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Introduction

Rhodium-catalysed selective C–C bond activation and borylation of cyclopropanes[†]

Yandong Wang,‡ Jingyi Bai,‡ Youqing Yang,‡ Wenxuan Zhao, Yong Liang,
^{*} Di Wang, Yue Zhao [•] and Zhuangzhi Shi[•]

Transition metal (TM)-catalysed directed hydroboration of aliphatic internal olefins which facilitates the construction of complex alkylboronates is an essential synthetic methodology. Here, an efficient method for the borylation of cyclopropanes involving TM-catalysed directed C–C activation has been developed. Upon exposure to neutral Rh(i)-catalyst systems, *N*-Piv-substituted cyclopropylamines (CPAs) undergo proximal-selective hydroboration with HBpin to provide valuable γ -amino boronates in one step which are otherwise difficult to synthesize by known methods. The enantioenriched substrates can deliver chiral products without erosion of the enantioselectivities. Versatile synthetic utility of the obtained γ -amino boronates is also demonstrated. Experimental and computational mechanistic studies showed the preferred pathway and the origin of this selectivity. This study will enable the further use of CPAs as valuable building blocks for the tunable generation of C–heteroatom or C–C bonds through selective C–C bond activation.

Alkylboronic esters1 are highly important in all facets of chemical science, ranging from synthetic chemistry² to materials science³ to drug discovery.⁴ Traditionally, these compounds can be prepared by the well-known Brown hydroboration, in which the *syn*-addition of hydroboranes to alkenes follows an anti-Markovnikov manner.5 During the past decades, the TM-catalysed hydroboration of aliphatic internal olefins with tethered directing groups has shown interesting chemo-, regio-, and diastereoselectivity, which greatly complements the classical noncatalytic process (Fig. 1A).6 The coordination of directing groups with the TM catalysts can deliver hydroboration products with controllable distal⁷ and/or proximal⁸ selectivity. Recently, directed borylation of aliphatic C-H bonds with proximal9 and distal10 selectivity has also gained momentum. The functionalization of the C-C bond is one of the most attractive strategies in organic chemistry. Despite the significant advances, transformations that allow the selective borylation of single C-C bonds remain effectively unknown.

CPAs are among the most prevalent molecules found in feedstock chemicals, drug molecules, and natural products.¹¹ In

‡ These authors contributed equally to this work.

this context, they have been exploited as robust building blocks in organic synthesis, including C-H and C-C bond functionalizations.12 Seminal work by Itami and Yamaguchi described the N-Piv-directed C-H borylation of CPAs with cis selectivity using phenanthroline-based ligands and a catalytic amount of an iridium salt (Fig. 1B).13 Inspired by the success of directed borylation of alkenes and alkanes, we envisioned that the borvlation of CPAs by chelation-assisted C-C bond activation¹⁴ could provide a unique opportunity to access amino-containing boronates as potent protease inhibitors.¹⁵ During the preparation of this paper, Yamaguchi and Yokogawa reported a related transformation achieved by C-C hydroboration of CPAs using an iridium catalyst and ^tBu-Quinox as the ligand (Fig. 1C).¹⁶ Notably, this elegant chemistry exhibited excellent distal selectivity, resulting in β -methyl alkylboronates. In contrast, herein, we describe a simple and practical method to achieve hydroboration of CPAs via proximal C-C bond cleavage affording yamino boronates in the presence of a rhodium complex and a simple phosphine ligand (Fig. 1D). A series of mechanistic experiments and DFT calculations revealed the preferred pathway of this transformation, which provides high proximalselectivity by the slightly catalytic system.

Results and discussion

Reaction design

Examination and optimisation of the reaction parameters were explored using CPA **1a** as the substrate and pinacolborane (HBpin) as the borylating source in conjunction with various TM catalysts and ligands (Table 1). As reported,¹⁶ when the

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State Key Laboratory of Coordination Chemistry, Chemistry and Biomedicine Innovation Center (ChemBIC), School of Chemistry and Chemical Engineering, Nanjing University, Nanjing 210093, China. E-mail: shiz@nju.edu.cn; yongliang@ nju.edu.cn

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Fig. 1 TM-catalysed borylation of olefins and CPAs. (A) The development of TM-catalysed directed borylation. (B) Ir-catalysed directed C-H borylation of CPAs. (C) Ir-catalysed distal-selective hydroboration of CPAs. (D) Rh-catalysed proximal-selective hydroboration of CPAs.

Table 1 Optimisation of reaction conditions for the proximal-selective hydroboration of CPA 1a ^a



Entry	[M] (mol%)	Ligand (mol%)	Solvent	$2a/3a^b$	Yield of $2\mathbf{a}^{b}$ (%)
1	$[Ir(cod)OMe]_2(5)$	^t Bu-Quinox (2.5)	Toluene	5/95	Trace
2	$[Ir(cod)OMe]_2(1)$	$PPh_3(4)$	Toluene	>99/1	42
3	$[Rh(CO)_2Cl]_2(1)$	$PPh_3(4)$	Toluene	>99/1	47
4	$[Rh(coe)_2Cl]_2(1)$	$PPh_3(4)$	Toluene	>99/1	52
5	$[Rh(cod)Cl]_2(1)$	PPh_3 (4)	Toluene	>99/1	99 (91) ^c
6	$Rh(PPh_3)_3Cl(1)$	_	Toluene	>99/1	65
7	$[Rh(OAc)_2]_2$ (1)	$PPh_3(4)$	Toluene	>99/1	4
8	$[Rh(cod)Cl]_2(1)$	$PPh_3(4)$	^c Hexane	>99/1	76
9	$[Rh(cod)Cl]_2$ (1)	$PPh_3(4)$	THF	>99/1	70
10^d	$[Rh(cod)Cl]_2$ (1)	$PPh_3(4)$	Toluene	>99/1	73
11^e	$[Rh(cod)Cl]_2$ (1)	$PPh_3(4)$	Toluene	_	0
12	$[Rh(cod)Cl]_2$ (0.5)	$PPh_3(2)$	Toluene	>99/1	85
13	$[Rh(cod)Cl]_2$ (1)	$PPh_3(2)$	Toluene	>99/1	79
14	$[Rh(cod)Cl]_2$ (1)	_	Toluene	_	8
15		$PPh_3(4)$	Toluene		0

^{*a*} Reaction conditions: cat [M] (0.5–5 mol%), L (2–4 mol%), 1a (0.20 mmol), HBpin (0.60 mmol) in solvent (1.0 mL), 24 h, 130 °C, under Ar, in sealed Schlenk tubes. ^{*b*} Determined by GC-MS analysis. ^{*c*} Yield of purified compounds after chromatography. ^{*d*} At 120 °C. ^{*e*} Using B₂pin₂ instead of HBpin.

reactions were performed with 5 mol% $[Ir(cod)OMe]_2$ as the catalyst and 2.5 mol% ^tBu-Quinox as the ligand at 130 °C for 24 h in toluene, branched boronate 3a was formed in 52% yield together with a small amount of the linear byproduct 2a (2a/3a = 5/95, entry 1). In this study, we focused our attention on product 2a. Completely inverse selectivity (2a/3a > 99/1) was observed with 1 mol% Ir catalyst and 4 mol% PPh₃, in which the

desired product **2a** was obtained in 42% yield (entry 2). Moreover, we found that Rh(1) complexes were more effective than the Ir catalyst. Upon switching the catalyst to $[Rh(CO)_2Cl]_2$, the yield of **2a** further increased to 47% (entry 3). Other Rh(1) complexes, such as $[Rh(coe)_2Cl]_2$, provided the corresponding product **2a** in 52% yield (entry 4), and $[Rh(cod)Cl]_2$ was the most efficient, giving a 99% yield of **2a** (entry 5). When Wilkinson's



Scheme 1 Evaluating different N-substituents

Substrate scope^a Table 2

catalyst (Rh(PPh₃)₃Cl) was used instead of [Rh(cod)Cl]₂ and PPh₃, the reaction provided 2a in a much lower yield (entry 6). Under these reaction conditions, upon switching to a Rh(II) catalyst, the reaction became very sluggish (entry 7). The replacement of toluene by other solvents such as ^chexane and THF provided acceptable yields of 2a (entries 8 and 9). The effect of temperature was also examined, and 120 °C was found to be less optimal (entry 10). Furthermore, the use of HBpin is crucial to this borylation process; other classes of borylating sources such as B₂pin₂ did not afford any borylation product (entry 11). Furthermore, decreasing the Rh catalyst loading to 0.5 mol%



^a Reaction conditions: [Rh(cod)Cl]₂ (1 mol%), PPh₃ (4 mol%), 1 (0.20 mmol), HBpin (0.6 mmol), in toluene (1.0 mL), 24 h, 130 °C, under argon, in sealed Schlenk tubes, yield of purified compounds after chromatography. ^b Using [Rh(cod)Cl]₂ (2.5 mol%), PPh₃ (10 mol%), 36 h, 130 °C. ^c Using [Rh(cod)Cl]₂ (2.5 mol%), PPh₃ (10 mol%), 36 h, 150 °C.

(entry 12) or ligand loading to 2 mol% still resulted in good yields (entry 13). Finally, control experiments showed that PPh_3 was required for high reactivity and the inhibition of side reactions (entry 14), and no product was observed in the absence of a Rh catalyst (entry 15).

We next evaluated the influence of the steric hindrance of substituents on directing groups (Scheme 1). Changing the directing group to *N*-Ac (**4a**) displayed no reactivity for ringopening. The directing group bearing a methylcyclohexane motif (**4b**) afforded the corresponding product **5b** in 73% yield. When a sample of **4c** containing a sterically hindered *N*-Ad moiety was subjected to the reaction conditions, the desired product **5c** was formed in a much lower yield. CPAs bearing other directing groups such as *N*-Boc (**4d**) and *N*-SO^tBu (**4e**), and substrate **4f** without a directing group completely failed. These results indicate that the appropriate selection of the *N*-Piv directing group ensures high reactivity and selectivity for this transformation.

Scope of the methodology

With the optimised conditions in hand, the scope of this methodology was subsequently explored (Table 2). The alkyl chains of the amide substituent R^1 , including methyl (1b), *n*butyl (1c), isopropyl (1d), cyclohexyl (1e) and benzyl (1f), were converted into the corresponding boronic esters 2b-2f in good to high yields. Phenyl (1g) and other aryl substituents bearing methyl (1h), methoxy (1i), fluoro (1j-1l), chloro (1m), cyano (1n) and ester (10) were compatible at the R^1 position. Among them, product 2l was subjected to X-ray crystallographic analysis. Beyond the use of aryl-containing CPAs, we examined naphthyl and heterocyclic substrates. Naphthyl compound 1p was readily transformed, as was thiophene substrate 1q. The use of the CPA 1r with an O-Piv motif was also tested and the desired boronate 2r was obtained in 84% yield. Furthermore, substrate 1s bearing a hydroxyl group still resulted in the desired product 2s in 56% yield. In addition, this process was not limited to substrates bearing diverse R¹ substituents, and CPAs with methyl (1t) and different aryl (1u-1w) groups at the R² position were also suitable starting materials to effectively afford the desired products 2t-2w efficiently. Notably, when the substrate 1x containing a terminal olefin motif was employed for the reaction, olefin hydroboration product 2x' was obtained in 54% yield and the ring-opening product 2x was not observed.

Synthetic applications

To showcase the synthetic utility of this proximal-selective hydroboration protocol, a 5.0 mmol reaction of CPA **1g** was first conducted, enabling the gram-scale synthesis of the linear product **2g** in 79% isolated yield (Scheme 2). The formed boronate group is an extremely versatile intermediate in organic synthesis, which can be converted into other functional substituents. Compound **2g** could be transformed into potassium trifluoroborate salt **4** using KHF₂ in an excellent yield. Subsequently, mild oxidation of **2g** provided the corresponding alcohol **5** in 78% yield. The Cu-catalysed Chan–Lam coupling¹⁷ of **2g** with 4-fluorobenzamide allowed the formation of **1**,3-



Scheme 2 Gram-scale synthesis and the follow-up transformation of compound 2g. Reagents and conditions: (a) aq. KHF₂, MeOH, 3 h, rt; (b) NaBO₃·4H₂O (4.0 equiv.), H₂O/THF, 6 h, rt; (c) 4-fluorobenzamide (1.0 equiv.), Cu(OAc)₂ (10 mol%), 1,10-phen (10 mol%), NaOSiMe₃ (1.1 equiv.), DTBP (3.0 equiv.), 4 Å MS, ^tBuOH, 24 h, 75 °C; (d) PhI (1.5 equiv.), PdCl₂(dppf)·CH₂Cl₂ (8 mol%), Ag₂O (1.5 equiv.), K₂CO₃ (3.0 equiv.), H₂O/THF, 33 h, 80 °C to rt; (e) 2-bromopyridine (1.0 equiv.), Pd(OAc)₂ (4 mol%), Ad₂PⁿBu (12 mol%), LiO^tBu (6.0 equiv.), H₂O/THF, 24 h, 100 °C; (f) C₂H₃MgBr (6.0 equiv.), 0 °C to rt, 1 h; then I₂ (6.0 equiv.) in MeOH, -78 °C, 30 min; then NaOMe (6.0 equiv.) in MeOH, (4.0 equiv.), H₂O/DCM, 5 h, 50 °C.

diamide derivative **6** in 70% yield. Furthermore, Pd-catalysed Suzuki–Miyaura coupling between substrate **2g** and iodobenzene or 2-bromopyridine gave the desired products 7 and **8** efficiently.¹⁸ The Zweifel olefination of **2g** with vinylmagnesium bromide furnished the terminal alkene product **9**.¹⁹ In addition, the Ag-catalysed fluorination of **2g** with Selectfluor® could give the monofluoroalkane **10** in a good yield based on a protocol developed by Li and co-workers.²⁰

Mechanistic studies

Based upon the above results and the reported studies, we proposed that the transformation proceeds through four possible pathways (Fig. 2). The Rh-catalysed C–H borylation proceeds first,¹⁶ followed by C–C activation and hydrogenation (pathway I). The Rh-catalysed C–C activation and β -H elimination generate an allylic amide, which further undergoes hydroboration (pathway II).²¹ Oxidative addition of Rh species to CPAs and β -H elimination generate an enamide, and subsequently olefin migration and hydroboration (pathway III) take place.²² The Rh-catalysed C–C activation and hydroboration deliver the final products directly (pathway IV).

Several plausible intermediates were then investigated to provide insight about the potential mechanism of this transformation (Scheme 3). The C–H borylation product **11** failed to afford the desired product **2a**, excluding pathway **I** (Scheme 3A-1). The reaction between pre-generated *N*-allylpivalamide (**12**) and HBpin could produce hydroboration product **2a** in



a modest yield (Scheme 3A-2). Impressively, treatment of substrate **1a** in the system without the addition of HBpin was found to be applicable to the formation of enamide **13** through Rh oxidative addition to the hindered proximal C–C bond (Scheme 3A-3). Moreover, this compound **13** can convert into the corresponding product **2a** by olefin isomerization and hydroboration (Scheme 3A-4). To further understand how these proximal-selective products formed, we investigated the reaction using the enantioenriched CPAs (Scheme 3B). The enantiomers (1*S*,2*R*)-**1u** (>99% ee) and (1*R*,2*S*)-**1u** (>99% ee) were, respectively, treated with HBpin under the standard conditions, giving the desired products (*R*)-**2u** and (*S*)-**2u** without erosion of

the enantioselectivities. These results clearly excluded the insertion of Rh–H species onto the olefin intermediates through pathways **II** and **III**, suggesting that pathway **IV** involving direct C–C cleavage and hydroboration is possible. In order to support this proposal, we tried to isolate and characterize the key metallacyclic intermediate *via* Rh-mediated C–C bond activation of CPAs (Scheme 3C). Treatment of several CPAs with [Rh(cod)Cl]₂ failed to provide any stable intermediate, but fortunately, heating CPA **14** with stoichiometric [Rh(CO)₂Cl]₂ under neat conditions could successfully generate a dimeric rhodacyclopentanone **15**.²³ When characterized by X-ray diffraction, the Rh(1) centre in the structure of complex **15** shows axial chelation



Scheme 3 Mechanistic experiments. (A) Investigation of the plausible intermediates. (B) Hydroboration of enantioenriched CPAs. (C) Synthesis of rhodacyclo complex 15.

assisted by the *N*-Piv directing group, leading to proximal selective C–C bond cleavage and CO insertion. This establishes that Rh(I) salts are effective for insertion into the proximal C–C bond of CPAs without the assistance of HBpin.

We also performed a series of density functional theory (DFT) calculations to obtain further mechanistic information of this transformation (for computational details, see the ESI†). Fig. 3 shows the Gibbs free energy profile of this reaction. The catalytic cycle begins with a ligand exchange that replaces a cod ligand in the stable intermediate **A** with substrate **1a**. Directed by the amide oxygen, a subsequent proximal C–C bond cleavage of **1a** takes place which affords **C**. This is the rate-limiting step in the catalytic cycle ($\Delta G^{\neq} = 35.5$ kcal mol⁻¹). In this C–C bond cleavage step, we found that the non-directed transition states for both proximal and distal cleavage are unfavourable (for details, see the ESI†). Then we computed two possible pathways for hydroboration of **C**, oxidative addition and σ -complex assisted metathesis (σ -CAM).²⁴ It was found that H–B oxidative

addition to form Rh(v) species K is endergonic by 39.5 kcal mol⁻¹. Alternatively, the σ -CAM pathway is much more favourable. The free energy of the four-membered cyclic σ-CAM transition state TS2 for the formation of intermediate D from intermediate C and Hbpin is 23.7 kcal mol⁻¹. We also calculated other possible σ -CAM transition states (TS7-TS9), which would afford other hydroboration products, and found them to be higher in energy. From D, the re-coordination of the cod ligand leads to E, which reductively eliminates the C-H bond and forms the product-coordinated F. We also located a side pathway that corresponds to the formation of alkene products. From C, the directing group may dissociate from the metal centre to form intermediate G. Then, a β-H elimination (TS4) takes place which forms π -allyl-Rh(III) complex H. The C-H bond subsequently undergoes reductive elimination via TS5 or TS6 to produce I or J, respectively. The enamide I is the kinetically favoured product, and its negative free energy $(-2.1 \text{ kcal mol}^{-1})$ supports the observation of 13 in Scheme 3A-



Fig. 3 Free energy diagram of hydroboration of CPAs using PPh₃ as the ligand and DFT-computed four transition states for the σ -CAM pathways. Energies are in kcal mol⁻¹.



Fig. 4 DFT-computed free energies for the two competitive pathways without PPh_3 as the ligand. The zero point is $[Rh(cod)Cl]_2$. Energies are in kcal mol^{-1} .

3, where HBpin is not added. This also indicated that, under standard conditions, alkene **12** and **13** can be reversely converted into C (for **12**, from J *via* **TS6** and **TS4** with a barrier of 26.5 kcal mol⁻¹; for **13**, from I *via* **TS5** and **TS4** with a barrier of 31.6 kcal mol⁻¹), and then they enter the hydroboration catalytic cycle to afford **2a**, as observed in Scheme 3A-3 and A-4.

It is noted that ligand PPh₃ plays an important role in regulating the competition of σ -CAM and β -H elimination from the C–C bond cleavaged intermediate. As shown in Fig. 3, the free energy of the β -H elimination pathway *via* **TS4** is 5.8 kcal mol⁻¹ higher than that of the σ -CAM process through **TS2** (29.5 *versus* 23.7 kcal mol⁻¹). However, in the absence of PPh₃, the β -H elimination transition state **TS11** is even 0.1 kcal mol⁻¹ lower in energy than the corresponding σ -CAM transition state **TS10** (30.0 *versus* 30.1 kcal mol⁻¹, Fig. 4). As a result, the efficiency of the desired hydroboration was very poor with many side reactions (Table 1, entry 14). These results imply that the electron-rich nature of the PPh₃ ligand facilitates the σ -CAM process, inhibiting the formation of alkene byproducts.

Finally, we investigated the opposite regioselectivity of hydroboration of CPAs in two Ir catalytic systems (Table 1, entries 1 and 2). Previous computations by Yamaguchi and Yokogawa showed that the real catalyst is Ir(III) species when [Ir(cod)OMe]₂ and ligand ^tBu-Quinox are employed.¹⁶ As shown in Fig. 5A, our calculations indicate that the free energy of ringopening transition state TS12 at the distal C-C bond is 2.2 kcal mol⁻¹ lower than that of **TS13** at the proximal C-C bond. However, the hydroboration of CPAs using [Ir(cod)Cl]₂ and PPh₃ involves Ir(1) species as the catalyst (Fig. 5B). The free energy of **TS15** with proximal selectivity is 4.1 kcal mol⁻¹ lower than that of TS14 with distal selectivity. These computational results are in good agreement with the observed selectivities in experiments. We also applied the distortion/interaction analysis on these transition states to understand the origins of the regioselectivity.²⁵ In the Ir(III) system, the distortion, particularly



Fig. 5 DFT-computed energies and distortion/interaction analysis of C-C activation transition states. (A) $[Ir(cod)OMe]_2$ and tBu -Quinox as ligands. (B) $[Ir(cod)Cl]_2$ and PPh₃ as ligands. Energies are in kcal mol⁻¹.

the distortion of the catalyst, controls the regioselectivity. To activate the sterically more hindered proximal C-C bond via TS13, the trigonal bipyramidal Ir(III) catalyst requires a distortion energy of 11.4 kcal mol^{-1} , which is 3.0 kcal mol^{-1} higher than that in TS12 to break the less hindered distal C-C bond (11.4 versus 8.4 kcal mol⁻¹, Fig. 5A), leading to the distalselectivity. By contrast, the favorable interaction between 1a and the catalyst determines the regioselectivity in the Ir(1)system (Fig. 5B), which favors proximal-selective C-C bond activation significantly $(-50.5 \text{ kcal mol}^{-1} \text{ for TS15 versus})$ -39.6 kcal mol⁻¹ for **TS14**). The tetrahedral structure of the Ir(1) catalyst makes the Ir center sterically less crowded compared with the trigonal bipyramidal Ir(III) catalyst. This favors the formation of the thermodynamically more stable intermediate **O** (versus **N**) with proximal-selectivity in the Ir(I) system, while less stable intermediate L (versus M) with distal-selectivity is preferred in the case of the Ir(III) catalyst.

Conclusions

In summary, we developed the first proximal-selective C–C bond activation followed by hydroboration of CPAs with HBpin

Chemical Science

leading to interesting γ -amino boronates enabled by the rhodium catalyst. The key to the high chemoselectivity and regioselectivity is the selection of PPh₃ as the ligand. Significantly, experimental and computational mechanistic studies greatly provided insight into the reactivity and selectivity of this transformation. Further investigations of enantioselective variants and different types of C–C functionalizations of CPAs with tunable distal and proximal selectivity are currently underway in our laboratory.

Conflicts of interest

There are no conflicts to declare.

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