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Photoinduced boryl-oximation of alkenes via NHC-boryl radical generation

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Boryl-functionalization of C–C multiple bonds offers a powerful route to organoboron compounds. Traditional transition-metal-catalyzed methods face challenges such as toxic residues and poor functional group tolerance. Furthermore, the recently reported photoredox-catalyzed boryl-functionalizations remain dependent on tailored photocatalysts, including $Ir^{III}(bpy)_3$, $Ir^{III}(bpy)_2(dtbbpy)PF_6$ and 4-CzIPN. *N*-Heterocyclic carbene (NHC) boranes, as versatile precursors, enable C–B bond formation *via* photochemical boryl radical generation. Herein, we describe a metal-free radical chain transfer strategy for NHC-boryl radical generation through thiol-catalyzed hydrogen atom transfer, achieving boryl-oximation of alkenes with a broad substrate scope and high selectivity. The resulting α -boryloximes serve as versatile synthetic intermediates, enabling late-stage functionalization of pharmaceutical molecules and further product transformations. Mechanistic studies highlight the critical role of light irradiation in sustaining the radical process.

Introduction

Organoboron compounds have garnered wide application in the fields of agricultural chemistry, pharmaceutical chemistry and materials science, due to their good stability and versatile reactivity (e.g., Suzuki cross-coupling reactions and carbonheteroatom bond forming reactions). Consequently, the development of C-B bond-forming reactions has long been of interest in organic synthesis.2 Of the known synthetic methods for organoboron compounds, transition-metal-catalyzed borylfunctionalization of alkenes and alkynes, involving representative processes such as diboration, silaboration, borylstannation,⁵ cyanoboration,⁶ alkynylboration,⁷ and aminoboration,⁸ is of significant importance because it enables the simultaneous incorporation of a boron atom and other valuable functional groups into the organic molecules. In such reactions, boron reagents are usually employed to form metal-boron active intermediates, which then undergo organometallic elementary reactions to obtain the target product. Despite the above advances in this field, several limitations remain, including the troublesome handling of boron sources, metal residue in pharmaceuticals, excessive use of ligands/bases and poor functional group tolerance.3-8 Therefore, it is highly

Recently, N-heterocyclic carbene (NHC) boranes have received increasing attention owing to their air stability and ease of handling compared to traditional boron reagents. Notably, the relatively low dissociation energies of the B-H bonds in NHCboranes (BDE = $70-80 \text{ kcal mol}^{-1}$), compared to that in BH₃ (BDE = 111.7 kcal mol⁻¹), facilitate their homolysis under thermal or photoredox conditions, leading to the generation of boryl radicals.9 Therefore, a series of radical borylation reactions of C-C multiple bonds have been explored successively using NHC-boranes (Scheme 1a).10 To our knowledge, there are only a handful of examples of catalytic boryl-functionalization of alkenes and alkynes with NHC-boranes,11-14 most of which require the involvement of photocatalysts. Despite the advances in boryl-functionalization, side reactions such as hydroboration15 and radical dimerization of alkenes are often unavoidable.16 In 2020-2025, Wang12 and Xuan14 reported visible light-induced arylboration and borylacylation of alkenes using NHC-boranes with [Ir^{III}] as the photocatalyst (Scheme 1b). Mechanistic studies showed that a NHC-boryl radical and an arene radical anion (or a ketyl radical) are generated concurrently via a photoredox-catalyzed single-electron transfer (SET) process. Subsequently, the NHC-boryl radical undergoes radical addition to alkenes, followed by radical cross-coupling with the arene radical anion (or ketyl radical), affording products. Moreover, the Xie group¹³ in 2024 achieved a rare alkynylborylation reaction of alkenes with NHC-boranes and alkynyl bromides by using Ir(ppy)₂(dtbbpy)PF₆ (Scheme 1b). In this work, the authors propose that radical addition to carbon-carbon triple bonds plays a key role in the introduction of alkynyl groups. Although certain

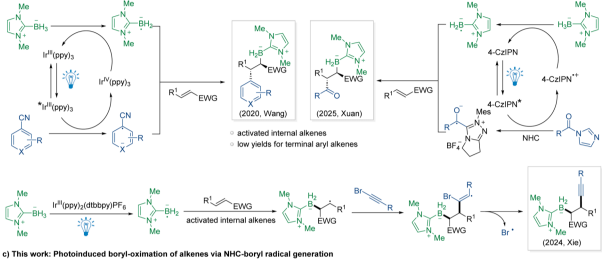
desirable to address these challenges encountered in the aforementioned boryl-functionalization reactions.

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a) Radical borylation reactions of C-C multiple bonds, using NHC-boranes

b) Photoredox-catalyzed radical boryl-functionalization of alkenes using NHC-boranes



Scheme 1 Strategies for the borylation/boryl-functionalization of C-C multiple bonds.

progress has been made in the boryl-functionalization of alkenes using NHC-boranes, the limited substrate scope and functional group type highlight the need for further investigation into borylfunctionalization reactions.

In recent years, significant progress has been made in the visible-light-induced radical functional-oximation of alkenes, particularly through the utilization of organonitroso compounds (*BuO-NO and R1R2N-NO) as efficient NO radical sources.17 Inspired by these breakthroughs, we proposed a photoinduced cascade radical chain transfer strategy involving NHC-boranes, thiols, and nitroso compounds for the boryl-functionalization of alkenes. This approach enables the generation of boryl and NO

radicals via homolytic N-N/N-O bond cleavage, thereby facilitating subsequent boryl-oximation of alkenes and allowing for the efficient synthesis of functionalized organoboron compounds (Scheme 1c). Notably, the newly established boryl-oximation proceeds well in the absence of any photocatalysts and exhibits excellent functional group tolerance, broad substrate scope, and compatibility with late-stage functionalization.

Results and discussion

Initially, methyl 4-vinylbenzoate 1aa, borane adduct 2 and Nnitrosopiperidine 3a were chosen as the model substrates to

Table 1 Selected optimization results

Entry	Deviation from optimized conditions a	4aa ^b (yield, %)	5aa ^b (yield, %)	Conv. ^b (%)
1	None	93 (88) ^c	<5	>95
2^d	376-400 nm LEDs	24	6	>95
3^d	406-443 nm LEDs	42	12	>95
4^d	420-480 nm LEDs	32	13	>95
5^d	437-471 nm LEDs	52	10	81
6^e	THF	56	16	>95
7^e	Dioxane	54	13	93
8^e	DMF	55	13	>95
9^e	EtOH	45	<5	75
10^e	ⁱ PrOH	55	<5	85
11^e	Acetone	52	12	>95
12^e	DCM	43	10	87
13 ^e	Toluene	65	8	94
14	No 1,2-ethanedithiol	<5	<5	<5
15	In the dark	n.d	n.d	0
16	Air	30	7	>95

^a Reaction conditions: **1aa** (0.3 mmol), **2** (0.9 mmol), **3a** (0.6 mmol), **1,2**-ethanedithiol (20 mol%), toluene (3.0 mL), 20 W 437–471 nm LEDs, rt, 40 h, and under N₂. ^b Determined by ¹H NMR using **1,3,5**-trimethoxybenzene as an internal standard. ^c The data in parentheses are isolated yields. ^d **1aa** (0.3 mmol), **2** (0.3 mmol), **3a** (0.45 mmol), **1,2**-ethanedithiol (20 mol%), CH₃CN (3.0 mL), rt, 30 h, and under N₂. ^e **1aa** (0.3 mmol), **2** (0.3 mmol), **3a** (0.45 mmol), **1,2**-ethanedithiol (20 mol%), solvent (3.0 mL), 20 W 437–471 nm LEDs, rt, 30 h, and under N₂.

optimize the reaction conditions, and the results are listed in Table 1. After the systematic screening, the best reaction conditions were obtained as follows: 1aa (1.0 equiv.), 2 (3.0 equiv.), 3a (2.0 equiv.), and 1,2-ethanedithiol (20 mol%) in a solution of toluene at room temperature under blue LED (20 W, 437–471 nm) irradiation for 40 h, producing the desired product 4aa in 88% isolated yield (entry 1). The observed stereoselectivity is further supported by the single-crystal structure of the product 4ai (CCDC number: 2448501).¹8 In addition, the wavelength of the light source significantly influences the reactivity and selectivity of the model reaction. Remarkably, the screening of light sources showed that the use of 437–471 nm LEDs resulted in the formation of 4aa in 52% yield and 81% conv. (entries 2–5).

Interestingly, during the course of optimization, a byproduct 5aa was isolated and well characterized (also see Scheme 3d). We further found that the choice of solvent significantly influenced the reaction efficiency. Under irradiation with blue LEDs (20 W, 437–471 nm), most solvents afforded product 4aa in approximately 50% yield, while toluene serving as the optimal solvent enhanced the yield to 65% (entries 6–13). Control experiments showed that the reaction almost could not proceed in the absence of the thiol catalyst or light irradiation (entries 14–15). The reaction proceeded under ambient air, however, the 4aa was achieved with only 30% isolated yield (entry 16). We next focused on evaluating some other reaction parameters (see Table S1 for details). Thiol catalysts with different substituents afforded product 4aa with comparable yields, while using

thiophenol as a catalyst only resulted in a 10% yield. Alkyl-substituted nitrosamines showed favorable results for the formation of the main product, whereas aryl-substituted nitrosamines and ^tBuONO (TBN) provided lower yields. Finally, the yield could be further improved by adjusting the material ratio to an excessive amount of 2 and 3a.

Subsequently, we investigated the substrate scope under the optimal conditions listed in Table 1 (Table 2). First, we studied the substituent effect on the aromatic rings of styrene derivatives. The reaction demonstrated good functional group compatibility, tolerating a variety of substituents, including ester, cyano, acylamino, sulfonyl, trifluoromethyl, acetal, and halogen groups. Most of the arylalkenes with an electronwithdrawing group at the para-position afforded boryloximation products (4aa-4ae, 4ag-4aj, and 4al) in moderate to excellent yields (52-91%). Individually, the reaction of 2-(4vinylphenyl)-1,3-dioxolane was sluggish, affording the desired product 4af only in 43% yield. Notably, when PhCN was used as the solvent instead of PhMe, both 4-phenylstyrene and styrene participated in the reaction, affording the corresponding products 4an and 4ao in 51% and 41% yields, respectively. Styrenes with electron-donating groups and internal olefins exhibited relatively poor reactivity. As a result, 4-(dimethylamino)styrene, 4-methoxystyrene and methyl-4-(prop-1en-1-yl)benzoate yielded the corresponding products (4ak, 4am, and 4aa-Me) only in trace to 18% yields, with most of the alkene substrates remaining unreacted.

Table 2 Photoinduced boryl-oximation of alkenes: substrate scope a,b,c,d,e

^a Unless otherwise noted, the reaction conditions were the same as those described in Table 1, entry 1, 1 (0.5 mmol), 2 (1.5 mmol), 3a (1.0 mmol), 1,2-ethanedithiol (20 mol%), toluene (5.0 mL), 20 W 437–471 nm LEDs, rt, 40 h, and under N₂. The isomer ratios were determined by ¹H NMR. ^b Used ^bBuSH (50 mol%) as the catalyst. ^c Used PhCN (5.0 mL) as the solvent. ^d Used ^bBuONO (TBN, 1.0 mmol) as the NO radical source. ^e 1 (1.0 mmol), 2 (3.0 mmol), 3a (2.0 mmol), 1,2-ethanedithiol (20 mol%), PhCN (5.0 mL), 20 W 437–471 nm LEDs, rt, 40 h, and under N₂.

We next found that *meta*-substituted substrates delivered the desired products in good yields (4ap-4as, 65-90%), whereas *ortho*-substituted counterparts showed reduced reactivity (4at-4av, 45-57%), indicating a mild steric effect in the boryloximation of alkenes. In addition to styrene and its derivatives, a fused-ring-substituted ethylene (*i.e.*, 2-naphthyl ethylene, corresponding to 4aw, 80%) and furanyl and pyridyl

ethylenes (corresponding to **4ax-4ay**, 63–81%) could also provide favorable results. Subsequently, we randomly evaluated the applicability of an aliphatic olefin **1az** in this reaction. Unfortunately, **1az** did not react with **2** and **3a** under the standard conditions, and no desired product was observed. Remarkably, by fine-tuning the standard conditions (using ¹BuONO instead of **3a** or employing 50 mol% ¹BuSH as

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a catalyst), the reaction could be further extended to functionalized alkenes including ester-, amide-, cyano- and sulfonyl-substituted ethylenes. Specifically, ethyl acrylate, benzyl acrylate, and phenyl vinyl sulfone gave the corresponding products (4ba-4bb and 4bh) in moderate yields with high selectivities. *N*-substituted acrylamides bearing phenyl, benzyl or alkyl reacted smoothly to afford 4bc-4bf in moderate yields. Finally, acrylonitrile demonstrated excellent reactivity, delivering the boryloximation product 4bg in 87% yield, albeit with relatively lower stereoselectivity.

Subsequently, we demonstrated the practicality of this method through gram-scale synthesis and late-stage functionalization of bioactive molecules. First, the model reaction was readily scaled up to a gram scale, affording compound **4aa** with well retained yield (Scheme 2a, 87% yield, 1.05 g of product **4aa** was obtained). Next, we evaluated the applicability of this protocol to the late-stage modification of multifunctionalized alkene-containing pharmaceutical derivatives (Scheme 2b). The results showed that derivatives of podophyllotoxin, atomoxetine, and cholesterol proceeded well to afford boryloximation products **4ca-4cc** in 56–87% yields. Given the wide-spread presence of oxime ether scaffolds in pharmaceuticals¹⁹—which demonstrate diverse biological activities such as

antifungal, antibacterial, antiviral and antidepressive effects—along with their versatile reactivity in synthetic chemistry, 20 we further explored the transformation of product **4aa** (Scheme 2c). Specifically, **4aa** could smoothly undergo a Michael addition with ethyl acrylate using triphenylphosphine as a catalyst, 21 leading to the formation of α -boryl oxime ether **6** in 95% yield with retention of configuration. The oxidative cleavage of the C–B bond in **6** yielded α -hydroxy oxime ether **7**, which was further reduced with NaCNBH₃ to afford amino alcohol **8** in 79% yield.

To understand the possible reaction mechanism, several control experiments were designed and conducted (Scheme 3). A radical-trapping experiment was performed by adding the radical scavenger TEMPO to the reaction system (Scheme 3a). The experimental results revealed that no target 4aa was formed, but both the adducts 9 and 10 were detected by high-resolution mass spectrometry, indicating that boryl radicals were involved therein. In addition, the absorption of 1aa, 2 and 3a (ranging from 380 to 440 nm) was clearly observed with UV-vis absorbance, showing that the peaks of 3a are mainly located near the UV region (see Fig. S5 for details). Next, a deuterium labeling experiment with the reaction of 1aa, 2-d₃ and 3d was carried out under the standard conditions (Scheme 3b). We found that the deuteration rate of the product 4aa-d₂ was

(a) Gram-scale synthesis

(b) Late-stage modification of pharmaceuticals

^a Unless otherwise noted, the reaction conditions were the same with those described in Table 1, entry 1, **1** (0.4 mmol), **2** (1.2 mmol), **3a** (0.8 mmol), 1,2-ethanedithiol (EDT, 20 mol %), toluene (4.0 mL), 20 W 454 nm LEDs, rt, 40 h, under N₂. ^b Used ^tBuSH (50 mol %) as catalyst, ^tBuONO (TBN, 1.0 mmol) as NO radical source.

$$\begin{bmatrix} B \end{bmatrix} = \begin{bmatrix} A_2 & Me \\ A_1 & N \\ N & N \\ Me & + \end{bmatrix}$$

(c) Transformations of the product 4aa

Me
$$H_2$$
 OEt OEt OEt OEt OEt OEt OEt OEt OO_2Me OO_2Me

Reaction conditions: (a) PPh₃ (30 mol %), ethyl acrylate (2.0 equiv), CH₃CN, 65 $^{\circ}$ C , 48 h. (b) H₂O₂ (10.0 equiv), CH₃CN/MeOH, 35 $^{\circ}$ C, 1 h. (c) NaCNBH₃ (2.0 equiv), AcOH, rt, 12 h.

Scheme 2 Applications of the boryl-oximation protocol and transformations of the products.

3a

Scheme 3 Mechanistic studies.

1aa

consistent with that of the borane adduct 2-d3, indicating that the B-D bonds in both the reactant and product did not undergo H-D exchange during the reaction. Notably, 1phenylpiperazine 11 was also obtained in 88% yield, which provides further evidence for the proposed reaction mechanism. Subsequently, the kinetic isotope effect (KIE) value of borane was determined through an intermolecular competition and two parallel reactions, with both results being close to 1.0. These experimental results suggest that the generation of boryl radicals might not be the rate-determining step (Scheme 3b). To

further corroborate the homolytic cleavage mechanism of Nnitrosamine, a control experiment was conducted in the absence of compound 2 (Scheme 3c). The reaction of benzyl acrylate (1bb) with 1-nitroso-4-phenylpiperazine (3d) afforded the hydroamination product 12 in 10% yield. Trace amounts of the aminoximation product 13 and 1-phenylpiperazine 11 were also detected by high-resolution mass spectrometry. These results further support that N-nitrosamine generates amino and nitrosyl radicals upon photoirradiation. Finally, we explored the origin of the byproduct 5aa observed in the reaction (Scheme

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Scheme 4 Proposed mechanism.

3d). Experiments showed that the photoinduced secondary boryl-oximation reaction of **1aa** with **4aa** and **3a** smoothly occurred to generate the byproduct **5aa** in 48% yield. This unexpected discovery provides new access to realizing multifunctionalization of NHC-boranes. ^{15h}

Based on the above control experiments and related reports,17,22 a plausible reaction mechanism for this borylfunctionalization of alkenes is proposed and listed in Scheme 4. First, the nitroso compound undergoes homolytic cleavage to form nitrosyl radical I and amino radical II (or alkoxyl radical II') upon light irradiation. Subsequently, the amino radical (or alkoxyl radical) undergoes a cascade radical chain transfer process (two-step HAT) to generate the NHC-boryl radical IV. Moreover, light on-off experiments (see Fig. S10 for details) demonstrated that the radical chain transfer process is transient and persistent irradiation is essential to run these cascades. Next, the NHC-boryl radical undergoes addition to the alkene, generating radical intermediate V, which subsequently couples with the nitroso radical I to form the unstable intermediate 4'. This intermediate then undergoes rearrangement to furnish the target product 4.

Conclusions

In summary, we have developed a new method that does not rely on the use of transition metals or tailored photocatalysts for the boryl-oximation of alkenes, affording a number of useful $\alpha\text{-boryl}$ oxime compounds in moderate to good yields. Both the latestage modification of various pharmaceutical molecules and the product transformations demonstrate its synthetic applications in the field of synthetic chemistry. Mechanistic experiments including radical-trapping experiments, deuterium-labeling experiments and KIE experiments disclose the possible pathway for this boryl-oximation of alkenes, in which a radical chain transfer is key for the generation of the boryl radical. The photoinduced activation mode of NHC-boranes offers an ideal platform for the bifunctionalization of alkenes. Our laboratory will focus on expanding this boryl radical-based strategy to address more challenging chemical transformations.

Author contributions

Qiang Huang (investigation, methodology, funding acquisition, and writing – original draft); Na Li (investigation and

methodology); Panke Zhang (supervision and funding acquisition); Hongji Li (conceptualization, supervision, funding acquisition, and writing – review & editing).

Conflicts of interest

The authors declare no competing financial interest.

Data availability

CCDC 2448501 contains the supplementary crystallographic data for this paper.²³

The detailed experimental process and additional data associated with this work are available in the SI. See DOI: https://doi.org/10.1039/d5sc03708e.

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