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Enantioselective Suzuki cross-coupling of 1,2-diboryl cyclopropanes†

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Herein, we describe the catalytic enantioselective cross-coupling of 1,2-bisboronic esters. Prior work on group specific cross coupling is limited to the use of geminal bis-boronates. This desymmetrization provides a novel approach to prepare enantioenriched cyclopropyl boronates with three contiguous stereocenters, that could be further derivatized through selective functionalization of the carbon-boron bond. Our results suggest that transmetallation, which is the enantiodetermining step, takes place with retention of stereochemistry at carbon.

Introduction

The Suzuki–Miyaura cross-coupling reaction is one of the main tools used in the pharmaceutical industry to forge carbon–carbon bonds.¹ A large variety of libraries of compounds are synthesized every year using this cross-coupling as a key point of diversification. In fact, a recent analysis showed that this transformation is the second most commonly used reaction in pharma, placed after amide coupling.² The robustness of the Suzuki cross-coupling has also made possible its use in automated settings³ and in the synthesis of DNA encoded libraries.⁴

As the pharmaceutical industry is shifting from compounds with a strong sp² character to libraries of compounds with increased three-dimensionality,⁵ the need to develop robust stereoselective Suzuki cross-couplings to provide compounds with an increased sp³ character becomes apparent.⁶ In this context, the development of enantioselective Suzuki–Miyaura cross-coupling reactions using prochiral bis-boronates remains largely unexplored.^{7,8} Morken⁹ and Hall¹⁰ reported the enantioselective Suzuki cross-coupling reaction of symmetric geminal bis-boronates to prepare enantiomerically enriched boronic esters (Fig. 1a). However, despite the elegant strategies developed to selectively functionalize 1,2-bis-boronates,^{11,12} the enantioselective desymmetrization of these species is still an unmet challenge (Fig. 1b). The degree of complexity in the

Following our interest in the functionalization of small rings¹³ and inspired by the relevance of cyclopropanes in synthetic methodology and pharmaceutical industry,¹⁴ we envisioned that symmetric bis-boryl cyclopropanes I could offer an ideal scenario to test this transformation (Fig. 1c). The products would be enantiomerically enriched cyclopropyl boronates,¹⁵ with three substituents in a hindered *cis* orientation and a handle for further stereospecific C–B functionalization.¹⁶

We realized from the outset that the proposed desymmetrization was a challenging transformation from the point of view of asymmetric catalysis (Fig. 2). Under the basic conditions needed for the Suzuki reaction, the bis-boronic ester **I** is likely to be in equilibrium with the racemic mixture of monoacids **II** and

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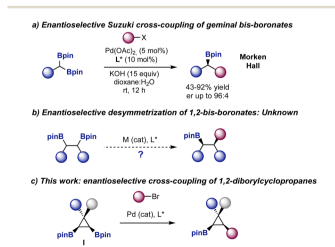


Fig. 1 Enantioselective Suzuki cross-coupling of prochiral bis-boronates.

products, relative to those prepared from geminal bisboronates, would be higher as two contiguous stereocenters are generated in the process.

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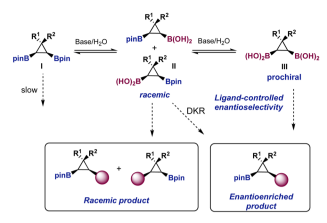


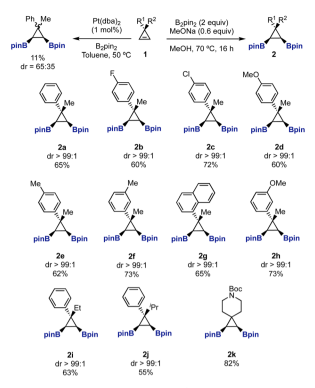
Fig. 2 Challenges involved in the proposed desymmetrization.

the prochiral bis-boronic acid **III**. If hydrolysis is a prerequisite for transmetallation, as it was proposed for geminal bis-boronates, ^{9,10} the chiral L*PdAr(X) complex formed after oxidative addition would be exposed to mixtures of **II** and **III**. Although a dynamic kinetic resolution (DKR) could not be discarded, the reaction of a L*PdAr(X) complex with racemic mixture **II** would likely provide racemic products, having a detrimental effect on the overall enantiocontrol. Therefore, we hypothesized that to maximize a potential ligand-controlled enantioselectivity the equilibriums in Fig. 2 should be displaced towards bis-boronic acid **III**.

Results and discussion

When we started our study, we realized that bis-boryl cyclopropanes such as 2 had not been prepared in the literature before. 17,18 We envisioned that diboration of readily available cyclopropenes 1 could provide easy access to these intermediates. Our first attempts using Pt-catalyzed diboration conditions 19,20 afforded a complex crude product from which we isolated a 65: 35 mixture of diborylated diastereomers 2a/2a' in 11% yield. Although we did not identify any other products, the dimerization of the cyclopropene under the reaction conditions could be a potential undesired reaction. 13b,21 Then, we turned our attention to the use of transition-metal free borylation conditions. We were pleased to find that heating cyclopropanes 1 in MeOH, in the presence of MeONa and B_2pin_2 , provided exclusively 1,2-syndiboryl cyclopropanes 2a-2k in good yields as single diastereomers (Scheme 1). 22,23

With a method to prepare diboryl cyclopropanes in hand, we started to explore the feasibility of the enantioselective Suzuki cross-coupling using cyclopropane **2a**, bromobenzene, and NaOH as the base, in the presence of 5 mol% of Pd(OAc)₂. Chiral bidentate ligands commonly used in palladium-catalyzed asymmetric transformations (not shown) consistently provided a racemic mixture of **3a** in variable yields.²⁴ A key observation was that in the absence of an added ligand, we observed exclusive formation of the protodeboronation product (Table 1, entry 1).²⁵ Indeed, we soon realized that the inhibition of this background reaction was one of the main challenges in this cross-coupling



Scheme 1 Base-promoted diboration of cyclopropenes.

reaction. We then moved to explore the reactivity of monodentate ligands, hoping that three-coordinate LPdAr(X) complexes would show enhanced selectivity.

After extensive experimentation, we found that TADDOL derived phosphoramidites (L₁-L₇, Table 1) showed encouraging results. Using bromobenzene, 5 mol% of Pd(OAc)2, NaOH (8 equiv.) and chiral ligand L₁, at 60 °C, cross-coupling product 3a was obtained in 66% yield with a promising enantiomeric ratio (er = 83:17, Table 1, entry 2). It is known that the addition of a fluoride source helps to increase the rate of the Suzuki crosscoupling reaction.26 In our case, the addition of KHF2 (Table 1, entry 3) had a positive effect on controlling the protodeboronation and, therefore, increasing the yield up to 78%. Using 6 equiv. of KHF2 the yield was further improved to 85% without reducing the enantiomeric ratio (er = 83:17). Less bulky groups in the para position of the aromatic rings of the ligands provided lower stereocontrol (L2-L3, Table 1, entries 5-6). Tuning of the substituents at nitrogen (L4-L5, Table 1, entries 7-8) and the acetal backbone (L₆-L₇, Table 1, entries 9-10) did not improve the results obtained with L₁. While at room temperature the cross-coupling did not occur, at 40 °C compound 3a was obtained with higher enantioselectivity (er = 86:14, Table 1, entry 11). Additionally, the use of 20 mol% of L1 had a beneficial effect on both the yield and the enantiomeric ratio (89%, er = 87:13, Table 1, entry 12). This is probably due to the inhibition of the background protodeboronation. According to this result, the use of 5 mol% of L₁ was detrimental to the yield and the enantioselectivity (Table 1, entry 13). Aryl iodides (Table 1, entry 14) afforded compound 3a with poorer results. Finally, entries 15-17 show that the use of a large excess of base was necessary. Using 1

Table 1 Optimization of the enantioselective Suzuki cross-coupling

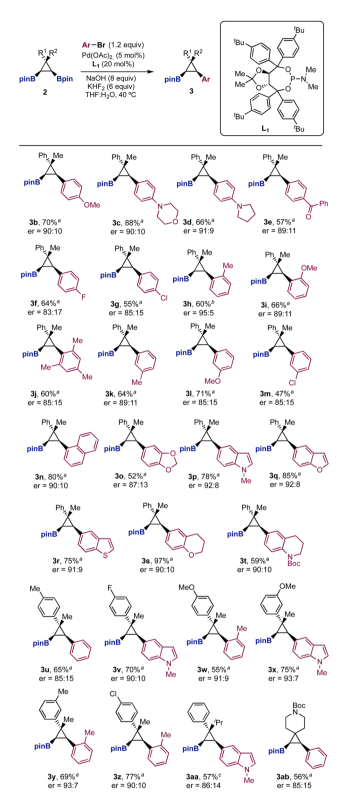
Entry	L	Equiv. NaOH	Equiv. KHF_2	T	Yield $3a^b$ (%)	er^c
1	_	8	_	60	_	_
2^a	L_1	8	_	60	66	83:17
3^a	L_1	8	3	60	78	82:18
4^a	$\mathbf{L_1}$	8	6	60	85	83:17
5 ^a	L_2	8	6	60	72	70:30
6 ^a	L_3	8	6	60	65	79:21
7^a	L_4	8	6	60	54	80:20
8 ^a	L_5	8	6	60	59	73:27
9^a	L_6	8	6	60	74	83:17
10^a	L_7	8	6	60	66	84:16
11^a	L_1	8	6	40	75	86:14
12^d	L_1	8	6	40	89	87:13
13^e	L_1	8	6	40	68	84:16
14^f	L_1	8	6	40	46	72:28
15^g	L_1	1	6	40	NR	_
16^g	L_1	4	6	40	Traces	_
17 ^g	$\mathbf{L_1}$	15	6	40	65	88:12

 a Reaction conditions: **2a** (0.1 mmol), PhBr (1.2 equiv.), Pd(OAc)₂ (5 mol%), L* (10 mol%), NaOH (8 equiv.), KHF₂ (0–6 equiv.), THF: H₂O (10:1, 0.1 M), and 16 h. b Yield calculated by 1 H NMR using an internal standard. c Enantiomeric ratio determined by chiral-phase HPLC. d 20 mol% of L₁ used. e 5 mol% L₁ used. f PhI was used instead of PhBr. g Reaction conditions [a] except for the equiv. of NaOH.

and 4 equiv. of NaOH we did not observe product formation. Increasing the amount of base to 15 equiv. maintained the level of stereocontrol (er = 88:12) but also increased the protodeboronation, lowering the yield of 3a to 65% (Table 1, entry 17). The use of other bases, Pd precatalysts or solvents were not found to be beneficial.²⁴

It should be highlighted that compound **3a** was obtained as a single *syn*-diastereomer. Assuming that the reductive elimination step occurs with retention, this result suggests that the transmetallation step takes place with retention of stereochemistry at carbon. This is in contrast with the results observed in the enantioselective cross-coupling with **1,1**-diboronates, in which the transmetallation seems to proceed with inversion.⁹

We next studied the structural scope of the enantioselective Suzuki cross-coupling reaction (Scheme 2). We were pleased to find that the reaction was quite general for different aryl



Scheme 2 Scope of the enantioselective desymmetrization. a Reaction conditions: 2 (0.2 mmol), PhBr (1.2 equiv.), Pd(OAc)₂ (5 mol%), L* (20 mol%), NaOH (8 equiv.), KHF₂ (6 equiv.), THF: H₂O (10:1, 0.1 M), 40 °C, and 16 h. b 15 equiv. of NaOH were used. c 8 equiv. of NaOH were used at 60 °C.

bromides (3b-3t) and cyclopropanes (3u-3ab). Importantly, in many cases the enantioselectivity was higher than that observed for bromobenzene. Aryl bromides with electron donating groups in the para and ortho positions provided the crosscoupling products (3b-3d and 3h-3l) in good yields and high enantiomeric ratios (up to 95:5 for 3h). Aryl bromides with electron-withdrawing substituents afforded arylated boryl cyclopropanes with similar stereoselectivites (3e-3g). Electrophiles with substituents in the meta position (3k-3m), sterically hindered aryl bromides (3j), as well as naphthalene (3n) and methylenedioxy derivatives (30) were well tolerated. Heterocycles such as indole (3p), benzofuran (3q), thiobenzofuran (3r), chromane (3s) and tetrahydroquinoline (3t) worked particularly well, providing cross-coupling products in good yields and enantiomeric ratios higher to that observed for bromobenzene. Finally, the method allowed structural modifications on the aryl and methyl groups of the diboryl cyclopropane framework (3u-**3ab**). In those cases, the cross-coupling products were prepared with comparable efficiencies to those shown with the Ph/Me substitution pattern.

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To gain insight into the boron species formed under the reaction conditions, we mixed pinacol ester 2a with (+)-pinanediol with KHF2 (6 equiv.) and NaOH (8 equiv.) at 40 °C, in the absence of Pd(OAc)₂ and bromobenzene (Scheme 3). We observed immediate disappearance of compound 2a by TLC and transesterified pinanediol derivative 21 was obtained in 86% yield. It is known that pinacol boronic esters form potassium trifluoroborate salts in the presence of KHF2, and that these salts are hydrolyzed to boronic acids27 in the basic aqueous media used in Suzuki cross-coupling reactions. Therefore, the experiment in Scheme 3 supports the in situ formation of bis-boronic acids but does not rule out the formation of transient trifluoroborate salts that could also participate in the equilibriums shown in Fig. 2. According to our results, the use of KHF2 has a positive effect on controlling the protodeboronation without compromising the enantioselectivity. Although the exact role of KHF2 is not clear at this point, the in situ formation of trifluoroborate salts could provide a slower release of the bis-boronic acid28 and fluoride26 under the basic aqueous conditions, minimizing the protodeboronation and increasing the rate of the cross-coupling. Bis-ester 21 did not react under the optimized conditions. Since pinanediol boronic esters are more robust towards hydrolysis, this result further supports the

a)

Ph Me
NaOH (8 equiv)

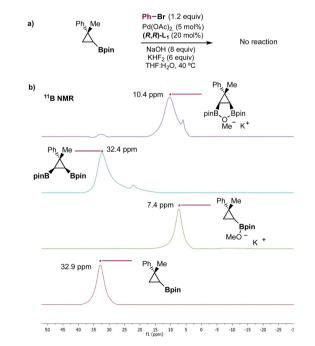
Ph Me

Ph Br (1.2 equiv)
Pd(OAc)₂ (5 mol%)
(R,R)-1- (20 mol%)
NaOH (8 equiv)

Scheme 3 Support for the in situ hydrolysis of diboronic ester 2a.

participation of a bis-boronic acid \mathbf{H} (Fig. 2) as an intermediate in the reaction.

Another question that emerged from our study was the role played by the second boryl moiety. Morken has reported that the presence of a vicinal pinacol boronic ester may have an activating effect towards transmetalation in Suzuki cross-coupling reactions.12g More recently, Morken has also shown that terminal 1,2diboronic esters, in the presence of potassium methoxide, form a 5-membered heterocycle with a single oxygen atom bridging both boron atoms.29 This chelated cyclic ate complex seemed to play a key role in the transmetallation with a copper complex. We were intrigued to compare the behavior of our 1,2-bisboryl cyclopropanes with that observed by Morken with terminal 1,2-bisboronates. We checked the reactivity of a monoborylated derivative (Scheme 4a) under the optimized reaction conditions and no product was observed after 16 hours. This result indicates that the presence of the adjacent boryl moiety is necessary for the transmetallation to take place. Additionally, we studied by 11B NMR spectroscopy the alkoxide complexation of mono- and diborylated cyclopropane (Scheme 4b). The treatment of the monoborylated derivative (32.9 ppm) with 1 equivalent of KOMe resulted in complete conversion to a compound with an upfield shifted resonance (7.4 ppm), which is in agreement with the formation of a sp3 hybridized borate complex. When we treated diborylated cyclopropane 2a (32.4 ppm) with 1 equivalent of KOMe, we observed an almost quantitative upfield shift of the ¹¹B NMR peak to 10.4 ppm. The smaller peak at 7.4 ppm corresponds to the borate complex of the residual protodeboronation product. Considering that we are using 0.5 equivalents of KOMe relative to the total amount of boron, the signal at 10.4 ppm suggests a chelation similar to that proposed by Morken for acyclic terminal 1,2-diboronates. Although the conditions used for the ¹¹B NMR



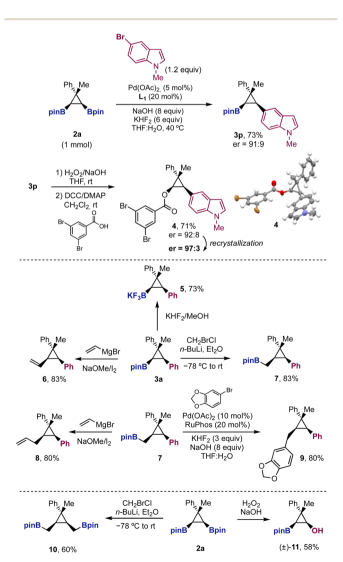
Scheme 4 Role of the vicinal borvl moietv.

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experiments (THF and KOMe) are not those used in the Suzuki cross coupling reaction (THF: H_2O , NaOH, and KHF₂), these results suggest that chelated cyclic ate complexes could potentially participate in our catalytic cycle. These observations could open the door to the design of further desymmetrizations.

The reaction was scaled up to 1 mmol with diboryl cyclopropane 2a and 5-bromoindole, affording cyclopropane 3p in similar yield and enantioselectivity to those observed before (Scheme 5). From boronate 3p, we performed the oxidation of the C-B bond followed by benzoylation to obtain benzoate 4. The recrystallization of 4 provided crystals of high enantiopurity (er = 97:3), which allowed us to assign the absolute configuration of the products in Table 1 and Scheme 2.³⁰

Finally, we showed the synthetic potential of the Suzuki cross-coupling products 3 with further stereospecific functionalization of the remaining boryl unit (Scheme 5). The transformation of the pinacol ester in the potassium trifluoroborate salt took place in high yield. Zweifel and Matteson homologations provided the desired products (6 and 7) in excellent yields, even though the



Scheme 5 Assignment of the absolute stereochemistry and C–B bond functionalizations.

boryl unit is placed in a crowded environment, surrounded by two substituents in *syn* relative distribution. The Matteson homologation product 7 is especially interesting as it still preserves the boryl unit for further transformations. A second sp²–sp³ Suzuki cross-coupling from 7 afforded trisubstituted cyclopropane 9 in 80% yield. Additionally, Zweifel homologation from 7 provided allylcyclopropane 8 in high yield. Finally, we briefly explored the synthetic potential of diboryl cyclopropanes 2 with reactions different than the Suzuki cross-coupling. Selective monooxidation of 2a afforded boryl cyclopropanol 11 as a single diastereomer. Moreover, double Matteson homologation provided symmetric bisboronate 10, which could be used to explore further desymmetrizations.

Conclusions

In summary, we have developed the first enantioselective desymmetrization of a 1,2-bisboronic ester. Prior work on group specific cross coupling has focused on geminal bis-boronates. Therefore, our study might pave the road to develop further desymmetrizations of prochiral 1,2-bis-boronates. This strategy allows for the preparation of highly functionalized enantioenriched boryl cyclopropanes with three stereocenters, one of them being quaternary. The products still preserve one boryl unit that can be used to design further stereospecific transformations. We believe that this methodology provides a useful tool for industrial and medicinal chemists to introduce functionalized three-membered rings into libraries of compounds.

Data availability

All the data supporting the findings of this study are available in the ESI.† Crystallographic data for compound 4 has been deposited in the Cambridge Crystallographic Data Centre under accession number CCDC 21910290.

Author contributions

J. T., M. V., R. F., A. V. and B. L. performed the experiments, wrote the ESI† and participated in discussions. M. T. conceived and directed the project and wrote the manuscript.

Conflicts of interest

There are no conflicts to declare.

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