

Hosomi–Sakurai Reaction

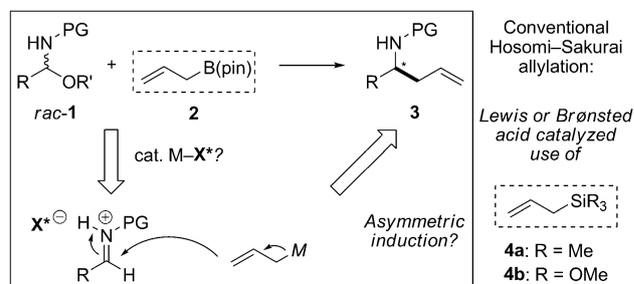
A Catalytic Asymmetric Borono Variant of Hosomi–Sakurai Reactions with N,O-Aminals**

Yi-Yong Huang, Ananya Chakrabarti, Naohide Morita, Uwe Schneider, and Shū Kobayashi*

Catalytic intermolecular cross-couplings between C_{sp}³ centers, such as O,O-acetals and N,O-aminals, and allyl species are challenging, but provide convenient access to various important substance classes such as homoallyl ethers and amines.^[1] For these transformations, Hosomi–Sakurai reactions using allyl silicon-based reagents are generally employed.^[2,3] These carbon–carbon bond formations proceed through either Lewis acid or Brønsted acid activation of the electrophile to generate a stabilized carbenium ion intermediate that can react with a silicon-based nucleophile. However, catalytic asymmetric Hosomi–Sakurai allylations of C_{sp}³ centers have proved to be challenging.^[2fj,4]

Allyl boronates are typically employed for additions to C_{sp}² centers,^[5] although a few C_{sp}²–C_{sp}³ cross-couplings have been reported.^[6,7] These nontoxic reagents are intrinsically less nucleophilic than silicon-based compounds and have been neglected in the context of allylation of C_{sp}³ centers. However, allyl boronates may offer significant advantages such as superior stability and unique reactivity and selectivity. During a project initially aimed at the catalytic activation of allyl boronates for selective C–C coupling with more complex electrophiles, we observed a peculiar reactivity with C_{sp}³ intermediates such as N,O-aminals. These electrophiles are abundant in natural products^[8a] and play an important role in organic synthesis.^[2f,8b–e] Thus, the development of catalytic asymmetric carbon–carbon bond formations with N,O-aminals is worthwhile.^[1] Intrigued by the unexpected reactivity, we started more detailed investigations with a view towards asymmetric catalysis. We report herein an approach to address the challenge of catalytic asymmetric Hosomi–Sakurai reactions involving C_{sp}³ centers by employing boronates instead of silicon-based reagents.

In an initial screen of various Lewis and Brønsted acids for the reaction of N,O-aminal *rac*-**1a** with allyl boronate **2** indium(I) triflate^[6a] was identified as the best catalyst for the formation of homoallyl amide *rac*-**3a** (Scheme 1; R = phenyl, PG = benzoyl, R' = methyl).^[9] In contrast, the corresponding



Scheme 1. Asymmetric borono variant of Hosomi–Sakurai reactions? B(pin) = pinacoloboron, PG = protecting group.

Hosomi–Sakurai allylations with silicon-based reagents **4a** and **4b** barely proceeded,^[9] which stands in sharp contrast to our earlier study.^[6a] The substantially higher reactivity of **2** over **4** under mildly Lewis acidic conditions constitutes a prerequisite for asymmetric catalysis. We postulated a dual catalytic activation^[6a] of *rac*-**1a** and **2** to generate iminium ion and allyl indium(I) intermediates (Scheme 1), thus we screened potential indium(I) catalysts bearing chiral counteranions^[9] rather than chiral ligands.^[10] In these experiments the combination of indium(I) chloride and chiral silver binol phosphate (*R*)-**5a**-Ag^[11] was found to be the most promising chiral catalyst system for the formation of product (*R*)-**3a** (e.r. = 88:12).^[9] Also, allyl silane **4a** proved to be substantially less effective than allyl boronate **2** in terms of both reactivity and selectivity.^[9] Thus, these results demonstrate the viability of our originally envisaged asymmetric concept, in which we proposed the use of boronates instead of silanes. At this stage, we explain the success of **2** based on its higher propensity to undergo transmetalation, thereby forming a more reactive allyl indium species.

Next, we further optimized the reaction conditions (Table 1). A screen of silver binol phosphates identified (*R*)-**5b**-Ag as the best chiral source (Table 1, entries 1–5). The use of an apolar cosolvent (cyclopentylmethyl ether) and a slight excess of the chiral silver salt further improved the asymmetric induction even at a lower catalyst loading (Table 1, entries 6–8). We then conducted several control experiments. In the absence of indium(I) chloride, (*R*)-**5b**-Ag displayed low reactivity and low asymmetric induction (Table 1, entry 10). The use of chiral Brønsted acid (*R*)-**5b**-H, which may be generated in situ under the present catalysis conditions, did not lead to any reaction (Table 1, entry 11).^[4g,8c] The combination of indium(I) chloride and (*R*)-**5b**-H provided low asymmetric induction (Table 1, entry 12), thereby demonstrating that an achiral metal salt and a chiral Brønsted

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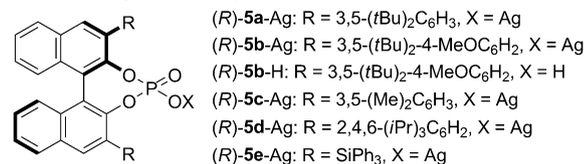
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Table 1: Optimization study and control experiments .

Entry	InCl [mol %]	5 [mol %]	Cosolvent	Yield ^[a] [%]	e.r. ^[b]
1 ^[c]	10	(<i>R</i>)-5 a-Ag (10)	–	81	88:12
2 ^[c]	10	(<i>R</i>)-5 b-Ag (10)	–	96	94.5:5.5
3 ^[c]	10	(<i>R</i>)-5 c-Ag (10)	–	90	44.5:55.5
4 ^[c]	10	(<i>R</i>)-5 d-Ag (10)	–	91	49.5:50.5
5 ^[c]	10	(<i>R</i>)-5 e-Ag (10)	–	94	49.5:50.5
6 ^[c]	10	(<i>R</i>)-5 b-Ag (10)	CPME	96	95.5:4.5
7 ^[c]	10	(<i>R</i>)-5 b-Ag (13)	CPME	98	98.5:1.5
8 ^[c,d]	5	(<i>R</i>)-5 b-Ag (6.5)	CPME	96	97.5:2.5
9 ^[d]	5	–	CPME	1	–
10 ^[d]	–	(<i>R</i>)-5 b-Ag (6.5)	CPME	5	57:43
11 ^[d]	–	(<i>R</i>)-5 b-H (6.5)	CPME	NR ^[e]	–
12 ^[c,d]	5	(<i>R</i>)-5 b-H (6.5)	CPME	88	62.5:37.5

[a] Yields of isolated (*R*)-3a after purification by preparative TLC on silica gel. [b] Enantiomeric ratios were determined by HPLC on a chiral stationary phase. [c] The chiral catalyst was performed in toluene at RT. [d] Reaction time: 18 h. [e] NR=no reaction (detected by ¹H NMR spectroscopy). CPME=Cyclopentyl methyl ether.



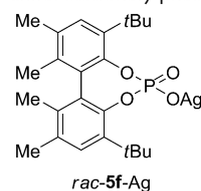
acid are ineffective.^[12,13] Importantly, we confirmed that redox disproportionation of indium(I), which would generate indium(0) and indium(III) in situ,^[14] did not occur in the present catalysis.^[15] Thus, the combination of indium(I) chloride and (*R*)-5b-Ag was shown to be crucial for the highly enantioselective formation of homoallyl amide (*R*)-3a under mild reaction conditions. The results of our control experiments (Table 1, entries 9–12)^[13,15] suggest the in situ generation of a chiral low-oxidation-state indium species as the active catalyst.

Next, we carried out a control experiment to investigate the reaction mechanism (Table 2). We employed the optically enriched aminal (*R*)-1a (e.r. = > 99.9:0.1) and allyl boronate 2 under standard reaction conditions using indium(I) chloride (10 mol %) combined with racemic silver phosphate *rac*-5f-Ag (13 mol %) as the catalyst system. This experiment was carefully analyzed over time by determining yields and enantiomeric ratios for both the generated product 3a and the recovered substrate 1a (15–640 min). Although the starting aminal (*R*)-1a was optically pure, the product 3a proved to be racemic at all stages of the reaction. At the same time, the racemization of (*R*)-1a proceeded relatively slowly under mildly Lewis acidic conditions. For example, we isolated product 3a in 15 % yield as a racemate (e.r. = 50:50) after a reaction time of 60 min, while substrate (*R*)-1a was recovered in 82 % yield with high optical purity (e.r. = 91:9). These results strongly indicate an iminium ion intermediate for this reaction (S_N1 pathway). In turn, these data

Table 2: Mechanistic control experiment.

t [min]	Yield [%] ^[a]	e.r. ^[b]	Yield [%] ^[a]	e.r. ^[b]
15	1	50:50	95	97.5:2.5
60	15	50:50	82	91:9
120	28	50:50	67	80:20
180	39	50:50	57	68:32
300	52	50:50	43	56.5:43.5
480	80	50:50	16	50:50
640	93	50:50	5	50:50

[a] Yields of isolated *rac*-3a and 1a after purification by preparative TLC on silica gel. [b] Enantiomeric ratios were determined by HPLC on a chiral stationary phase. Bz = benzoyl.



provide proof that the catalytic asymmetric C–C bond formation (see, Table 1) proceeds by the postulated S_N1 mechanism with an iminium ion species as a key intermediate, thus confirming the critical role of the chiral counteranion (see, Scheme 1).^[10,16,17] Overall, the present C–C bond-forming method relies on the generation of a chirally modified electrophile (acyclic transition state), and represents therefore an orthogonal approach compared with our related earlier study, in which we proposed a chirally modified nucleophile as a key intermediate (cyclic transition state).^[10]

We then examined the scope of this catalytic asymmetric transformation (Table 3). Under the optimized reaction conditions the reactions between substituted aromatic or heteroaromatic aminals *rac*-1a–k and allyl boronate 2 proceeded smoothly to provide the desired products (*R*)-3a–k with high asymmetric induction (Table 3, entries 1–12). In addition, even the challenging aliphatic aminals *rac*-1l and *rac*-1m proved to be good substrates for this new catalytic asymmetric method. Product (*R*)-3l was formed with excellent asymmetric induction (Table 3, entry 13), which demonstrates a dramatic improvement compared with our earlier related study.^[10] Product (*S*)-3m was obtained in high yield albeit with lower asymmetric induction (Table 3, entry 14). Overall, we consider these results remarkable as the levels of asymmetric induction exceed^[4c,e,g] or equal^[18b,c] even those of the corresponding catalytic asymmetric allylations of unactivated aldimines (C_{sp²} centers) with boronate 4 or silicon-based reagents 2.^[18]

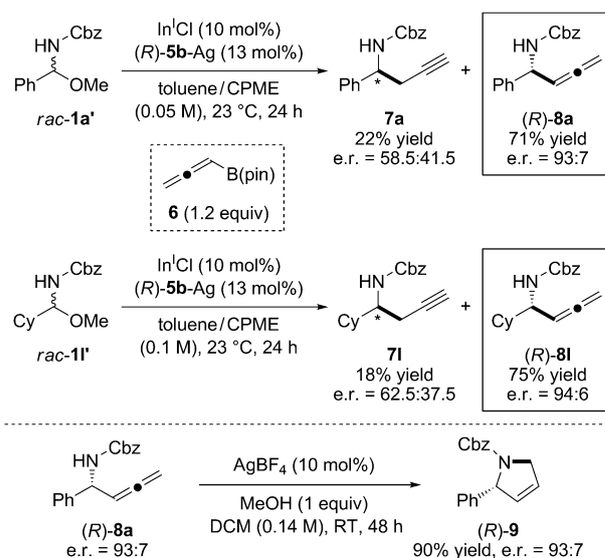
In addition, we were pleased to find that the novel chiral catalyst system was applicable to asymmetric allenylation

Table 3: Scope for the catalytic asymmetric N,O-aminal allylation.

Entry ^[a]	Product (R)-3	Yield [%] ^[b]	e.r. ^[c]
1		96	97.5:2.5
2		96 ^[d]	98.5:1.5 ^[d]
3		98	97.5:2.5
4		98	98:2
5		96	97:3
6		99	98:2
7		94	97:3
8		96	96.5:3.5
9		98	95:5
10		99	97.5:2.5
11		98	95:5
12		99	97.5:2.5
13		96	98:2
14		88	86:14

[a] Reaction conditions: *rac*-**1a–m** (1 equiv), InCl (5 mol%), (*R*)-**5b-Ag** (6.5 mol%), **2** (1.2 equiv), toluene/CPME (0.2 M), 23 °C, 18 h. [b] Yields of isolated (*R*)-**3a–l** and (*S*)-**3m** after purification by preparative TLC on silica gel. [c] Enantiomeric ratios were determined by HPLC on a chiral stationary phase. [d] Modified reaction conditions: InCl (10 mol%), (*R*)-**5b-Ag** (13 mol%), 12 h.

(Scheme 2). The reactions of aromatic and aliphatic aminals *rac*-**1a'** and *rac*-**1l'** with allenyl boronate **6** afforded the homoallenyl carbamates (*R*)-**8a** and (*R*)-**8l** in 71% and 75% yields, respectively, with high asymmetric induction (e.r. = 93:7 and 94:6, respectively). The homopropargyl carbamates **7a** and **7l** were obtained as the minor regioisomers, which were separated from (*R*)-**8a** and (*R*)-**8l** by chromatography. The observed regioselectivity is unprecedented for the use of allenyl boronate **6** in asymmetric catalysis. Thus, our work is clearly distinct from related studies.^[6a,19] The utility of highly functionalized compounds of type **8** was demonstrated by a catalytic intramolecular hydroamination with (*R*)-**8a**^[20] to generate azaheterocycle (*R*)-**9**^[21] (Scheme 2). This *5-endo-trig* cyclization occurred smoothly without loss of optical purity (e.r. = 93:7), and constitutes a straightforward method to access optically enriched 2-substituted 2,5-dihydropyrroles.


Scheme 2. Catalytic asymmetric allenylation and subsequent cyclization. Cbz = benzyloxycarbonyl, Cy = Cyclohexyl.

This report features several notable characteristics: 1) Under mildly Lewis acidic conditions, boronates proved to be dramatically more reactive and selective than classic silicon-based reagents. 2) The described transformations represent the first highly enantioselective Hosomi–Sakurai reactions with C_{sp^3} centers.^[2f,j] 3) This study also constitutes the first main-group-metal-catalyzed activation of allyl boronates for asymmetric C–C bond formation with C_{sp^3} centers.^[6] 4) Chiral Brønsted acid catalysis either with^[12] or without^[4g,8c] achiral metal salts proved to be inefficient. 5) In the context of asymmetric intermolecular C–C bond formation, the chemistry presented herein is a rare example not only of chiral-counteranion-directed metal catalysis,^[10,12] but also of dynamic kinetic resolution.^[22] Current investigations include elucidation of the catalyst structure and application to other reactions.

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