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Nickel(0)-catalyzed linear-selective hydroarylation of unactivated alkenes and styrenes with aryl boronic acids†

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Herein, we describe the first linear-selective hydroarylation reaction of unactivated alkenes and styrenes with aryl boronic acids, which was achieved by introducing a directing group on the alkenes. This efficient, scalable reaction serves as a method for modular assembly of structurally diverse alkyl arenes, including γ -aryl butyric acid derivatives, which are widely utilized as chemical building blocks for the synthesis of various drugs and other biologically active compounds.

Introduction

Transition-metal-catalyzed alkene hydroarylation involving metal hydrides (M–H)¹ has become one of the most widely used strategies for the synthesis of alkyl arenes.² To date, three types of aryl sources have been used for this purpose: (1) simple arenes, (2) aryl halides, and (3) aryl metals (*i.e.*, aryl stannanes or boronic acids). The simple arenes are usually restricted to heteroarenes and arenes with a directing group because of the challenge of activating the inert C–H bonds of simple arenes.³ Aryl halides and aryl metals as the aryl sources require the use of O₂ or an excess of a reductant (such as a silane or an alkyl halide), which limits the practicality of alkene hydroarylations with these two sources (Scheme 1a).^{4,5} A significant breakthrough was made by Zhou *et al.*, who reported a highly selective Ni-catalyzed hydroarylation of styrenes and 1,3-dienes with aryl boron compounds under redox-neutral conditions (Scheme 1b).⁶ However, this method still suffers from some limitations: (1) the reaction is efficient only with styrene and 1,3-diene substrates, and (2) only branched products can be achieved.

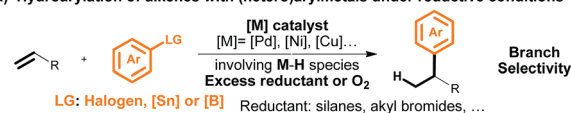
Inspired by recent work on olefin functionalization reactions in which the reactivity and regioselectivity are controlled *via* the introduction of a coordinating group,^{7,8} we have developed the first method for directing-group-controlled linear-selective hydroarylation of unactivated alkenes and styrenes with aryl boronic acids under redox-neutral conditions (Scheme 1c). This efficient, scalable reaction serves as a method for modular assembly of structurally diverse alkyl arenes, including γ -aryl butyric acid derivatives, which are widely utilized as chemical

building blocks for the synthesis of various drugs and other biologically active compounds (Scheme 1). γ -Aryl butyric acids are traditionally prepared by Friedel–Crafts reactions between arenes and butyrolactones,⁹ but this method suffers from disadvantages such as harsh reaction conditions, limited substrate scope (it is suitable only for electron-rich arenes), and poor regioselectivity.

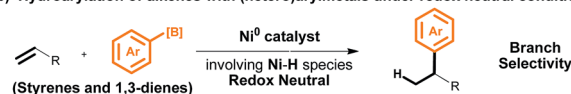
Results and discussion

In a preliminary experiment, we found that treatment of 3-butenic acid derivative **1a** (0.2 mmol), which bears an 8-

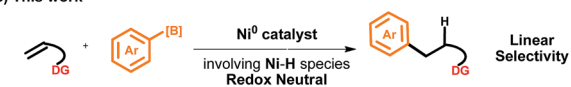
a) Hydroarylation of alkenes with (hetero)arylmets under reductive conditions



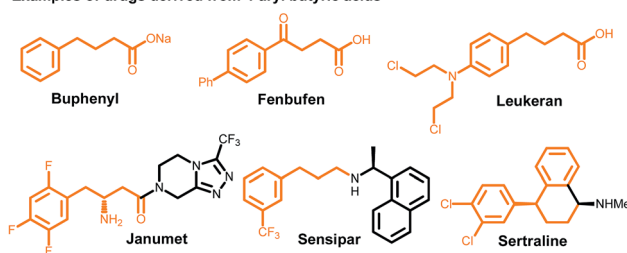
b) Hydroarylation of alkenes with (hetero)arylmets under redox neutral conditions



c) This work



Examples of drugs derived from 4-aryl butyric acids



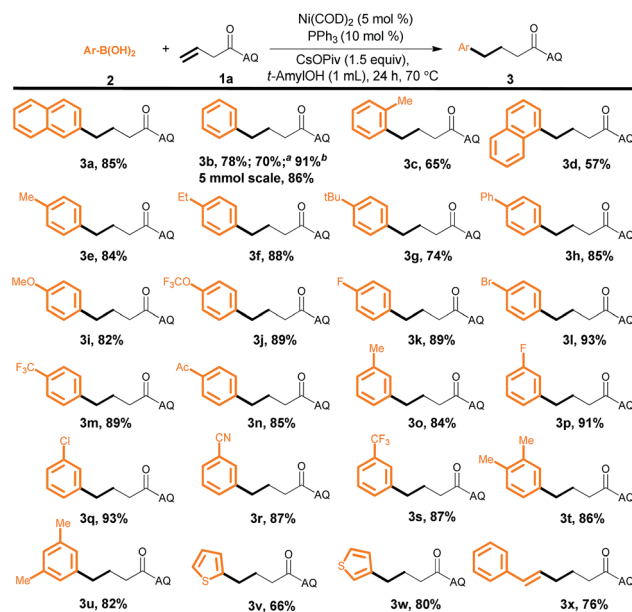
Scheme 1 Transition-metal-catalyzed hydroarylation of alkenes involving M–H and examples of drugs derived from 4-aryl butyric acids.

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aminoquinoline (AQ) directing group, with $\text{Ni}(\text{COD})_2$ (5 mol%), PCy_3 (10 mol%), and 2-naphthalenylboronic acid **2a** (0.3 mmol) in MeOH (1 mL) at 70 °C for 24 h gave linear product **3a** in 21% yield (Table 1, entry 1). Replacing MeOH with *t*-amyl alcohol slightly improved the yield (entry 2), and adding 1.5 equiv. of K_2CO_3 improved the yield even further (to 38%, entry 3). Evaluation of different bases revealed that CsOPiv was the best choice (entries 3–7). We also found that changing the **1a**/**2a** molar ratio to 1 : 2 increased the yield to 61% (entry 8). Finally, various phosphine ligands were investigated (entries 9–12); switching from PCy_3 to PPh_3 dramatically improved the yield (to 85%, entry 9). A control experiment confirmed that the reaction did not occur in the absence of $\text{Ni}(\text{COD})_2$ (entry 13).

Next, we investigated the substrate scope with respect to the aryl boronic acid by carrying out reactions with 3-butenic acid derivative **1a** as the alkene (Scheme 2). To our delight, electron-rich, electron-poor, and sterically hindered aryl boronic acids all afforded the desired products in good to excellent yields (**3a–3u**). Heteroaryl boronic acids were also reactive (**3v** and **3w**), as was an alkenyl boronic acid, which selectively afforded (*E*)-6-phenylhexenoic acid derivative **3x** under the optimized conditions. In addition, we found that a high yield of **3b** could be obtained by hydroarylation of **1a** with an aryl boroxine or an aryl boronic ester as the aryl source. It is important to stress that these reaction conditions were compatible with a remarkable variety of functional groups on the aryl boronic acids, including halogens (F, Cl, and Br) and acetyl, cyano, and methoxy groups, yielding products (**3i–3s**) that could be subjected to further



Scheme 2 The reactions of diverse aryl boronic acids **2** with 3-butenic acid derivative **1a**. Yield of isolated product is given. ^a 5,5-Dimethyl-2-phenyl-1,3,2-dioxaborinane was used as the aryl source. ^b Triphenylboroxine was used as the aryl source.

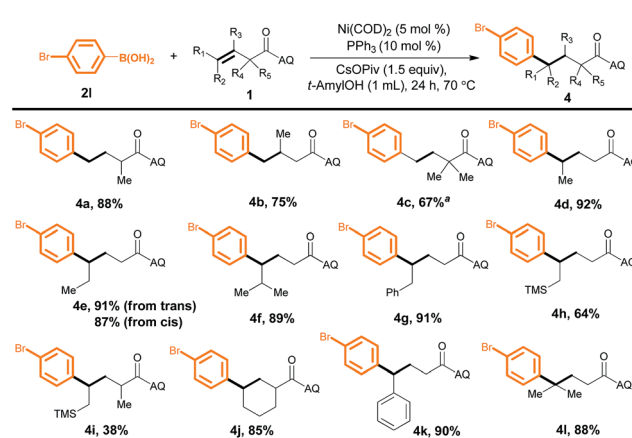
synthetic transformations. In addition, the reaction could be carried out on gram scale without a decrease in the yield (1.5 g of **3b**, 86%). The structure of **3b** was confirmed by X-ray diffraction analysis of a single crystal.¹⁰

We next evaluated the utility of this method for various unactivated alkenes (Scheme 3). Terminal alkenes bearing a single substituent at the α - or β -position reacted smoothly to afford the desired products in good to excellent yields (**4a** and **4b**). A sterically congested α,α -disubstituted terminal alkene was also reactive, furnishing desired product **4c** in moderate yield, although a higher temperature (125 °C) and a higher loading (10 mol%) of a different ligand, PhPMe_2 , were required. Furthermore, internal alkenes bearing a variety of substituents at the γ -position could also be efficiently converted into the

Table 1 Optimization of directed hydroarylation of unactivated alkene **1a** with 2-naphthalenylboronic acid **2a**^a

Entry	Ligand	Base	Solvent (H source)	Yield ^b [%]
1	PCy_3	—	MeOH	21
2	PCy_3	—	<i>t</i> -AmylOH	29
3	PCy_3	K_2CO_3	<i>t</i> -AmylOH	38
4	PCy_3	KHCO_3	<i>t</i> -AmylOH	23
5	PCy_3	K_3PO_4	<i>t</i> -AmylOH	43
6	PCy_3	NaOtBu	<i>t</i> -AmylOH	NR
7	PCy_3	CsOPiv	<i>t</i> -AmylOH	53
8 ^c	PCy_3	CsOPiv	<i>t</i> -AmylOH	61
9 ^c	PPh_3	CsOPiv	<i>t</i> -AmylOH	86(85) ^d
10 ^c	CyPPH_2	CsOPiv	<i>t</i> -AmylOH	77
11 ^c	S-Phos	CsOPiv	<i>t</i> -AmylOH	33
12 ^c	XantPhos	CsOPiv	<i>t</i> -AmylOH	59
13 ^e	PPh_3	CsOPiv	<i>t</i> -AmylOH	NR

^a Reactions were carried out with $\text{Ni}(\text{COD})_2$ (5 mol%), ligand (10 mol%), base (0 or 1.5 equiv.), **1a** (0.2 mmol), and 2-naphthalenylboronic acid (**2a**, 0.3 mmol) in solvent (1 mL) for 48 h at 25 °C under a N_2 atmosphere. ^b Yields were determined by NMR spectroscopy with an internal standard. NR = no reaction. ^c 2.0 equiv. of **2a** was used. ^d The isolated yield is given in parenthesis. ^e Reaction without $\text{Ni}(\text{COD})_2$.



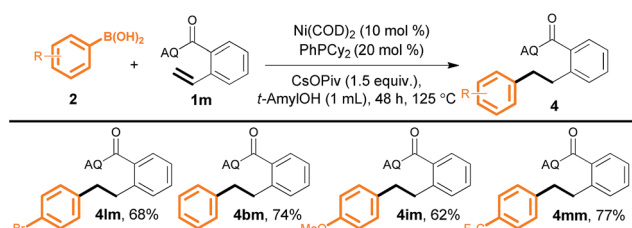
Scheme 3 Reactions of unactivated alkenes with aryl boronic acid **2l**. Yield of isolated product is given. ^a PhPMe_2 (10 mol%) as the ligand at 125 °C.



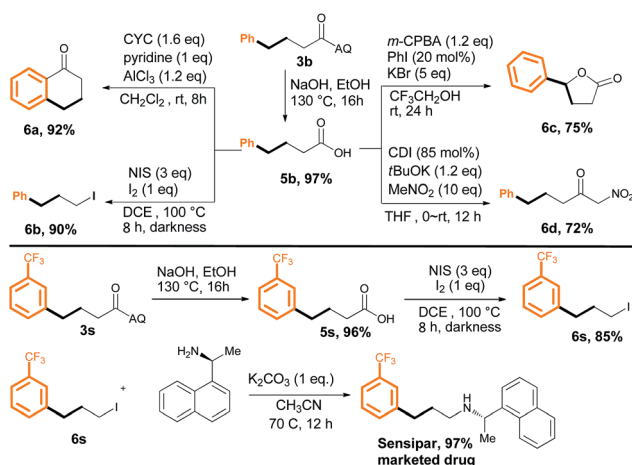
corresponding products (**4d–4l**); the steric and electronic effects of the substituents appeared to be negligible. Notably, the yields of **4e** from *trans*- and *cis*-3-hexenoic acid were essentially the same. In addition, α,γ -disubstituted alkenes gave desired products **4i** and **4j**. Surprisingly, a γ,γ -disubstituted alkene could also be hydroarylated to produce compound **4l** (88%), which has a quaternary carbon center.

We wondered whether the introduction of the AQ directing group at the *ortho*-position of styrene would result in linear selectivity rather than the branched selectivity reported by Sigman *et al.* and Zhou *et al.* for hydroarylation of styrenes with aryl metals.^{4,6} After extensive investigation, we found, to our delight, that reactions of styrene derivative **1m** with various aryl boronic acids occurred smoothly with completely linear selectivity and in moderate yield by employment of PhPCy₂ as the ligand and higher catalyst loading (10 mol% Ni(COD)₂), higher temperature (125 °C), and longer reaction time (Scheme 4). This result clearly proved that the directing group controlled the regioselectivity of the hydrometalation.

The products obtained by the method described herein could provide access to structurally diverse synthetic building blocks by means of additional transformations (Scheme 5, upper panel). Specifically, removal of the AQ directing group of **3b** smoothly provided 4-phenylbutanoic acid (**5b**) in 97% yield. Acid **5b** could undergo intramolecular cyclization to yield tetralin-derivative **6a**, or the carboxyl group could be replaced with an iodine atom (**6b**). Five-membered lactone **6c** could be obtained by treatment of **5b** with a hypervalent iodine reagent, or the carboxyl group could be replaced with a nitro group (**6d**). Five-membered lactone **6c** could be obtained by treatment of **5b** with a hypervalent iodine reagent,



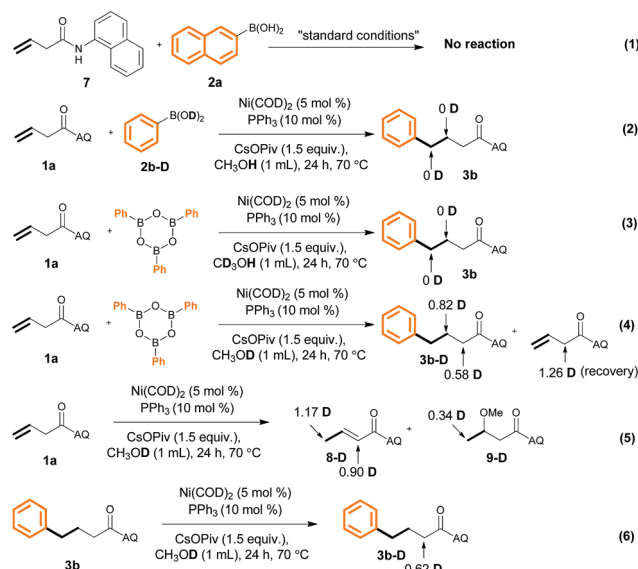
Scheme 4 Reactions of styrene derivative **1m** with aryl boronic acids.



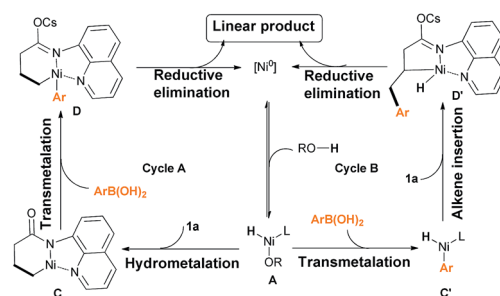
Scheme 5 Transformations of hydroarylation products **3b** and **3s**.

and **5b** readily underwent attack by CH₃NO₂ to afford corresponding ketone derivative **6d**. To further demonstrate the synthetic usefulness of our method, we synthesized the drug Sensipar from **3s** (Scheme 5, lower panel). First, free acid **5s** was synthesized by removal of the AQ directing group, and then the acid was transformed into the corresponding alkyl iodide (**6s**) by treatment with NIS/I₂ in DCE. Simple amination of **6s** gave Sensipar in 97% yield.

To probe the reaction mechanism, we carried out a control experiment and a series of deuteration experiments (Scheme 6). We found that no reaction occurred when *N*-(naphthalenyl) butenamide **7** and 2-naphthalenylboronic acid (**1a**) were used as the substrates (eqn (1)). This result indicates that the presence of the AQ directing group on the unactivated alkene was indispensable. Treatment of PhB(OD)₂ **2b-D** with alkene **1a** in MeOH or treatment of (PhBO)₃ with **1a** in CD₃OH gave undeuterated product **3b** (eqn (2) and (3)). However, the use of MeOD as the solvent led to the formation of **3b** that was only partially deuterated at both the α - and β -positions (eqn (4)), which reveals that the hydrogen atom of the Ni-H intermediate **A** (Scheme 7) came from the methanol O-H group and



Scheme 6 Control experiment and deuteration experiments.



Scheme 7 Proposed mechanism of the Ni(0)-catalyzed directing-group-controlled hydroarylation of unactivated alkenes with aryl boronic acids.

hydrometalation of Ni–H is irreversible in the presence of aryl boronic acid. Further experiment suggests that hydrometalation of starting material **1a** via Ni–H species would also happen to form 5-membered nickelacycle and is reversible in the absence of aryl boronic acid under our reaction conditions (eqn (5) and (6)). Additionally, the deuteration of α -positions might be because of CsOPiv base.

On the basis of literature reports and our own mechanistic experiments, we propose two catalytic cycles that starts with the reversible formation of Ni–H species **A** (Scheme 7). For path A, coordination and subsequent hydrometalation of the C=C bond affords the six-membered nickelacyclic intermediate **C**, and then transmetalation of the aryl boronic acid generates intermediate **D**.¹¹ Finally, reductive elimination gives the desired product and regenerates the Ni(0) catalyst. Alternatively, intermediate **A** could also undergo transmetalation first to form Aryl–Ni–H species **C'**, and then alkene insertion to form five-membered nickelacyclic intermediate **D'**, which would involve reductive elimination to yield the same linear product. Note, that we have detected the production of naphthalene under our reaction condition. It indicated that cycle B might be the real process. However, at this stage, we still can't rule out cycle A.

Conclusions

In conclusion, we have described the first method for Ni(0)-catalyzed directing-group-controlled linear-selective hydroarylation of unactivated alkenes and styrenes with aryl boronic acids under redox-neutral conditions. Our efficient, scalable method provides a general route to structurally diverse alkyl arene derivatives, including γ -aryl butyric acid derivatives, which are widely utilized as chemical feedstocks for the preparation of drugs and other biologically active compounds.

Conflicts of interest

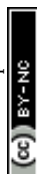
There are no conflicts to declare.

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