Nickel-catalyzed reductive cleavage of aryl alkyl ethers to arenes in absence of external reductant†

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The reductive cleavage of the C–O bonds of aryl ethers has great potential in organic synthesis. Although several catalysts that can promote the reductive cleavage of aryl ethers have been reported, all such systems require the use of an external reductant, e.g., hydrosilane or hydrogen. Here, we report the development of a new nickel-based catalytic system that can cleave the C–O bonds of ethers in the absence of an external reductant. The hydrogen atom required in this new reductive cleavage reaction is provided by the alkoxy group of the substrate, which serves as an internal reductant. The absence of an external reductant enables the unique chemoselectivity, i.e., the selective reduction of an alkoxy group over aikenes and ketones.

Introduction

The reductive deoxygenation of phenols and their derivatives is a fundamental synthetic transformation that has been widely used in various areas of organic synthesis; for example, reductive deoxygenation enables the use of oxygen-based functionalities as removable activating and directing groups in arene functionalization reactions. 1 Reductive deoxygenation is also a potentially useful strategy for the synthesis of deoxygenated analogs of phenol-based natural products.2 The reductive deoxygenation of phenols has traditionally been achieved by converting them to activated sulfonates such as triflates, followed by catalytic reduction of the activated species in the presence of a hydride donor.3 Recent progress in nickel-catalyzed4 C–O bond activation reactions5 has enabled the direct use of unactivated phenol derivatives. One of the most important achievements in this area has been the development of a catalytic process for the reduction of inert aryl ethers;4c,t this not only expands the substrate scope of reductive deoxygenation reactions, but also provides a platform for the development of new catalytic methods for the conversion of alkyl ether-containing biomass to alternative sources of fuel and chemicals, including the depolymerization of lignin.4a However, it is worth noting that these nickel-catalyzed reductive cleavage reactions of aryl ethers only proceed in the presence of a stoichiometric amount of an external reductant, e.g., hydrosilane or hydrogen.4b Here, we report the development of a nickel-catalyzed reductive cleavage reaction for the conversion of aryl alkyl ethers to the parent arenes in the absence of an external reductant.

Results and discussion

One possible mechanism for the reductive cleavage of aryl methyl ethers with an external hydride reagent involves oxidative addition of the C(aryl)–O bond to form nickel(i)-methoxide A, followed by reduction of an aryl–nickel bond (Scheme 1a). Although this mechanism seems plausible in view of the reports of nickel-catalyzed cross-couplings of aryl ethers with other nucleophiles, the key oxidative addition of a C(aryl)–OMe bond has never been verified experimentally. Gómez-Bengoa and Martin proposed a new mechanism for the nickel-catalyzed reductive cleavage of aryl ethers by hydrosilane (Scheme 1b). Spectroscopic and computational studies indicate that silylnickel(i) species can be generated under these conditions and mediate C(aryl)–OMe bond cleavage via migratory insertion of an arene to form benzylnickel species B, followed by concerted elimination of MeOSiR3. Polyaromatic compounds such as naphthyl ethers serve as good substrates, but anisole derivatives do not react unless they contain a suitable directing group; these results are consistent with the proposed mechanism involving a dearomatized intermediate B. It was envisaged that if oxidative addition of a C(aryl)–O bond occurs to form intermediate A, β-hydrogen elimination would lead to the formation of the same reductive cleavage product via C (Scheme 1c). This mechanistic variant therefore enables the reductive cleavage to proceed without an external reductant. Although the relevance of β-hydrogen elimination of nickel-alkoxide species in catalytic reductive cleavage of aryl ethers has been suggested before, catalytic systems that can effect such a reaction in the absence of an external reductant have not been reported.
To achieve the catalytic reaction shown in Scheme 1c, we evaluated the effect of ligands in the nickel-catalyzed reduction of 2-methoxynaphthalene (1a) to give naphthalene (2). As previously reported, the addition of PCy₃ and SiPr, which are the most effective ligands for the reductive deoxygenation of 1a with hydrosilane and hydrogen, respectively, did not lead to the formation of appreciable amounts of the reduced product 2 in the absence of an external reductant (Table 1, entries 1 and 2). Systematic screening of several other N-heterocyclic carbene (NHC)-based ligands revealed that those bearing 1-adamantyl [I(1-Ad)] and cyclohexyl (ICy) groups on their imidazole nitrogen atoms promoted the desired reaction to provide 2 in modest yields (entries 7 and 8). The highest yield of 2 was obtained with an NHC containing 2-adamantyl groups [I(2-Ad)] (entry 9). Although it was possible to reduce the amount of NaO'Bu (80% yield with 50 mol% NaO'Bu), the use of 2 equivalents resulted in a better yield of 2 (see ESI† for details). When the catalyst loading was reduced to 10 mol%, the yield of 2 was decreased to 52% (the catalyst loading could be reduced with other substrates. See the ESI† for details). The reductive cleavage proceeded in the presence of excess mercury; therefore we suggest that the soluble nickel species is responsible for the catalysis.

These newly developed conditions were also used with the labeled substrate 2-CD₃O-naphthalene (1a-d₂). The results of this experiment showed that the hydrogen delivered to the reduced product was primarily derived from the methoxy group of the substrate (eqn (1)); this supports our proposed mechanism shown in Scheme 1c.

![Scheme 1](image)

**Scheme 1** Mechanistic diversity of reductive cleavage of aryl methyl ethers.

The effect of the alkoxy group on the reductive cleavage reaction was then examined using an I(2-Ad) ligand (Table 2). Primary alkoxy groups, including EtO and the longer "C₆H₄O groups, were efficiently removed under these conditions, as in the reactions of 1b and 1c, but the bulkier "PrO groups was completely unreactive. When a substrate bearing a PhCH₂O group (i.e., 1e) was used, Caryl-O bond cleavage was accompanied by cleavage of C(benzyl)-O bonds. A PhO group remained virtually intact under these conditions, which is consistent with our proposed mechanism (Scheme 1c). Alkoxy groups on polyaromatic scaffolds such as naphthalenes 4–11 and phenanthrene 3 successfully underwent nickel-catalyzed reductive cleavage under these conditions to provide the parent arenes. Functional groups such as silyl (5), and amide (8 and 9) were compatible. Notably, a pendant alkene moiety, which could be reduced under the previously reported conditions using hydrosilane or hydrogen, remained intact under the current conditions, as shown by the reactions of 10 and 11. Heteroarenes, namely the electron-deficient quinoline 12 and electron-rich carbazole 13, were good substrates. In addition, this catalytic reaction could also be applied to benzylic C-O bonds (14–16). Unfortunately, benzylic C-O bond activation is limited to polyaromatic substrates, and simple benzylic alkoxy ethers were unreactive under the current conditions.

For comparison, the reductive cleavage of 14 was also examined in the presence of an external reductant (Scheme 2). The reported method using hydrosilane was far less effective (4% yield). The protocol using dihydrogen was also less effective in the presence of absence of AlMe₃. These results highlight the superior catalytic activity of the Ni/I(2-Ad) system for reductive cleavage reactions.

The lower reactivity of substrates lacking a fused aromatic ring is a common problem in nickel-catalyzed transformations of inert phenol derivatives. Similarly, 4-tert-butylandisole (15) was found to be much less reactive under the current conditions (Table 3). However, the introduction of a phenyl group, as in 16,
significantly enhanced the reactivity, and the reductive cleavage product was obtained in 51% yield. Note that compound 16 is completely unreactive under previously reported nickel-catalyzed reductive cleavage conditions using hydrosilane.

Replacement of the methoxy group with an ethoxy group further improved the efficiency of this catalysis. Ethoxybenzenes bearing a phenyl group at the para and meta positions, i.e., 17 and 18, respectively, therefore underwent the reaction to give good yields. In the case of methoxy arene substrates, hydrogen elimination of the postulated intermediate A generates formaldehyde complex C (see Scheme 1c), which readily

### Table 2 Ni/I(2-Ad)-catalyzed reductive cleavage of naphthyl and naphthylmethyl ethers

<table>
<thead>
<tr>
<th>0</th>
<th>OR</th>
<th>R'N'</th>
<th>R2N' = NP2 (8)</th>
<th>morpholin (9)</th>
</tr>
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<tr>
<td>11</td>
<td>99%</td>
<td>12</td>
<td>84%</td>
<td>13</td>
</tr>
</tbody>
</table>

**Scheme 2** Comparison with reported catalytic systems.

**Table 3 Ni/I(2-Ad)-catalyzed reductive deoxygenation of anisole derivatives**

<table>
<thead>
<tr>
<th>0</th>
<th>OR</th>
<th>R'N'</th>
<th>R2N' = NP2 (8)</th>
<th>morpholin (9)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>77%</td>
<td>16</td>
<td>51%</td>
<td>17</td>
</tr>
</tbody>
</table>

**Table 4** TEPa and %Vburb values for selected NHCs used in this study

<table>
<thead>
<tr>
<th>Entry</th>
<th>Ligand</th>
<th>TEPa</th>
<th>%Vburb</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>PCy3</td>
<td>2060</td>
<td>(35.7)</td>
</tr>
<tr>
<td>2</td>
<td>IMes</td>
<td>2050.7</td>
<td>33.8</td>
</tr>
<tr>
<td>3</td>
<td>IPr</td>
<td>2051.5</td>
<td>36.9</td>
</tr>
<tr>
<td>4</td>
<td>ICy</td>
<td>2049.6</td>
<td>28.1</td>
</tr>
<tr>
<td>5</td>
<td>I(1-Ad)</td>
<td>2048.3</td>
<td>37.3</td>
</tr>
<tr>
<td>6</td>
<td>I(2-Ad)</td>
<td>2049.4</td>
<td>33.5</td>
</tr>
</tbody>
</table>

**Scheme 2** Comparison with reported catalytic systems.
decomposes to give an inactive Ni(CO) species via decarbonylation. Although some Ni(CO) species can generate an active Ni(0) species by the dissociation of a CO ligand under the current conditions, Ni(CO) formation would be expected to decrease the efficiency of catalyst turnover. The higher reactivities of ethoxyarenes compared with methoxyarenes can be attributed to the lower tendency of the acetaldehyde complex, which is generated from the nickel-ethoxide intermediate, to undergo the undesired decarbonylation. As well as a phenyl group, several electron-withdrawing groups can accelerate the corresponding arenes. ortho-Substituted anisoles, such as 2-ethoxy-1,1’-biphenyl, failed to react under the current conditions.

The key to a successful reaction is the use of a new NHC ligand I(2-Ad). The electronic and steric properties of I(2-Ad) were investigated to gain insights that will help in the future design of better ligands for catalytic C–O bond activation processes (Table 4). In terms of the electronic properties, the Tolman electronic parameter (TEP) for the new I(2-Ad) ligand was determined to be 2049, by synthesizing [IrCl(I(2-Ad))(CO)2]. This value is similar to the TEP values obtained for ICy and I(1-Ad), and I(2-Ad) is a significantly stronger σ-donor than PCy3, the commonest ligand used in nickel-mediated C–O bond activation. The steric properties were determined by calculating the buried volume (%V_{bur}), based on the crystal structure of [IrCl(I(2-Ad))(cod)]. The %V_{bur} of I(2-Ad) (33.5) was found to be close to that for IMes (33.8), exerting a significantly larger steric demand than ICy, but not as large as those of I(1-Ad) and IPr. The catalytic activity in this study clearly cannot be explained solely by these two static parameters, and other factors such as dynamic behavior of the substituents should be considered for a complete understanding of the structure–activity relationships.

Conclusions

In summary, we have developed a novel procedure for the nickel-catalyzed reductive deoxygenation of aryl ethers. This new reaction proceeds in the absence of an external stoichiometric reductant via a mechanism involving oxidative addition of the C–O bond followed by β-hydrogen elimination, which has not previously been exploited in a catalytic setting. The present catalytic system not only provides an atomically economic process by avoiding the use of an external reductant, but also provides a significantly wider substrate scope. These results also show that the elusive oxidative addition of a C(aryl)–OME bond is a reliable elementary step in nickel-mediated C–O bond transformations of aryl ethers, particularly when an NHC-based ligand is used. Further studies of nickel-catalyzed transformations of aryl ethers using a Ni/NHC system are currently underway in our laboratory.

Acknowledgements

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Notes and references


7 Selected works on reductive cleavage of aryl ethers under heterogeneous catalysis: (a) A. G. Sergeev, J. D. Webb and J. F. Hartwig, J. Am. Chem. Soc., 2012, 134, 20226; (b) Y. Ren, M. Yan, J. Wang, Z. C. Zhang and K. Yao, Angew. Chem., Int. Ed., 2013, 52, 12674; (c) M. Chatterjee,


12 In fact, stoichiometric studies by Martin (ref. 6d) reported successful isolation of an acetaldehyde-bound nickel(0) complex, while the attempted synthesis of the corresponding formaldehyde complex resulted in the formation of a carbonyl–nickel(0) complex via a rapid decarbonylation process.

