall, these experiments showed that deuterium incorporation (product $3 \cdot d_1$) was only observed when deuterated borane **B1**- d_3 was exploited as an XAT agent, while the use of **B1** resulted in the formation of product 3 exclusively.

These results reveal that the hydroalkylated product is obtained upon HAT from another molecule of **B1** rather than the solvent, thus pointing toward a radical chain mechanism. We next wondered if the latter could be the rate-determining step of the transformation; therefore, we set off to evaluate the kinetic isotope effect (KIE) of the reaction (Figure 4d). First, we measured the KIE through a competition experiment. Hereto, we performed our benchmark reaction in the presence of an equimolar mixture of **B1** and **B1**-*d*₃ (5 equiv each) and a KIE value of 3.5 was found. However, when we calculated this value with the parallel reactions method by performing independent reactions under optimized conditions, one containing **B1** and one containing **B1**-*d*₃, we found a KIE of



Figure 4. (A) Electron paramagnetic resonance (EPR) spectrum (black) obtained upon irradiation ($\lambda = 460$ nm) of a deoxygenated 0.0125 M benzene solution of phenyl *N-tert*-butylnitrone containing PC1 (0.05 M) and B1 (0.05 M). Simulated profiles for acridine radical generated from PC1 upon SET (red) and of the PBN*NHCBH₂• (blue) are also shown (see Section 6.1 in the Supporting Information for experimental and simulation parameters). (B) UV–vis spectra of a deoxygenated CH₃CN solution of PC1 and B1 (both 10⁻⁵ M) irradiated over 4.25 min. (C) Deuteration experiments. (D) Determination of the kinetic isotope effect (KIE). (E) Computational investigation.

only 1.09. Taken together, these experiments suggest that the HAT step is not involved in the rate-determining step of the reaction.²⁶ Finally, we determined a quantum yield of 2%. Such a modest value is in accordance with a process being supported by either short-lived radical chain propagations, an inefficient initiation process²⁷ or decomposition of the photocatalyst

(further mechanistic insights are reported in Section 6 of the Supporting Information).²⁸

We also performed a computational investigation intended to model the entire reaction profile through the simulation of all the key steps, including some possible parasitic pathways. Thus, we adopted DFT at the ω B97xD/def2TZVP level of theory to optimize the relevant stationary points, also including



Figure 5. Proposed mechanism for the photoinduced XAT by N-heterocarbene boryl radicals for C–C bond formation. Redox potentials are reported vs SCE.

the effect of the solvent through an implicit model (Figure 4e; see also Section 10 of the Supporting Information for further details). We started by considering LBR I reacting with iodocyclohexane 1a through TS1 ($\Delta G^{\ddagger} = +11.6 \text{ kcal} \cdot \text{mol}^{-1}$) to afford cyclohexyl radical II. This nucleophilic radical (II) adds subsequently onto dimethyl maleate 2b through TS2 ($\Delta G^{\ddagger} = +15.5 \text{ kcal} \cdot \text{mol}^{-1}$) to deliver radical adduct III. Next, the targeted hydroalkylated product 5 is formed through reaction of III with NHC-ligated borane B1 via TS3 ($\Delta G^{\ddagger} = +12.4 \text{ kcal} \cdot \text{mol}^{-1}$). Notably, all these steps are exergonic in nature, with ΔG values in the $-11.9 \text{ to } -17.4 \text{ kcal} \cdot \text{mol}^{-1}$ range.

In this intricate ballet of fleeting radical intermediates, we realized that a careful balance between the XAT, the radical addition, and the final HAT steps was crucial to avoid parasitic reaction pathways.^{11a,b,15b} Accordingly, we also evaluated the possibility for these intermediates to undergo competitive, yet nonproductive pathways, including the direct addition of LBR I onto dimethyl maleate **2b** with formation of a new B–C bond (through TS4) and the reduction of the cyclohexyl radical II to cyclohexane (through TS5). However, our computational analysis revealed that both steps occur with higher activation energies and less negative energy gains compared to those describing the desired process. Intrigued by the lack of reactivity of organic bromides in our reaction, we also computed ΔG and ΔG^{\ddagger} for the XAT step by LBR I for bromocyclohexane.

By comparing the results with those obtained for the iodo analogue 1a, it seems that the difference in reactivity can be mainly attributed to kinetic factors ($\Delta G^{\ddagger}_{\text{R-I}} = +11.6 \text{ kcal} \cdot \text{mol}^{-1}$) vs $\Delta G^{\ddagger}_{\text{R-Br}} = +17.9 \text{ kcal} \cdot \text{mol}^{-1}$), as the process shows similar driving forces for both halides ($\Delta G_{\text{R-I}} = -13.0 \text{ kcal} \cdot \text{mol}^{-1}$ vs $\Delta G_{\text{R-Br}} = -13.8 \text{ kcal} \cdot \text{mol}^{-1}$).

Finally, we were interested in comparing quantitatively LBR I with other commonly used halogen abstractors, including α -aminoalkyl and (tris(trimethylsilyl)silyl radicals and a conventional tin-based XAT reagent (Me₃Sn[•]). As depicted in Figure 4e, a clear trend emerges. On the one hand, the XAT step shows very low activation energies in the case of metalloidyl radicals (i.e., R₃Si[•] and R₃Sn[•]; for the tin-based abstractor it is barrierless), while LBR I and the α -aminoalkyl radical display significant ΔG^{\ddagger} values (+11.6 and 15.0 kcal·mol⁻¹, respectively). On the other hand, from a thermodynamic point of

view, the XAT process is highly exergonic for metalloidyl radicals, moderately exergonic for LBR I, and essentially thermoneutral for the α -amino radical. In the latter case, the formation of an iminium ion resulting from the elimination of iodide was invoked as the driving force for the whole process.^{19a}

With both experimental and computational insights considered together, a mechanistic scenario is proposed in Figure 5. PC1 absorbs light resulting in formation of the corresponding highly oxidizing excited state $(E(PC^*/PC_{red}) =$ +2.06 V vs SCE).²⁹ This excited state can react with B1 $(E_{va}(B1^{\bullet+}/B1) = +0.89 \text{ V vs} (SCE) \text{ via single-electron transfer}$ to afford LBR I upon deprotonation. The latter intermediate is entrusted with the desired XAT step, which is expected to be relatively fast $(k \approx 10^8 \text{ M}^{-1} \text{ s}^{-1})$,³⁰ thus yielding the alkyl radical II. This radical can be subsequently trapped by the electron-poor olefin to give adduct III. The observed inhibition effect of O_2 (Table 1, entries 10-11) can be explained by taking into account that LBR I is known to react even faster with molecular oxygen $(k > 10^8 \text{ M}^{-1} \text{ s}^{-1})$,³⁰ thus confiscating this crucial intermediate for the XAT event. Alternatively, it has been reported that molecular oxygen can also quench the excited state of PC1 at diffusion controlled rates to form singlet oxygen ($k = 2 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$).³¹ Next, as shown by the deuterium labeling experiments, III abstracts a hydrogen atom from B1 in a polarity-matched step to give product 3 and subsequently kicks off the chain propagation.

CONCLUSIONS

In conclusion, we have shown that *N*-heterocyclic carbene (NHC) borane **B1** is an efficient XAT agent to sustain the radical hydroalkylation of olefins. This is a significant advancement to previous reports where NHC-ligated boryl radicals were mainly used as radical chain carriers for radical reductions. Our method shows remarkable generality, robustness, and versatility, as it does not rely on any interaction between the ligated borane and the organic halide to generate nucleophilic alkyl radicals under visible light irradiation. Due to the mild reaction conditions, it is applicable to a vast array of substrates, including biologically active compounds. And

finally, continuous-flow technology can be exploited to accelerate and scale our methodology.

A detailed experimental and spectroscopic mechanistic investigation describes the key role of the NHC-based boryl radicals in the operative reaction mechanism. This is further corroborated by computational analysis, indicating that the described process is the most favorable one with respect to possible competing pathways.

While NHC-ligated boryl radicals have been only recently exploited in radical chemistry, this investigation represents an important step toward the appreciation and the exploitation of the properties and the reactivity of these boryl species. Hence, we are confident that this work will stimulate further research into the use of LBRs for radical-based synthetic transformations.

ASSOCIATED CONTENT

1 Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.2c10444.

Experimental procedures, characterization data of synthesized compounds, copies of NMR spectra, complete mechanistic investigation. The primary NMR FID files for starting materials, compounds **3–40**, as well as optimized geometries of species displayed in Figure 4e, IRC and PES profiles are available in the FigShare repository at https://www.doi.org/10.21942/uva. 20459517. (PDF)

AUTHOR INFORMATION

Corresponding Author

Timothy Noël – Flow Chemistry Group, van 't Hoff Institute for Molecular Sciences (HIMS), University of Amsterdam, 1098 XH Amsterdam, The Netherlands; orcid.org/0000-0002-3107-6927; Email: t.noel@uva.nl

Authors

- Ting Wan Flow Chemistry Group, van 't Hoff Institute for Molecular Sciences (HIMS), University of Amsterdam, 1098 XH Amsterdam, The Netherlands
- Luca Capaldo Flow Chemistry Group, van 't Hoff Institute for Molecular Sciences (HIMS), University of Amsterdam, 1098 XH Amsterdam, The Netherlands; orcid.org/0000-0001-7114-267X
- Davide Ravelli PhotoGreen Lab, Department of Chemistry, University of Pavia, 27100 Pavia, Italy; @ orcid.org/0000-0003-2201-4828
- Walter Vitullo Flow Chemistry Group, van 't Hoff Institute for Molecular Sciences (HIMS), University of Amsterdam, 1098 XH Amsterdam, The Netherlands
- Felix J. de Zwart Homogeneous, Supramolecular and Bioinspired Catalysis Group (HomKat), van't Hoff Institute for Molecular Sciences (HIMS), Universiteit van Amsterdam (UvA), 1098 XH Amsterdam, The Netherlands;
 orcid.org/0000-0002-0981-1120
- Bas de Bruin Homogeneous, Supramolecular and Bioinspired Catalysis Group (HomKat), van't Hoff Institute for Molecular Sciences (HIMS), Universiteit van Amsterdam (UvA), 1098 XH Amsterdam, The Netherlands;
 ◎ orcid.org/0000-0002-3482-7669

Complete contact information is available at: https://pubs.acs.org/10.1021/jacs.2c10444

^{II}T.W. and L.C. contributed equally to this work.

Fundina

Author Contributions

China Scholarship Council (CSC) and European Union's Horizon 2020 research and innovation program

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

T.W. has received support from the China Scholarship Council (CSC) for her PhD studies. L.C. acknowledges European Union's Horizon 2020 research and innovation programme under the Marie Skłodowska-Curie Grant Agreement No. 101023615 (project name: HAT-TRICK). Calculations were carried out at the CINECA Supercomputer Center (Italy), with computer time granted by ISCRA projects (codes: HP10CIH7D5 and HP10CXZVO3).

REFERENCES

(1) (a) Stephenson, C.; Yoon, T.; MacMillan, D. W. C. Visible Light Photocatalysis in Organic Chemistry; Wiley-VCH Verlag GmbH & Co. KGaA: Weinheim, Germany, 2018. DOI: 10.1002/9783527674145. (b) Capaldo, L.; Ravelli, D.; Fagnoni, M. Direct Photocatalyzed Hydrogen Atom Transfer (HAT) for Aliphatic C-H Bonds Elaboration. Chem. Rev. 2022, 122 (2), 1875-1924. (c) Chan, A. Y.; Perry, I. B.; Bissonnette, N. B.; Buksh, B. F.; Edwards, G. A.; Frye, L. I.; Garry, O. L.; Lavagnino, M. N.; Li, B. X.; Liang, Y.; et al. Metallaphotoredox: The Merger of Photoredox and Transition Metal Catalysis. Chem. Rev. 2022, 122 (2), 1485-1542. (d) Holmberg-Douglas, N.; Nicewicz, D. A. Photoredox-Catalyzed C-H Functionalization Reactions. Chem. Rev. 2022, 122 (2), 1925-2016. (e) Murray, P. R. D.; Cox, J. H.; Chiappini, N. D.; Roos, C. B.; McLoughlin, E. A.; Hejna, B. G.; Nguyen, S. T.; Ripberger, H. H.; Ganley, J. M.; Tsui, E.; et al. Photochemical and Electrochemical Applications of Proton-Coupled Electron Transfer in Organic Synthesis. Chem. Rev. 2022, 122 (2), 2017-2291. (f) Witzel, S.; Hashmi, A. S. K.; Xie, J. Light in Gold Catalysis. Chem. Rev. 2021, 121 (14), 8868-8925.

(2) Huang, C.-Y.; Li, J.; Li, C.-J. Photocatalytic $C(sp^3)$ Radical Generation via C–H, C–C, and C–X Bond Cleavage. *Chem. Sci.* **2022**, 13 (19), 5465–5504.

(3) Kärkäs, M. D. Photochemical Generation of Nitrogen-Centered Amidyl, Hydrazonyl, and Imidyl Radicals: Methodology Developments and Catalytic Applications. *ACS Catal.* **2017**, 7 (8), 4999–5022.

(4) Chang, L.; An, Q.; Duan, L.; Feng, K.; Zuo, Z. Alkoxy Radicals See the Light: New Paradigms of Photochemical Synthesis. *Chem. Rev.* **2022**, *122* (2), 2429–2486.

(5) Bonciolini, S.; Noël, T.; Capaldo, L. Synthetic Applications of Photocatalyzed Halogen-Radical Mediated Hydrogen Atom Transfer for C-H Bond Functionalization. *Eur. J. Org. Chem.* **2022**, 2022, No. e202200417.

(6) Noël, T.; Zysman-Colman, E. The Promise and Pitfalls of Photocatalysis for Organic Synthesis. *Chem. Catal.* **2022**, *2* (3), 468–476.

(7) (a) Su, Y.; Kinjo, R. Boron-Containing Radical Species. *Coord. Chem. Rev.* 2017, 352, 346–378. (b) Renaud, P. Boron in Radical Chemistry. In *Encyclopedia of Radicals in Chemistry, Biology and Materials*; John Wiley & Sons, Ltd.: Chichester, U.K., 2012. DOI: 10.1002/9781119953678.rad020.

(8) Capaldo, L.; Noël, T.; Ravelli, D. Photocatalytic Generation of Ligated Boryl Radicals from Tertiary Amine-Borane Complexes: An Emerging Tool in Organic Synthesis. *Chem. Catal.* **2022**, *2* (5), 957–966.

(9) For representative works on ligated boryl radicals (LBRs), see: (a) Baban, J. A.; Roberts, B. P. An Electron Spin Resonance Study of Phosphine-Boryl Radicals; Their Structures and Reactions with Alkyl Halides. J. Chem. Soc. Perk. T. 1984, 10, 1717. (b) Baban, J. A.; Roberts, B. P. An e.s.r. Study of Amine-Boryl Radicals (R₃N-BH₂•) in Solution. J. Chem. Soc., Chem. Commun. 1983, 21, 1224-1226. (c) Baban, J. A.; Marti, V. P. J.; Roberts, B. P. Spin-Trapping of Ligated Boryl Radicals. Tetrahedron Lett. 1985, 26 (10), 1349-1352. (d) Baban, J. A.; Marti, V. P. J.; Roberts, B. P. Ligated Boryl Radicals. Part 2. Electron Spin Resonance Studies of Trialkylamine-Boryl Radicals. J. Chem. Soc. Perk. T. 2 1985, 11, 1723-1733. (e) Lu, D.; Wu, C.; Li, P. Synergistic Effects of Lewis Bases and Substituents on the Electronic Structure and Reactivity of Boryl Radicals. Chem.-Eur. J. 2014, 20 (6), 1630-1637. (f) Paul, V.; Roberts, B. P. Polarity Reversal Catalysis of Hydrogen Atom Abstraction Reactions. J. Chem. Soc., Chem. Commun. 1987, 17, 1322-1324. (g) Paul, V.; Roberts, B. P. Homolytic Reactions of Ligated Boranes. Part 10. Electron Spin Resonance Studies of Radicals Derived from Ligated Arylboranes. J. Chem. Soc., Perk. T. 2 1988, 10, 1895-1901. (h) Paul, V.; Roberts, B. P. Homolytic Reactions of Ligated Boranes. Part 8. Electron Spin Resonance Studies of Radicals Derived from Ligated Alkylboranes. J. Chem. Soc., Perk. T. 2 1988, 7, 1183-1193. (i) Baban, J. A.; Roberts, B. P. Homolytic Reactions of Ligated Boranes. Part 9. Overall Addition of Alkanes to Electron-Deficient Alkenes by a Radical Chain Mechanism. J. Chem. Soc., Perk. T. 2 1988, 7, 1195-1200.

(10) For recent synthetic application of LBRs, see: (a) Zhang, Z.-Q.; Sang, Y.-Q.; Wang, C.-Q.; Dai, P.; Xue, X.-S.; Piper, J. L.; Peng, Z.-H.; Ma, J.-A.; Zhang, F.-G.; Wu, J. Difluoromethylation of Unactivated Alkenes Using Freon-22 through Tertiary Amine-Borane-Triggered Halogen Atom Transfer. J. Am. Chem. Soc. 2022, 144 (31), 14288– 14296. (b) Lei, G.; Xu, M.; Chang, R.; Funes-Ardoiz, I.; Ye, J. Hydroalkylation of Unactivated Olefins via Visible-Light-Driven Dual Hydrogen Atom Transfer Catalysis. J. Am. Chem. Soc. 2021, 143 (29), 11251–11261. (c) Kim, J. H.; Constantin, T.; Simonetti, M.; Llaveria, J.; Sheikh, N. S.; Leonori, D. A Radical Approach for the Selective C– H Borylation of Azines. Nature 2021, 595 (7869), 677–683.

(11) (a) Ueng, S.-H.; Solovyev, A.; Yuan, X.; Geib, S. J.; Fensterbank, L.; Lacôte, E.; Malacria, M.; Newcomb, M.; Walton, J. C.; Curran, D. P. N -Heterocyclic Carbene Boryl Radicals: A New Class of Boron-Centered Radical. J. Am. Chem. Soc. 2009, 131 (31), 11256–11262. (b) Ueng, S.-H.; Makhlouf Brahmi, M.; Derat, E.; Fensterbank, L.; Lacôte, E.; Malacria, M.; Curran, D. P. Complexes of Borane and N-Heterocyclic Carbenes: A New Class of Radical Hydrogen Atom Donor. J. Am. Chem. Soc. 2008, 130 (31), 10082– 10083. (c) Curran, D. P.; Solovyev, A.; Makhlouf Brahmi, M.; Fensterbank, L.; Malacria, M.; Lacôte, E. Synthesis and Reactions of N-Heterocyclic Carbene Boranes. Angew. Chem., Int. Ed. 2011, 50 (44), 10294–10317.

(12) Taniguchi, T. Advances in Chemistry of N-Heterocyclic Carbene Boryl Radicals. *Chem. Soc. Rev.* 2021, 50 (16), 8995–9021. (13) For reviews about radical addition of LBRs onto unsaturated systems, see: (a) Lai, D.; Ghosh, S.; Hajra, A. Light-Induced Borylation: Developments and Mechanistic Insights. *Org. Biomol. Chem.* 2021, 19 (20), 4397–4428. (b) Taniguchi, T. Boryl Radical Addition to Multiple Bonds in Organic Synthesis. *Eur. J. Org. Chem.* 2019, 2019 (37), 6308–6319.

(14) For selected works about radical addition of boryl radicals onto unsaturated systems, see: (a) Zhu, C.; Dong, J.; Liu, X.; Gao, L.; Zhao, Y.; Xie, J.; Li, S.; Zhu, C. Photoredox-Controlled β -Regioselective Radical Hydroboration of Activated Alkenes with NHC-Boranes. Angew. Chem., Int. Ed. 2020, 59 (31), 12817-12821. (b) Li, G.; Huang, G.; Sun, R.; Curran, D. P.; Dai, W. Regioselective Radical Borylation of $\alpha_{,\beta}$ -Unsaturated Esters and Related Compounds by Visible Light Irradiation with an Organic Photocatalyst. Org. Lett. 2021, 23 (11), 4353-4357. (c) Qi, J.; Zhang, F.; Jin, J.; Zhao, Q.; Li, B.; Liu, L.; Wang, Y. New Radical Borylation Pathways for Organoboron Synthesis Enabled by Photoredox Catalysis. Angew. Chem., Int. Ed. 2020, 59 (31), 12876-12884. (d) Xia, P. J.; Song, D.; Ye, Z. P.; Hu, Y. Z.; Xiao, J. A.; Xiang, H. Y.; Chen, X. Q.; Yang, H. Photoinduced Single-Electron Transfer as an Enabling Principle in the Radical Borylation of Alkenes with NHC-Borane. Angew. Chem., Int. Ed. 2020, 59 (17), 6706-6710. (e) Xu, W.; Jiang, H.; Leng, J.; Ong,

H. W.; Wu, J. Visible-Light-Induced Selective Defluoroborylation of Polyfluoroarenes, Gem-Difluoroalkenes, and Trifluoromethylalkenes. *Angew. Chem., Int. Ed.* **2020**, *59* (10), 4009–4016. (f) Zhou, N.; Yuan, X. A.; Zhao, Y.; Xie, J.; Zhu, C. Synergistic Photoredox Catalysis and Organocatalysis for Inverse Hydroboration of Imines. *Angew. Chem., Int. Ed.* **2018**, *57* (15), 3990–3994. (g) Shi, Q.; Xu, M.; Chang, R.; Ramanathan, D.; Peñin, B.; Funes-Ardoiz, I.; Ye, J. Visible-Light Mediated Catalytic Asymmetric Radical Deuteration at Non-Benzylic Positions. *Nat. Commun.* **2022**, *13* (1), 4453.

(15) (a) Pan, X.; Lacôte, E.; Lalevée, J.; Curran, D. P. Polarity Reversal Catalysis in Radical Reductions of Halides by N-Heterocyclic Carbene Boranes. J. Am. Chem. Soc. 2012, 134 (12), 5669-5674.
(b) Ueng, S.-H.; Fensterbank, L.; Lacôte, E.; Malacria, M.; Curran, D. P. Radical Reductions of Alkyl Halides Bearing Electron Withdrawing Groups with N-Heterocyclic Carbene Boranes. Org. Biomol. Chem. 2011, 9 (9), 3415-3420.

(16) Supranovich, V. I.; Levin, V. v.; Struchkova, M. I.; Korlyukov, A. A.; Dilman, A. D. Radical Silyldifluoromethylation of Electron-Deficient Alkenes. *Org. Lett.* **2017**, *19* (12), 3215–3218.

(17) Juliá, F.; Constantin, T.; Leonori, D. Applications of Halogen-Atom Transfer (XAT) for the Generation of Carbon Radicals in Synthetic Photochemistry and Photocatalysis. *Chem. Rev.* **2022**, *122* (2), 2292–2352.

(18) (a) Le, C.; Chen, T. Q.; Liang, T.; Zhang, P.; MacMillan, D. W. C. A Radical Approach to the Copper Oxidative Addition Problem: Trifluoromethylation of Bromoarenes. Science 2018, 360 (6392), 1010-1014. (b) Lovett, G. H.; Chen, S.; Xue, X.-S.; Houk, K. N.; MacMillan, D. W. C. Open-Shell Fluorination of Alkyl Bromides: Unexpected Selectivity in a Silyl Radical-Mediated Chain Process. J. Am. Chem. Soc. 2019, 141 (51), 20031-20036. (c) Smith, R. T.; Zhang, X.; Rincón, J. A.; Agejas, J.; Mateos, C.; Barberis, M.; García-Cerrada, S.; de Frutos, O.; MacMillan, D. W. C. Metallaphotoredox-Catalyzed Cross-Electrophile $C(sp^3)-(Csp^3)$ Coupling of Aliphatic Bromides. J. Am. Chem. Soc. 2018, 140 (50), 17433-17438. (d) Brill, Z. G.; Ritts, C. B.; Mansoor, U. F.; Sciammetta, N. Continuous Flow Enables Metallaphotoredox Catalysis in a Medicinal Chemistry Setting: Accelerated Optimization and Library Execution of a Reductive Coupling between Benzylic Chlorides and Aryl Bromides. Org. Lett. 2020, 22 (2), 410-416. (e) Devery, J. J.; Nguyen, J. D.; Dai, C.; Stephenson, C. R. J. Light-Mediated Reductive Debromination of Unactivated Alkyl and Aryl Bromides. ACS Catal. 2016, 6 (9), 5962-5967. (f) Zhang, P.; Le, C.; MacMillan, D. W. C. Silyl Radical Activation of Alkyl Halides in Metallaphotoredox Catalysis: A Unique Pathway for Cross-Electrophile Coupling. J. Am. Chem. Soc. 2016, 138 (26), 8084-8087. (g) ElMarrouni, A.; Ritts, C. B.; Balsells, J. Silyl-Mediated Photoredox-Catalyzed Giese Reaction: Addition of Non-Activated Alkyl Bromides. Chem. Sci. 2018, 9 (32), 6639-6646. (h) Dong, J.; Lyu, X.; Wang, Z.; Wang, X.; Song, H.; Liu, Y.; Wang, Q. Visible-Light-Mediated Minisci C-H Alkylation of Heteroarenes with Unactivated Alkyl Halides Using O2 as an Oxidant. Chem. Sci. 2019, 10 (4), 976-982. (i) Sakai, H. A.; Liu, W.; Le, C.; MacMillan, D. W. C. Cross-Electrophile Coupling of Unactivated Alkyl Chlorides. J. Am. Chem. Soc. 2020, 142 (27), 11691-11697. (j) Luridiana, A.; Mazzarella, D.; Capaldo, L.; Rincón, J. A.; García-Losada, P.; Mateos, C.; Frederick, M. O.; Nuño, M.; Jan Buma, W.; Noël, T. The Merger of Benzophenone HAT Photocatalysis and Silyl Radical-Induced XAT Enables Both Nickel-Catalyzed Cross-Electrophile Coupling and 1,2-Dicarbofunctionalization of Olefins. ACS Catal. 2022, 12 (18), 11216-11225.

(19) (a) Constantin, T.; Zanini, M.; Regni, A.; Sheikh, N. S.; Juliá, F.; Leonori, D. Aminoalkyl Radicals as Halogen-Atom Transfer Agents for Activation of Alkyl and Aryl Halides. *Science* **2020**, 367 (6481), 1021–1026. (b) Constantin, T.; Juliá, F.; Sheikh, N. S.; Leonori, D. A Case of Chain Propagation: α -Aminoalkyl Radicals as Initiators for Aryl Radical Chemistry. *Chem. Sci.* **2020**, 11 (47), 12822–12828. (c) Zhao, H.; McMillan, A. J.; Constantin, T.; Mykura, R. C.; Juliá, F.; Leonori, D. Merging Halogen-Atom Transfer (XAT) and Cobalt Catalysis to Override E2-Selectivity in the Elimination of Alkyl Halides: A Mild Route toward *Contra* -Thermodynamic Olefins. J. Am. Chem. Soc. 2021, 143 (36), 14806–14813. (d) Caiger, L.; Sinton, C.; Constantin, T.; Douglas, J. J.; Sheikh, N. S.; Juliá, F.; Leonori, D. Radical Hydroxymethylation of Alkyl Iodides Using Formaldehyde as a C1 Synthon. Chem. Sci. 2021, 12 (31), 10448– 10454.

(20) Qi, J.; Zhang, F.; Jin, J.; Zhao, Q.; Li, B.; Liu, L.; Wang, Y. New Radical Borylation Pathways for Organoboron Synthesis Enabled by Photoredox Catalysis. *Angew. Chem., Int. Ed.* **2020**, *59* (31), 12876–12884.

(21) (a) Wan, T.; Wen, Z.; Laudadio, G.; Capaldo, L.; Lammers, R.; Rincón, J. A.; García-Losada, P.; Mateos, C.; Frederick, M. O.; Broersma, R.; Noël, T. Accelerated and Scalable C(sp³)-H Amination via Decatungstate Photocatalysis Using a Flow Photoreactor Equipped with High-Intensity LEDs. ACS Centr. Sci. 2022, 8 (1), 51-56. (b) Buglioni, L.; Raymenants, F.; Slattery, A.; Zondag, S. D. A.; Noël, T. Technological Innovations in Photochemistry for Organic Synthesis: Flow Chemistry, High-Throughput Experimentation, Scale-up, and Photoelectrochemistry. Chem. Rev. 2022, 122 (2), 2752-2906. (c) Mazzarella, D.; Pulcinella, A.; Bovy, L.; Broersma, R.; Noël, T. Rapid and Direct Photocatalytic C(sp³)-H Acylation and Arylation in Flow. Angew. Chem. In. Ed. 2021, 60 (39), 21277-21282. (d) Sambiagio, C.; Noël, T. Flow Photochemistry: Shine Some Light on Those Tubes! Trends Chem. 2020, 2 (2), 92-106. (e) Rehm, T. H. Flow Photochemistry as a Tool in Organic Synthesis. Chem.-Eur. J. 2020, 26 (71), 16952-16974.

(22) Zhang, L.; Si, X.; Rominger, F.; Hashmi, A. S. K. Visible-Light-Induced Radical Carbo-Cyclization/gem-Diborylation through Triplet Energy Transfer between a Gold Catalyst and Aryl Iodides. *J. Am. Chem. Soc.* **2020**, *142* (23), 10485–10493.

(23) Tsudaka, T.; Kotani, H.; Ohkubo, K.; Nakagawa, T.; Tkachenko, N. v.; Lemmetyinen, H.; Fukuzumi, S. Photoinduced Electron Transfer in 9-Substituted 10-Methylacridinium Ions. *Chem.—Eur. J.* **2017**, *23* (6), 1306–1317.

(24) Walton, J. C.; Brahmi, M. M.; Fensterbank, L.; Lacôte, E.; Malacria, M.; Chu, Q.; Ueng, S.-H.; Solovyev, A.; Curran, D. P. EPR Studies of the Generation, Structure, and Reactivity of N-Heterocyclic Carbene Borane Radicals. *J. Am. Chem. Soc.* **2010**, *132* (7), 2350– 2358.

(25) MacKenzie, I. A.; Wang, L.; Onuska, N. P. R.; Williams, O. F.; Begam, K.; Moran, A. M.; Dunietz, B. D.; Nicewicz, D. A. Discovery and Characterization of an Acridine Radical Photoreductant. *Nature* **2020**, 580 (7801), 76–80.

(26) Simmons, E. M.; Hartwig, J. F. On the Interpretation of Deuterium Kinetic Isotope Effects in C–H Bond Functionalizations by Transition-Metal Complexes. *Angew. Chem., Int. Ed.* **2012**, *51* (13), 3066–3072.

(27) Pitre, S. P.; McTiernan, C. D.; Vine, W.; DiPucchio, R.; Grenier, M.; Scaiano, J. C. Visible-Light Actinometry and Intermittent Illumination as Convenient Tools to Study $Ru(bpy)_3Cl_2$ Mediated Photoredox Transformations. *Sci. Rep.-UK* **2015**, *5* (1), 16397.

(28) Tlili, A.; Lakhdar, S. Acridinium Salts and Cyanoarenes as Powerful Photocatalysts: Opportunities in Organic Synthesis. *Angew. Chem., Int. Ed.* **2021**, *60* (36), 19526–19549.

(29) Joshi-Pangu, A.; Lévesque, F.; Roth, H. G.; Oliver, S. V.; Campeau, L.-C.; Nicewicz, N.; DiRocco, D. A. Acridinium-Based Photocatalysts: A Sustainable Option in Photoredox Catalysis. *J. Org. Chem.* **2016**, *81* (16), 7244–7249.

(30) Pan, X.; Vallet, A.-L.; Schweizer, S.; Dahbi, K.; Delpech, B.; Blanchard, N.; Graff, B.; Geib, S. J.; Curran, D. P.; Lalevée, J.; Lacôte, E. Mechanistic and Preparative Studies of Radical Chain Homolytic Substitution Reactions of *N*-Heterocyclic Carbene Boranes and Disulfides. *J. Am. Chem. Soc.* **2013**, *135* (28), 10484–10491.

(31) Benniston, A. C.; Harriman, A.; Li, P.; Rostron, J. P.; van Ramesdonk, H. J.; Groeneveld, M. M.; Zhang, H.; Verhoeven, J. W. Charge Shift and Triplet State Formation in the 9-Mesityl-10-Methylacridinium Cation. *J. Am. Chem. Soc.* **2005**, *127* (46), 16054–16064.

(32) When monitoring the reaction over time, we detected *N*,*N*-dimethylimidazolium iodide (via ¹H NMR) and boric acid (via ¹¹B-

NMR). These byproducts are formed upon decomposition of NHC-BH₂I in an aqueous environment as reported in the literature (ref 15b, see also Section 6.8 in the Supporting Information). Notably, when the reaction was performed in dry C_6D_6 , the formation of NHC-BH₂I could be observed via ¹¹B-NMR spectroscopy (see Section 6.8 in the Supporting Information), which supports again the occurrence of the XAT step.

Recommended by ACS

Intermolecular Organophotocatalytic Cyclopropanation of Unactivated Olefins

David M. Fischer, Erick M. Carreira, et al.

JANUARY	06,	2023
IOURNAL	OF	THE AMERICAN CHEMICAL SOCIETY

Photochemical Organocatalytic Functionalization of Pyridines via Pyridinyl Radicals

Emilien Le Saux, Paolo Melchiorre, et al. DECEMBER 27, 2022

JOURNAL OF THE AMERICAN CHEMICAL SOCIETY

Asymmetric C-Alkylation of Nitroalkanes via Enzymatic Photoredox Catalysis

Haigen Fu, Todd K. Hyster, et al.	
JANUARY 06, 2023	
JOURNAL OF THE AMERICAN CHEMICAL SOCIETY	READ 🗹

Late-Stage C(*sp*³)-H Methylation of Drug Molecules

Edna Mao and David W.C. MacMillan

ANUARY 25, 2023	
OURNAL OF THE AMERICAN CHEMICAL SOCIETY	READ 🗹

Get More Suggestions >

READ 🗹

READ 🗹

J,