Synthesis of Nickel(I)–Bromide Complexes via Oxidation and Ligand Displacement: Evaluation of Ligand Effects on Speciation and Reactivity

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ABSTRACT: Nickel's +1 oxidation state has received much interest due to its varied and often enigmatic behavior in increasingly popular catalytic methods. In part, the lack of understanding about Ni¹ results from common synthetic strategies limiting the breadth of complexes that are accessible for mechanistic study and catalyst design. We report an oxidative approach using tribromide salts that allows for the generation of a well-defined precursor, [Ni¹(COD)Br]₂, as well as several new Ni¹ complexes. Included among them are complexes bearing bulky monophosphines, for which structure–speciation relationships are established and catalytic reactivity in a Suzuki-Miyaura coupling (SMC) is investigated. Notably, these routes also allow for the synthesis of well-defined monomeric ^{t-Bu}bpy-bound Ni¹ complexes, which has not previously been achieved. These complexes, which react with aryl halides, can enable previously challenging mechanistic investigations and present new opportunities for catalysis and synthesis.

INTRODUCTION

Nickel-catalyzed cross-couplings have emerged as broadly useful methodologies in synthetic organic chemistry—in large part due to the complementary reactivity of Ni to precious metals like Pd.^{1,2} In addition to Ni's ability to facilitate challenging oxidative addition reactions,^{3,4} its relative propensity⁵ to access open-shell electron configurations (i.e., Ni¹ & Ni¹¹¹) underlies many modern cross-coupling reactions (e.g., Ni/photoredox,⁶ Ni/electrocatalysis,⁷ crosselectrophile coupling⁸). Conversely, this same property can attenuate turnover in traditional Ni⁰/Ni¹¹ cycles, where Ni¹ species have been identified as off-cycle.⁹ It is therefore of great interest to understand the nuanced factors that control the formation, reactivity, and speciation of Ni¹ intermediates.^{10,11}

Stoichiometric synthesis of well-defined Ni¹ complexes presents an opportunity to investigate these nuances and characterize species that are relevant in catalysis. Examples of such investigations have examined several ligand classes on Ni¹ centers: Matsubara¹² and Schoenebeck¹³ have studied N-heterocyclic carbene (NHC) ligands, Hazari9,14-16 and Schoenebeck¹⁷ bisphosphine ligands, and Hazari^{18,19} and Martin^{19,20} phenanthroline ligands. For each ligand class, tractable conclusions about the role(s) of Ni¹ species in the respective catalytic mechanisms have been put forth. However, existing strategies for synthesizing Ni¹ species (Figure 1A) limit the breadth of complexes that can be accessed with catalytically relevant ligands. We postulate two contributing factors therein: (1) common preparations for Ni¹ species necessitate the formation of L_nNi¹¹X₂ complexes that are intractable for bulky ligands²¹ and (2) components of [Ni⁰] and [Ni¹¹] precursors used for Ni¹ synthesis can induce undesired redox equilibria and speciation effects.^{20,22} These limitations are borne out in a literature survey of Ni¹

complexes: certain structural motifs are well-explored while others remain elusive (Figure 1B).

Bulky monophosphines are among the ligand classes that have scarcely been characterized on Ni¹ centers. Notably, small monophosphine ligands have been known to form monomeric (PR₃)_nNi¹X (n = 2 or 3) species via comproportionation and oxidative addition,^{23,24} but dimeric [(PR₃)Ni¹X]₂ complexes bearing larger phosphine congeners were unknown prior to a 2022 report from our group.^{25,26} This is in stark contrast to precedent for Pd¹, for which complexes of the type [(PR₃)Pd¹X]₂ are long-known²⁷ and have been thoroughly studied by Schoenebeck and coworkers.^{28,29} As precedent for bisligated (PR₃)₂Ni¹ species has shown that such species are capable of catalytic cycle (re)entry, ^{9,14-16} the structure and speciation of monoligated (PR₃)Ni¹ complexes warrants further investigation.

Moreover, while 4,4'-di-*tert*-butylbipyridine (^{t-Bu}bpy) is the ligand of choice for many Ni-catalyzed cross-couplings that invoke open-shell intermediates,³⁰⁻³³ only two examples of formal (^{t-Bu}bpy)Ni¹ species have been structurally characterized: Hazari's [(^{t-Bu}bpy)Ni¹Cl]₂ and Nocera's [(^{t-Bu}bpy)Ni^{1.5+}(quinuclidine)Cl]₂Cl.^{18,34} Importantly, these complexes do not replicate the behavior of (^{t-Bu}bpy)Ni¹ species in catalysis due to irreversible dimerization of reactive (^{t-Bu}bpy)Ni¹X. While *in situ* generation of monomeric (^{t-Bu}bpy)Ni¹ species has clarified the reactivity of important catalytic intermediates,^{35,36} these approaches are more challenging to apply to future catalyst design and mechanistic studies than stoichiometric synthesis.

Strategies to enable the synthesis of previously unknown Ni¹ complexes create downstream opportunities for more effective Ni catalysis. Contributions from Morandi and coworkers, who recently reported a phenoxide-bound Ni¹ precursor,³⁷ have already led to mechanistic insight and development in Ni-catalyzed methodologies.³⁸ Further goals for the synthetic organometallic community in this area include: the complexation of catalytically relevant ligands to structurally novel Ni¹ centers, the incorporation of easily derivatized X-type ligands such as halides, and application of synthetic discoveries to catalytic systems.

Herein, we present the synthesis of over 20 previously unidentified Ni¹ complexes bearing olefins and catalytically relevant monophosphines and bipyridine ligands. We access these complexes through a mild one-pot oxidation or via ligand displacement of well-defined Ni¹ precursors. For the bulky monophosphine class, we conduct a structurespeciation analysis and catalytic studies to elucidate these ligands' behavior on Ni¹ centers. For the bipyridine class, we identify the first examples of monomeric (*t*-Bubpy)Ni¹ complexes and investigate their electronic structure and stoichiometric reactivity with aryl halides.



Figure 1. (A) Ni¹ complex synthetic strategies and their prevalence. (B) Examples of structurally characterized Ni¹ complexes for (I) phosphine ligands, (II) bipyridine/phenanthroline ligands, and (III) olefin ligands.

RESULTS AND DISCUSSION

In our recent study of dialklylbiaryl phosphine ligands in Ni catalysis, we serendipitously identified the structure of a unique monophosphine-bound Ni¹ halide dimer.²⁵ This species, [(CyJohnPhos)Ni¹Cl]₂ (**1-Cl**), formed in trace amounts following oxidative addition of 2-chlorotoluene to

(CyJohnPhos)₂Ni⁰. To the best of our knowledge, **1-Cl** was the first structurally characterized example of a [(PR₃)Ni¹X]₂ dimer. To further investigate the role of **1** in the catalytic system of interest, we sought to synthesize this complex independently by comproportionation. However, comproportionation attempts with several Ni⁰ and Ni^{II} precursors failed, perhaps due to challenges forming phosphine-bound Ni^{II}Cl₂ species. Taking inspiration from studies from Fout³⁹ and Uyeda⁴⁰, we were able to isolate [(CyJohnPhos)Ni^IBr]₂ (**1-Br**) by oxidizing (CyJohnPhos)₂Ni⁰ with trimethylphenylammonium tribromide (TMPhATB), a commercially available, easily manipulated solid (Figure 2B). **1-Br** assumes an analogous solid-state structure to **1-Cl**, with κ^{1} -P phosphine binding, μ_2 -Br ligand bridging, and a Ni–Ni bonding interaction (2.5955(4) Å).



Figure 2. (A) Attempted syntheses of **1-Cl** and **1-Br** via comproportionation (X = Cl or Br). (2) Initial discovery of tribromide oxidation to generate **1-Br**. (3) One-pot oxidation from Ni(COD)₂ to give **2**. (4) Solid state structures of **1-Br** and **2**, with thermal ellipsoids displayed at 50% probability and hydrogen atoms omitted for clarity. Selected solid-state bond distances are tabulated. EPR simulation parameters: $g_1 = 2.306$, $g_2 = 2.138$, $g_3 = 2.039$. For simulated hyperfine splitting and strain, see SI.

Seeking to extend this oxidative approach to bulkier Buchwald phosphines, we treated $Ni(COD)_2$ (COD = 1,5-cyclooctadiene) with TMPhATB in the presence of XPhos, generating (XPhos)Ni^IBr (2). In the solid state, 2 is monomeric, with XPhos adopting a κ^{1} -P, η^{2} -C_{arene} binding mode. This speciation contrasts with both 1-Cl and 1-Br, in which CyJohnPhos binds κ^{1} -P only in a dimeric complex. EPR measurements in glassy toluene at 77 K suggest that the solid-state speciation is conserved in the solution phase: no signal is observed for 1-Br, while spectra of 2 feature a rhombic signal with large hyperfine splitting from the ³¹P nucleus of XPhos (Figure 2D). The low solubility of 1-Br in organic solvents prevented the measurement of its solution magnetic moment. However, 2 was observed to have a solution-phase magnetic moment of $1.73 \mu_B$, consistent with a monomeric d⁹ species.⁴¹

Having identified tribromide oxidation as a tenable route to Ni¹ species, we sought to evaluate the scope of ligands that could be complexed to Ni¹ centers. We were able to generate Ni¹ halide complexes for a variety of catalytically important ligand classes including mono- and bisphosphine, *N*-heterocyclic carbene (NHC), and polypyridyl (Table 1). In most cases, yields are comparable to other reported syntheses,^{4,13,42-44} but the unique generality of Br₃⁻ oxidation is highly enabling.

Table 1. Ligand generality of tribromide oxidation

4		Br ^{_B}	⊖ ^r .Br ⊃⊳⊳∖⊕ + L –	→ 1/m [L _n Ni ^l Br] _n
4		(0.5 e	quiv) (<i>n</i> equiv)	rt, 10 min
	Ligand	n	Ni ^I complex	Br ₃ ⁻ oxidation yield (isolated)
	PPh ₃	3	L₃Ni ^l Br	75%
	PCy ₃	2	L₂Ni ^I Br	67%
	dppf	1	LNi ^I Br/[LNi ^I Br] ₂	77%
	<i>t</i> -BuXantPhos	1	L Ni ^I Br	48%
	IPr	1	[LNi ^I Br] ₂	69%
	tpy	1	L Ni ^I Br	42%
	dtbbpy	1	[LNi ^l Br] ₂	66% ^a

^aNMR yield using displaced COD as an internal standard

However, challenges arose in the isolation of Ni¹ complexes bearing two ligand classes: bipyridines and bulky trialkylphosphines. The former case will be discussed in detail (*vide infra*). Our challenges with the latter case are best summarized with the product mixture resulting from the oxidation of a Ni(COD)₂/P(*t*-Bu)₃ mixture. The desired product, a dimeric [(P(*t*-Bu)₃)Ni¹Br]₂ complex, was not detected. Instead, this reaction generated two unanticipated P(*t*-Bu)₃-bound species: an over-oxidized Ni^{1.5+} dimer, [(P(*t*-Bu)₃)₂Ni^{1.5+}₂Br₃] (**3**), and a nickelate complex with an outersphere ammonium cation, [(P(*t*-Bu)₃)Ni¹Br₂]Me₃PhN (**4**) (Figure 3). **3** likely forms as a product along with Ni⁰ black in a redox equilibrium with the expected Ni¹ dimer, while **4** is presumably the product of ammonium bromide coordination to a (P(*t*-Bu)₃)Ni¹Br species.

While complexes **3** and **4** are interesting from a structural and electronic perspective, we sought an alternative route to the desired $[(P(t-Bu)_3)Ni^{1}Br]_2$ complex that did not

employ ammonium salts. To this end, we identified that the formation of complexes **2**, **3**, and **4** implies a reactive intermediate from which ligation of these bulky phosphines is relatively facile. Both XPhos and $P(t-Bu)_3$ are too sterically encumbered to displace COD from Ni(COD)₂ directly,²⁵ but both ligate to Ni following tribromide oxidation. We therefore aimed to isolate this putative intermediate, which could serve as an ammonium-free Ni¹ precursor.



Figure 3. Synthesis and solid state structures of **3** and **4**, with thermal ellipsoids displayed at 50% probability and hydrogen atoms omitted for clarity. Selected bond distances (Å) for **3**: Ni1–Ni2: 2.378(1); Ni1–P1: 2.306(1); Ni1–Br1: 2.4637(1). Selected bond distances (Å) for **4**: Ni3–P3: 2.209(1); Ni3–Br4: 2.4096(7).

Following tribromide oxidation of a Ni(COD)₂ solution in THF- d_8 , we observed the formation of a single paramagnetic species by ¹H NMR. Use of a more soluble tribromide oxidant, tetrabutylammonium tribromide (TBATB),⁴⁵ cooling, and addition of solvent-quantity free COD to the oxidation mixture stabilized the resulting complex. With these modifications, we were able to isolate and characterize [Ni¹(COD)Br]₂ (**5**), a golden yellow solid that is stable at –35 °C under inert atmosphere (Figure 4).^{46,47} **5** is a rare example of a Ni¹–olefin complex and a tractable precursor to a variety of Ni¹–Br complexes (*vide infra*).



Figure 4. Synthesis, selected interatomic distances, and solid state structure of **5**. Thermal ellipsoids are displayed at 50% probability and hydrogen atoms are omitted for clarity. Selected solid-state bond distances are tabulated.

Structure-Speciation Relationships of Monodentate Phosphines with [Ni¹(COD)Br]₂. With access to [Ni¹(COD)Br]₂, we studied its ability to serve as a precursor to generate L_nNi¹ complexes with bulky monophosphines. Unlike with Ni(COD)₂,²⁵ Buchwald-type ligands of all steric profiles were capable of displacing the COD ligands of [Ni¹(COD)Br]₂ to generate the desired Ni¹ complex. In addition to **1-Br** and **2** with CyJohnPhos and XPhos, respectively, Ni¹ complexes with SPhos, DavePhos, JohnPhos, and t-BuBrettPhos formed readily from 5; each of these complexes was characterized by SCXRD (see SI). For Buchwaldtype ligands, three distinct classes of L1Nil species were observed in the solid-state: (i) dimers with Ni-Ni bonds and no η^2 -Carene interaction, (ii) dimers with no Ni-Ni bond (distances >3.0 Å) and a strong η^2 -Carene interaction between each Ni and the ligand B ring, and (iii) monomers with a strong η^2 -C_{arene} interaction to the B ring (Figure 5). The ligand structural feature that most clearly leads to the observed speciation outcome is the presence of a 4-*i*-Pr group on the B ring, which prevents dimer formation altogether. The structural feature(s) that distinguishes the two classes of dimers is less clear, though only the smaller Buchwaldtype ligands formed species with Ni-Ni bonds.



Figure 5. (A) Buchwald ligand-bound Ni¹ complex binding modes in the solid state. (B) Solid-state structure of [(JohnPhos)Ni¹Br]₂ (6) with thermal ellipsoids displayed at 50% probability and hydrogen atoms omitted for clarity.

Given the success of bulky Buchwald-type ligands at displacing COD from **5**, we reexamined the synthesis of bulky trialkylphosphine-bound Ni¹ complexes using the NR₄Br salt-free precursor (Figure 6). We were pleased to find that reaction of **5** with 2 equiv of P(*t*-Bu)₃ (1:1 L:Ni) led to the formation of [(P(*t*-Bu)₃)Ni¹Br]₂ (**7**). The X-ray crystal structure of **7** confirmed its identity as a Ni¹ dimer, with a Ni–Ni bond distance (2.6005(6) Å) similar to that observed in **1-Br**. We found that P(*t*-Bu)₃:Ni ratios >1:1 did not affect the speciation of the resulting complexes, with only the L₁Ni¹ dimer complex observed.

Recent work by our lab and the Sigman lab has found that minimum percent buried volume ($%V_{bur}$ (*min*))—a steric quantification of the smallest energetically accessible conformation of a ligand within 3.5 Å of the metal center—enables the discovery of structure–speciation relationships of phosphine ligand/metal complexes in cross-coupling.⁴⁸ With the demonstrated success of $%V_{bur}$ (*min*) in rationalizing phosphine ligand effects at Ni⁰/Ni¹¹, we were curious to investigate similar effects at Ni¹.

For ligands with %*V*_{bur} (*min*) values⁴⁹ slightly lower than $P(t-Bu)_3$ (36.3%), such as $CyP(t-Bu)_2$ (34.3%) and $Cy_2P(t-Bu)_3$ Bu) (32.0%), both the L_1Ni^1 dimer⁵⁰ (8 (L = CyP(t-Bu)₂) and 9 (L = Cy₂P(t-Bu))) and L₂Ni¹ monomer species (10 (L = $CvP(t-Bu)_2$ and **11** (L = $Cv_2P(t-Bu)$) could be generated and were structurally characterized, with the L:Ni stoichiometric ratio controlling the outcome.⁵¹ While these ligands normally do not form L2Ni complexes at Ni⁰ or Ni^{II}, the relatively small size of the single halide ligand leaves the majority of Ni's coordination sphere unencumbered, allowing the coordination of large phosphines. The crystal structure of $[(CyP(t-Bu)_2)_2Ni^{I}Br]$ revealed a nearly T-shaped complex, as the high amount of steric pressure between the bulky phosphines distorted the complex from the ideal trigonal geometry. For ligands smaller than $P(t-Bu)_3$ where more than one Ni¹ species can form, L:Ni stoichiometry controlled the outcome for isolated material in the solid-state. These trends were generally conserved in solution-state NMR characterization, though a small amount of phosphine dissociation and dimerization was observed for L_2Ni^1 monomer **10** with $CyP(t-Bu)_2$ (see SI).

We also found that PCy₃ (%*V*_{bur} (*min*) = 30.2 %) could form a L₁Ni¹ dimer (**12** (L = PCy₃)) from **5** with a 1:1 ratio of phosphine:Ni. In the presence of excess COD, the PCy₃ complex is more prone to disproportionation—generating Ni(COD)₂ and (PCy₃)₂Ni^{II}Br₂—than ligands with higher %*V*_{bur} (*min*) values. This is unsurprising given that stable L₂Ni⁰ and L₂Ni^{II} complexes can readily form with PCy₃. Nonetheless, the ability to synthesize [(PCy₃)Ni^IBr]₂ differs from the ligand structure–speciation trends observed in generation of the analogous Pd^I dimers.²⁹ Like CyP(*t*-Bu)₂ and Cy₂P(*t*-Bu), the L₂Ni^{IB}r monomer (**13** (L = PCy₃)) was readily obtained with PCy₃.

For even smaller PPh₃ (% V_{bur} (min) = 28.2 %), Ni¹ species were obtainable for 3:1, 2:1, and 1:1 ratios of phosphine:Ni. Unlike the examined phosphines with greater % V_{bur} (min) values, a L₃Ni¹ (**14** (L = PPh₃)) monomer is sterically accessible for PPh₃. The L₂Ni¹ (**15** (L = PPh₃)) monomer is also isolable. Uniquely, treatment of **5** with only one equivalent of PPh₃ relative to Ni resulted in the formation of a rare monomeric COD-bound Ni¹ complex, (PPh₃)Ni¹(COD)Br (**16**) (Figure 6B).⁵² Similar to COD-bound **5**, complex **16** rapidly decomposes in solution in the absence of added COD or ligand.



Figure 6. (A) Monophosphine structure–speciation relationships. (B) Synthesis and characterization of (PPh₃)Ni^I(COD)Br. EPR simulation parameters: $g_1 = 2.593$, $g_2 = 2.075$, $g_3 = 2.027$. For simulated hyperfine splitting and strain, see SI. (C) Solid-state structures of selected L₁Ni^I dimers, with thermal ellipsoids displayed at 50% probability and hydrogen atoms omitted for clarity. (D) Solid-state structures of selected L₂Ni^I monomers, with thermal ellipsoids displayed at 50% probability and hydrogen atoms omitted for clarity.

Overall, we have found that $[Ni^{l}(COD)Br]_{2}$ is a versatile precursor for synthesizing monophosphine-bound Ni¹ complexes. The high lability of COD in **5** enables formation of otherwise challenging-to-access complexes with bulky phosphines such as Buchwald-type ligands and P(*t*-Bu)₃. Furthermore, the excellent stoichiometric control on product outcome when using **5** as a precursor allows for a detailed understanding of phosphine structural effects on Ni¹ speciation (Figure 6).

Implications of Bulky Monophosphine Speciation at Ni^I in Cross-Coupling. Recently, in a collaborative study with the Sigman group, we identified phosphine ligand reactivity thresholds in Ni- and Pd-catalyzed cross-coupling datasets;48 these reactivity thresholds were linked to the %V_{bur} (min) steric descriptor.⁴⁹ For Ni, only monodentate phosphines with %Vbur (min) values less than 32% were effective at catalyzing the studied Suzuki-Miyaura coupling (SMC) reactions. This value corresponded to the region of chemical space where two phosphines could bind and stabilize Ni⁰ and Ni^{II} complexes; these even oxidation states of Ni are believed to be the catalytically relevant species in SMC reactions.9 Given Ni's propensity to engage in unproductive side pathways, we hypothesized that attaining bisligated Ni⁰/Ni¹¹ complexes was necessary to stabilize these on-cycle species.53 The inability of bulky monodentate phosphines (i.e., with $\% V_{\text{bur}}$ (*min*) values greater than 32%) to adequately stabilize these species would leave Ni more prone to falling into off-cycle thermodynamic sinks. However, this hypothesis has eluded testing due to previous synthetic challenges in accessing Ni complexes of any oxidation

state with bulky monodentate phosphines. Indeed, ligands like P(*t*-Bu)₃ do not displace olefin ligands from Ni⁰ precursors like Ni(COD)₂,²⁵ nor do they ligate Ni^{II}X₂ salts.²¹

Given the demonstrated ability of well-defined, bidentate phosphine-bound Ni¹ complexes to re-enter Ni⁰/Ni¹¹ catalytic cycles,¹⁵ we were interested to determine if monodentate phosphine-bound Ni¹ complexes could also reenter the cycle and if the $\% V_{\text{bur}}$ (*min*) threshold behavior was retained. In order to evaluate the behavior of these monophosphine Ni¹ species in a catalytic system, [Ni¹(COD)Br]₂ was employed as a precatalyst in two SMC reactions (Figure 7A). For the three phosphines tested with $\% V_{\text{bur}}$ (*min*) values less than 32%, the observed yields are lower when [Ni¹(COD)Br]₂ is used as a precursor relative to Ni(COD)₂, in line with previous precedent with dppf/Ni¹ species.

However, moderate product formation did occur for these three ligands, indicating reentry to the Ni^{0/II} cycle. Notably, a precipitous decline in yield occurs for monophosphines possessing $\% V_{bur}$ (min) values >32%, analogous to the reactivity threshold we recently reported with Ni(COD)₂ as a precursor. Even in the absence of COD, **7** was an ineffective precatalyst, giving trace yields in both reactions.

These data suggest that bulky monophosphine-bound Ni¹ complexes are thermodynamic sinks that are recalcitrant towards Ni^{0/II} catalytic cycle (re)entry. Specifically, the ability to coordinate two or more phosphine ligands to substrate bound Ni⁰ or Ni^{II} complexes appears to be necessary for stability of these even oxidation states in catalysis. The unique geometric and electronic structure of Ni^I monomer and dimer species appears to be ideal for supporting bulky ligands, whereas typical coordination spheres of Ni⁰ and/or Ni^{II} complexes are not amenable to coordination of multiple bulky phosphines (Figure 7B).



Figure 7. (A) Catalytic reactivity studies comparing **5** and $Ni(COD)_2$ as precatalysts in an SMC reaction. (B) Rationalization of catalytic reactivity by phosphine steric environment across oxidation states.

Synthesis of Ni^I Complexes with Bipyridine Ligands. Motivated by the catalytic importance of ^{t-Bu}bpy, we sought to access (^{t-Bu}bpy)Ni^I complexes using the synthetic methods discussed thus far. Tribromide oxidation of a Ni(COD)₂/ ^{t-Bu}bpy mixture did not afford isolable Ni^I species; bidentate, π -accepting COD induced the formation of COD-bound Ni⁰ and L_nNiBr₂ species upon concentration (Table 1).²² Likewise, reaction of ^{t-Bu}bpy with the well-defined precursor **5** did not afford isolable [(^{t-Bu}bpy)Ni^IBr]₂. With these results in hand, we envisioned that reaction of ^{t-Bu}bpy with precursors that contain supporting ligands other than COD would enable access to Ni^I complexes of catalytic interest.

First, we investigated bulky monophosphine-bound Ni¹ dimers as synthetic precursors of (^{t-Bu}bpy)Ni¹ complexes. Matsubara and coworkers have previously demonstrated that unsubstituted 2,2'-bipyridine (bpy) could displace the bridging μ -Cl/Br interactions of NHC-bound Ni¹ halide dimers to give monomeric (NHC)(bpy)Ni¹X complexes.¹² We evaluated an analogous route from **7** and were able to isolate (^{t-Bu}bpy)(Pt-Bu₃)Ni¹Br (**17**) by facile ligand displacement. By varying bipyridine ligand identity, it was also

possible to identify $(^{CO2Et}bpy)(P(t-Bu)_3)Ni^{I}Br$ (see SI, $^{CO2Et}bpy = diethyl 2,2'-bipyridine-4,4'-dicarboxylate)$. Indeed, the affinity of large phosphines for the steric environment of Ni^I species (*vide supra*) seems to favor the formation of monomeric species in heteroleptic complexes with bipyridine ligands. It is exciting to recognize the synergy of this relationship: monomeric bipyridine-ligated Ni^I systems are coveted. Furthermore, this class of compounds may suggest previously unexplored mechanistic possibilities for methodologies in which both (poly)pyridyl and bulky monophosphine ligands are employed in one pot.^{54,55}

However, we foresaw potential limitations of a (t Bu bpy)Ni¹ complex bearing a strongly σ -donating phosphine ligand. Such a mixed-ligand system may lead to ambiguity about which ancillary ligand is responsible for reactivity, among other shortcomings. A more general monomeric (t Bu bpy)(L)Ni¹X precursor would fill its coordination sphere with a highly labile ligand, such as an olefin. To this end, we attempted a tribromide oxidation from the recently reported Ni⁰ precursor, Ni⁰(stb)₃ (stb = (E)-stilbene).⁵⁶ Upon treatment of a Ni⁰(stb)₃/ ${}^{t-Bu}$ bpy mixture with 0.5 equiv TBATB, we observed the formation of a previously undetected paramagnetic species by ¹H NMR. Layering the resulting THF solution with pentane and cooling to -35 °C afforded red-black crystals suitable for SCXRD, which identified the complex (${}^{t-Bu}$ bpy)Ni¹(stb)Br (**18**).

Solid-state structural analysis of 17 and 18 reveals subtle differences in the coordination sphere of the metal center (Figure 8D). The *t*-Bubpy ligand of **17** is positioned closer to Ni than in 18, while 18 features a more tightly bound bromo ligand. The Ni-P(t-Bu)₃ bond of **17** is longer than for bpyfree congeners **4** and **10**, consistent with greater steric hindrance around the metal center. Bond metrics on the *t*-Bubpy ligand are within error for the two complexes, suggesting a similar extent of donation from the Ni¹ center. The C-C bonds in the backbones of the *t*-Bubpy ligands for **17** and **18** are significantly longer than in $(t-Bubpy)_2Ni^0$ (C-C_{bpy} = 1.439(6) Å, see SI). This observation is consistent with a relatively π-basic metal center endowing greater t-Bubpy-character for Ni⁰ species. Also in accordance with π -basicity, 18 is observed to activate the olefin of stilbene to a lesser extent than its Ni⁰ congener, (t-Bubpy)Ni⁰(stb) (19).

The electronic structure of these complexes was studied with EPR spectroscopy (Figure 8E). Continuous-wave spectra of 17 and 18 in glassy frozen solutions afforded rhombic signals. The spectrum of a frozen toluene solution containing 17 features no resolved hyperfine splitting and an observed value of g_{avg} = 2.261, which is similar to previous reports of well-defined monomeric (bpy/phen)Ni¹(halide) complexes ($g_{avg} = 2.19 - 2.24$).^{19,22,57,58} For **18**, a color change from dark green to red was observed upon freezing in both 2-MeTHF and toluene, and spectra exhibited abnormal lineshape (see SI). Thorough investigation of a chemical process that may be occur ring upon freezing has not been conducted, as spin relaxation for **18** is sufficiently long to observe a room-temperature spectrum. The observed g_{iso} value is 2.211, which is also consistent with prior reports.^{19,22,57,58} Overall, both EPR spectra are consistent with monomeric d⁹ Ni¹ complexes as observed by SCXRD.



Figure 8. (A) Precedented dimeric (*t*-Bubpy)Ni¹ halide complex and reactivity. (B) Monomeric (*t*-Bubpy)(L)Ni¹Br complexes accessed in this work. (C) Synthesis of complex **18** via complex **19**. Conditions (a): 1.35 equiv (*E*, *E*, *E*)-1,5,9-cyclododecatriene, 2.3 equiv Al(OEt)Et₂, Et₂O, -35 °C to rt, 16 h. Conditions (b): 1 equiv *t*-Bubpy, Et₂O. (D) Solid-state structures of **17** and **18**, with thermal ellipsoids displayed at 30% probability and hydrogen atoms omitted for clarity. Selected solid-state bond distances (Å) for **17**: Ni1-P1: 2.249(1); Ni1-Br1: 2.4438(7); Ni1-N1: 1.983(3); N1-C1: 1.365(3); C1-C2: 1.472(3). Selected solid-state bond distances (Å) for **18**: Ni2-Br2: 2.4062(6); Ni2-N3: 1.997(3) N3-C3: 1.351(4); C3-C4: 1.471(4); Ni2-C5: 2.057(5); C5-C6: 1.406(5). (E) X-band EPR spectra of **17** (toluene glass, 77 K) and **18** (THF, 298 K). Simulation parameters (**17**): *g*₁ = 2.557, *g*₁ = 2.148, *g*₃ = 2.077. Simulation parameters (**18**): *g*_{1so} = 2.211. Synthesis of **17** from **7** not shown. See SI for more information.

Complexes 17 and 18 are the first structurally characterized monomeric t-Bubpy-bound Ni¹ complexes. The nuclearity of similar complexes is observed to be crucial to their reactivity: dimeric [(t-Bubpy)Ni^IX]₂ species (X = Cl, Br) cannot kinetically disaggregate and therefore are inert towards aryl halides (Figure 8A),¹⁸ whereas in situ generated monomeric (t-Bubpy)Ni^IX species have been shown to activate even aryl chlorides.³⁶ Indeed, 17 and 18 are capable of activating aryl halide bonds (Figure 8B). Furthermore, complex **18** is an especially useful Ni¹ model system for mechanistic and stoichiometric studies: it has a direct Ni⁰ analog, **19**, against which its reactivity can be evaluated. A three-step, two-pot synthesis from Ni(acac)2-which proceeds via 19—ultimately provided 18 cleanly (Figure 8C). Efforts to prepare 8 through an analogous pathway were unsuccessful, likely due to the instability of a putatively trigonal planar (t-Bubpy)(P(t-Bu)3)Ni⁰ complex.

Interested in evaluating the reactivity of these monomeric (^{t-Bu}bpy)(L)Ni^IBr complexes, we undertook stoichiometric oxidative addition experiments. Both **17** and **18** are observed to react with 1,4-bromofluorobenzene, affording a 1:1 mixture of (^{t-Bu}bpy)Ni^{II}Br₂ and (^{t-Bu}bpy)Ni^{II}(4-fluorophenyl)Br (Figure 9). An analogous product mixture has been observed in other stoichiometric oxidative studies with (bpy/phen)Ni^I complexes; it is proposed to arise from rapid comproportionation of a putative Ni^{III} oxidative adduct with remaining Ni^I.^{22,55}

Complexes **17** and **18** are also observed to activate the $C(sp^2)$ -Cl bond of 1,4-chlorofluorbenzene (Figure 9). While the oxidative addition of aryl chlorides to (bpy/phen)Ni¹

species has been invoked in methodologies,^{59,60} it has not previously been demonstrated for a well-defined, isolable (bpy/phen)Ni¹ complex.^{22,34,55} Complex **18** is observed to convert the anticipated amount of aryl chloride within minutes, while **17** requires several hours of reaction time for the expected 50% conversion. This is perhaps due to the increased lability of olefinic stilbene relative to P(*t*-Bu)₃. Rigorous kinetic and mechanistic investigations into the reactivity profile of these complexes is beyond the scope of this study, but work is underway in our laboratory to further interrogate the behavior of this unique class of compounds.



Figure 9. Stoichiometric reactivity with aryl halides for **17** and **18**. Conversion measured after 4 h.

CONCLUSIONS

In summary, we have identified a mild oxidative approach and precursor to a variety of Ni¹ complexes bearing catalytically relevant ligands. These strategies enabled access to previously elusive complexes, including L₁Ni monophosphine dimers and the first examples of monomeric ^{t-} ^{Bu}bpy-bound Ni¹ species. For the monophosphine ligand

class, we have elucidated structure-speciation relationships at Ni¹ and conducted catalytic studies to relate established steric relationships to observed trends for Ni¹ complexes. For the bipyridine ligand class, we have synthesized and characterized well-defined monomeric complexes that are capable of activating aryl bromides and chlorides. We anticipate that our findings will enable future mechanistic studies and catalyst design for Ni-catalyzed cross-coupling reactions.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website.

Experimental procedures, experimental data, and characterization and spectral data for new compounds (PDF)

Accession Codes

CCDC 2204445–2204452, 2204454–2204455, 2204457, 2204459, 2245704–2245705, 2245707–2245709, 2245759, and 2262986–2262993 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc. cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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Notes

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ACKNOWLEDGMENT

We thank Dr. Robert Taylor (UCLA) and Dr. Paul Oyala (Caltech) for assistance with EPR measurements and Dr. Saeed Khan (UCLA) for assistance with X-ray diffraction data collection. We thank Prof. Franziska Schoenebeck for helpful discussions. Financial support for this work was provided by the NIGMS (R35 GM126986). These studies were supported by shared instrumentation grants from the National Science Foundation under equipment grants CHE-1048804 and 2117480, along with the NIH Office of Research Infrastructure Programs under grant S100D028644.

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(51) For ligands smaller than $P(t-Bu)_3$ where more than one Ni¹ species can form, L:Ni stoichiometry controlled the outcome for isolated material in the solid state. Speciation was generally conserved in solution-state NMR characterization (C₆D₆), though a

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