

Allylation of Nitrosobenzene with
Pinacol Allylboronates. A Regioselective
Complement to Peroxide Oxidation

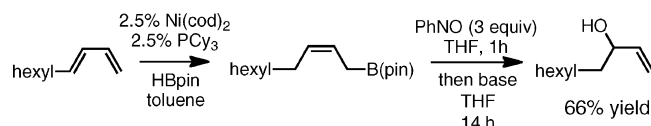
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ABSTRACT



Addition of nitrosobenzene to pinacol allylboronates leads to oxidation of the organoboron with concomitant rearrangement of the substrate alkene. This reaction appears to proceed by allylboration of the nitroso group in analogy to carbonyl and imine allylation reactions. Remarkably, the N–O bond is cleaved during the reaction such that simple alcohols are the final reaction product.

Because of the utility of allylboron reagents in organic synthesis, their preparation has been studied intensely.¹ Along these lines, recent efforts from our laboratory have focused on the development of the catalytic hydroboration of dienes² and the catalytic diboration of both allenes³ and dienes.⁴ These reactions convert simple hydrocarbon building blocks into substituted pinacolato allylboronates. These types of allylmetal reagents participate in a wide range of allylations with carbonyl and imine derivatives.⁵ However, the only other reactions that have been developed for these species are a narrow range of oxidation,⁶ cross-coupling,⁷ conjugate

addition,⁸ and homologation reactions.⁹ To expand the utility of allyl boronates in organic synthesis, we have begun to study other reactions that might apply to these reagents. Considering the isoelectronic relationship between the nitroso group and carbonyl groups, we were prompted to study the reaction between nitrosobenzene and allyl boronates. While a single example by Bubnov describes the reaction between nitrosobenzene and highly reactive triallylborane, a number of critical issues remain unaddressed.¹⁰ First, it is not clear whether the diminished electrophilicity of boronic esters

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(5) For a review of allylboration of carbonyl compounds, see: Hall, D. G. *Pure Appl. Chem.* **2008**, *80*, 913. See also: (b) Zhang, P.; Morken, J. P. *J. Am. Chem. Soc.* **2009**, *131*, 12550. For the allylboration of imines, see: (c) Sugiura, M.; Hirano, K.; Kobayashi, S. *Org. Synth.* **2006**, *83*, 170. (d) Sieber, J. D.; Morken, J. P. *J. Am. Chem. Soc.* **2006**, *128*, 74. (e) Elford, T. G.; Hall, D. G. *Tetrahedron Lett.* **2008**, *49*, 6995.

(6) Review of oxidation of organoboron compounds: Brown, H. C.; Snyder, C.; Rao, B. C. S.; Zweifel, G. *Tetrahedron* **1986**, *42*, 5505.

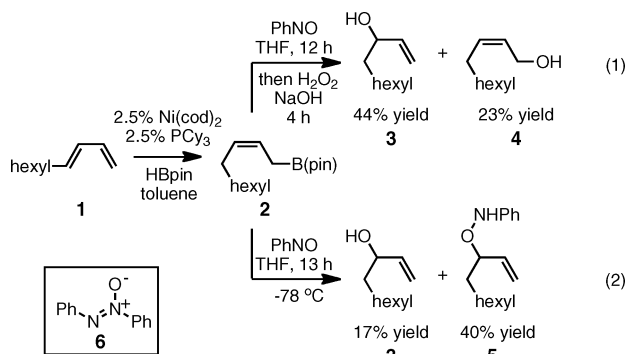
(7) Selected examples: (a) Nilsson, K.; Hallberg, A. *Acta Chem. Scand. B* **1987**, *41*, 569. (b) Kalinin, V. N.; Denisov, F. S.; Bubnov, Y. N. *Mendeleev Commun.* **1996**, 206. (c) Kotha, S.; Behera, M.; Shah, V. R. *Synlett* **2005**, *12*, 1877. (d) Yamamoto, Y.; Takada, S.; Miyaura, N. *Chem. Lett.* **2006**, *35*, 704. (e) Kotha, S.; Shah, V. R.; Mandal, K. *Adv. Synth. Catal.* **2007**, *349*, 1159. (f) Kotha, S.; Shah, V. R. *Eur. J. Org. Chem.* **2008**, *6*, 1054. (g) Gerbino, D. C.; Mandolesi, S. D.; Schmalz, H.; Podestá, J. C. *Eur. J. Org. Chem.* **2009**, *23*, 3964. (h) Flegeau, E. F.; Schneider, U.; Kobayashi, S. *Chem.—Eur. J.* **2009**, *15*, 12247.

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relative to boranes will impede the reaction. Second, with substituted allylic boronates, it is not readily apparent whether the reaction with nitrosobenzene will occur with allylic transposition as occurs with carbonyls or by coordination and subsequent 1,2 alkyl migration as occurs in the reaction between organoboranes and many reagents.¹¹ Lastly, although Bubnov has demonstrated that triallylborane and nitrosobenzene react with relatively nonselective formation of both N–O and C–O bonds, the issue of N- versus O-allylation with allylic boronates was uncertain.¹²

Two preliminary experiments revealed much about the reaction between allylboronates and nitrosobenzene. In the first (eq 1, Scheme 1), 1,3-decadiene (**1**) was subjected to

Scheme 1. Reaction of Nitrosobenzene with Allylboronate **2**

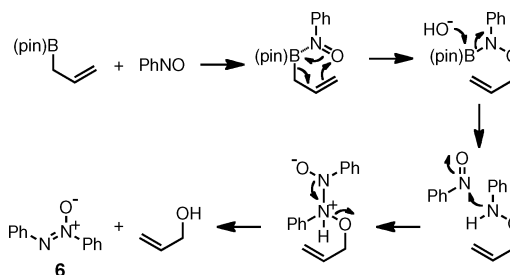


Ni-catalyzed 1,4-hydroboration, a reaction that delivers *cis* allyl boronate **2**.^{2a} After the reaction mixture was diluted with THF, it was treated with 1.05 equiv of nitrosobenzene. Subsequent oxidative treatment furnished allylic alcohols **3** and **4** in 44% and 23% yield, respectively. In the second experiment (eq 2, Scheme 1), intermediate allyl boronate **2** was treated with nitrosobenzene for 13 h at $-78\text{ }^{\circ}\text{C}$ prior to workup with brine and passage through a short silica gel plug. This reaction did not provide any of the terminal alcohol **4** but did produce internal allylic alcohol **3** and alkoxyamine **5**. The observation that allylic alcohol **4** is produced only when oxidative workup is employed suggests that **4** arises from unreacted **2**. The observation that regioisomers **3** and **5**, but not **4**, are produced in the absence of hydrogen peroxide suggests that the predominant reaction

pathway for allylic boronate **2** and nitrosobenzene occurs by addition of the allylboron to the oxygen atom and with allylic rearrangement.¹³

Considering the robustness of the N–O bond, it is somewhat surprising that N–O bond-cleaved compound **3** is produced in eq 2 (Scheme 1). A clue to its formation can be found in the fact that compound **5** is not isolated from eq 1 and that mass spectral analysis of the unpurified reaction mixture from eq 2 revealed the presence of a compound with an *m/z* ratio corresponding to compound **6** (Scheme 1). ¹H NMR analysis also reveals the presence of **6**. In line with these observations, it was surmised that a second molecule of nitrosobenzene and the Brønsted base might conspire to cleave the N–O bond of the initial allylation product in a fashion such as that depicted in Scheme 2. This type of

Scheme 2. Proposed Mechanism for N–O Bond Cleavage



reaction has been observed by Barbas and appears consistent with the reaction outcome.^{14,15}

In view of the mechanism depicted in Scheme 2, it might be anticipated that additional nitrosobenzene and alternate bases would provide an improved yield of allylation product **3**. As depicted in Table 1 (entry 1), addition of 3 equiv of

Table 1. Tandem Diene Hydroboration/Nitrosobenzene Oxidation

entry	base	% yield ^a
1	NaOH/H ₂ O ₂	69
2	NaOH	61
3	none	37
4	CsOH	55
5	LiOH	62
6	KOH	59
7	NH ₄ OH	67

^a Isolated yield of purified product.

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(12) For recent examples that use PhNO as an electrophile, see: (a) Momiyama, N.; Yamamoto, H. *J. Am. Chem. Soc.* **2003**, 125, 6038. (b) Brown, S. P.; Brochu, M. P.; Sinz, C. J.; MacMillan, D. W. C. *J. Am. Chem. Soc.* **2003**, 125, 10808. (c) Yamamoto, Y.; Momiyama, N.; Yamamoto, H. *J. Am. Chem. Soc.* **2004**, 126, 5962. (d) Momiyama, N.; Yamamoto, H. *J. Am. Chem. Soc.* **2005**, 127, 1080. (e) Simmons, B.; Walji, A. M.; MacMillan, D. W. C. *Angew. Chem., Int. Ed.* **2009**, 48, 4349.

Table 2. Substrate Scope for Tandem Diene Hydroboration/Oxidation

entry	diene	method A product/yield (%) ^a	method B product/yield (%) ^a
1	hexyl-CH=CH-CH=CH2	hexyl-CH(OH)-CH=CH2 66	hexyl-CH=CH-CH2OH 85
2	Cy-CH=CH-CH=CH2	Cy-CH(OH)-CH=CH2 62	Cy-CH=CH-CH2OH 81
3	TBDPSO-CH2-CH=CH-CH=CH2	TBDPSO-CH2-CH(OH)-CH=CH2 57	TBDPSO-CH2-CH=CH-CH2OH 56
4	Ph-CH=CH-CH=CH2	Ph-CH(OH)-CH=CH2 64	Ph-CH=CH-CH2OH 91
5	BnO-C(Me)2-CH=CH-CH=CH2	BnO-C(Me)2-CH(OH)-CH=CH2 44	BnO-C(Me)2-CH=CH-CH2OH 89
6	TBDPSO-CH2-CH2-CH=CH-CH=CH2	TBDPSO-CH2-CH2-CH(OH)-CH=CH2 58	TBDPSO-CH2-CH2-CH=CH-CH2OH 95
7	BnO-CH2-CH2-CH=CH-CH=CH2	BnO-CH2-CH2-CH(OH)-CH=CH2 63	BnO-CH2-CH2-CH=CH-CH2OH 93
8	hexyl-CH=CH-C(Me)=CH2	hexyl-CH(OH)-CH2-C(Me)=CH2 33	hexyl-CH=CH-C(Me)=CH2 93
9	pentyl-CH=CH-C(Me)=CH2	pentyl-CH(OH)-CH2-C(Me)=CH2 58	pentyl-CH=CH-C(Me)=CH2 81

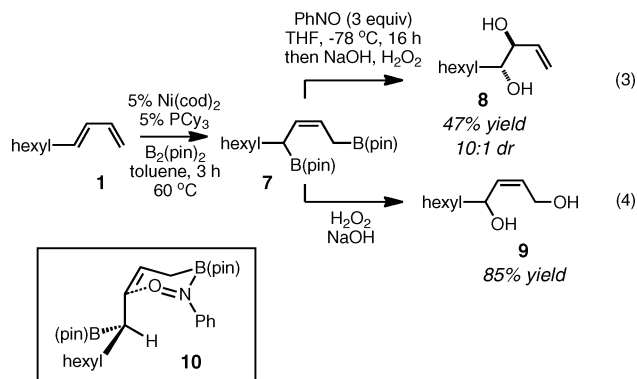
^a Isolated yield of purified product. Value is an average of two experiments.

nitrosobenzene, as opposed to 1 equiv (Scheme 1), results in a significantly enhanced yield of the secondary allylic alcohol **3**, and with oxidative workup, none of the terminal allylic alcohol **4** could be detected. According to the mechanism in Scheme 2, hydrogen peroxide is not required for the reaction, and the experiment in entry 2 (Table 1) bears this out. In the absence of base, however, the reaction yield

is significantly diminished (entry 3). With the requirement for addition of H₂O₂ apparently obviated, we examined workup under basic, nonoxidative conditions. As depicted in Table 1, a number of bases suffice for the nitrosobenzene-mediated oxidation of allylboronates. In all cases reasonable yields of secondary allylic alcohol **3** were isolated.

To study the generality of the nitrosobenzene-mediated allylboronate oxidation, a number of substrates were explored in the tandem hydroboration/oxidation sequence, and both nitrosobenzene/NH₄OH (method A) and NaOH/H₂O₂ (method B) oxidation procedures were examined. As depicted in Table 2, a number of different terminally substituted butadienes participate in the reaction and deliver moderate yields of the secondary allylic alcohol from method A. Whereas protected oxygen functional groups (entries 3, 5–7) and steric encumbrance (entries 2 and 5) at the diene terminus are tolerated, substitution at the 2 position of the diene leads to substantially diminished reactivity (entry 8). As noted in Table 2, oxidation with H₂O₂/NaOH consistently delivers the terminal allylic alcohol **4**. The fact that method B is efficient with all substrates in Table 2 suggests that the diminished reactivity observed with method A in entry 8 arises due to inefficient reaction of the intermediate allylic boronate with nitrosobenzene. Surprisingly, substitution at the 3 position is tolerated in the nitrosobenzene reaction, and the derived tertiary alcohol was isolated in good yield (entry 9).

With achiral allylboronates such as **2** the nitrosobenzene oxidation products **3** are racemic. However, stereogenicity in the allylic boronate may impact the oxidation if the reacting alkene is prochiral. To study the capacity for diastereoselection in the nitrosobenzene-mediated oxidation reaction, the chiral 1,4-diboryl-2-alkene **7** (Scheme 3) was

Scheme 3. Diastereoselection in the Oxidation of Chiral Allylboronate **6**

(13) The N-allylation product was not detected in these reactions. This observation is in contrast to that of Bubnov,¹⁰ who notes that a 2:1 mixture of O-allylation and N-allylation products is obtained from nitrosobenzene and triallylborane. In the reaction of tricrytylborane, Bubnov also notes allylic rearrangement.

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(15) Bubnov also notes N–O bond cleavage but claims a different mechanism that is light-promoted. We find that the reactions of allyl boronates proceed equally well in the dark.

generated by Ni-catalyzed diene diboration and subjected to the nitrosobenzene allylation reaction. Whereas reaction with nitrosobenzene at room temperature delivered diol **8** in 2.6:1 diastereoselection (data not shown), when nitrosobenzene was added to diboronate **7** at –78 °C and followed by oxidative workup, diol **8** was isolated in 10:1 *anti:syn* stereoselection and in moderate yield (eq 3, Scheme 3). For

comparison, the direct oxidation with hydrogen peroxide furnishes regioisomeric 1,4-diol **9** in 85% yield (eq 4).

The fact that the allylation of nitrosobenzene proceeds with allylic rearrangement suggests that a cyclic transition structure operates. This hypothesis was further supported by subjecting octylB(pin) to nitrosobenzene and NH_4OH ; less than 5% of 1-octanol could be detected. With these observations in mind, the stereochemical outcome of the reaction depicted in eq 3 may be rationalized by considering transition structure **10** as the predominating reaction pathway. Presumably, the enhanced basicity of nitrogen relative to oxygen results in N–B bonded complex **10** wherein nitrosobenzene has coordinated to the least hindered boronate. The stereogenic carbon atom in transition structure **10** is oriented in a manner where the electron-rich C–B bond is aligned with the π -system of the reacting alkene in a manner that should enhance π nucleophilicity. If the small hydrogen is placed inside with respect to the alkene to minimize A[1,3] strain, the model correctly predicts the stereochemical outcome of

the reaction. Similar features account for the stereochemistry of hydroboration of allylic silanes and boranes.¹⁶

In conclusion, treatment of simple allylboronates with nitrosobenzene and base can be considered a reliable strategy for oxidation with allylic rearrangement. Future studies will expand upon this reactivity pattern and will be reported in due course.

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Supporting Information Available: Complete experimental procedures and characterization data (^1H and ^{13}C NMR, IR, and mass spectrometry). This material is free of charge via the Internet at <http://pubs.acs.org>.

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