Nickel-catalyzed C–N bond activation: activated primary amines as alkylation reagents in reductive cross-coupling†

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Nickel-catalyzed reductive cross coupling of activated primary amines with aryl halides under mild reaction conditions has been achieved for the first time. Due to the avoidance of stoichiometric organometallic reagents and external bases, the scope regarding both coupling partners is broad. Thus, a wide range of substrates, natural products and drugs with diverse functional groups are tolerated. Moreover, experimental mechanistic investigations and density functional theory (DFT) calculations in combination with wavefunction analysis have been performed to understand the catalytic cycle in more detail.

1. Introduction

Transition-metal-catalyzed cross-coupling reactions play an important role in modern organic synthesis.1 Among them, reductive cross-coupling which uses two electrophiles as coupling partners represents an attractive catalytic platform for the formation of diverse chemical bonds.2 Due to the avoidance of stoichiometric organometallic reagents this transformation often possesses a broader substrate scope. Traditionally, alkyl halides,3 which may have limited stability and availability, have been used as alkyl electrophiles. Recently, N-hydroxypyridinethalimide esters, anhydrides, benzyl oxalates, tosylates and mesylates have been reported as alkyl coupling partners.4 However, the use of primary amines as readily available, cheap, green and stable alkylation reagents in reductive cross-coupling has not been described. Recently, bench-stable pyridinium salts have been reported to be attractive substrates in deaminative cross-coupling with boronic acids.5 In this transformation, the radical is generated via a single electron transfer (SET) pathway between the pyridinium salt and the nickel catalyst. In addition, an alkyl radical can be formed by a photoinduced SET process6 or by an elegant metal-free formation of a boron-based electron-donor–acceptor (EDA) complex7 under blue light irradiation. Also, a Lewis base promoted C–N borylation has been reported.8 As part of our continuing efforts in metal catalyzed functional group interconversion and the activation of inert bonds,9 we herein describe the first Ni-catalyzed deaminative reductive cross-coupling of activated primary amines with aryl halides, providing a versatile method for the transformation of amino groups to aryl motifs under mild conditions (Scheme 1).

2. Results and discussion

We started our investigation by evaluating the reductive cross-coupling of cyclohexylamine derived pyridinium salt 1a and iodobenzene 2a using NiCl₂-dme as the catalyst, bipyridine ligand L1 as the ligand and a reducing agent (Table 1). Since solvents typically play an important role in cross-coupling reactions, we first tested several solvents (entries 1–5) and found DMA to be the optimal solvent. Subsequently the ratio of reactants 1a and 2a was evaluated (entries 6 and 7) and the yield was improved to 61%. Bidentate as well as tridentate N-containing ligands L1–L4 can be applied (entries 8–10) and L1, L2,
and L4 showed a similar efficiency, providing the desired product in about 60% yield. With the readily available and cheap bipyridine L2, we examined a series of nickel catalysts, including NiBr₂·dme, Ni(OAc)₂·4H₂O, Ni(acac)₂, and NiCl₂·6H₂O. However lower yields were observed (entries 11–14). The yield was significantly improved to 99% when Mn powder was used as a reductant (entry 16), whereas the utilization of Mg powder gave the desired product in only 8% yield (entry 15). Besides aryliodides, the protocol was also applied to aryl bromides. When bromobenzene was used, the desired product was also obtained in a good yield (entry 17). Control experiments demonstrated that the nickel catalyst, ligand, and reductant are all essential for the success of this transformation (entries 18–20).

With the optimized reaction conditions in hand, the scope of aryl halides was first evaluated (Table 2a). A wide variety of aryl halides bearing electron-donating, electron-neutral, and electron-withdrawing functional groups could be successfully converted into the corresponding products in good to excellent yields. For example, phenyl and diphenyl iodides and bromides underwent this reaction smoothly, giving the corresponding products in excellent yields (3a and 3b).

In addition, a wide range of functional groups including ketone (3c and 3d), trifluoromethyl (3e), trifluoromethoxy (3f), trifluoromethylthio (3g), tosyl (3h), cyano (3i and 3m), methoxy (3j), methylthio (3k), -butyl (3l), fluoro (3m, 3r and 5k), and ester (5j) were well tolerated under the mild reaction conditions, highlighting the high chemoselectivity of this newly developed deaminative reductive cross-coupling reaction. Use of disubstituted aryl bromide and bicyclic substrates including phtha-lides and naphthyl halides also gave the products 3m–3p in good yields. It is noteworthy that pharmaceutically relevant 3- and 4-bromopyridines could be applied to this protocol with good to high efficiency (3q–3s, 5n). Next, the scope of pyridinium salts was explored. A wide range of structurally diverse pyridinium salts were suitable substrates for this transformation. Cyclic and acyclic secondary amine substrates could undergo this deaminative arylation reaction in good to excellent yield (4a–4g) and the same applies for N-heterocyclic pyridinium salts (4h and 4i) (Table 2b).

It should be mentioned that when we applied primary alkyl pyridinium salts to this protocol, the reaction did not occur. However, simply switching the ligand from L2 to L4 and slightly raising the reaction temperature to 60 °C allowed this transformation to occur smoothly. A series of primary alkyl pyridinium salts bearing diverse functional groups such as amine, acetal, dioxy, cyclohexenyl, thiophene, and pyridine were suitable coupling partners for this deaminative reductive cross-coupling, leading to products 4n–4s. Notably, chloro (4k), unprotected OH and indole NH groups were also tolerated (4t and 5f), providing the option for further functionalization. Moreover, methylation reaction, which is challenging in reductive cross-coupling, was also realized via the utilization of methyl pyridinium salts (Table 2c). Importantly, our newly developed protocol could also be readily extended to a wide range of complex molecules derived from natural products and drugs. As such pregnenolone, galactopyranose, probenecid, adamantane carboxylic acid, and cholesterol derivatives could be transformed to the corresponding products 5a–5e in good to excellent yield. Moreover, a series of pyridinium salts derived from drugs or drug intermediates, including tryptamine, mexiteline, amphetamine, Lipitor intermediate, and dopamine, all underwent the mild coupling protocol with good to excellent efficiency (5f–5n). Use of Mosapride derived pyridinium salts gave product 5o in a lower yield (Table 2d).

Additionally, a gram-scale reaction was successfully conducted using 1a and 4-iodobiphenyl 2b in the presence of only 5 mol% nickel catalyst and the desired product 3b was obtained in 96% yield (Scheme 2a), demonstrating the practicability of our newly developed deaminative reductive cross-coupling methodology. Also, byproduct 6, which is potentially a useful organic base, could be isolated in 85% yield. To shed light on the mechanism of this transformation, an experiment was conducted with the radical trapping reagent TEMPO (2,2,6,6-tetramethyl-1-piperidinyloxy, 2 equiv.). The reaction was suppressed and no product 5m was detected (Scheme 2b).
Table 2  Scope of substrates

\[ \text{Reaction conditions: pyridinium salt 1 (0.30 mmol), aryl halide 2 (0.20 mmol), NiCl}_2 \text{ dme (0.02 mmol), L2 (0.02 mmol, for secondary alkyl) or L4 (0.02 mmol, for primary alkyl), Mn powder (0.40 mmol) in 1.0 ml DMA at rt (for secondary alkyl) or 60 °C (for primary alkyl); yields after purification.} \]

\[ \text{GC yield.} \]

\[ \text{The ratio of pyridinium salt to aryl halide 1 : 3, 100 °C, NMR yield.} \]
Also, ring-opened product 9 was generated in 55% yield when a substrate bearing a cyclopropane motif 8 was used (Scheme 2c). Both these results suggest the involvement of an alkyl radical in this transformation. When ligand L5 which is effective for chain-walking reductive cross-coupling was used in our catalytic system, non-walking and chain-walking products (4e and 4e’, 5h and 5h’) were obtained with a ratio of 4 : 1, suggesting an oxidative addition of aryl halide to Ni0 to give a NiII intermediate prior to alkyl radical generation. Since the NiII intermediate generated from addition of the alkyl radical to the NiII intermediate is less likely to undergo the chain-walking step due to steric hindrance the non-chain-walking product is the major product.

Furthermore, detailed DFT calculations were performed to rationalize our newly designed catalytic reaction (Scheme 3; computational methods, see ESI†). As a model system, we investigated the reaction of phenyl bromide with A1 in the presence of NiCl2·dme, bpy as the ligand and Mn as the reducing agent. The reaction starts with the complexation of the ligand bpy to the NiII precatalyst, followed by reduction to form the active Ni0 catalyst B (Scheme S1 in ESI†). The catalytic process is initiated by oxidative addition of phenyl bromide to Ni0 via transition state B-TS with an energy barrier of 11.7 kcal mol⁻¹. The formed NiII intermediate C is reduced by Mn, leading to intermediate D with a free energy gain of 6.9 kcal mol⁻¹. In the next step, A1 is coordinated to D, followed by SET reduction of A1 to generate radical A2 and NiII intermediate F. The radical A2 is prone to undergo C–N bond cleavage with an energy barrier of 19.9 kcal mol⁻¹, liberating the alkyl radical A3 and the aromatic pyridine A4. At this point, alkyl
radical \(A_3\) adds to \(\text{Ni}^{\text{II}}\) intermediate \(F\) to form \(\text{Ni}^{\text{III}}\) intermediate \(G\). Subsequently, the C-C bond cross-coupling product \(A_5\) is formed \textit{via} reductive elimination from \(\text{Ni}^{\text{III}}\) with an energy barrier of 4.9 kcal mol\(^{-1}\).

Finally, the \(\text{Ni}^{\text{II}}\) intermediate \(H\) is further reduced by Mn to regenerate the \(\text{Ni}^{\text{I}}\) active catalyst \(B\) and initiate the next catalytic cycle.

Subsequently, we focused on the origin of the SET reduction of the pyridinium salt and the generation of the alkyl radical.\(^{11}\) The molecular orbital plots (Fig. 1a) show that the SOMO of \(E\) corresponds to the singly occupied MO predominantly localized on the \(d_z\)-orbital of \(\text{Ni}\). At the same time, the LUMO of \(E\) corresponds to a \(\pi\)-orbital delocalized around the central nitrogen-containing aromatic ring of the pyridinium salt. However, after \(A_2\) is displaced away from the \(\text{Ni}\) species, the SOMO becomes localized on the central aromatic ring of the pyridinium salt. This indicates that upon separation of the \(\text{Ni}\) and pyridyl fragments the unpaired electron transfers from \(\text{Ni}\) to the pyridinium salt and delocalizes around the central aromatic ring. Spin density analysis further supported this process. The unpaired electron density is localized on the \(\text{Ni}\)-center when the pyridinium salt is coordinated to the \(\text{Ni}\)-complex, as in \(E\) (Fig. 1b, left), while it is transferred to the central pyridine ring as the pyridine moiety \(A_2\) dissociates from the Ni complex (Fig. 1b, middle). Subsequently, the C-N bond dissociates and the spin density is further transferred to the sp\(^3\)-carbon atom, indicating generation of the alkyl radical (Fig. 1b, right). Additionally, cyclic voltammetry (CV) measurements of the pyridinium salt were conducted (see ESI†) and reversible peaks at \(-0.94\) \(V\) vs. \(\text{SCE}\) were observed, suggesting the existence of stable radical intermediate \(A_2\). In order to understand the C-N bond dissociation, we performed localized orbital locator (LOL) - \(\pi\) analysis\(^{22}\) and multi-center bond order calculations \(\langle I_{\text{ring}} \rangle\) \(^{13}\) for the whole process (Fig. 1c). Before the SET reduction, the central pyridine ring has full aromaticity \(\langle I_{\text{ring}} \rangle = 0.05\) in \(A_1\), for comparison, \(I_{\text{ring}} = 0.05\) in benzene and pyridine\(^{16}\). After the reduction the aromaticity is lost and the free energy is 3.0 kcal mol\(^{-1}\) higher \(\langle I_{\text{ring}} \rangle = 0.05\) in \(A_2\). With the C-N bond dissociating, the aromaticity is partially regained (c.f. \(I_{\text{ring}} = 0.01\) in transition state\(A_2\)). After the generation of alkyl radicals, the aromaticity of \(A_3\) is fully restored \(\langle I_{\text{ring}} \rangle = 0.05\) in \(A_3\) with a free energy gain of 5.5 kcal mol\(^{-1}\).

3. Conclusions

In summary, we have developed a new deaminative reductive cross-coupling of aryl halides with pyridinium salts derived from readily available alkyl amines. Due to the avoidance of stoichiometric organometallic reagents and external bases, the scope is rather broad as demonstrated using the many different substrates employed (>55 examples). In addition, the chemoselectivity of this protocol is good and functional groups, including cyano, methoxy, methylthio, fluoro as well as chloro, unprotected OH and indole NH groups, are well tolerated. Importantly, the cross-coupling reaction can be scaled-up using a lower amount of nickel catalyst without diminishing the yield, demonstrating the practicability of this protocol. Furthermore, experimental mechanistic investigations and DFT calculations combined with wavefunction analysis have been conducted to gain insight into the catalytic process.

Conflicts of interest

There are no conflicts to declare.

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Notes and references


