## Citation


## As Published

http://dx.doi.org/10.1021/jacs.8b01696

## Publisher

American Chemical Society (ACS)

## Version

Author’s final manuscript

## Citable link

https://hdl.handle.net/1721.1/125953

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Breaking the Base Barrier: An Electron-Deficient Palladium Catalyst Enables the Use of a Common Soluble Base in C–N Coupling

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Abstract

Due to the low intrinsic acidity of amines, palladium-catalyzed C–N cross-coupling has been plagued continuously by the necessity to employ strong, inorganic, insoluble bases. To surmount the many practical obstacles associated with these reagents, we utilized a commercially available dialkyl triarylmonophosphine-supported palladium catalyst that facilitates a broad range of C–N coupling reactions in the presence of weak, soluble bases. The mild and general reaction conditions show extraordinary tolerance for even highly base-sensitive functional groups. Additionally, insightful heteronuclear NMR studies using 15N-labeled amine complexes provide evidence for the key acidifying effect of the cationic palladium center.

Graphical Abstract

INTRODUCTION

Palladium-catalyzed carbon-nitrogen (C–N) bond formation has become a valuable tool in the modern synthesis of structurally complex target molecules. Over the last two decades, numerous methods have been established for the coupling of aryl halide and pseudohalide electrophiles with several classes of amine nucleophiles in the presence of base.† A number of improvements in the development of ligands‡ and precatalysts§ have taken place. Despite considerable effort, there has been no general and practical solution to allow for the use of...
amine bases in Pd-catalyzed C–N cross-coupling. Because the majority of current catalytic systems utilize inorganic bases, modern amination methods suffer from restrictions with regard to functional group compatibility or other operational limitations due to insolubility (Figure 1). For instance, metal alkoxide and lithium amide bases are moisture-sensitive and are incompatible with functional groups that are prone to nucleophilic attack either by base or by base-activated amine. As an alternative, inorganic bases such as Cs₂CO₃, K₃PO₄, and K₂CO₃ are frequently employed, but because of their insolubility, reactions often require a phase transfer agent or high catalyst loadings, as well as elevated reaction temperatures. Furthermore, because deprotonation likely takes place on the particle surface of the base in heterogeneous systems, the use of these bases introduces reproducibility issues on large scales when stirring methods or particle sizes are varied. Finally, due to their insolubility in most organic solvents, these bases also pose challenges for continuous flow chemistry and in reagent dosing for robotic miniaturization methods, which has stifled usage of industrial technologies. Thus, the discovery of a catalyst system that facilitates the coupling of multiple amine classes in the presence of a weak, amine base would provide an important solution to this long-standing synthetic challenge. Herein, we report the first general solution to this problem. Moreover, we have uncovered the factors that enable the success of such a system, which should help guide the future discovery of even more active catalytic systems.

Organic bases are employed infrequently in C–N coupling methodologies despite their widespread use in synthesis, broad pKₐ⁺ range, suitability for continuous flow chemistry, and compatibility with electrophilic functional groups (Figure 2). In one example, organic phosphazene bases such as P₂-Et have been used in nanomolar, robotic high-throughput screening to couple a variety of nucleophiles. While effective, phosphazene bases are expensive and must be stored in an inert atmosphere. Weaker organic amine bases such as 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU) and 7-methyl-1,5,7-triazabicyclo[4.4.0]dec-5-ene (MTBD) have been employed in C–N coupling of aryl nonaflate under microwave radiation. However, only simple anilines could be coupled using DBU while heteroaryl amines required the moisture-sensitive and expensive MTBD. Utilizing photoredox catalysis, C–N coupling has been performed with 1,4-diazabicyclo[2.2.2]octane (DABCO) as the base. Despite the mild nature of this method, C–N couplings facilitated by photoredox processes are primarily focused on aliphatic amine nucleophiles. Finally, a few examples of electrochemically-mediated, nickel-catalyzed amidation and amination of aryl bromides have been reported with DBU as the base. In hopes of overcoming current limitations of these methodologies, we aimed to develop a system that utilizes commercially available ligands and a weak organic base to couple a variety of amine classes.

RESULTS AND DISCUSSION

Our efforts focused on whether commonly used dialkyl biarylmonophosphine ligands could facilitate the coupling of aryl halides or triflates with amines using soluble organic bases. Initial stoichiometric experiments using oxidative addition (OA) complexes (See Supporting Information) indicated that weaker bases such as triethylamine (TEA) and DABCO were outperformed by DBU in procedures using many different ligands when coupling aryl amines, amides, and primary amines. Experiments utilizing p-tolyl-OA complexes (Figure 1, H) as catalysts (1.0 mol%) to couple p-tolyl triflate with aniline revealed that multiple
ligands are capable of facilitating the desired reaction (Table 1). Precatalyst P6 bearing L6 (AlPhos)\(^{16}\) and P5 (AdBrettPhos) provided nearly quantitative cross-coupling product at room temperature. P4 (t-BuBrettPhos) provided the desired product in moderate yields,\(^{17}\) and ligands bearing cyclohexyl (Cy) groups on the phosphine, including P3 (BrettPhos) and L1 (XPhos) failed to yield any of the desired product. While P5 is highly reactive, the ligand from which it is derived is known to undergo an in situ ligand modification in the presence of aryl (pseudo)halides, generating different active catalytic species for each ArX substrate.\(^ {18}\) As a result, the commercially available, air-stable, L6-supported precatalyst, COD(L6-Pd)\(_2\), was selected for further investigation in C–N coupling with amines of varying pKa and nucleophilicity. When coupling benzamide with the aryl triflate, 76% yield was obtained, with unreacted electrophile making up the mass balance. Upon raising the temperature to 60 °C, the yield was increased to 97%. Similar reactivity was observed with benzyl amine, demonstrating that this ligand facilitates the coupling of different amine classes.

To explore the scope of the methodology, a variety of aryl halides, triflates, and nucleophiles were tested (Table 2). Both five- and six-membered heterocyclic amines, including a pyrazine (1a), a thiazole (1e), and an oxazole (1i) were coupled in high yield under the optimized reaction conditions. Substrates bearing acidic functional groups, such as an unprotected pyrazole (1d, 1f), an indole (1b, 1h, 3d, 3f), and a phenol (1c, 3c), are also compatible with the weakly basic reaction conditions when an excess of base is used. Importantly, five-membered heterocyclic bromides are competent in the reaction, as well as highly functionalized electrophiles such as the Merck informer compound, X18 (1f). Using 0.2 mol% of Pd, 1g was prepared at room temperature in 95% yield after crystallization on a 10 mmol scale. Amide-type coupling partners, including an oxazolidinone (2b), a primary urea (2g), and a secondary cyclic amide (2d) underwent the desired arylation reaction in good to excellent yields. Heterocyclic amides, including pyridines (2c, 2e), a furan (2h), and a thiophene (2f) are tolerated, though these substrates required diluted reaction conditions due to their low solubility. Similarly, heterocyclic products derived from aliphatic amine coupling partners are obtained in high yields. In cases where standard reaction conditions show low conversion to product, excess base and amine were employed (3e). Hindered aliphatic amines, including secondary aliphatic amines, proved to be difficult substrates under the reaction conditions, most likely due to unfavorable steric interactions near the Pd center. Relative to COD(L6-Pd)\(_2\), t-BuBrettPhos-G3 provides 1d and 1f in comparable yields (Table 2, red numbers) with 1.0 mol% of Pd, while 1a, and 2e require higher loadings. We note that the L4-supported precatalyst fails to couple primary amines under these conditions.

Due to the mildness of the reaction conditions, we envisioned that this methodology would be amenable to substrates that would readily undergo undesired elimination and substitution reactions in the presence of strong bases. For example, Hartwig has shown that fluoroalkylamines participate in base-promoted decomposition reactions.\(^ {19}\) In our system, these amines are readily coupled at room temperature without decomposition (Table 3, 4c). Moreover, elimination-prone coupling partners bearing alkyl halide functional groups, including unhindered primary bromides (4a) and chlorides (4b, 4d), are suitable coupling partners. When subjected to previously developed conditions, these coupling partners and
products were found to undergo elimination reactions to yield the corresponding alkenes; however, under our reaction conditions, the alkyl halide provides opportunities for subsequent synthetic elaboration.

In both stoichiometric and catalytic experiments, we observed that ligands bearing large alkyl groups performed better than smaller variants. Inspired by the work of Tyler, we hypothesized that the size of the alkyl groups on the phosphine might affect the electrophilicity of palladium through geometric distortion at the phosphorus atom. In turn, these size effects should influence the acidity of Pd-bound amines. Specifically, the magnitude of this acidifying effect directly affects the ease of the key deprotonation step and consequently dictates the strength of base that is required. The structure of an amine-bound OA complex (Figure 1, III) was elucidated by single crystal X-ray diffraction, which indicated that the amine was bound trans to the phosphine (Figure 3). Analogous complexes bearing a 15N-enriched n-butyl amine were used to study the effect of ligand bulkiness on the central Pd atom charge (Table 4). NMR spectroscopic analyses of the two-bond 31P–15N coupling for AlPhos and BrettPhos ligand families indicated that the ligands bearing larger alkyl groups exhibited smaller coupling constants (Ad < t-Bu < Cy). The magnitude of the trans 31P–15N 2J constant is directly influenced by the electron-donation to the Pd atom: specifically, more cationic (at Pd) amine-bound complexes exhibit less negative 2J values (see Supporting Information for discussion). Thus, the trend we observed indicates that the Ad ligands are the least electron-donating, a conclusion that is consistent with our DFT calculated Hirshfeld charges at Pd. Of the intermediates analyzed, the smallest 2J was observed with AlPhos (L6) as the supporting ligand, suggesting that the amine bound to this complex is most easily deprotonated. We propose that this explains the superior performance of L6 in reactions promoted by weak bases.

CONCLUSION

In summary, we have developed a catalyst system that utilizes a weak organic amine base in the coupling reaction of aryl (pseudo)halides with a variety of amine and amide partners. The electrophile scope includes five- and six-membered heterocycles, as well as alkyl halide functional groups that were not tolerated by previously developed conditions. While L6 is the most generally effective ligand, we found that more traditional ligands such as L4 can be economical substitutes for some aryl amine and amide couplings. We show that the size of the phosphine substituents modulates the charge on the Pd atom, with L6 generating the most cationic, and thus most active, amine–Pd complex. Ongoing work is focused on a more detailed understanding of the reaction mechanism.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

This material is based upon work supported by the NSF Graduate Research Fellowship Program under Grant No. 1122374 (J.M.D.), an NIH Postdoctoral Fellowship under Grant No. 1F32GM120847-01 (N.A.W.), and the NIH (Grant Nos. GM122483 and GM58160). Any opinions, findings, conclusions, or recommendations expressed in this material are those of the authors and do not necessarily reflect the views of the NSF or NIH. R.Y.L. thanks Bristol-
Myers Squibb for a Fellowship in Synthetic Organic Chemistry. We acknowledge several companies for donations of chemicals used in this work: Sigma-Aldrich for BrettPhos and t-BuBrettPhos, Nippon Chemical for cyclohexyl chlorophosphine, FMC Lithium for tert-butyl chlorophosphine, and Merck for informer compound X18. We are grateful to Dr. Peter Müller for crystallographic analysis, and we acknowledge Dr. Andy Thomas and Dr. Christine Nguyen for assistance in the preparation of this manuscript.

References


17. In high-throughput reaction screening, this ligand was found to be suitable for amide couplings facilitated by DBU. See reference 11a.


21. Preliminary mechanistic studies have indicated that there is a positive order in [DBU] for the coupling of p-tolyl triflate and propyl amine. This suggests that the base is involved in the turnover-limiting step.


23. The variation in $^{2}J$ values (~1.0 Hz) is in the range of those observed in $^{2}J$ coupling constants of $^{13}$C–Au–$^{31}$P complexes bearing trialkyl and triaryl phosphine ligands. For studies on the effect of phosphine size on two-bond coupling, see: Isab AA, Hussain MS, Akhtar MN, Wazeer MIM, Al-Arfaj AR. Polyhedron. 1999; 18:1401–1409.
Figure 1.
(A) The proposed catalytic cycle for the palladium-catalyzed coupling of aryl halides with amines: I, monoligated Pd(0); II, oxidative addition (OA) complex; III, amine-bound OA complex; IV, amido complex. (B) Synthetic considerations for bases employed in C–N coupling.
Figure 2.
Current metal-catalyzed C–N coupling methodologies that utilize soluble, amine bases.
Figure 3.
Propyl-amine-bound L6-supported OA complex. Thermal ellipsoids are shown at 50% probability. Hydrogen atoms and residual toluene molecules are omitted for clarity.
### Table 1
Comparison of Ligands in Pd-Catalyzed Amination Facilitated by DBU<sup>a</sup>

<table>
<thead>
<tr>
<th>Precatalyst</th>
<th>Nucleophile</th>
<th>Temp, (°C)</th>
<th>Time</th>
<th>Yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P1</td>
<td>Aniline</td>
<td>RT</td>
<td>20 min</td>
<td>0</td>
</tr>
<tr>
<td>P2</td>
<td>Aniline</td>
<td>RT</td>
<td>20 min</td>
<td>30</td>
</tr>
<tr>
<td>P3</td>
<td>Aniline</td>
<td>RT</td>
<td>20 min</td>
<td>0</td>
</tr>
<tr>
<td>P4</td>
<td>Aniline</td>
<td>RT</td>
<td>20 min</td>
<td>61</td>
</tr>
<tr>
<td>P5</td>
<td>Aniline</td>
<td>RT</td>
<td>20 min</td>
<td>99</td>
</tr>
<tr>
<td>P6</td>
<td>Aniline</td>
<td>RT</td>
<td>20 min</td>
<td>99</td>
</tr>
<tr>
<td>COD(L6-Pd)₂</td>
<td>Aniline</td>
<td>RT</td>
<td>3 h</td>
<td>99</td>
</tr>
<tr>
<td>COD(L6-Pd)₂</td>
<td>Benzamide</td>
<td>RT</td>
<td>16 h</td>
<td>76</td>
</tr>
<tr>
<td>COD(L6-Pd)₂</td>
<td>Benzamide</td>
<td>60</td>
<td>16 h</td>
<td>97</td>
</tr>
<tr>
<td>COD(L6-Pd)₂</td>
<td>Benzyl amine</td>
<td>60</td>
<td>16 h</td>
<td>98</td>
</tr>
</tbody>
</table>

<sup>a</sup>Reaction conditions: aryl triflate (0.25 mmol) nucleophile (0.30 mmol), DBU (0.50 mmol), precatalyst (1% Pd; 2.5 μmol or 1.25 μmol dimer), MTBE (0.25 mL). GC and ¹H NMR yields referenced to hexamethylbenzene as an internal standard. MTBE = methyl tert-butyl ether.
Table 2

Amination and Amidation of Aryl Triflates, Bromides, and Chlorides

<table>
<thead>
<tr>
<th>DBU (2.0 equiv)</th>
<th>COD(L6-Pd)2 (1% Pd)</th>
<th>MTBE (1.0 M), 60 °C, 16 h</th>
<th>L4-G3</th>
<th>Aryl Amin</th>
<th>Acyl Amin</th>
<th>Primary Amin</th>
</tr>
</thead>
<tbody>
<tr>
<td>X = C(O)N</td>
<td>X = C(O)N</td>
<td>X = C(O)N</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Isolated yields are reported as the average of two runs. Unless noted, standard reaction conditions: aryl(pseudo)halide (1.0 mmol), nucleophile (1.2 mmol), COD(L6-Pd)2 (0.005 mmol, 1% Pd), MTBE (1.0 mL, 1 M), 60 °C for 16 h. While these conditions are suitable for each substrate, in many cases milder conditions can be used. Red yields indicate reactions performed with L4-G3 under substrate-specific conditions.

b3% L4-G3

c4.0 equiv of base.

d3 h at RT, 4.0 equiv of DBU.

e1% L4-G3.

f0.50 mmol, 0.5 M in THF.

gh3 h at RT.

i10 mmol scale, 0.2% Pd, 16 h, RT.

j0.25 M, 0.95 equiv of nucleophile.

k0.25 M.

h3% Pd, 4.0 equiv of DBU.

l4.0 equiv of DBU and amine, 50 °C.
Table 3

Base-Sensitive Coupling Partners

<table>
<thead>
<tr>
<th>X = OTf, Br (1.2 eq.)</th>
<th>DBU (2.0 equiv)</th>
<th>COD(L6-Pd)2 (1% Pd)</th>
<th>H-NR2</th>
<th>MTBE, 1.0 M, RT, 16 h</th>
<th>X = Cl, Br, N</th>
</tr>
</thead>
<tbody>
<tr>
<td>4a</td>
<td>87%b From ArBr</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4b</td>
<td>98% From ArBr</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4c</td>
<td>74% From ArBr</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4d</td>
<td>90% From ArOTf</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

a Isolated yields are reported as the average of two runs. Unless noted, standard reaction conditions: aryl(pseudo)halide (1.0 mmol), nucleophile (1.2 mmol), DBU (2.0 mmol), COD(L6-Pd)2 (0.005 mmol, 1% Pd), MTBE (1.0 mL, 1 M), RT for 16 h.

b 1.0 mmol of DBU and 1.05 mmol of nucleophile.
Table 4

$^{31}$P–$^{15}$N $^{2}J$ Coupling Constants$^{a}$

<table>
<thead>
<tr>
<th>Ligand</th>
<th>Alkyl Group ($R^{1}$)</th>
<th>$^{2}J$($^{31}$P–$^{15}$N) (Hz)</th>
<th>Pd Charge (Hirshfeld, 10$^{-3}$e)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BrettPhos (L3)</td>
<td>Cy</td>
<td>$-$37.0</td>
<td>+186</td>
</tr>
<tr>
<td>t-BuBrettPhos (L4)</td>
<td>t-Bu</td>
<td>$-$36.1</td>
<td>+191</td>
</tr>
<tr>
<td>AdBrettPhos (L5)</td>
<td>Ad</td>
<td>$-$35.2</td>
<td>+195</td>
</tr>
<tr>
<td>CyAIPhos (L8)</td>
<td>Cy</td>
<td>$-$35.9</td>
<td>+194</td>
</tr>
<tr>
<td>t-BuAIPhos (L7)</td>
<td>t-Bu</td>
<td>$-$35.1</td>
<td>+203</td>
</tr>
<tr>
<td>AIPhos (L6)</td>
<td>Ad</td>
<td>$-$34.2</td>
<td>+203</td>
</tr>
</tbody>
</table>

$^{a}$Hirshfeld charges were computed at the B3LYP/6-31G(d)-SDD(SMD(THF)) level of theory with geometries optimized at the same level.