

Enantioselective Rhodium(III)-Catalyzed Markovnikov Hydroboration of Unactivated Terminal Alkenes

James R. Smith,[†] Beatrice S. L. Collins,[†] Matthew J. Hesse,[†] Mark A. Graham,[‡] Eddie L. Myers,[†] and Varinder K. Aggarwal^{*,†,‡}[†]School of Chemistry, University of Bristol, Cantock's Close, Bristol BS8 1TS, United Kingdom[‡]Pharmaceutical Technology and Development, AstraZeneca, Silk Road Business Park, Charter Way, Macclesfield SK10 2NA, United Kingdom

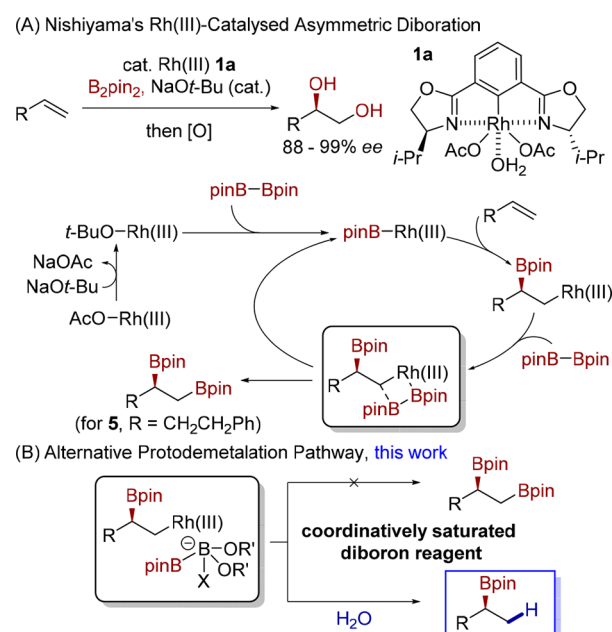
Supporting Information

ABSTRACT: We report the first enantioselective Rh-catalyzed Markovnikov hydroboration of unactivated terminal alkenes. Using a novel sp^2 – sp^3 hybridized diboron reagent and water as a proton source, a broad range of alkenes undergo hydroboration to provide secondary boronic esters with high regio- and enantiocontrol.

The H. C. Brown asymmetric hydroboration of alkenes, reported in 1961, holds an important position in the history of asymmetric synthesis as the first example of a chemical transformation in which high enantioselectivity was conferred by a small molecule.¹ Previously, high selectivity was the sole preserve of macromolecular structures, like enzymes. Although historically significant, and practical at the time, this method has been largely superseded by methods involving asymmetric metal catalysis, principally rhodium- and copper-catalyzed processes.² Certain classes of alkenes, e.g., styrenes,^{3,4} electronically activated alkenes^{5–8} or alkenes bearing directing groups,⁹ give high regio- and enantiocontrol, but archetypal aliphatic terminal alkenes have not succumbed to asymmetric hydroboration through either metal-catalyzed or noncatalyzed processes. Although aliphatic terminal alkenes usually give the linear alkylboronic ester,¹⁰ recently disclosed copper-catalyzed hydroboration processes, employing bulky phosphine or NHC ligands, give the branched Markovnikov product.^{11,12} The process, however, has yet to be rendered asymmetric. A general catalytic asymmetric method for the generation of secondary alkylboronic esters from the abundant feedstock of aliphatic terminal alkenes¹³ remains an unmet challenge, which is now addressed in this paper.

We sought a process for adding a metal–boron bond across an alkene, placing a bulky metal at the less hindered terminal carbon atom. We were attracted to Nishiyama's diboration reaction,^{14,15} which is postulated to proceed via a rhodium(III)–boryl species that undergoes insertion into the alkene, installing a secondary carbon–boron bond and generating a terminal rhodium(III)–alkyl species. The introduction of the second boron moiety then occurs through σ -bond metathesis (Scheme 1A). We surmised that if we could prevent the introduction of the second boron moiety, and protodemetalate instead, we could access the desired Markovnikov hydroboration products. However, introduction of a proton source

Scheme 1. Diverting Diboration into Hydroboration

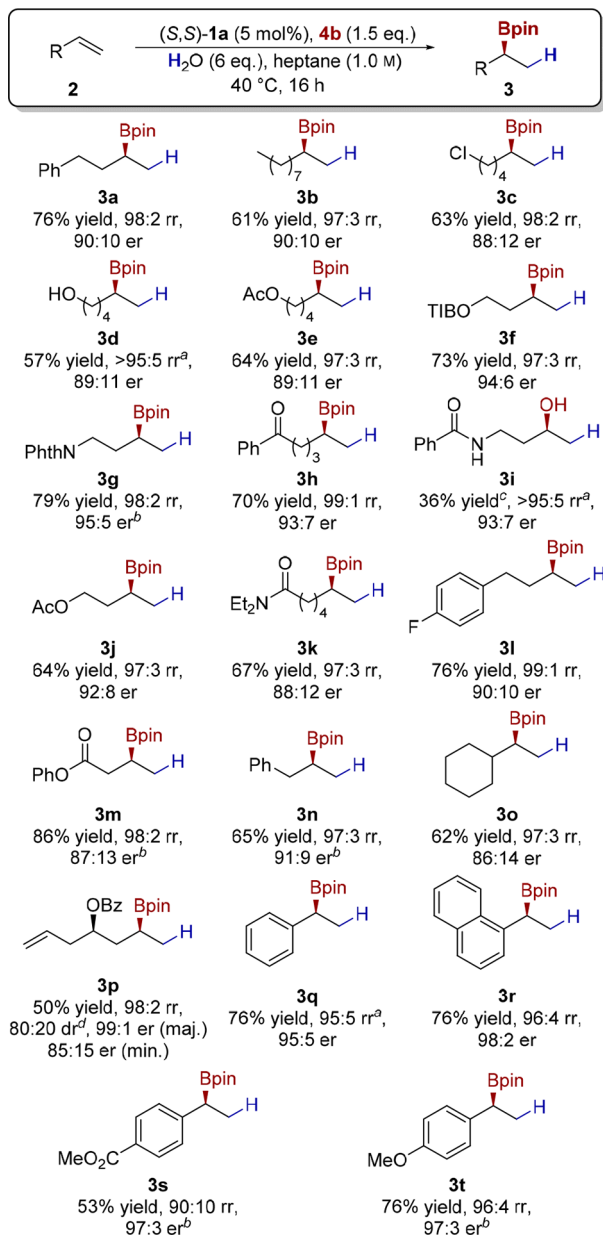


was not sufficient to favor a protodemetalation pathway: addition of isopropyl alcohol to the standard Nishiyama diboration conditions with 4-phenyl-1-butene (**2a**) as substrate did not lead to the desired hydroboration product and diboration product **5** was formed exclusively (86%, 98:2 er, Table 1, Entry 1). We reasoned that if one of the boron centers of the diboron reagent was coordinatively saturated, the terminating σ -bond metathesis would be inhibited, allowing for the desired protodemetalation (Scheme 1B). For this, we envisioned using a mixed sp^2 – sp^3 hybridized diboron species, in which one boron atom is bound to an amino diol ligand.¹⁶ The use of these “preactivated” diboron reagents would also enable the direct transfer of the sp^2 boron to the rhodium(III) center, obviating the need for external base.

We thus treated alkene **2a** with Santos's diboron reagent **4a**^{16b} in the presence of Nishiyama's [(*S,S*)-Rh(Phebox-*i*-Pr)(OAc)₂H₂O] catalyst **1a** and isopropyl alcohol (Table 1,

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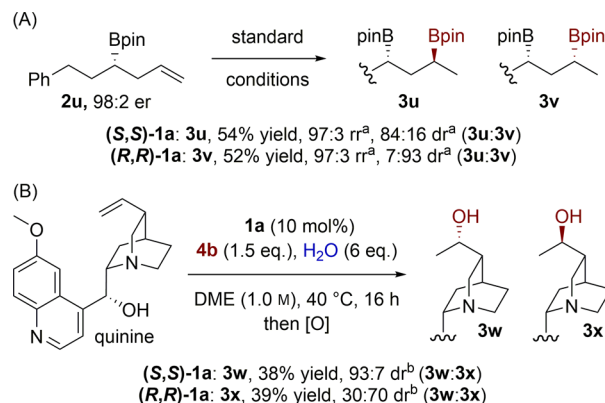
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Scheme 2. Scope of the Hydroboration Reaction^a

^aReactions were conducted with 0.38 mmol of **2**. Quoted yields are those of isolated product and are based on an average of values obtained from two experiments. Regioselectivity (rr) was determined by GC–MS analysis of the crude reaction mixtures, unless otherwise stated. Enantioselectivity (er) was determined by either chiral HPLC, SFC or GC analysis following oxidation (and in some cases further derivatization—see SI) of the isolated products (**3**), unless otherwise stated. ^bDetermined by ¹H NMR analysis of the crude reaction mixture. ^cDetermined by chiral SFC or HPLC analysis of the boronic ester (**3**). ^dAlcohol **3i** was obtained following an oxidative work up using H₂O₂/NaOH. ^eDetermined by LCMS analysis of the crude reaction mixture.

of the alkene into the rhodium–boron bond is the first irreversible step of the catalytic cycle, followed by rapid protodemetalation, or alternatively, reversible migratory insertion occurs before a rate-determining binding of a water molecule to the rhodium center, followed by rapid intramolecular protodemetalation. To differentiate between these two pathways, the reaction was conducted under the standard

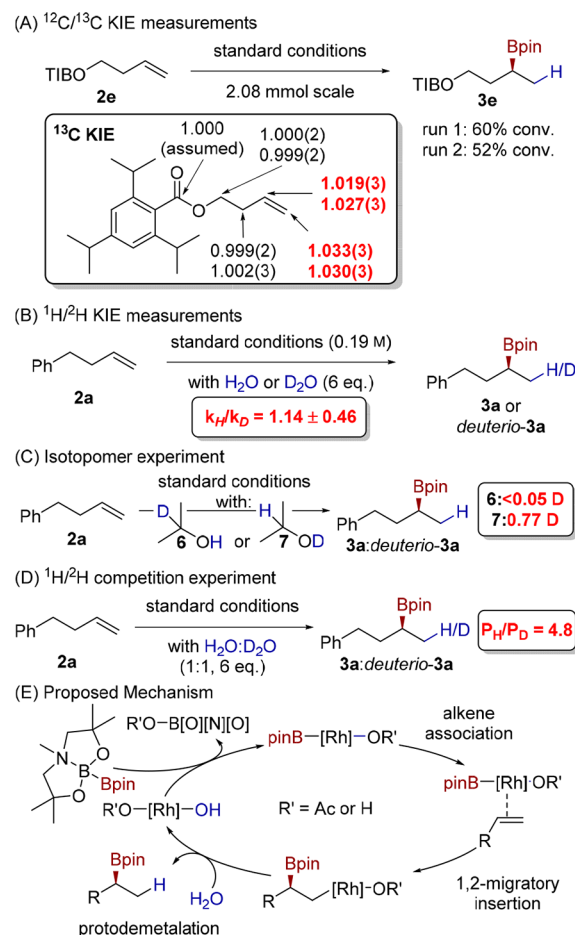
Scheme 3. Hydroboration of an Enantioenriched Substrate



^aDetermined by GCMS analysis of the crude reaction mixture.

^bDetermined by ¹³C NMR analysis of isolated material.

Scheme 4. Mechanistic Studies



reaction conditions using a 1:1 mixture of H₂O and D₂O (Scheme 4D). The product was obtained with 83% hydrogen incorporation, ruling out a nonreversible binding event of the water molecule prior to protodemetalation. We thus propose the mechanism outlined in Scheme 4E. The rhodium(III) catalyst undergoes transmetalation with the diboron reagent, which is activated by internal nitrogen coordination. Following alkene coordination, migratory insertion of the alkene into the rhodium–boron bond generates a primary rhodium–alkyl species with the boron moiety installed at the secondary

position. This is the first irreversible step of the catalytic cycle. Subsequent protodemetalation involving a water molecule then provides the secondary alkyl boronic ester product and regenerates the active rhodium(III) catalyst. We propose that the Markovnikov selectivity derives from regioselective migratory insertion and is controlled by a combination of steric and electronic factors. Isomerization and reduction side-products likely arise from the presence of small quantities of rhodium–hydride species (perhaps formed from competing β -hydride elimination of the β -boron rhodium–alkyl intermediate). The absolute configuration of the hydroboration products was the same as that observed in Nishiyama's diboration reaction. The slightly lower levels of enantioselectivity observed in the hydroboration reaction compared to the diboration reaction most likely results from the different ligand attached to the rhodium center (hydroxide or acetate versus *tert*-butoxide in Nishiyama's system).

In summary, we report the first asymmetric hydroboration of unactivated terminal alkenes. Secondary alkyl boronic esters are formed in good yields and high levels of enantioselectivity. Very high levels of regioselectivity are obtained without the need for directing groups or electronic biasing of the alkene substrates. Efforts to probe further the mechanism of this novel hydroboration method are currently underway.

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.7b05149.

X-ray crystallographic data for **4b** (CCDC reference number: 1550415) (CIF)

Experimental details and characterization data (PDF)

■ AUTHOR INFORMATION

Corresponding Author

*v.aggarwal@bristol.ac.uk

ORCID

Eddie L. Myers: 0000-0001-7742-4934

Varinder K. Aggarwal: 0000-0003-0344-6430

Notes

The authors declare no competing financial interest.

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