

Ni(II) Salts and 2-Propanol Effect Catalytic Reductive Coupling of Epoxides and Alkynes

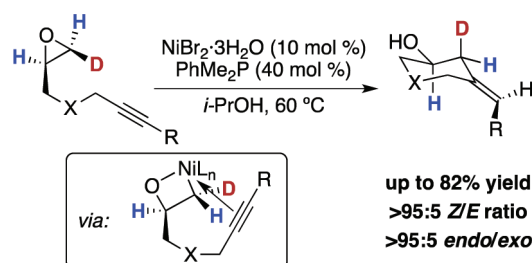
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ABSTRACT



A Ni-catalyzed reductive coupling of alkynes and epoxides using Ni(II) salts and simple alcohol reducing agents is described. Whereas previously reported conditions relied on Ni(cod)₂ and Et₃B, this system has several advantages including the use of air-stable and inexpensive Ni(II) precatalysts (e.g., NiBr₂·3H₂O) as the source of Ni(0) and simple alcohols (e.g., 2-propanol) as the reducing agent. Deuterium-labeling experiments are consistent with oxidative addition of an epoxide C–O bond that occurs with inversion of configuration.

The development of methods for transition-metal-catalyzed C–C bond formation using air-stable and inexpensive

reagents continues to offer a challenge to synthetic chemists. Recent improvements in the scope and utility of Ni-catalyzed C–C bond forming reactions present a significant advance toward this goal.^{1,2} The advent of methods employing Ni(II) precatalysts, for example, has obviated the need to handle air-sensitive Ni(0) sources.³ In many cases, however, Ni-catalyzed coupling protocols are limited by the inclusion of air- and moisture-sensitive reducing agents. Herein, we describe a practical method for the Ni-catalyzed reductive coupling of alkynes and epoxides using air-stable and inexpensive Ni(II) salts as the precatalysts and 2-propanol (*i*-PrOH) as the reducing agent.^{4,5} Mechanistic analysis of the reductive coupling process via

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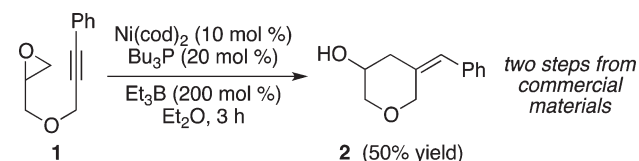
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deuterium-labeling studies provides evidence that oxidative addition of the epoxide C–O bond occurs with inversion of configuration.

Our laboratory has previously demonstrated that alkynes and epoxides undergo reductive coupling in the presence of $\text{Ni}(\text{cod})_2$, Bu_3P , and Et_3B .^{6–8} For example, Ni-catalyzed reductive coupling of epoxide **1** provided homoallylic alcohol **2** in 50% yield (Scheme 1). The combination of $\text{Ni}(\text{cod})_2$ and Bu_3P , both of which are exceedingly prone to air oxidation, was found to provide optimal results. In addition, the inclusion of Et_3B , a highly pyrophoric reducing agent, was critical for the desired transformation to proceed. We hypothesize that Et_3B may serve as a Lewis acid to facilitate oxidative addition of the epoxide, in addition to its standard role as a reducing agent.

Scheme 1. Previously Reported Conditions for the Ni-Catalyzed Reductive Coupling of Alkynes and Epoxides⁶



In light of our recent studies involving regioselective epoxide-opening cascades promoted by hydroxylic solvents,⁹ we envisioned that alcohol solvents might promote the Ni-catalyzed reductive coupling of alkynes and epoxides. In this setting, the role of the alcohol solvent would be twofold. First, oxidative addition of the epoxide C–O bond could be facilitated by hydrogen-bonding interactions between the epoxide oxygen and the alcohol solvent.^{9b,10} Second, the appropriate alcohol could serve as a reducing agent to provide the desired homoallylic alcohol product (**2**) and regenerate the active $\text{Ni}(0)$ catalyst (*vide infra*, Scheme 2).^{4,5,11}

In this manner, simple alcohols (e.g., *i*-PrOH) could serve as mild, air-stable, and inexpensive alternatives to Et_3B .

Table 1. Influence of Reaction Parameters on the Ni-Catalyzed Reductive Coupling of an Alkyne and Epoxide

1a (Z)-2a

entry	variation from standard conditions	yield (%) ^a	Z/E ^b
1	none ^c	82	90:10
2	$\text{Ni}(\text{cod})_2$ instead of $\text{NiBr}_2 \cdot 3\text{H}_2\text{O}$ ^d	82	95:5
3	$\text{NiBr}_2 \cdot \text{diglyme}$ instead of $\text{NiBr}_2 \cdot 3\text{H}_2\text{O}$	76	88:12
4	$\text{NiCl}_2 \cdot 6\text{H}_2\text{O}$ instead of $\text{NiBr}_2 \cdot 3\text{H}_2\text{O}$	74	89:11
5	5 mol % $\text{NiBr}_2 \cdot 3\text{H}_2\text{O}$ and 20 mol % PhMe_2P instead of standard catalyst/phosphine loading	75	95:5
7	THF instead of <i>i</i> -PrOH	<5 ^e	---
8	no $\text{NiBr}_2 \cdot 3\text{H}_2\text{O}$	<5 ^e	---
9	30 mol % PhMe_2P instead of 40 mol %	76	90:10
10	20 mol % PhMe_2P instead of 40 mol %	52	91:9
11	no PhMe_2P	<5 ^e	---

^a Isolated yield of the mixture of olefin isomers. ^b Determined by ¹H NMR spectroscopy of the crude reaction mixture. ^c Complete conversion of starting material observed within 3 h. ^d Phosphine loading was 20 mol % instead of 40 mol %. ^e Determined by ¹H NMR spectroscopy relative to mesitylene as an internal standard.

Our investigations revealed that *i*-PrOH was an effective reducing agent for the Ni-catalyzed reductive coupling of alkynes and epoxides. For example, employing *i*-PrOH as the solvent in the presence of PhMe_2P and either $\text{NiBr}_2 \cdot 3\text{H}_2\text{O}$ (Table 1, entry 1) or $\text{Ni}(\text{cod})_2$ (entry 2) provided high yields of the desired homoallylic alcohol **2a**.^{12–14} The use of air-stable $\text{Ni}(\text{II})$ precatalysts and reducing agents (i.e., $\text{NiBr}_2 \cdot 3\text{H}_2\text{O}$ and *i*-PrOH, as opposed to previous conditions employing $\text{Ni}(\text{cod})_2$ and Et_3B),⁶ allowed us to setup and perform the coupling reactions outside of a glovebox. Furthermore, all reactions listed in Table 1 could be performed without drying of the reagents or glassware. Of note, reactions employing inexpensive $\text{NiX}_2 \cdot \text{hydrate}$ salts (entries 1 and 4) were of comparable efficiency to reactions performed with

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(12) Details of substrate synthesis are provided as Supporting Information.

(13) Subjecting enantioenriched epoxide **1a** (88% ee) to the standard conditions for Ni-catalyzed reductive coupling provided homoallylic alcohol **2a** (88% ee) with complete preservation of optical purity. See Supporting Information for details.

(14) Preliminary investigation of the reaction conditions indicated that *i*-PrOH was the optimal alcohol reducing agent. See Supporting Information for additional details of reaction optimization including alcohol reducing agent, $\text{Ni}(\text{II})$ precatalyst, and temperature.

anhydrous $\text{NiBr}_2 \cdot \text{diglyme}$ (entry 3). Decreasing the catalyst loading to 5 mol % was tolerated and provided the desired homoallylic alcohol **2a** in only slightly lower yields (entry 5). No product formation was observed in the absence of *i*-PrOH, the nickel catalyst, or phosphine ligand (entries 7, 8, 11).

Increased phosphine loadings were found to provide the highest yields of homoallylic alcohol **2a** for reactions in which NiX_2 salts were employed as the catalyst. Previously, a 2:1 ($\text{R}_3\text{P}/\text{Ni}$) ratio was determined to be the optimal loading for reductive coupling reactions of epoxides and alkynes using $\text{Ni}(\text{cod})_2$ (Table 1, entry 2).⁶ When $\text{NiBr}_2 \cdot 3\text{H}_2\text{O}$ was employed as the catalyst, however, a phosphine loading of 20 mol % did not result in complete conversion of starting material (entry 10). Increasing the phosphine loading to 30 mol % led to complete consumption of the starting material and increased yields (entry 9). Finally, a phosphine loading of 40 mol % was found to provide the highest yield of homoallylic alcohol **2a** (entry 1). We attribute the requirement for excess phosphine ligand to its involvement in the reduction of the $\text{Ni}(\text{II})$ precatalyst to the active $\text{Ni}(0)$ species, which has been observed previously for both $\text{Ni}(\text{II})$ ¹⁵ and $\text{Pd}(\text{II})$ ¹⁶ precatalysts. Alternatively, excess phosphine may be necessary to suppress the formation of catalytically inactive $\text{Ni}(\text{O}i\text{-Pr})_2$ species.¹⁷

In all cases examined, reductive coupling of epoxide **1a** proceeded to provide the desired tetrahydropyran product (**2a**) as a single regioisomer with respect to both the epoxide and the alkyne. Stereospecific *cis* addition to the alkyne occurred to provide the trisubstituted olefin (**Z-2a**),⁶ however, isomerization of the olefin geometry was observed to a limited extent under the standard conditions for reductive coupling. The amount of undesired isomerization product (**E-2a**) could be minimized by judicious choice of catalyst identity and loading (entries 1, 2, and 5).

The mild conditions for Ni-catalyzed reductive coupling were found to effect product formation for substrates containing variation within the epoxide–alkyne tether. Both substituted cyclohexane (Table 2, entries 1 and 2) and piperidine (entry 3) derivatives could be prepared in this manner.¹²

Varying the substituent on the alkyne had a more significant impact on the efficiency of reductive coupling. Alkyl-substitution was tolerated and provided the desired homoallylic alcohol (**2e**) in yields comparable to those observed for phenyl-substituted alkynes (Table 2, entries 1–4). Conversely, subjecting alkynoate **1f** to the standard

conditions for reductive coupling did not result in formation of the desired product (entry 5).¹⁸ Terminal alkynes are a particularly difficult class of substrates in $\text{Ni}(0)$ -catalyzed coupling reactions, as this moiety often succumbs to deleterious cyclotrimerization.¹⁹ Under these conditions for reductive coupling, however, terminal alkyne **1g** underwent reductive coupling in 55% yield (entry 6).

Table 2. Investigation of Reaction Scope

entry	X	R	product	yield (%) ^a	Z/E ^b
1	CH ₂	Ph	2b	70	88:12
2	C(CO ₂ Me) ₂	Ph	2c	76	>95:5
3	NBn	Ph	2d	74	88:12
4	O	<i>n</i> -C ₆ H ₁₁	2e	75	>95:5
5	O	CO ₂ Me	2f	<5 ^c	---
6	O	H	2g	55 ^d	---

^a Isolated yield of the mixture of olefin isomers. ^b Determined by ¹H NMR spectroscopy of the crude reaction mixture. ^c Determined by ¹H NMR spectroscopy relative to mesitylene as an internal standard. ^d Dropwise addition of substrate via syringe pump.

The proposed mechanism for Ni-catalyzed reductive coupling of epoxides and alkynes involves initial oxidative addition of the epoxide C–O bond to form nickella(II)-oxetane **5** (Scheme 2).⁶ Previous reports have demonstrated that oxidative addition of epoxides with group 10 metals occurs with either inversion (Pd and Pt)^{20,21} or scrambling (Ni)²² of configuration, corresponding to $\text{S}_{\text{N}}2$

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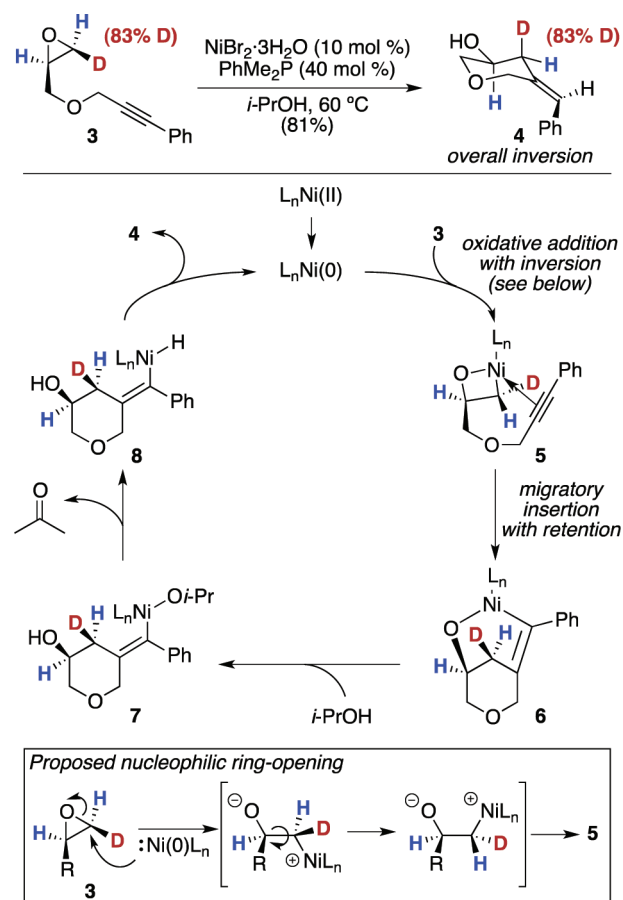
ring opening or homolytic C–O bond cleavage, respectively. To the best of our knowledge, there is no experimental evidence describing oxidative addition of an epoxide C–O bond with Ni(0) that occurs with inversion of configuration.²³

To gain insight into the oxidative-addition process, we prepared monodeuterated epoxide **3** and observed the relative configuration of the deuterated carbon center after subjection to the standard conditions for Ni-catalyzed reductive coupling (Scheme 2). Monodeuterated epoxide **3** was prepared with 83% deuterium incorporation at the terminal epoxide site bearing a cis relationship to the alkyne tether.¹² Subjecting epoxide **3** to the standard conditions for reductive coupling provided homoallylic alcohol **4** with complete inversion of configuration at the deuterated carbon center.²⁴ This result is consistent with inversion of configuration in the oxidative addition step to afford the nickella(II)oxetane intermediate (**5**), followed by migratory insertion that occurs with retention of configuration.²⁵ We propose that inversion of configuration in the oxidative addition step occurs via S_N2 attack by the nucleophilic Ni(0) species facilitated by hydrogen-bond activation of the epoxide.¹⁰ Importantly, identical results were obtained when epoxide **3** was subjected to the previously described conditions for reductive coupling (Scheme 1, PhMe₂P instead of Bu₃P) indicating that the active Ni(0) catalyst is responsible for the inversion process.

Following C–C bond formation, *i*-PrOH serves as a mild reducing agent to complete the catalytic cycle. Protonolysis of the Ni–O bond of strained bicyclic oxanickellacycle **6** results in formation of the L_nNi(O*i*-Pr) species (**7**) necessary for catalytic turnover. Subsequent β-H elimination occurs to liberate acetone, which is inert under the reaction conditions, followed by reductive elimination to afford homoallylic alcohol **4** and regenerate the active Ni(0) catalyst.

In summary, we have developed novel conditions for the Ni-catalyzed reductive coupling of alkynes and epoxides that employ air-stable and inexpensive reagents. These reaction conditions are significantly milder and more convenient than those previously used for similar reductive coupling reactions in which a combination of Ni(cod)₂ and Et₃B was necessary to effect C–C bond formation. Further investigation of the electrophile/π-nucleophile scope is warranted in an effort to improve the efficiency of previously developed methods and for the invention of novel reductive coupling procedures.

Scheme 2. Deuterium-Labeling Experiment and Proposed Mechanism for Ni-Catalyzed Reductive Coupling



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Supporting Information Available. Complete experimental procedures and characterization data of novel compounds. This material is available free of charge via the Internet at <http://pubs.acs.org>.