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# Communication

# The Application of Pulse Radiolysis to the Study of Ni(I) Intermediates in Ni-Catalyzed Cross-Coupling Reactions

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**ABSTRACT:** Here we report the use of pulse radiolysis and spectroelectrochemistry to generate low-valent nickel intermediates relevant to synthetically important Ni-catalyzed cross-coupling reactions and interrogate their reactivities toward comproportionation and oxidative addition processes. Pulse radiolysis provided a direct means to generate singly reduced [(dtbbpy)NiBr], enabling the identification of a rapid Ni(0)/Ni(II) comproportionation process taking place under synthetically relevant electrolysis conditions. This approach also permitted the direct measurement of Ni(I) oxidative addition rates with electronically differentiated aryl iodide electrophiles ( $k_{OA} = 1.3 \times 10^4 - 2.4 \times 10^5 \text{ M}^{-1} \text{ s}^{-1}$ ), an elementary organometallic step often proposed in nickel-catalyzed cross-coupling reactions. Together, these results hold implications for a number of Ni-catalyzed cross-coupling processes.

ver the past 10 years, significant advances in nickelmediated cross-coupling technology have elevated this mode of catalytic bond formation to widespread use in both academic and industrial settings.<sup>1,2</sup> At the same time, the development of dual-catalytic Ni/photoredox<sup>3-5</sup> and Ni/ electrocatalysis<sup>6–8</sup> platforms, as well as further developments in Zn- and Mn-mediated reductive coupling reactions," has enabled a wide array of challenging and/or unique C-C and C-heteroatom couplings that were previously unknown. In practice, these advances have led to the wide-scale adoption of nickel cross-coupling reactions across many industrial sectors, perhaps most notably the medicinal chemistry community.<sup>10,11</sup> While the development and deployment of these technologies has taken place at an impressive rate, a detailed understanding of the underlying mechanisms responsible for these new bondforming processes has lagged behind in comparison.<sup>12</sup>

The use of photoredox and electrocatalysis has proven particularly valuable in modulating nickel oxidation states and generating transient high-energy radical intermediates that, in turn, can readily participate in previously elusive bond-forming processes.<sup>6,13,14</sup> While such reaction pathways provide a means to forge otherwise challenging C-C and C-heteroatom bonds, the fleeting nature of these catalytic Ni species makes them accordingly difficult to study. Electroanalytical techniques, such as cyclic voltammetry, can be used to probe the propensity for catalyst oxidation and reduction as well as the fate of resulting electrogenerated transition metal intermediates. For dual-catalytic Ni/photoredox reactions, Stern-Volmer quenching experiments are routinely utilized to measure rapid photoinduced electron-transfer rates, and transient absorption spectroscopy can provide insight into ensuing elementary steps.<sup>15-21</sup> However, while electroanalytical<sup>22-</sup> and spectroscopic techniques have provided invaluable insight into many open-shell processes, the use of these experiments is less informative for reactions involving intermediates that exhibit non-reversible redox behavior or which participate in low quantum efficiency steps. Moreover,

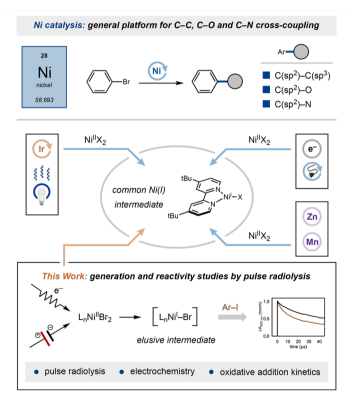
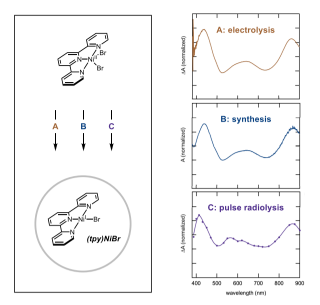


Figure 1. Approach to studying elusive Ni(I) intermediates.

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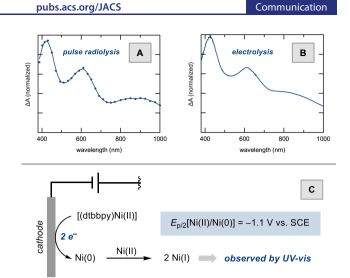


**Figure 2.** Normalized  $\Delta A/A$  data for [(tpy)NiBr] generated by electrolysis ( $E_{app} = -750 \text{ mV vs SCE}$ ) (A), stoichiometric synthesis (B), and pulse radiolysis (t = 100 ns) (C).

multiple recent observations of radical chain mechanisms involving photogenerated Ni(I)/Ni(III) species further underscore the need to examine catalytic intermediates produced with low quantum yields that nonetheless are responsible for significant levels of product formation.  $^{16,17,19}$ 

While significant mechanistic complexity is a hallmark of nickel-catalyzed cross-coupling reactions, initial single-electron reduction of the employed Ni(II) dihalide precatalyst is a commonly proposed initiation event across many of these transformations (Figure 1).<sup>12,16,26–28</sup> The relative instability of the corresponding three-coordinate Ni(I) halide complexes resulting from reduction of the respective Ni(II) precatalyst makes studying these catalytic intermediates challenging. Such Ni(I) complexes can be generated with sterically encumbered ligands;<sup>29–31</sup> however, the cross-coupling relevant bipyridine-ligated analogs form stable and unreactive halide-bridged dimers or Ni(I)–Ni(II) mixed-valent bimetallic species.<sup>16,20,32</sup> For these reasons, the study of catalytically active, coordinatively unsaturated Ni(I) intermediates has remained largely unexplored.

Pulse radiolysis has a rich history of successfully probing reactivities of short-lived catalytic intermediates,<sup>33,34</sup> interrogating biological redox reactions,<sup>35</sup> and measuring fast electron-transfer rates.<sup>36</sup> Enabled by MeV electron pulses, this technique induces rapid solvent ionization, followed by electron or hole transfer to the solute of interest. When coupled to an appropriate light source and detector, timeresolved optical absorption measurements can be made on the transiently generated oxidized or reduced intermediates.<sup>3</sup> Importantly, the judicious choice of an appropriate organic solvent can ensure the selective generation of solvated electrons, thereby providing access to short-lived reduced intermediates in a catalytically relevant setting. Pulse radiolysis is particularly well-suited to studying short-lived Ni(I) intermediates, as the rapid injection of a low concentration of charges enables (i) selective one-electron reduction, (ii) generation of monomeric intermediates, and (iii) time-resolved measurements of ns- $\mu$ s time scale follow-on reactions. Given these parameters, we were optimistic that pulse radiolysis could



**Figure 3.** Normalized  $\Delta A$  data for [(dtbbpy)NiBr] generated by pulse radiolysis (see SI, pages S4 and S16, for details) (A),  $\Delta A$  data obtained upon electrolysis of [(dtbbpy)NiBr<sub>2</sub>] (B), and the proposed mechanism of electrolytic Ni(I) generation (C).

be used to generate highly reactive Ni(I) species relevant to recently developed Ni-catalyzed cross-coupling reactions. If successful, the outlined approach would provide a platform for undertaking a wide range of reactivity studies with short-lived Ni(I) intermediates which would otherwise be difficult to study.

To initiate our study of transiently generated Ni(I) intermediates, we first turned our attention to validating the use of pulse radiolysis to generate the proposed Ni(I) bipyridine species. To this end, terpyridine-ligated Ni(I) bromide, previously reported by Vicic and co-workers,<sup>38</sup> was employed as a model system. Because this four-coordinate Ni(I) complex is sufficiently stable to permit traditional stoichiometric organometallic studies,<sup>39</sup> we reasoned that independent synthesis and spectroelectrochemical generation could serve as a means to verify putative Ni(I) signals generated by reductive pulse radiolysis of the corresponding terpyridine-ligated Ni(II)Br<sub>2</sub> precursor,  $[(tpy)NiBr_2]$  (tpy = 2,2':6',2"-terpyridine). For these measurements, N,N-dimethylformamide (DMF) was selected as an optimal solvent, as it is commonly employed in Ni-catalyzed cross-coupling reactions<sup>9,13,28</sup> and generates solvated electrons upon pulse radiolytic ionization (Figure S11).40 Spectroelectrochemical measurements with a solution of [(tpy)NiBr<sub>2</sub>] in DMF showed pronounced differences in absorption features ( $\Delta A$  postelectrolysis vs pre-electrolysis, Figures 2A and S29) at 430, 600, and 860 nm, which align well with the analogous absorbance spectrum obtained from a synthetic sample of [(tpy)NiBr] (Figure 2B). Using the 9 MeV Laser Electron Accelerator Facility (LEAF),<sup>37</sup> a solution of [(tpy)NiBr<sub>2</sub>] (20 mM in DMF) was then irradiated under pulse radiolytic conditions (see Supporting Information (SI), pages S4 and S16, for details), generating  $\Delta A$  features mirroring those observed under electrolytic and synthetic conditions (Figure 2C), validating the use of pulse radiolysis to generate and spectroscopically investigate Ni(I) intermediates.

The electrochemical reduction of [(dtbbpy)NiBr<sub>2</sub>] (dtbbpy = 4,4'-di-*tert*-butyl-2,2'-bipyridine) is irreversible by cyclic voltammetry at moderate scan rates (100 mV/s, Figure S33), and previous studies have revealed that the corresponding

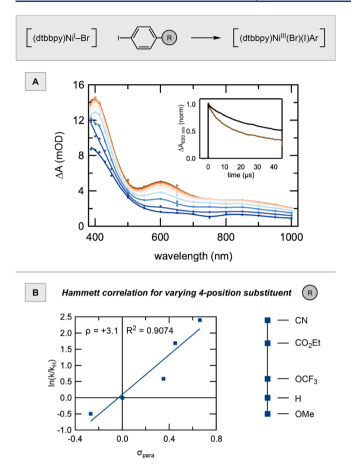


Figure 4. Spectral evolution for Ni(I) signal in the presence of ethyl 4-iodobenzoate (500 mM) (A) and Hammett correlation demonstrating oxidative addition rate sensitivity to aryl iodide electronics (B).

cathodic wave represents a two-electron reduction event  $(E_{p/2}[Ni^{II}/Ni^{0}] = -1.1 V \text{ vs SCE, Figure S33})^{23,41}$  Under preparative electrocatalytic conditions, a Ni(II)/Ni(0) comproportionation step following the initial cathodic reduction has been proposed to generate a catalytically active Ni(I) intermediate relevant to C-N cross-coupling.<sup>23</sup> Additionally, this elementary comproportionation step has been proposed in the context of Ni-catalyzed C-SCF<sub>3</sub> coupling.<sup>42</sup> With these studies in mind, we recognized that pulse radiolysis provided an opportunity to selectively generate single-electron-reduced [(dtbbpy)NiBr], as the pulsed electron beam generates a low total concentration of solvated electrons (~1  $\mu$ M).<sup>37</sup> Meanwhile, spectroelectrochemical investigation of [(dtbbpy)NiBr<sub>2</sub>] reduction would enable spectroscopic measurement of the bulk electrolysis products relevant to the electrocatalytic setting. When irradiated in a manner analogous to that described previously, [(dtbbpy)NiBr<sub>2</sub>] reduction produces a long-lived three-peaked signal with  $\Delta A$  maxima at 430, 620, and 860 nm (Figure 3A). When [(dtbbpy)NiBr<sub>2</sub>] reduction was monitored spectroelectrochemically, the resulting  $\Delta A$  spectrum matched well with the pulse radiolysis data (Figure 3B), indicating that indeed Ni(0)/Ni(II) comproportionation to produce [(dtbbpy)NiBr] is rapid under synthetically relevant electrolysis conditions (Figure 3C). This finding holds implications for Ni-catalyzed cross-coupling processes, as the relevant nickel catalyst oxidation state can have a dramatic impact on feasible elementary organometallic steps.<sup>1,12</sup> Moreover, this

experimental approach provides a method to interrogate the effects that additives in specific Ni-catalyzed reactions have on Ni(0)/Ni(II) comproportionation.

We next turned our attention to investigating the reactivity of [(dtbbpy)NiBr] generated through electrolysis and pulse radiolysis. Oxidative addition between reductively generated bipyridine-ligated Ni(I) intermediates and aryl halides is a commonly invoked elementary step across many nickelcatalyzed arylation reactions.<sup>12</sup> While the use of cyclic voltammetry and spectroelectrochemistry can provide some insight into this step, the two-electron nature of [(dtbbpy)-NiBr<sub>2</sub>] cathodic reduction complicates the interpretation of these results. Nonetheless, reductive generation of low-valent dtbbpy-ligated nickel was carried out in the presence of 4bromobenzotrifluoride, and its subsequent oxidative addition could be monitored by cyclic voltammetry and UV-vis absorption measurements (Figures S1-S4). Unfortunately, the possibility of [(dtbbpy)Ni] aryl halide oxidative addition competing with Ni(0)/Ni(II) comproportionation to generate [(dtbbpy)NiBr] makes it challenging to assign this reactivity to a Ni(0)/Ni(II) or Ni(I)/Ni(III) oxidative addition process (SI, page S5). Pulse radiolysis was then employed to directly probe the rate of aryl halide oxidative addition with [(dtbbpy)NiBr]. Examining the reactivity of 4-bromobenzotrifluoride with [(dtbbpy)NiBr] revealed no significant lifetime change in the Ni(I) signal by pulse radiolysis, providing an upper limit on the oxidative addition rate ( $k_{OA} < 10^4 \text{ M}^{-1} \text{ s}^{-1}$ , Figures S7 and S8). However, moving to the more reactive ethyl 4-iodobenzoate showed significant quenching of the Ni(I) signal ( $k_{OA} = 1.18 (\pm 0.05) \times 10^5 \text{ M}^{-1} \text{ s}^{-1}$ , Figure 4A), consistent with previously measured rate constants for Ni(I) aryl iodide oxidative addition.<sup>17</sup> To ensure efficient Ni(I) generation in quenching experiments involving high concentrations of aryl iodide, 1,3-dicyanobenzene (500 mM) was included as an electron-transfer mediator (see SI, page S12, for details). Rapid initial decay of the 620 nm [(dtbbpy)NiBr] signal leaves behind a long-lived spectrum which can be tentatively assigned to the corresponding Ni(III) aryl oxidative addition adduct (see SI, page S20, for discussion and Figure S16 for Ni(I) data without aryl iodide, showing no spectral evolution on the same time scale).

In order to probe the sensitivity of the Ni(I) oxidative addition rate to substrate electronics, a panel of electronically differentiated aryl iodides were examined by pulse radiolysis. Ni(I) signal decay kinetics were measured in the presence of up to 1000 mM 4-iodoanisole, iodobenzene, 4-(trifluoromethoxy)iodobenzene, ethyl 4-iodobenzoate, or 4-iodobenzonitrile. A plot of  $\ln(k_{OA}/k_{OA}(H))$  vs  $\sigma_p$  for the corresponding 4-substituted iodoarenes reveals a strong dependence of the oxidative addition rate on arene electronics ( $\rho = +3.1$ , Figure 4B), with an 18-fold increase in rate over the studied range ( $\sigma = -0.27$  to +0.66). The rapid reaction of iodobenzene with [(dtbbpy)NiBr] is of particular note, as the corresponding chloro-bridged dimer [(dtbbpy)NiCl]<sub>2</sub> was recently shown by Hazari and co-workers to be unreactive toward iodobenzene oxidative addition.<sup>32</sup>

In order to assess the influence of dimer formation on the rate of oxidative addition by [(dtbbpy)NiBr],  $[(dtbbpy)NiBr]_2$  was independently synthesized, exposed to iodobenzene, and monitored by <sup>1</sup>H NMR analysis (Figure S38). Consistent with results obtained by Hazari and co-workers for the chloro-variant, <sup>32</sup>  $[(dtbbpy)NiBr]_2$  shows no reactivity toward iodobenzene at room temperature up to 16 h. This reactivity

contrasts dramatically with the rate measured for the monomeric form by pulse radiolysis ( $k_{OA} = 2.2 (\pm 0.3) \times 10^4 M^{-1} s^{-1}$ , Figure 5). Spectroelectrochemical measurements

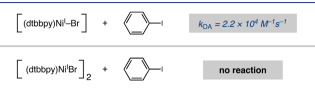
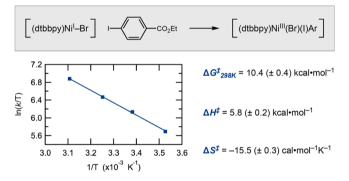


Figure 5. Relative rates of oxidative addition with iodobenzene for monomeric and dimeric bipyridine-ligated Ni(I)Br.

(Figure S9) show that [(dtbbpy)NiBr] dimerization occurs no faster than the minute time scale at catalytically relevant nickel concentrations. Together, these results indicate the importance of considering the immediate reduction products of nickel precatalysts in addition to complexes which may be thermodynamically favored.

To quantitatively assess the barrier to Ni(I) aryl iodide oxidative addition, Ni(I) quenching studies were carried out at various temperatures. Following the method of Eyring, a plot of  $\ln(k_{OA}/T)$  vs 1/T was used to determine the enthalpic  $(\Delta H^{\ddagger} = 5.8 \pm 0.2 \text{ kcal} \cdot \text{mol}^{-1})$  and entropic  $(\Delta S^{\ddagger} = -15.5 \pm 0.3 \text{ cal} \cdot \text{mol}^{-1} \cdot \text{K}^{-1})$  contributions to the overall reaction barrier  $(\Delta G^{\ddagger}_{298 \text{ K}} = 10.4 \pm 0.4 \text{ kcal} \cdot \text{mol}^{-1}$ , Figure 6). This data may



**Figure 6.** Eyring analysis of the oxidative addition between [(dtbbpy)NiBr] and ethyl 4-iodobenzoate.

serve as a useful experimental benchmark for computational investigations of Ni/photoredox-based cross-coupling reactions which have primarily relied on DFT-based oxidative addition barrier heights.<sup>43–45</sup>

The combined use of pulse radiolysis and spectroelectrochemistry has provided insights into fundamental elementary steps relevant to low-valent Ni intermediates commonly encountered in Ni-catalyzed cross-coupling reactions (Figure 7). The finding that Ni(0)/Ni(II) comproportionation to

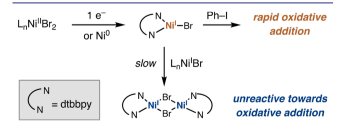


Figure 7. Overview of mechanisms for [(dtbbpy)NiBr] generation and reactivity.

generate bipyridine-ligated Ni(I) is rapid under synthetically relevant conditions holds relevance to the overall mechanism of multiple Ni-catalyzed C-C and C-X (X = N, O) crosscoupling reactions. Additionally, the investigation of Ni(I) oxidative addition by pulse radiolysis revealed the influence of aryl halide electronics on Ni(I) oxidative addition rates and the profound impact that dimer formation has on this elementary step. In particular, transiently generated [(dtbbpy)NiBr] rapidly reacts with iodobenzene, while the dimeric form, [(dtbbpy)NiBr]<sub>2</sub>, is virtually unreactive. This information is especially relevant to reactions wherein high concentrations of low-valent Ni are thought to be deleterious, given the kinetic favorability of dimer formation under such conditions.<sup>46</sup> Taken together, these studies have important implications for reaction design in the area of Ni-catalyzed cross-coupling. More broadly, pulse radiolysis has been demonstrated to serve as a powerful tool for interrogating short-lived catalytic intermediates and measuring fast radical-based reactions relevant to photoredox and electrocatalysis cross-coupling platforms.

## ASSOCIATED CONTENT

## **Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.1c04652.

Experimental procedures, spectroscopic and electroanalytical data, and further discussions (PDF)

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#### Notes

The authors declare no competing financial interest.

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