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# **Bimetallic C–C Bond Forming Reductive Elimination from Nickel**

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**ABSTRACT:** Ni-catalyzed cross-coupling reactions have found important applications in organic synthesis. The fundamental characterization of the key steps in cross-coupling reactions, including C–C bond forming reductive elimination, represents a significant challenge. Bimolecular pathways were invoked in early proposals, but the experimental evidence was quite limited. We present the preparation of well-defined (pyridine-pyrrolyl)Ni mono-methyl and phenyl complexes that allow the direct observation of bimolecular reductive elimination to generate ethane and biphenyl, respectively. The sp<sup>3</sup>-sp<sup>3</sup> and sp<sup>2</sup>-sp<sup>2</sup> couplings proceed via two distinct pathways. Oxidants promote the fast formation of Ni(III) from (pyridine-pyrrolyl)Ni-methyl, which dimerizes to afford a bimetallic Ni(III) intermediate. Our data is most consistent with the subsequent methyl coupling from the bimetallic Ni(III) to generate ethane as the rate-determining step. In contrast, the formation of biphenyl is facilitated by the coordination of a bidentate donor ligand.

#### Introduction

Nickel complexes have emerged as appealing catalysts for cross-coupling reactions (eq 1).<sup>1</sup> While contemporary Ni catalysts employing chelating ligands have started to solve important problems in organic synthesis,<sup>2</sup> catalyst design will benefit from advanced understanding of the reaction mechanism to achieve more widespread utility.<sup>3,4</sup> The diversity of pathways, depending on the ligand framework,

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highlights the challenges associated with characterizing the mechanism for Ni-catalyzed cross-coupling reactions. The two prevailing mechanisms include the Pd-like "closed-shell" redox pathway via Ni(0)/Ni(II) intermediates<sup>5</sup> and the "open-shell" pathway featuring Ni(I)/Ni(III) intermediates.<sup>4,6</sup>

$$\begin{array}{c} X \\ R^1 \stackrel{}{\searrow} R^2 + R^3 - M \xrightarrow{\text{Ni catalyst}} R^3 \\ R^1 \stackrel{}{\searrow} R^2 \\ R^1 \stackrel{}{\swarrow} R^3 \\ R^2 \\ M^2 B, Zn, Mg, Si, Zr, etc. \end{array}$$
(1)

In parallel with catalytic studies, historical and contemporary efforts have focused on stoichiometric characterization of the fundamental steps of nickel-mediated C–C bond coupling. Several Ni(II) and Ni(III) systems have been identified to undergo reductive elimination,<sup>6,7</sup> whereas C–C bond formation from isolated Ni(IV) has just been reported recently.<sup>8</sup> Reductive elimination engaging two metal centers has been determined to promote C–C coupling with Pd and Pt complexes,<sup>9</sup> but such a bimetallic mechanism with Ni has seldom been considered.<sup>10</sup> Yamamoto observed transmetalation between two Ni(II) intermediates prior to a "closed-shell" reductive elimination of biphenyl from Ni(II) to Ni(0) (Scheme 1A).<sup>7a-d</sup> In contrast, early proposals by Kochi and Hegedus described a group transfer between Ni(III) and Ni(II) as the key step, followed by reductive elimination from the Ni(III) intermediate (Scheme 1B).<sup>6,11</sup> This "open-shell" transmetalation was supported by minimal experimental evidence.<sup>12</sup> In light of the fundamental interest and potential applications of bimolecular mechanisms in Ni catalysis, herein we present the direct observation of two distinct bimetallic pathways for Ni-mediated sp<sup>3</sup>-sp<sup>3</sup> and sp<sup>2</sup>-sp<sup>2</sup> C–C coupling. Spectroscopic and kinetic studies, in combination with DFT calculations, provide evidence for high-valent Ni intermediates. Our discovery of new bimetallic mechanisms for Ni-mediated C–C bond formation provides valuable insight for catalyst design in cross-coupling reactions.

# Scheme 1. Proposed Bimetallic Pathways for Ni-Mediated C-C Bond Formation

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(A) "Closed-Shell" Transmetalation (Yamamoto)
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$$2 (bpy)NiII \begin{pmatrix} Br \\ Ph \end{pmatrix} \xrightarrow{- (bpy)NiBr_2} (bpy)NiII \begin{pmatrix} Ph \\ Ph \end{pmatrix} \xrightarrow{- Ni^0} Ph -Ph$$

(B) "Open-Shell" Group Transfer (Kochi and Hegedus)

 $\left[ArNi^{III}Br_{2}\right] + ArNi^{II}Br \xrightarrow{-Ni^{II}Br_{2}} \left[Ar_{2}Ni^{III}Br\right] \xrightarrow{-Ni^{I}Br} Ar-Ar$ 

#### **Results and Discussion**

**Synthesis of mono-Methyl and mono-Phenyl Ni Complexes.** LX-type bidentate ligands are beneficial to Ni-catalyzed cross-coupling reactions,<sup>13</sup> but fundamental understanding of the ligand effect on C–C bond formation has been limited by the challenges associated with isolating Ni-carbyl intermediates.<sup>14</sup> We prepared (py-<sup>Me</sup>pyrr)Ni(acac), **1** (py-<sup>Me</sup>pyrr = 3,5-dimethyl-2-(2-pyridyl)pyrrole, acac = acetylacetonate), which underwent methylation and phenylation with Grignard reagents to afford *cis* and *trans* mixtures of (py-<sup>Me</sup>pyrr)Ni(CH<sub>3</sub>)(lut) **2** and (py-<sup>Me</sup>pyrr)Ni(Ph)(lut) **3** (lut = 2,4-lutidine) (Scheme 2). The *cis* and *trans*-isomers of **2** co-crystallized into a highly disordered packing (Figure S1(B)). Compound **3** did not form single crystals, so we prepared the alternative (py-<sup>Ph</sup>pyrr)Ni(acac) **7** and (py-<sup>Ph</sup>pyrr)Ni(Ph)(lut) **8**. Single crystal X-ray diffraction established a square planar geometry for **2** and **8** (Figure S1 and S2). This geometry implies a low-spin state for Ni and is consistent with the observed diamagnetic <sup>1</sup>H NMR spectra. In solution, the assignments of *cis*- and *trans*-isomers were confirmed by NOESY and COSY experiments (Figure S4). The *cis/trans* ratio remained constant over time, indicating that the preference for the *trans*-geometry was a thermodynamic effect.<sup>15</sup> In the following studies, the isomers were not separated, but used as a mixture.

Scheme 2. Syntheses of (py-<sup>Me</sup>pyrr)Ni Complexes



**Oxidant-Promoted Ethane Formation.** The isolation of **2** has allowed us to carry out fundamental studies on C–C bond formation from the well-defined mono-methyl complex. We initiated our study by evaluating **2** under Yamamoto conditions, where transmetalation followed by biphenyl coupling was reported for (bpy)Ni(Ph)(Br) (bpy = bipyridine) in DMF.<sup>7c</sup> In contrast to this precedent, **2** remained stable in DMF- $d_7$  and C<sub>6</sub>D<sub>6</sub> (Table 1, entries 1-2). We continued to explore the effect of external donor ligands and oxidants. Immediate decomposition of **2** to CH<sub>4</sub> was observed in the presence of five equivalents of

bpy (entry 3).<sup>16</sup> Addition of CH<sub>3</sub>I to **2** resulted in minor conversion upon heating (entry 4). Introducing O<sub>2</sub> to **2** immediately furnished ethane in 19% yield (maximum theoretical yield = 50%) (entry 5).<sup>17</sup> The yield of ethane was increased to 43% with I<sub>2</sub> as the oxidant, with concomitant formation of 10% CH<sub>3</sub>I (entry 6). Addition of NBS (*N*-bromosuccinimide) also led to high yield of ethane (entry 7). Notably, no methane was observed under these oxidative conditions.

 Table 1. Formation of Ethane from 2

_		3 C <sub>6</sub> D <sub>6</sub> oxidant/add	→ CH <sub>4</sub>	+ CH <sub>3</sub> Cł	H <sub>3</sub>
Entry	Oxidant or	Temp (°C)	Time (h)	Yield	d (%) <sup>a</sup>
	auunive			$CH_4$	CH <sub>3</sub> CH <sub>3</sub>
1 <i>b</i>	none	22	12	0	0
2	none	50	4	0	0
3	bpy (5 equiv)	50	4	60	trace
4	CH <sub>3</sub> I (1 equiv)	50	4	4	4
5	O <sub>2</sub> (5 equiv)	22	0.1	0	19 ± 2
6	l <sub>2</sub> (0.5 equiv)	22	0.1	0	43 <sup>c</sup>
7	NBS (1 equiv)	22	0.1	<1	45 <sup>d</sup>

<sup>*a*</sup> Internal standard = hexamethyldisiloxane (TMS<sub>2</sub>O). The yield of ethane is based on molar concentration (the maximum yield is 50%). Methane and ethane in the gas phase are calculated using Henry's law. <sup>*b*</sup> In DMF- $d_{7}$ .

<sup>c</sup> CH<sub>3</sub>I was observed in 10% yield.

<sup>d</sup> CH<sub>3</sub>Br was observed in 2% yield.

The selective ethane formation from **2** under oxidative conditions provided preliminary support for a bimolecular reductive elimination process. We continued our investigation to obtain further evidence for a bimolecular pathway. A crossover experiment was performed in which a mixture of **2** and **2**-*d*<sub>3</sub> in a 1:1 ratio reacted with I<sub>2</sub> to generate a mixture of CH<sub>3</sub>CH<sub>3</sub>, CH<sub>3</sub>CD<sub>3</sub>, and CD<sub>3</sub>CD<sub>3</sub> in a statistical distribution (eq 2). The ratio of ethane to CH<sub>3</sub>I exhibited a linear dependence on [**2**] with a positive slope (Figure 1). This influence of [**2**] on the product distribution reflects the requirement for two molecules of **2** to form ethane, whereas the formation of CH<sub>3</sub>I is uni-molecular in **2**.





Figure 1. The dependence of [ethane]/[CH<sub>3</sub>I] ratio on [2]. Reaction conditions: [2] = 5-40 mM,  $[I_2] = 20$  mM,  $C_6D_6 = 0.6$  mL, 22 °C, 1 h. Internal standard = TMS<sub>2</sub>O. Estimated standard deviation for individual points:  $\leq 5\%$ .

Experimental and Theoretical Characterization of Intermediates. (A) Spectroscopic Data. Our subsequent spectroscopic studies on I<sub>2</sub>-promoted ethane formation from 2 provided structural information on the Ni intermediates and shed some light on the reaction mechanism. The cyclic voltammetry of 2 exhibited two irreversible oxidation waves at 12 and 411 mV (vs. Fc/Fc<sup>+</sup>) (Figure S5). In a liquid N<sub>2</sub> filled cold well, 2 and I<sub>2</sub> were mixed and the reaction mixture was immediately analyzed by EPR spectroscopy. The EPR spectrum displayed a single species of an S = 1/2 state with a 1:1:1 superhyperfine splitting (Figure 2, red).<sup>18</sup> To determine the lifetime of this intermediate, we allowed the mixture to react at -20 °C for 10 minutes and examined it by EPR spectroscopy. The signal corresponding to the intermediate decayed to zero. When NBS was used to replace I<sub>2</sub> as the oxidant, an analogous intermediate was observed (Figure 2, black). The features and splitting patterns for the two intermediates, generated from I<sub>2</sub>

and NBS (red and black, respectively), are similar, but the g values are slightly different, reflecting two distinct complexes with a similar structure.



**Figure 2.** X-band EPR spectra of the intermediate generated from the reaction of **2** with I<sub>2</sub> (red) and the reaction of **2** with NBS (black) recorded in toluene glass at 10 K. The blue spectrum (top) is the simulation of the red spectrum. Spectroscopic parameters:  $g_x = 2.20$ ,  $g_y = 2.14$ ,  $g_z = 2.03$ ,  $A_{xx} = 7.7$  MHz,  $A_{yy} = 63$  MHz,  $A_{zz} = 55$  MHz. Microwave frequency = 9.380 GHz, power = 0.25 mW, modulation amplitude = 1 mT/100 kHz.

The short-lived Ni(III) intermediate detected by EPR spectroscopy prompted us to explore its structure by DFT calculations. Geometry optimization of possible Ni(III) intermediates converged to (py- $^{Me}$ pyrr)Ni<sup>III</sup>(CH<sub>3</sub>)(I)(lut), **4** (Figure 3). This structure features a square pyramidal geometry with pyridine in the axial position and iodide *trans* to pyrrole. The spin-density plot reveals a Ni-centered radical strongly coupled with the *N*-atom of pyridine, whereas its couplings with pyrrole and lutidine are minor. The EPR parameters of **4** were calculated using the ORCA package,<sup>19</sup> and the computational g values and A<sub>iso</sub> reproduced the experimental spectra shown in Figure 2 (Table S1).



Figure 3. Structures and Mulliken spin-density plots of 4 obtained from DFT calculations.

Variable temperature NMR experiments allowed us to identify the diamagnetic intermediates. In a liquid nitrogen-filled cold well, upon injection of I<sub>2</sub>, 2 rapidly evolved into a diamagnetic intermediate with the concomitant formation of CH<sub>3</sub>I in 10% yield and minor paramagnetic species (Figure S7A). While the complete conversion of this intermediate to ethane was fast at 22 °C (Figure S7B), the intermediate was stable on the NMR time scale at -20 °C (Figure S8). Multiple singlet resonances in the upfield region between 2.0 - 3.0 ppm are accompanied with several aromatic protons between 5.5 - 7.0ppm. COSY experiments elucidate that there are a number of resonances overlap with the residual toluene solvent (Figure S9). To determine the identity of this intermediate and assign the spectrum, we prepared deuterium labeled (py-<sup>Me</sup>pyrr)Ni-methyl complexes,  $2-d_3$ ,  $2-d_1$ , and  $2-d_7$  (Figure 4). Complex  $2-d_7$  is a mixture of isotopologues and isotopomers. The overall deuterium incorporation in  $2-d_7$  is 60% at each position. When 2-d<sub>3</sub> was treated with I<sub>2</sub>, four signals at 3.00, 2.94, 2.91, and 2.85 ppm disappeared (Figure 5A). HSQC experiments revealed that the carbon chemical shifts corresponding to these proton resonances were between 26.00 - 29.00 ppm (Figure S10). The concentration of these four proton resonances, in combination with CH<sub>3</sub>I, accounted for 95% of the Ni- $CH_3$  from the starting **2**. Moreover, the overall integration of these four signals is half the overall integration of eight singlet peaks between 2.30 and 2.60 ppm (Figure S8). The reaction of  $2-d_1$  led to the disappearance of two singlet <sup>1</sup>H resonances at 6.14 and 6.17 ppm, while all other resonances remained (Figure 5B). When  $2-d_7$  was submitted to the reaction conditions, all eight singlet resonances between 2.30 and 2.60 ppm diminished with small residuals due to the incomplete deuterium incorporation to  $2-d_7$  (Figure 5C).



Figure 4. Deuterium labeled 2 for structure assignment of the diamagnetic intermediate.







6.75	6.65	6.55	6.45	6.35	6.25	6.15	6.05	5.95	5.85	5.75
					it (bbiii)	/				



05 3.00 2.95 2.90 2.85 2.80 2.75 2.70 2.65 2.60 2.55 2.50 2.45 2.40 2.35 2.30 f1 (ppm)

**Figure 5.** Comparison of the <sup>1</sup>H NMR spectra of the intermediate generated from I<sub>2</sub> reacting with proteo and deuterio (py-pyrr)Ni-methyl complexes  $2-d_3(A)$ ,  $2-d_1(B)$ , and  $2-d_7(C)$ . Reaction conditions: [2] = 16 mM, [I<sub>2</sub>] = 16 mM, toluene- $d_8$  = 0.65 mL, -20 °C. Internal standard = 1,3,5-trimethoxybenzene.

The <sup>1</sup>H NMR resonances corresponding to the diamagnetic intermediate gradually decrease at  $-20 \,^{\circ}$ C, while a singlet resonance at 0.820 ppm, corresponding to ethane, increases. The concentration of CH<sub>3</sub>I remains constant. We were able to monitor the decrease of the intermediate and increase of ethane over 3–4 hours. The concentration of the Ni–CH<sub>3</sub> at 3.00 ppm exhibits an exponential decay (Figure 6A).<sup>20</sup> The concentration of this species, however, only accounts for a fraction of ethane. When all four Ni-CH<sub>3</sub> resonances identified in Figure 5A at 3.00, 2.94, 2.91, and 2.85 ppm are integrated, half of the total concentration of ethane.<sup>21</sup> Other models, including second order kinetics, were fully considered but failed to fit the experimental data. When the conversion of the intermediate to ethane was complete, the resulting Ni species were NMR silent (Figure S7B). The EPR spectrum revealed the absence of any radical species. Independently prepared (py-pyrr)<sub>2</sub>Ni<sup>II</sup> and the comparison of the spectra revealed that (py-pyrr)<sub>2</sub>Ni<sup>II</sup> is absent from the reaction (Scheme 3, *cf.* Figures S11-S12). Alternatively, tetrahedral (py-pyrr)Ni-iodides are possible products (Scheme 3). Our attempts to prepare (py-pyrr)Ni-iodides via multiple methods were unsuccessful, consistent with previous reports by Caulton.<sup>14b,22</sup>



**Figure 6.** Reaction time courses for the consumption of the Ni–CH<sub>3</sub> resonance at 3.00 ppm (2[5a]) (A) and the conversion of 5 to ethane (B). The concentration of 5 ([5] = [5a] + [5b] + [5c]) was calculated by dividing the total concentration of four Ni-CH<sub>3</sub> resonances at 3.0, 2.94, 2.91, and 2.85 ppm by 2 (see discussions below). Reaction conditions:  $[2]_0 = 17$  mM,  $[I_2]_0 = 15$  mM, toluene- $d_8 = 0.65$  mL, -20 °C. Internal standard = 1,3,5-trimethoxybenzene.

Scheme 3. Proposed Ni Products from the Reaction of 2 with I<sub>2</sub>



Proposed Ni product

(B) Structural Assignment of Intermediate 4. The irreversible CV of 2 at 12 and 411 mV (vs. Fc/Fc<sup>+</sup>) suggests that the high-valent Ni species generated from electrochemical oxidation are unstable.

Nevertheless, the data provides an estimation of the oxidation potentials for Ni(II)/Ni(III) and Ni(III)/Ni(IV) transitions. Previous reports indicate that the oxidation potential of  $I_2$  is 140 mV (vs. Fc/Fc<sup>+</sup>),<sup>23</sup> implying that  $I_2$  is suitable for oxidizing **2** to Ni(III) instead of Ni(IV). This analysis is in alignment with previous experimental and computational studies on the oxidation of Ni(II) to Ni(III) by Br<sub>2</sub>.<sup>24</sup> The EPR experiments provide further support for the oxidation of **2** to a Ni(III) intermediate upon addition of  $I_2$ . Replacing  $I_2$  with NBS leads to a minor shift of the g values while the overall shape of the EPR signal remains. This observation suggests that the Ni(III) intermediate possesses a halide ligand.<sup>25</sup> DFT calculations allow us to propose (py-<sup>Me</sup>pyrr)Ni<sup>III</sup>(I)(CH<sub>3</sub>)(lut) **4** as the possible structure (Figure 3). The square pyramidal geometry is consistent with a Ni(III)-halide previously isolated by van Koten.<sup>25b</sup> The calculated EPR parameters for **4** are in good agreement with the experimental values (Table S1). The observed 1:1:1 superhyperfine splitting is due to the strong coupling of the unpaired electron on Ni with the N of pyridine (I = 1) in the axial position. The lifetime of **4** at -20 °C is short, as its EPR signal has completely decayed after 10 minutes.

(C) Structural Assignment of Intermediate 5. Low-temperature <sup>1</sup>H NMR studies reveal a diamagnetic intermediate 5 with a lifetime of 3–4 hours at –20 °C. The assignment of the peaks is facilitated by the deuterium labeling studies, 2D NMR experiments, and literature precedents. The reaction of 2- $d_3$  leads to the disappearance of four peaks at 3.00, 2.94, 2.91, and 2.85 ppm (Figure 5A), suggesting four Ni– $CH_3$  resonances exist for intermediate 5. Consistently, the total concentration of these methyl groups, along with the small amount of CH<sub>3</sub>I by-product, accounts for 95% of the starting Ni- $CH_3$  from 2. The concentration of each individual Ni– $CH_3$  resonance (3.00 ppm, for instance) undergoes first order decay at –20 °C (Figure 6A). Half of the overall concentration of the four Ni– $CH_3$  resonances fit to a first order kinetic model, which accounts for the formation of ethane (Figure 6B). These kinetic data are consistent with two Ni– $CH_3$  groups in each molecule of 5, which couple in an intramolecular fashion to generate ethane.

The <sup>1</sup>H chemical shifts of Ni–CH<sub>3</sub> and the <sup>13</sup>C resonances of the corresponding Ni–CH<sub>3</sub> are unusually downfield for methyl resonances attached to metals (Figure 5A and Figure S10), implying an electrondeficient Ni center. Previous reports of high-valent Ni complexes display an analogous trend (Scheme 4).<sup>8,26</sup> The experiment with  $2-d_7$  reveals that all eight methyl peaks between 2.30 and 2.60 ppm belong to the py-pyrr ligand (Figure 5C). The integrations suggest that the ratio of Ni-CH<sub>3</sub>/py-pyrr is 1:1. These considerations, coupled with the kinetic data, lead us to propose the dimeric  $[(py-pyrr)Ni^{III}(CH_3)(\mu-I)]_2$  as the structure intermediate 5 (Scheme 5). The diamagnetic property of 5 arises from the antiferromagnetic coupling of low-spin Ni(III) ( $d^7$ , S = 1/2) centers.<sup>27</sup> The integrations of the <sup>1</sup>H NMR resonances and 2D NMR experiments suggest that three isomers exist. Duplicate experiments show different ratios of the isomers, suggesting a distribution originated from a kinetic effect instead of thermodynamic equilibrium. DFT calculations on possible stereoisomers converged to two symmetric structures that have the lowest energy, 5a and 5b (Scheme 5). The Ni centers favor a square pyramidal geometry with pyridine occupying the axial position. NMR data determined that the third isomer has two inequivalent Ni-CH<sub>3</sub> and two sets of py-pyrr ligands. DFT calculations converged to 5c, in which one Ni adapts a square pyramidal geometry and the other is trigonal bipyramidal with an iodide and a methyl located at the axial position (Scheme 5). The assignment of the <sup>1</sup>H resonances for each isomer is presented in Scheme 5. The assignments for 5c in brackets are arbitrary, since we do not have a way to distinguish between the two inequivalent py-pyrr ligands. The missing resonances are attributed to peaks underneath the residual toluene solvent. The calculated chemical shifts of Ni– $CH_3$  in each isomer are 3.75, 3.57, 2.87 and 2.77 ppm, respectively, in agreement with the experimental values.

# Scheme 4. Reported <sup>1</sup>H Chemical Shifts of High-Valent Ni Complexes



Scheme 5. Proposed Structures for Intermediate 5 and Assignments of <sup>1</sup>H NMR Resonances

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Alternatively, Ni(IV)-dimethyl complexes are considered as the structure of **5**, and can be ruled out by our data. (Py-pyrr)Ni<sup>IV</sup>(CH<sub>3</sub>)<sub>2</sub>(lut)(I) **5d** is a possible intermediate (Scheme 6). The integration of Ni– *CH<sub>3</sub>*/py-pyrr for **5d** is 2:1, inconsistent with the observed Ni–*CH<sub>3</sub>*/py-pyrr ratio of 1:1 by NMR labeling studies. (Py-pyrr)<sub>2</sub>Ni<sup>IV</sup>(CH<sub>3</sub>)<sub>2</sub> **5e** would fit the kinetic and NMR data. The reductive elimination from **5e** to form ethane is expected to generate (py-pyrr)<sub>2</sub>Ni<sup>II</sup> as the resulting Ni species. The lack of (py-pyrr)<sub>2</sub>Ni<sup>II</sup> after the reaction suggests that **5e** is absent (Scheme 3, *cf*. Figures S11-S12). In addition, the formation of **5e** requires the transfer of the py-pyrr ligand and methyl from one molecule of **4** to another. This process is highly unlikely. The mixed-valent (py-pyrr)Ni<sup>IV</sup>(CH<sub>3</sub>)<sub>2</sub>( $\mu$ -I)<sub>2</sub>Ni<sup>II</sup>(py-pyrr) **5f** would fit the kinetic data. But the py-pyrr ligands in any stereoisomers of **5f** are inequivalent, inconsistent with the symmetry of the three isomers determined by <sup>1</sup>H NMR spectroscopy. Moreover, we performed DFT calculations on **5f**, which converged to an octahedral Ni(IV) bridged with a tetrahedral Ni(II). The tetrahedral Ni(II) is highspin, contradictory to the observed diamagnetic NMR spectrum. The calculated Ni–*CH*<sub>3</sub> <sup>1</sup>H NMR chemical shifts are presented in Scheme 6, which are significantly more downfield compared to the experimental values.

# Scheme 6. Ni(IV) Complexes Ruled Out as the Structure of 5 and the Corresponding Ni– $CH_3$ <sup>1</sup>H NMR Chemical Shifts Predicted by DFT Calculations



**Mechanism of Oxidant-Promoted Ethane Formation.** In the studies presented above, we explored and characterized the C–C bond formation process from (py-pyrr)Ni mono-methyl complex **2**. The lack of reactivity of **2** in DMF contradicts Yamamoto's "closed-shell" pathway.<sup>7</sup> The absence of ethane in the presence of  $CH_3I$  reveals a pathway distinct from simple nucleophilic substitution<sup>28</sup> or oxidation of Ni(II) by  $CH_3I$ .<sup>2a</sup> Instead, **2** undergoes reductive elimination to generate ethane in the presence of oxidants. This reaction is highly selective, since methane is absent from the reaction. The I<sub>2</sub>-promoted ethane formation is bimolecular, evident in the crossover experiment and the increased [ethane]/[ $CH_3I$ ] ratio as a function of [**2**].

On the basis of the structural assignment of the intermediates above, we propose the mechanism for  $I_2$ -promoted ethane formation (Scheme 7). The process is initiated by the oxidation of **2** to afford a Ni(III) intermediate, assigned as (py-<sup>Me</sup>pyrr)Ni<sup>III</sup>(I)(lut)(CH<sub>3</sub>) **4** (step *i*) (Scheme 7). Intermediate **4** has a short lifetime and rapidly undergoes further conversion. Four possible pathways are considered for the subsequent ethane formation. In the free radical pathway (Scheme 7, pathway A), **4** undergoes a Ni– carbon bond homolysis to eject a methyl radical, which dimerizes to generate ethane. This pathway is inconsistent with the absence of methane after the reaction, since methane is an expected product from the ejection of a methyl radical.<sup>29</sup> Moreover, monitoring the reaction by <sup>1</sup>H NMR spectroscopy clearly reveals the conversion of organometallic Ni–CH<sub>3</sub> intermediates to ethane. The radical mechanism is inconsistent with the clean first order kinetics. In comparison, the reaction of **2** with bpy is speculated to form a

methyl radical, which results in methane as the major product (Table 1, entry 3) (see below for further discussion about ligand-promoted radical ejection).

Scheme 7. Proposed Mechanism for I2-Promoted Reductive Elimination of Ethane



An alternative mechanism involves a methyl transfer from **4** to **2** to afford a Ni(III)-dimethyl intermediate (pathway B). This pathway is a reminiscent of Kochi and Hegedus' proposal.<sup>6</sup> The rapid decay of paramagnetic intermediate **4** into EPR silent species is contradictory to a Ni(III)-dimethyl intermediate. Instead, a diamagnetic intermediate **5** converts to ethane over 3–4 hours at –20 °C. Pd(III) has been reported to undergo disproportionation to afford Pd(IV), followed by reductive elimination to form ethane.<sup>9</sup> We evaluated the analogous pathway (pathway C). The possible Ni(IV) complexes, generated from disproportionation, can be ruled out as the structure of intermediate **5** based on direct and circumstantial evidences presented above (Scheme 6).

The NMR and kinetic data fit well to  $[(py-pyrr)Ni^{III}(CH_3)(\mu-I)]_2$  **5**, which is generated from the dissociation of 2,4-lutidine from **4** (step *ii*, pathway D). The reductive elimination from **5** to furnish ethane is first order, and is the rate-determining step (step *iii*). The precise transition state for the C–C bond formation is unknown. We speculate that the dissociation of the  $\mu$ -iodide occurs prior to methyl coupling. The lack of S = 1/2 species after the reaction is evident from the EPR studies, which implies a Ni(II) as the resulting Ni product. The broad NMR signals suggest that the resulting Ni(II) is high-spin. The competing reductive elimination of CH<sub>3</sub>I from **4** (step *vi*) takes place in parallel with dimerization

(step *ii*). Once **5** is formed, the bridging  $\mu$ -iodide inhibits CH<sub>3</sub>I reductive elimination, consistent with the constant [CH<sub>3</sub>I] after the initial period of the reaction. The favorable formation of ethane relative to CH<sub>3</sub>I is a result of the fast dimerization (step *ii*).

**Donor Ligand-Promoted Bimolecular Biphenyl Formation**. (Py-<sup>Me</sup>pyrr)Ni(Ph)(lut), **3**, underwent C–C bond formation via a different pathway, relative to **2**. Complex **3** remained intact in DMF- $d_7$  and C<sub>6</sub>D<sub>6</sub> (Table 2, entries 1-2). Addition of bpy afforded biphenyl in 49% yield upon heating (maximum theoretical yield = 50%) (entry 3). Neither addition of CH<sub>3</sub>I (entry 4) nor O<sub>2</sub> (entry 5) to **3** afford biphenyl or toluene. Repeating the experiment in toluene- $d_8$  resulted in the absence of benzene and phenol. Addition of I<sub>2</sub> resulted in PhI in high yield (entry 6).

#### Table 2. Formation of Biphenyl from 3

		∠Ph ∑lut oxida	C <sub>6</sub> D <sub>6</sub>	► Ph–Ph + Ph	nl
Entry	Oxidant or	Temp (°C)	Time (h)	Yield (%) <sup>a</sup>	
	auullive			Ph-Ph Phi	PhCH <sub>3</sub>
1 <i><sup>b</sup></i>	none	22	12	0	
2	none	50	4	trace	
3	bpy (5 equiv)	50	4	49	
4	CH₃I (1 equiv)	50	4	0	0

<sup>*a*</sup> TMS<sub>2</sub>O was added as the internal standard. The yield of biphenyl is based on molar concentration (the maximum yield is 50%). <sup>*b*</sup> In DMF- $d_{7}$ .

0.1

0.1

The high yield of biphenyl in the presence of bpy prompted us to undertake a further characterization of the reaction pathway and the intermediate. The crossover experiment between **3** and  $(py-^{Me}pyrr)Ni(p-Tol)(lut)$  afforded a mixture of biphenyl, 4-phenyltoluene, and 4,4'-dimethylbiphenyl in a ratio comparable to the statistical distribution (eq 3). While one equivalent of phenanthroline was capable of fully displacing lutidine from **3** to afford the penta-coordinated  $(py-^{Me}pyrr)Ni(Ph)(phen)$  **10** (Figure S3(B)), ligand substitution between **3** and bpy favored dissociation (eq 4). The equilibrium constant was determined to be 0.019 ± 0.002 at 22 °C, by integrating the <sup>1</sup>H resonances of **3** and **6** in the presence of different concentrations of 2,4-lutidine. The full conversion of **3** to **6** was achieved by removing 2,4-

 $O_2$  (5 equiv)

 $I_2$  (0.5 equiv)

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lutidine under vacuum followed by re-dissolving in  $C_6D_6$  five times. At 22 °C, 6 evolved biphenyl in full conversion after 2 h.

$$3 + \binom{L}{X} Ni \stackrel{p-\text{Tol}}{\underset{\text{lut}}{}} \xrightarrow{\text{bpy}} Ph-Ph + Ph-Tol + Tol-Tol \qquad (3)$$

$$1: 2.7 \qquad \text{Statistical:} \qquad 1: 5.4: 7.3$$

$$\text{Experimental:} \quad 1: 5.0: 6.3$$

$$\mathbf{3} + bpy \xrightarrow[(22 \circ C)]{-lutidine, C_6D_6}_{K_{eq}} \underbrace{\mathbf{0.019 \pm 0.002}}_{(22 \circ C)} \underbrace{\mathbf{N}}_{\mathbf{6}} \underbrace{\mathbf{N}}_{\mathbf{N}} \underbrace{\mathbf{N}} \underbrace{\mathbf{N}} \underbrace{\mathbf{N}} \underbrace{\mathbf$$

EPR analyses provided additional insight to the mechanism of this transformation. The reaction of  $(py-^{Me}pyrr)Ni^{II}(Me)(lut)$ , **2**, with 1 equiv. of  $(bpy)Ni^{0}(cod)$  (eq 5) exhibited a rhombic signal corresponding to an S = 1/2 Ni species at 50 K in toluene glass (Figure 7, red spectrum). An organic radical is present, which could result from the decomposition of Ni(I). This paramagnetic species is assigned to  $(py-^{Me}pyrr)Ni^{I}(bpy)$ , generated in-situ by the comproportionation of Ni(II) with Ni(0). The EPR spectrum of the reaction mixture of **3** with bpy displayed the same species as  $(py-^{Me}pyrr)Ni^{I}(bpy)$  (Figure 7, blue spectrum).





**Figure 7.** X-band EPR spectra recorded in toluene glass at 50 K for the reaction mixture of **3** with bpy (blue) and the comproportionation reaction (eq 5) (red). The black spectrum is the simulation of the red spectrum:  $g_x = 2.27$ ,  $g_y = 2.11$ ,  $g_z = 2.04$ ,  $A_{xx} = 3.7$  MHz,  $A_{yy} = 26$  MHz,  $A_{zz} = 8.5$  MHz.

The statistical distribution of products from the crossover experiment (eq 3) provides crucial evidence for a bimolecular pathway for bpy-promoted biphenyl formation. Increasing the coordination number of a transition metal complex has been reported to trigger the ejection of alkyl/aryl radical via steric interactions.<sup>30</sup> Free phenyl radicals are not involved here, since benzene is not an observed product. Instead, a phenyl group transfer from **6** to **3** can be proposed to generate a Ni(III)-diphenyl intermediate, which undergoes reductive elimination. This hypothesis is consistent with  $(py-^{Me}pyrr)Ni^{I}(bpy)$  as the resulting Ni species; however, other pathways cannot be ruled out. For example, bpy may bridge between two molecules of **3** to facilitate the bimolecular reductive elimination. Our ongoing efforts focus on elucidation of the precise role of bpy in this reaction by engaging in DFT calculations.

### Conclusion

Collectively, we have established two bimolecular pathways for Ni-mediated C–C bond formation. The sp<sup>3</sup>-sp<sup>3</sup> coupling of Ni–CH<sub>3</sub> to form ethane is initiated by oxidation to generate Ni(III), which dimerizes to afford the diamagnetic, bimetallic Ni(III)–CH<sub>3</sub> intermediate. The resulting Ni(III)–CH<sub>3</sub> dimer undergoes rate-determining reductive elimination to afford ethane (Scheme 7). The bimolecular pathway is verified by crossover experiments and the increased [ethane]/[CH<sub>3</sub>I] ratio as increasing [Ni–CH<sub>3</sub>]. The initial Ni(III) intermediate is characterized by EPR spectroscopy, and evidence for the bimetallic [Ni(III)–CH<sub>3</sub>]<sub>2</sub> include <sup>1</sup>H NMR experiments and kinetic studies. The experimental data are in good agreement with DFT calculations. Biphenyl formation, in contrast, is triggered by the coordination of a bidentate donor ligand (eq 4). While EPR characterization of the resulting Ni(I) species supports a phenyl transfer between two Ni(II) intermediates to generate a Ni(III)-diphenyl intermediate, the precise mechanism requires further investigation. These two bimetallic reductive elimination pathways presented here highlight the complexity of Ni-mediated C–C bond formation reactions. The elucidation of the

bimolecular mechanism provides important insight for future design of active Ni catalysts for crosscoupling and homo-coupling reactions.

### ASSOCIATED CONTENT

**Supporting Information**. All experimental procedures, additional discussions and figures, details of DFT calculations, NMR spectra, and crystallographic data (PDF, CIF). This material is available free of charge via the Internet at http://pubs.acs.org.

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The authors declare no competing financial interest.

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