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Ligand-controlled diastereodivergent, enantio- and regioselective copper-catalyzed hydroxyalkylboration of 1,3-dienes with ketones†

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A copper-catalyzed three-component coupling of 1,3-dienes, bis(pinacolato)diboron, and ketones allows for the chemo-, regio-, diastereo- and enantioselective assembly of densely functionalized tertiary homoallylic alcohols. The relative configuration of the vicinal stereocenters is controlled by the chiral ligand employed. Subsequent transformations illustrate the versatility of these valuable chiral building blocks.

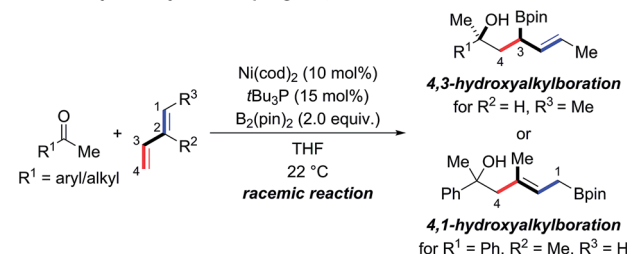
Introduction

The enantioselective synthesis of tertiary homoallylic alcohols¹ continues to attract attention as these are highly useful intermediates in complex molecule synthesis and for medicinal chemistry.² An established way to access that motif is by ketone allylation^{3–7} where enantiofacial discrimination and low reactivity are the key challenges compared to aldehydes as electrophiles.⁸ Many methods are based on preformed allylmetal reagents.^{3–6} An alternative to these nucleophiles is their *in situ* formation by hydrometalation of 1,3-dienes^{9,10} and allenes,¹⁰ and examples of transition-metal-catalyzed reductive couplings with ketones were recently achieved.^{10–12} A powerful variation of this approach is the borylmatalation of 1,3-dienes in the presence of a carbon electrophile.^{13–17} These and related stereoselective borylative coupling reactions of other π -systems form a carbon–boron and a carbon–carbon bond in a single operation.¹³ However, reactions involving ketones as electrophiles are scarce.^{14,17a,d–g} To the best of our knowledge, there are only three examples of the preparation of tertiary homoallylic alcohols by the borylative coupling strategy. Morken and co-workers reported a nickel-catalyzed three-component coupling of 1,3-dienes, bis(pinacolato)diboron, and ketones in racemic fashion (Scheme 1, top).¹⁴ The reaction outcome was dependent on the substitution pattern of the 1,3-diene; (*E*)-penta-1,3-diene converted into 4,3-hydroxyalkylboration products while isoprene (one example) afforded the 4,1-hydroxyalkylboration product. Starting from allenes as the precursor of the allylic nucleophiles, Hoveyda and co-workers realized enantioselective borylative couplings with carbonyl

compounds with *syn* selectivity but enantiocontrol was lower for ketones than for aldehydes (Scheme 1, middle).^{17a} Low enantioselectivity was found by Tian and Tao in an

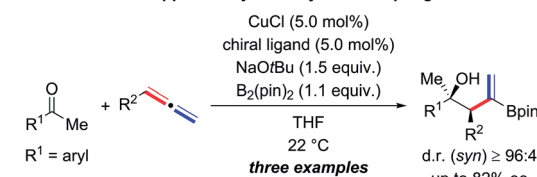
Morken (2011):

Nickel-catalyzed borylative coupling of 1,3-dienes and ketones



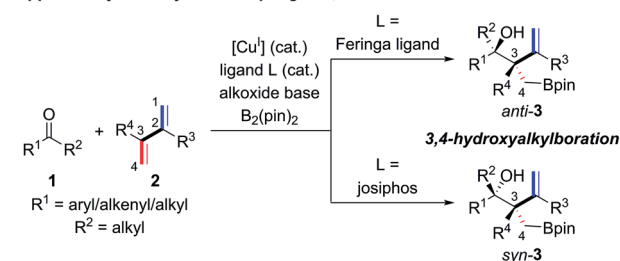
Hoveyda (2013):

Enantioselective copper-catalyzed borylative coupling of allenes and ketones



This work:

Diastereodivergent and enantioselective copper-catalyzed borylative coupling of 1,3-dienes and ketones

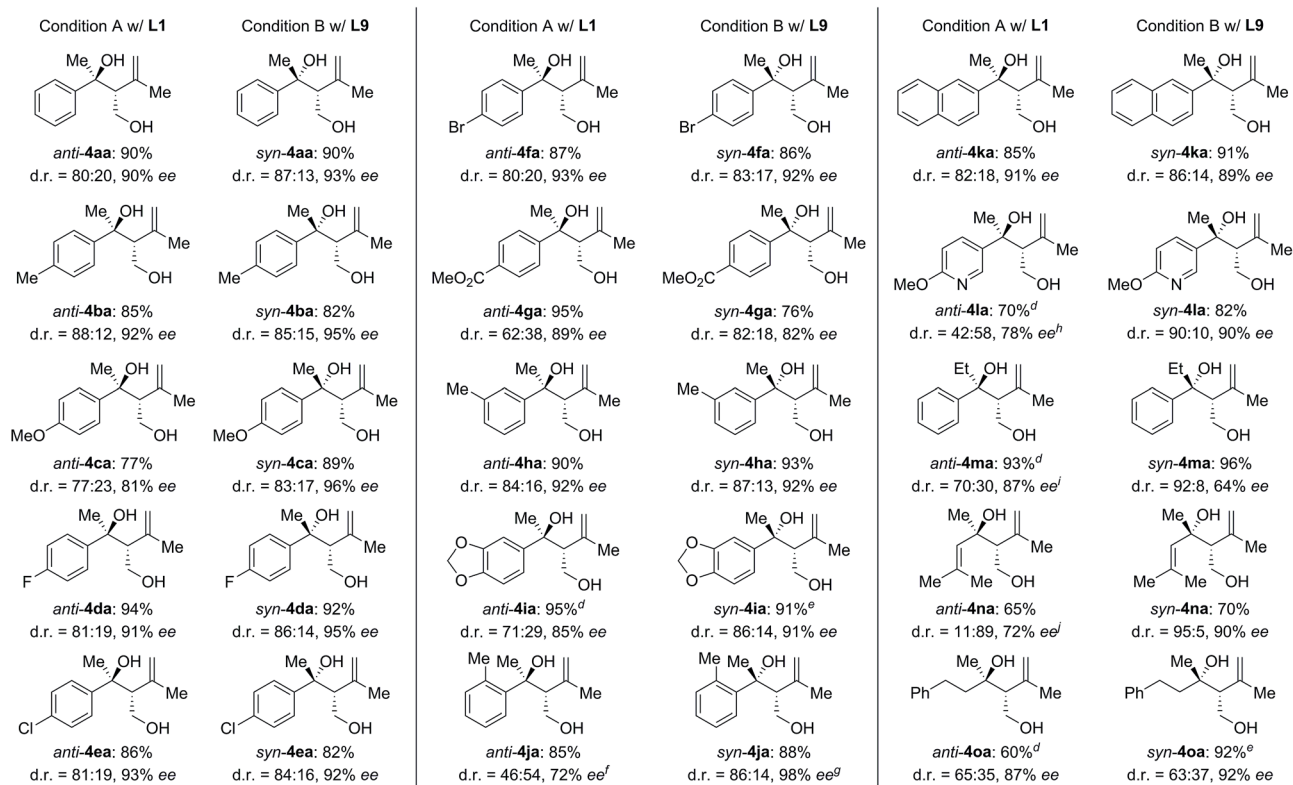
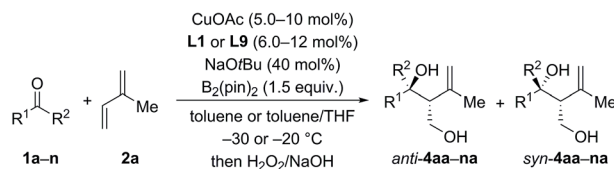


Scheme 1 Transition-metal-catalyzed intermolecular borylative coupling reactions of ketones for the construction of tertiary homoallylic alcohols. cod = cycloocta-1,5-diene, pin = pinacolato.

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Scheme 2 Scope I: variation the ketone.^{a-c} ^aCondition A: CuOAc (10 mol%), **L1** (12 mol%), NaOtBu (40 mol%), ketone **1** (0.20 mmol), isoprene (**2a**, 1.0 mmol), and B₂(pin)₂ (1.5 equiv.) in toluene (2 mL) at –30 °C. Condition B: CuOAc (5.0 mol%), **L9** (6.0 mol%), NaOtBu (40 mol%), ketone **1** (0.40 mmol), isoprene (**2a**, 2.0 mmol), and B₂(pin)₂ (1.5 equiv.) in toluene/THF = 8 : 2 (3.5 mL) at –20 °C. ^bYields are combined isolated material; diastereomers are usually separable by flash chromatography on silica gel. ^cThe enantiomeric excess of the major diastereomer was determined by HPLC analysis on chiral stationary phases. ^dCuOAc (15 mol%) and **L1** (18 mol%) were used. ^eCuOAc (10 mol%) and **L9** (12 mol%) were used. ^f*anti*-**4ja**: 29% ee. ^g*anti*-**4ja**: 80% ee. ^hee value of *anti*-**4la**. ⁱ*syn*-**4ma**: 78% ee. ^j*syn*-**4na**: 72% ee.

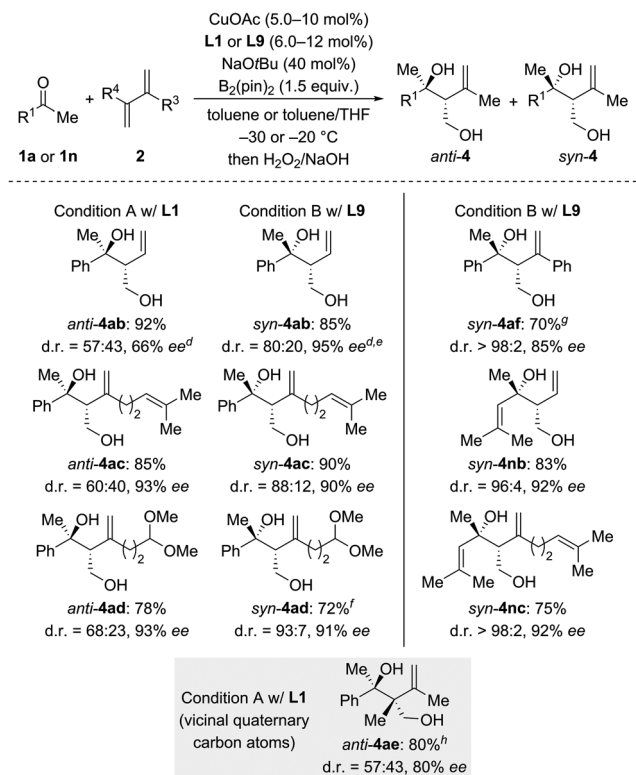
myrcene (**2c**), its functionalized derivative **2d**, and 2,3-dimethylbuta-1,3-diene (**2e**). Yields were generally good but stereoselectivities ranged from poor to good under Condition A. In contrast, good to excellent stereoselectivities were observed for these 1,3-dienes under Condition B, *e.g.*, d.r. = 96 : 4 and 92% ee for **1n** → *syn*-**4nb** and d.r. = 93 : 7 and 91% ee for **1a** → *syn*-**4ad**. In the case of 2-aryl-substituted 1,3-diene **1f**, diastereodivergency was not achieved. Subjecting **1f** to Condition A afforded *syn*-**4af** in low yield as a single *syn*-isomer (not shown). However, applying Condition B at −5 °C significantly improved the yield and furnished the *syn*-**4af** with d.r. > 98 : 2 and 85% ee.

To explore synthetic transformations of these tertiary homoallylic alcohols (Scheme 4), a scale-up synthesis of *syn-4aa* (1.0 mmol) under Condition B was done without any loss in efficiency and selectivity (see the ESI†). The primary alkyl borane generated by the multicomponent reaction was subjected to a Suzuki–Miyaura coupling to afford *syn-5* in 83%

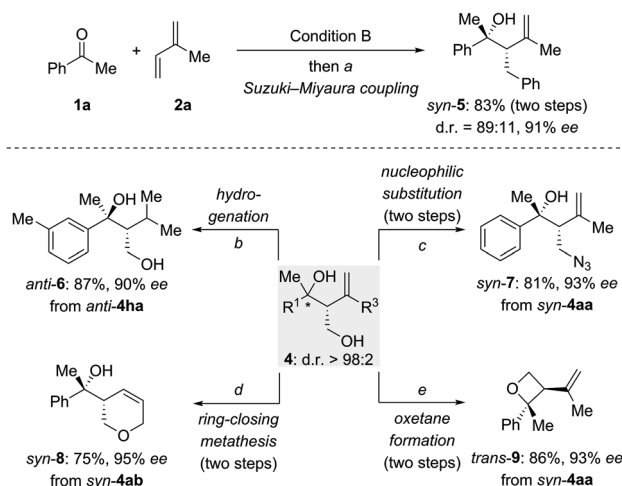
yield (Scheme 4, top). The versatility of the diol products **4** is illustrated for several transformations (Scheme 4, bottom). The 1,1-disubstituted double bond in *anti*-**4ha** was hydrogenated over Pd/C to produce *anti*-**6** in 87% yield. The hydroxy group in *syn*-**4aa** was replaced by an azide group through an S_N2 reaction of an intermediate mesylate with NaN₃ (*syn*-**4aa** → *syn*-**7**). Pyran *syn*-**8** was synthesized from *syn*-**4ab** by sequential alcohol allylation and ring-closing metathesis. Of note, a chemoselective tosylation of the primary alcohol in *syn*-**4aa** followed by a 4-*exo-tet* ring closure allowed for the construction of enantioenriched, trisubstituted oxetane *trans*-**9** in 86% yield.

Conclusion

In summary, we have developed an efficient copper-catalyzed diastereodivergent and enantioselective borylative coupling of 1,3-dienes and ketones. Using a Feringa-type ligand **L1**, the



Scheme 3 Scope II: variation of the 1,3-diene.^{a–c} For footnotes a–c, see Scheme 2. ^dThe absolute configuration was assigned by chemical correlation after separation of the diastereomers by flash chromatography on silica gel (see the ESI†). ^e*anti*-**4ab**: 84% ee. ^fCuOAc (8.0 mol%) and **L9** (10 mol%) were used. ^gRun at –5 °C with CuOAc (10 mol%), **L9** (12 mol%), NaOtBu (50 mol%), and B₂(pin)₂ (2.0 equiv.). ^hCuOAc (15 mol%) and **L1** (18 mol%) were used.



Scheme 4 Tertiary homoallylic alcohols as versatile building blocks. (a) PhBr (1.8 equiv.), Pd(OAc)₂ (5.0 mol%), RuPhos (10 mol%), KOtBu (3.0 equiv.), toluene/H₂O (10/1), 80 °C, 24 h; (b) Pd/C (10%), H₂ (1 atm), MeOH, rt, 26 h; (c) (i) MsCl (1.5 equiv.), Et₃N (1.5 equiv.), CH₂Cl₂, 0 °C to rt, 50 min; (ii) NaN₃ (2.0 equiv.), DMF/H₂O (10/1), 80 °C, 12 h; (d) (i) NaH (2.0 equiv.), allyl bromide (1.1 equiv.), THF, 0 °C to rt, 14 h; (ii) Hoveyda–Grubbs II (5.0 mol%), CH₂Cl₂, Δ, 12 h; (e) (i) TsCl (2.4 equiv.), pyridine, 0 °C to rt, 24 h; (ii) *n*BuLi (1.1 equiv.), −25 °C to rt, 15 h. Ms = methanesulfonyl.

reaction yielded *anti*-configured tertiary homoallylic alcohols while a switch to josiphos ligand **L9** resulted in *syn* selectivity (see the ESI† for a discussion of the reaction mechanism). This three-component coupling reaction represents a useful method for the preparation of stereochemically diverse tertiary alcohols bearing versatile alkenyl and boryl motifs from feedstock 1,3-dienes, ketones, and B₂(pin)₂. The synthetic utility of the reaction was showcased by several transformations.

Conflicts of interest

There are no conflicts to declare.

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