Insight into Transmetalation Enables Cobalt-Catalyzed Suzuki–Miyaura Cross Coupling

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ABSTRACT: Among the fundamental transformations that comprise a catalytic cycle for cross coupling, transmetalation from the nucleophile to the metal catalyst is perhaps the least understood. Optimizing this elementary step has enabled the first example of a cobalt-catalyzed Suzuki–Miyaura cross coupling between aryl triflate electrophiles and heteroaaryl boron nucleophiles. Key to this discovery was the preparation and characterization of a new class of tetrahedral, high-spin bis(phosphino)pyridine cobalt(1) alkoxide and aryloxide complexes, (tBuPNP)CoOR, and optimizing their reactivity with 2-benzofuranylBPin (Pin = pinacolate). Cobalt compounds with small alkoxide substituents such as R = methyl and ethyl underwent swift transmetalation at 23 °C but also proved kinetically unstable toward β−H elimination. Secondary alkoxides such as R = Pr or CH(Ph)Me balanced stability and reactivity. Isolation and structural characterization of the product following transmetalation, (tBuPNP)Co(2-benzofuranyl), established a planar, diamagnetic cobalt(1) complex, demonstrating the high- and low-spin states of cobalt(1) rapidly interconvert during this reaction. The insights from the studies in this elementary step guided selection of appropriate reaction conditions to enable the first examples of cobalt-catalyzed C−C bond formation between neutral boron nucleophiles and aryl triflate electrophiles, and a model for the successful transmetalation reactivity is proposed.

Transition metal-catalyzed cross coupling has revolutionized carbon−carbon formation by enabling selective and efficient reactions of organic electrophiles and various nucleophiles.1 Among these methods, the palladium-catalyzed Suzuki–Miyaura reaction of organic halides electrophiles and organoboron nucleophiles is one of the most commonly applied strategies for C−C bond formation in chemical synthesis.2−23 The safe handling, relative stability, and broad availability of boron-based nucleophiles distinguishes the Suzuki–Miyaura reaction from other cross coupling methods that rely on reactive organolithium, Grignard, or zinc nucleophiles that can be incompatible with functional groups found in late stage intermediates and are dangerous to handle on a large scale.2 As a result, Suzuki–Miyaura cross coupling has become one of the preferred methods for making C−C bonds in the pharmaceutical, agricultural, and fine chemicals industries.2−5 The distinction between boron and other nucleophiles is particularly apparent in a recent analysis of reactions employed by the pharmaceutical industry that found Suzuki–Miyaura coupling accounted for approximately 40% of C−C bond-forming reactions, while coupling with other nucleophiles represented less than 5%.6−8 Palladium compounds are state-of-the-art catalysts for the Suzuki–Miyaura cross coupling. Their widespread use raises concerns about the sustainability7 and economics of these processes, the latter being influenced not only by cost of goods but by the purification steps required to meet strict regulations (typically <5 ppm) for palladium content in active pharmaceutical ingredients.6−8 These practical motivations, coupled with opportunities for new modes of reactivity and potential expansion of reaction scope, have prompted exploration of Earth abundant, first row transition metals as potential catalysts for the Suzuki–Miyaura reaction. Considerable progress has been made with nickel-9−18 and copper-19−25 catalyzed variants, but reactivity with earlier first row transition metals such as iron and cobalt has remained elusive. State-of-the-art iron-26−30 and cobalt-31−35 catalyzed C−C cross coupling methods rely on more reactive nucleophiles rather than the preferable neutral boron reagents associated with Suzuki–Miyaura chemistry (Scheme 1). Specifically, several recent examples of C(sp2)−C(sp3) and C(sp3)−C(sp3) iron-catalyzed cross coupling rely on stoichiometric addition of organolithium reagents to form activated borinate nucleophiles because the corresponding neutral boron partners do not participate.31−38 As such, the advantages of neutral boron nucleophiles are lost, and the reliance on radical capture for C−C bond formation has limited the scope to principally pure hydrocarbyl products such as cycloheptylbenzene. Understanding transmetalation reactivity is therefore key for developing improved iron- and cobalt-catalysts for Suzuki–Miyaura cross coupling. Here we describe a cobalt pincer complex that promotes Suzuki–Miyaura cross coupling of aryl triflate electrophiles and organoboron nucleophiles. Stoichiometric reactions designed to mimic elementary steps of the catalytic reaction provided crucial insight about the optimal nucleophile, electrophile, base, and conditions for the desired...
reactivity. A combination of these observations ultimately enabled successful catalytic turnover.

Scheme 2 presents a plausible cycle for cobalt-catalyzed C(sp²)−C(sp²) Suzuki−Miyaura coupling. Guiding principals from this pathway are (i) transmetalation of an aryl group from a neutral boron reagent to a CoOR complex to generate a cobalt(I) aryl, (ii) interaction with the aryl electrophile to promote carbon−carbon bond formation and release of product, generating a cobalt(I) halide, and (iii) exchange of the halide ligand with exogenous base to regenerate the CoOR species. One notable difference with first row transition metals is the likelihood of variable spin states and coordination geometries and kinetically accessible one-electron chemistry.49 Depending on the field strength of the supporting ligands, cobalt(I) complexes like those shown in Scheme 2 may be high (S = 1) or low (S = 0) spin with tetrahedral or planar geometries, respectively. With these considerations in mind, the flexible diisopropyl-substituted bis(phosphino)pyridine pincer ligand (iPrPNP) was selected for this study due to its known ability to support both tetrahedral (X = Cl) and planar (X = alkyl, aryl) cobalt(I) complexes.50−54 Furthermore, the (iPrPNP)cobalt platform promotes two-electron oxidative addition53 and has been applied to catalytic C−H borylation.50,52

The requirement for highly reactive nucleophiles in current iron and cobalt-catalyzed C−C coupling methods suggests sluggish transmetalation prevents catalytic turnover with neutral boron reagents. While transmetalation from boron is well established55 and mechanistically understood56,57 for palladium, analogous reactivity with iron58,59 or cobalt60,61 has not been demonstrated.52 Precedent with palladium63−65 suggested a transmetalation study would be best approached with the corresponding (iPrPNP)CoOR derivatives, a new class of bis(phosphine)pyridine cobalt complexes with unknown spin states, geometries and substitution chemistry. Synthesis of the target (iPrPNP)CoOR complexes was accomplished by protonolysis of the cobalt(I) alkyl complex (iPrPNP)CoCH₂SiMe₃ with a stoichiometric quantity of the appropriate alcohol (ROH, Scheme 3). In this manner, a series of paramagnetic cobalt(I) aryloxides (iPrPNP)CoOR and alkoxides were prepared. The cobalt aryloxide complexes (R = Ph, C₆H₄(4-OMe) and C₆H₄(3-F)) were isolated as purple powders, and solid state magnetic measurements established S = 1 ground states for each, consistent with high spin Co(I) derivatives. Despite their paramagnetism, these compounds were reliably identified by ¹H NMR spectroscopy, exhibiting the number of peaks expected for compounds with C₂ᵥ molecular symmetries. The cobalt phenoxide, (iPrPNP)CoOPh, was also characterized by X-ray diffraction (Figure 1). The pseudotetrahedral geometry is consistent with the high spin cobalt(I) ground state. To accommodate this geometry, the pincer ligand is significantly distorted from planarity, as evidenced by a P−Co−P bond angle of 123.75(1)° and the N(1)−Co(1)−O(1) bond angle of 128.18(4)°. Accordingly, the cobalt is deviated by 1.032 Å from the idealized plane of the PNP pincer. Other metrical parameters are as expected for a tetrahedral cobalt(I) center (see Supporting Information for complete details). Several cobalt(I) alkoxides (iPrPNP)CoOR (R = CH(Ph)Me, iPr, Et and Me) were also prepared by protonolysis (Scheme 3). Attempts to prepare the tert-butoxide derivative (BuO)CoO(Bu) have been unsuccessful, resulting in recovery of the
starting cobalt alkyl. Like the aryloxides, the cobalt(I) alkoxides were paramagnetic but observable by $^1$H NMR spectroscopy. These complexes proved unstable in benzene-d$_6$ solution at 23 °C with relative rates of decomposition decreasing from R = Me > Et > iPr > CH(Me)Ph. Approximate times for decomposition at 23 °C vary from over the course of 2 h for R = Me to 4 h for Et, 24 h for iPr, and 72 h for CH(Me)Ph. These times are approximate because the decomposition reactions do not yield a single cobalt product, and the complexity of the reaction mixture precluded detection of the expected aldehydes or ketones. In general, the relative stability is consistent with decomposition of the alkoxide complex by β-hydride elimination, where small alkyl substituents with more β-hydrogens decompose more rapidly. The putative cobalt hydride product, (iPrPNP)CoH, is known to decompose by P−C bond cleavage and is likely the source of the product mixtures.53 The formation of (2bPNP)Co(PHiPr$_2$)$_2$H was in 25% yield by $^1$H NMR, accompanied by 50% free (2bPNP) ligand, by methanolysis of (2bPNP)CoCH$_2$SiMe$_3$ in the presence of HPPr$_2$ provides support for this hypothesis (Scheme 3).

Carboxylates were also explored as possible oxygen-based ligands for cobalt to promote transmetalation with boron reagents. These ligands were of interest due to precedent from our laboratory$^{66}$ and Nagashima’s$^{67}$ that cobalt carboxylate derivatives are bench-stable catalyst precursors. Addition of 1 equiv of benzoic acid (BzOH, Bz = C(O)Ph) to (2bPNP)-CoCH$_2$SiMe$_3$ at room temperature produced a complex mixture of cobalt products from which the cobalt(II) bis(carboxylate) (2bPNP)Co(OBz)$_2$ was identified as the major component. Addition of 2 equiv of BzOH to (2bPNP)-CoCH$_2$SiMe$_3$ resulted in clean formation of (2bPNP)Co(OBz)$_2$ isolated as a purple solid in 87% yield and was characterized by X-ray diffraction (see Figure S6). The target cobalt(I) benzoate (2bPNP)CoOBz was prepared by slow addition of BzOH to a thawing toluene solution of (2bPNP)CoCH$_2$SiMe$_3$ (see Supporting Information). The solid-state structure of this S = 1 complex ($\mu_{tot} = 2.8 \mu_B$, 23 °C) established a $\kappa_1$-benzoate ligand. The P–Co–P bond angle of 126.081(14)$^\circ$ and the N(1)–Co(1)–O(1) bond angle of 119.65(4)$^\circ$ as well as the deviation of the cobalt by 1.000 Å from the metal chelate plane establish an idealized tetrahedral geometry, similar to (2bPNP)-CoOPh (Figure 2).68

With a series of cobalt aryloxide, alkoxide, and carboxylate compounds in hand, transmetalation with neutral boron reagents was explored (Scheme 4). Addition of 1 equiv of phenylboronic acid pinacol ester (PhBPin) to a benzene-d$_6$ solution of (2bPNP)CoOPh at room temperature resulted in no reaction as judged by $^1$H NMR spectroscopy. By contrast, the corresponding reaction with 2-benzofuranylBPin resulted in gradual appearance of a new diamagnetic cobalt compound over the course of 24 h at 23 °C. Characterization by $^1$H, $^{13}$C, and $^{31}$P NMR spectroscopies as well as X-ray diffraction identified the product as (2bPNP)Co(2-benzofuranyl), arising from transmetalation of the heteroaryl group from boron to cobalt (Figure 3). The solid-state structure of (2bPNP)Co(2-benzofuranyl) was determined by X-ray diffraction and confirmed the idealized planar geometry about the metal center, consistent with the observed diamagnetic ground state. While the analogous reaction with (2bPNP)CoOBz generated an intractable mixture, clean formation of the desired organometallic complex was observed upon addition of 2-benzofuranylBPin to the cobalt alkoxides (2bPNP)CoOR (R =...
CH(Me)Ph, iPr, Et, and Me) generated in situ. In these cases, transmetalation occurred immediately, signaled by a color change to a red brown solution, and \(^1\text{H} \text{NMR} \) spectroscopy confirmed clean formation of \((^\text{P}^\text{NP})\text{Co}(2\text{-benzofuranyl})\) within the time required for \(^1\text{H} \text{NMR} \) analysis. Notably, transmetalation occurred rapidly from \(S = 1\) cobalt alkoxides to form diamagnetic cobalt heteroarene products, a difference from palladium chemistry where catalysis occurs predominantly on the \(S = 0\) spin surface. Use of 2-benzofuranyl boronic acid as the boron reagent in the reaction of \((^\text{iPrPNP})\text{CoOCH}(\text{Me})\text{Ph} \) led to partial formation of the desired product within a complex mixture, indicating boronic acids are not optimal organoboron nucleophiles for transmetalation in this system.

To gain further insight about this unprecedented reactivity, transmetalation of various heteroaryl boron reagents was explored (Scheme 5). Addition of a solution of 2-methylfuran-arylBPin or benzothiophenylBPin to \((^\text{iPrPNP})\text{CoOCH}(\text{Me})\text{Ph} \) led to partial formation of the desired product within the time required for analysis by \(^1\text{H} \text{NMR} \) spectroscopy, determined by analogy to the benzofuranyl complex (vide supra). When \(N\)-methylindolylBPin was used as the boron reagent, transmetalation did not proceed. These observations suggest the lone pair of the heteroaryl group is essential for transmetalation reactivity. This influence may arise from initial coordination of the lone pair to cobalt, bringing the boron atom in proximity of the alkoxide ligand and initiating the transmetalation process (Scheme 5). This coordination event may also induce a change in the spin state from \(S = 1\) to \(S = 0\), opening up a coordination site and hence a vacant cobalt orbital for enabling heteroaryl group transfer.

Given the observed difference in the reactivity of cobalt phenoxy and alkoxide complexes toward 2-benzofuranylBPin, experiments were conducted to establish the relative rates of transmetalation as a function of the oxygen substituent in \((^\text{P}^\text{NP})\text{CoOR} \) (Scheme 5). Cobalt aryloxides with varied electronic properties (\(R = \text{C}_6\text{H}_4(4\text{-OMe})\) and \(\text{C}_6\text{H}_4(3\text{-F})\)) were synthesized and subjected to transmetalation with 2-benzofuranylBPin. Relative rate constants \((k_{\text{rel}})\) were measured by comparing initial rates of these and the parent cobalt phenoxy complex (see Figure S1). This kinetic analysis established that the relative rate constant for transmetalation increases with increasing electron donation from the aryl substituent (\(\text{C}_6\text{H}_4(4\text{-OMe}) > \text{Ph} > \text{C}_6\text{H}_4(3\text{-F})\)). This behavior is consistent with accumulation of positive charge within the aryl oxide ligand during the transmetalation process. Additional kinetic experiments established an overall second-order reaction indicated by a linear slope of inverse product concentration versus time (see Figure S2).

In a typical cross coupling reaction, for example, as illustrated in Scheme 2, the aryl or alkoxide ligand \((\text{OR})\) enters the coordination sphere of the cobalt catalyst by salt metathesis rather than by protonation. Thus, the catalytically relevant route to the \((^\text{P}^\text{NP})\text{CoOR} \) species was explored using the known cobalt(I) chloride \((^\text{P}^\text{NP})\text{CoCl} \). Addition of 1 equiv of NaOPh in THF-\(d_8\) solution resulted in formation of \((^\text{P}^\text{NP})\text{CoOPh} \) as the major product observed by \(^1\text{H} \text{NMR} \) spectroscopy (Scheme 6). Addition of 2-benzofuranylBPin to the resulting THF-\(d_8\) solution produced a mixture containing \((^\text{P}^\text{NP})\text{CoCl} \) as the only cobalt species observable by \(^1\text{H} \text{NMR} \) spectroscopy. These results suggest a preference for the cobalt chloride and boronate species over the cobalt alkoxide and neutral boronic ester at room temperature, potentially complicating the combination of salt metathesis and transmetalation steps that is necessary for catalytic turnover. Indeed, addition of 1 equiv each of NaOCH(Ph)Me and 2-benzofuranylBPin in THF to \((^\text{P}^\text{NP})\text{CoCl} \) resulted in slow formation of \((^\text{P}^\text{NP})\text{Co}(2\text{-benzofuranyl}) \) at room temperature (incomplete conversion after 3 h). However, heating the same reaction mixture to 60 °C resulted in complete conversion to the desired organometallic complex after 90 min (Scheme 6).
Replacing the bulky alkoxide base with NaOMe resulted in a decreased yield of (iPrPNP)Co(2-benzofuranyl) (55% versus 75%), highlighting the importance of the steric profile of the alkoxide ligand substituent and providing key insight for base selection in the catalytic cross coupling reaction.

These observations provided an understanding of salt metathesis and transmetalation steps of the cobalt-catalyzed Suzuki–Miyaura cross coupling reaction proposed in Scheme 2. Aiming to complete the catalytic cycle, interaction of the organometallic complex (iPrPNP)Co(2-benzofuranyl) and potential electrophilic coupling partners was investigated (Scheme 7). Heating a mixture of 1 equiv of phenyl halide (PhX, X = Cl, Br, I) and (iPrPNP)Co(2-benzofuranyl) in benzene-d₆ resulted in partial conversion (<50%) to 2-phenylbenzofuran after 90 min at 60 °C. Formation of the anticipated cobalt(I) halide species (iPrPNP)CoX was not observed in any case. Performing the same reaction at 60 °C using phenyl triflate (X = OTf) as the electrophile generated a quantitative amount of 2-phenylbenzofuran and a new paramagnetic species (Scheme 7). Though difficulties in isolation prevented additional characterization, this compound was identified as (iPrPNP)CoOTf by ¹H NMR spectroscopy by analogy to the related cobalt(I) chloride complex. Oxidation with Ph₃CCl yielded the stable cobalt(II) species (iPrPNP)CoOTf(Cl) (see Supporting Information) that was characterized by X-ray diffraction (see Figure S9).

Results from these stoichiometric studies offered guidelines for realization of a successful Co-catalyzed Suzuki–Miyaura coupling method. The stability and transmetalation reactivity of (iPrPNP)CoOR complexes pointed to NaOCH(Ph)Me as the optimal base. Reactions of the organometallic complex and different electrophiles indicated that the use of aryl triflates, rather than aryl halides, would allow for catalytic turnover. Combining salt metathesis and transmetalation processes suggested the reaction might require elevated temperature, and entry to the catalytic cycle could be accomplished from the isolable complex (iPrPNP)CoCl. Thus, heating a 1:1:1 mixture of PhOTf, 2-benzofuranylBPin, and NaOCH(Ph)Me in the presence of 5 mol % (iPrPNP)CoCl in THF at 60 °C for 24 h resulted in formation of 2-phenylbenzofuran in 85% yield (Table 1). Analysis by GC and ¹H NMR spectroscopy indicated a trace amount (<4%) of bis(benzofuran) and no observable amount of biphenyl in the reaction mixture, amounting to >20:1 selectivity for the cross coupled product. Preliminary evaluation of the reaction scope established tolerance for CF₃, OMe, C(O)Me, and F functional groups on the aryl electrophile (Table 1). Heteroaryl cross coupling with 3-pyridinyl triflate was also successful, albeit in low yield.

The scope of nucleophilic partner was also examined (Table 2). Furanyl derivatives containing CHO and BPin groups in the 2-position underwent selective cross coupling with PhOTf. Only a trace amount of product was observed in the reaction of 2-methylfuranylBPin but use of the more electron-rich 4-pyrrolidinyl-(iPrPNP)CoCl complex allowed for catalytic turnover, suggesting that ligand modification may overcome current limitations with substrate scope. While benzothiophenylBPin participated in stoichiometric transmetalation with (iPrPNP)CoOR, catalytic reactivity was very low, suggesting

Table 1. Preliminary Aryl Triflate Scope

| Aryl Triflate (0.05 mmol), boron reagent (1 equiv), NaOCH(Ph)Me (1 equiv), and (iPrPNP)CoCl (5 mol %) in THF (1 mL) heated at 60 °C for 24 h. > 20:1 cross selectivity in all cases. Yields determined by ¹H NMR. |
|---|
| 85% yield |
| 90% yield |
| 75% yield |
| 80% yield |
| F |
| 50% yield |
| N |
| 30% yield |

*See Table 1.*

Table 2. Preliminary Boron Reagent Scope

| Boron Reagent (1 equiv) + PhOTf (1 equiv) |
|---|
| PhOCH(O)Me |
| PhOCHO |
| PhOBF |
| 95% yield |
| 50% yield |
| 60% yield |
| 35% yield |
| trace product |
| no reaction |

Additional PhOTf, base and catalyst added after 12 h. Formed as a 3:1 mixture of bis- and monophenyl products. Using 4-pyrro-(iPrPNP)CoCl as the catalyst.

Scheme 6. Salt Metathesis/Transmetalation Studies

Scheme 7. Electrophile-Dependent Reactivity of (iPrPNP)Co(2-benzofuranyl)
that interaction with the aryl triflate electrophile is also substrate dependent. Cross coupling of PhOTf and N-methylindolylBPin did not proceed, as expected due to the lack of transmetalation reactivity observed in the stoichiometric reaction with the nucleophilic partner.

Fundamental insights gained by stoichiometric reactions of proposed transition metal intermediates have enabled the first example of a cobalt-catalyzed C(sp²)–C(sp³) Suzuki–Miyaura cross coupling. Specifically addressing the transmetalation step in the catalytic cycle provided access to crucial information regarding turnover and base selection, ultimately enabling catalytic reactivity. The flexibility of the supporting PNP pincer allows rapid interconversion between high-spin tetrahedral and low-spin planar catalytic intermediates, a distinguishing feature for the first row transition metal over state-of-the-art palladium catalysts. Importantly, these findings provide insight into the interaction of Earth abundant first row transition metals and neutral boron reagents, and this mechanistic foundation may enable new catalytic reactions involving this important class of nucleophiles.

**EXPERIMENTAL SECTION**

**Preparation of (iPrPNP)CoOPh.** A scintillation vial was charged with (iPrPNP)CoCl₂ (82 mg, 0.17 mmol) in 1.7 mL of toluene, and a magnetic stir bar. A solution of phenol (16 mg, 0.032 mmol) in 0.28 mL of benzene-pentane was added to a scintillation vial containing (iPrPNP)CoCl (1.0 mg, 0.0025 mmol) and a magnetic stir bar. A solution of phenyltriﬂate (12 mg, 0.05 mmol) in 0.5 mL of THF was added to a scintillation vial containing (iPrPNP)CoCl (1.0 mg, 0.0025 mmol) and a magnetic stir bar. A solution of phenyl triﬂate (12 mg, 0.05 mmol) in 0.5 mL of THF was added to this mixture. The reaction was sealed and heated in a heating block at 60 °C for 24 h. Trimethoxybenzene was added as a standard, and the volatiles were removed in vacuo. The residue was dissolved in CDCl₃ and filtered through a glass frit, and analyzed by ¹H NMR spectroscopy. This data matched that previously reported for 2-phenylbenzoﬂuran.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscentsci.6b00283.

Complete experimental procedures and full characterization data (PDF)

Crystallographic data (CIF)

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**Notes**

The authors declare no competing financial interest.

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