Cobalt co-catalysis for cross-electrophile coupling: diarylmethanes from benzyl mesylates and aryl halides†

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The nickel-catalyzed cross-coupling of aryl halides with alkyl radicals derived from alkyl halides has recently been extended to couplings with carbon radicals generated by a co-catalyst. In this study, a new co-catalyst, cobalt phthalocyanine (Co(Pc)), is introduced and demonstrated to be effective for coupling substrates not prone to homolysis. This is because Co(Pc) reacts with electrophiles by an \( \text{S}_\text{N}^2 \) mechanism instead of by the electron-transfer or halogen abstraction mechanisms previously explored. Studies demonstrating the orthogonal reactivity of (bpy)Ni and Co(Pc), applying this selectivity to the coupling of benzyl mesylates with aryl halides, and the adaptation of these conditions to the less reactive benzyl phosphate ester and an enantioconvergent reaction are presented.

Introduction

Cross-coupling relies upon the selective, ordered activation of two different substrates. For the coupling of nucleophiles with electrophiles, a single catalyst reacts with the electrophile by oxidative addition and the nucleophile by transmetalation, resulting in high cross-selectivities. Cross-electrophile coupling,\(^1\) the union of two different electrophiles, achieves selectivity by different mechanisms. Specifically, we have recently shown that in nickel-catalyzed reactions electrophiles can be differentiated by heterolysis and homolysis (Fig. 1, entry 1).\(^2\) Selectivity arises because (L)\(\text{Ni}^0\) reacts with aryl halides faster than alkyl halides, but (L)\(\text{Ni}^1\) forms alkyl radicals faster than aryl radicals.\(^3\)

The key to successful cross-electrophile coupling is selective radical generation from R–Y (Fig. 1).\(^4\) In order to expand the types of substrates that can be coupled with aryl halides by this electrophile + radical mechanism, alternative methods of generating radicals must be developed. A key advance was that radical generation and coupling can be accomplished by two different catalysts (Fig. 1). Our group\(^5\) and the groups of Sanford,\(^6\) Molander,\(^7\) and MacMillan and Doyle\(^8\) have independently shown that a variety of co-catalysts can allow coupling of otherwise unreactive substrates under mild conditions (Fig. 1, entries 2–4).

All of the methods reported to date convert the substrate into a radical by single-electron oxidation or reduction (Fig. 1). As such, substrates must be easily oxidized or reduced. The development of co-catalysts that form radicals by different mechanisms

- **Mechanism**
- **R–Y**
- **Co-catalyst**
- **R+**

|   | SET halogen abstraction | Alkyl-X | \([\text{Ni}^0]\) | Alkyl+ |
|---|-------------------------|---------|-----------------|--------|
| 1 | SET reductive ring opening | O | \(\text{Cp}_2\text{Ti}^{\text{III}}\text{Cl}\) | \(\text{Cp}_2\text{Ti}^{\text{IV}}\) | \(\text{Cp}_2\text{Ti}^{\text{IV}}\) |
| 2 | photoredox SET oxidation | \(\text{Ar–N}_2^+\) | \(\text{Ru(bpy)}_3\text{Cl}_2\) | \(\text{Ar}^+\) |
| 3 | photoredox SET reduction | \(\text{Bn–BF}_3\text{K}\) | \(\text{Ar–NMe}_2\) | \(\text{Bn–}\) |
| 4 |   | Alkyl-CO\(_2^+\) | \(\text{Ar–NMe}_2\) | \(\text{Bn–}\) |
| 5 | SN2 followed by homolysis | \(\text{Bn–OMs}\) | \(\text{Co(Pc)}\) | \(\text{Bn–}\) |
|   | This Report | | | |

*Fig. 1* Comparison of radical co-generation methods in cross-coupling. An electrophile (Ar–X) reacts to form an arylmetal intermediate and the other substrate (R–Y) reacts to form a radical (R+).
would enable further expansion of substrate scope in this arylation strategy.

We report here that cobalt phthalocyanine (Co(Pc)) is an excellent, non-photochemical co-catalyst for radical generation that is compatible with nickel catalysis (Fig. 1, entry 5). Co(Pc) differs from previously described co-catalysts because it generates radicals after 2-electron nucleophilic substitution rather than single-electron transfer. This gives Co(Pc) different selectivity than previous approaches. For example, alkysulfonate and alkylphosphate esters are unreactive towards single-electron transfer due to the strength of the C–O bond, but react rapidly by nucleophilic substitution.

Results and discussion

To demonstrate the potential of Co(Pc), we applied this co-catalyst to the synthesis of diarylmethanes from two abundant electrophiles, a benzyl alcohol derivative and an aryl halide. Although a variety of approaches to diarylmethanes have been developed, their prominence in medicinal chemistry has justified continued attention. The majority of recent approaches involve the cross-coupling of benzylmetal reagents or arylmetal reagents, but the need to pre-form each organometallic reagent can be limiting. A major advance was the development of methods where organozinc reagents were generated and coupled concurrently, but 2–4 equiv. of the benzyl halide was still required for high yields and no examples with more abundant benzyl alcohol derivatives were reported. Gosmini reported one example of the coupling of benzyl chloride with ethyl 4-bromobenzoate under conditions that might not involve an organozinc intermediate, but the scope of this method has not been explored. Finally, Reisman reported the coupling of secondary benzyllic chlorides with vinyl bromides, but the use of aryl halides or benzyl alcohols was not reported. Compared to these known methods, our new approach avoids pre-formed nucelophiles, starts from benzyl alcohols instead of the less abundant benzyl halides, and does not use a large excess of one coupling partner.

The application of our reported conditions to the coupling of aryl halides with benzyl bromide resulted primarily in the formation of dibenzyl (Scheme 1A). This is due to the fact that benzyl bromide reacts with (L)Ni faster than aryl halides (Scheme 1B). For example, benzyl bromide is converted to dibenzyl and toluene in only 60 min (ESI Table S1†).

In order to restore selectivity, we sought to take advantage of the low reactivity of bipyridine-ligated nickel catalysts with alkyl sulfonate esters in order to prevent formation of benzyl(nickel) and favor formation of aryll nickel (Scheme 1C). Benzyl mesylates can be conveniently generated in situ from abundant benzyl alcohols. Although this approach prevented dibenzyl formation (Table S1†), it did not improve the yield of diarylmethane because nickel is slow to form a benzyl radical from a benzyl mesylate (Table 1, entry 1). The addition of Co(Pc) as a co-catalyst could generate the missing benzyl radical (Scheme 1C), separating the oxidative addition and radical generation steps.

Significantly, this optimized nickel and cobalt system allows for the coupling of benzyl mesylates with aryl bromides, which occurs in low yield in the absence of Co(Pc) (Table 1, entries 1–3). Furthermore, it appears that both (dtbbpy)NiBr2 and Co(Pc) are required for high selectivity as neither catalyst is effective without a ligand (entries 4 and 7). An alternative co-catalyst, sodium iodide, was not as effective as Co(Pc) (entry 5 and Table S1†). The different products formed arise from different mechanisms of co-catalysis: Co(Pc) selectively forms benzyl radicals from BnOMs while NaI converts BnOMs to BnI which reacts similarly to benzyl bromide (Scheme 1B).

For convenience, the aryl halide, ligand, Co(Pc), and zinc could be added with the alcohol, Ms2O, and EtN(i-Pr)2 or after mesylate formation was complete with little difference in yield. However, if nickel was added at the beginning, rapid reduction or dimerization of the starting materials resulted.

The nickel and cobalt co-catalytic strategy of synthesizing diarylmethanes was also successful when aryl iodides were employed as coupling partners. When benzyl mesylate was reacted with iodobenzene under the optimized reaction conditions, the diphenylmethane product was formed in good yield. Although product was formed in the absence of Co(Pc), the co-catalytic method improved yield and selectivity (Table 1, entries 9–12). Added NaI depressed the yield only slightly (entry 13 vs. entry 9), suggesting that PhI competes with in situ formed BnI...
more effectively than PhBr for oxidative addition to nickel (Scheme 1B).

Application of these conditions to a variety of different aryl bromides, aryl iodides, and benzyl alcohol demonstrated the generality of the method (Scheme 2). The reaction tolerates functional groups, such as aldehyde (3e) and methyl ketone (3d), which are susceptible to the nucleophilic and basic character of some organometallic reagents. Additionally, a boronic acid ester was coupled without chemoselectivity problems, demonstrating complementarity with existing cross-coupling methods (3f).13

The coupling of aryl bromides and aryl iodides with benzyl mesylate provided comparable yields. In examples where the selectivity over bibenzyl was high but the yield was low, hydrodehalogenation of the arenne was responsible for diminished yields.20 The aryliodoide conditions could be applied without further optimization to a vinyl bromide with reasonable success (3j). Both electron-rich and electron-poor benzyl alcohol coupled effectively and steric hindrance on the benzyl alcohol did not result in lower yields (3l, 3m, 3g', 3n).22 Finally, we were able to apply this approach to the synthesis of beclobrate (3o), a diaryl methane compound that can be used to alter lipid levels in humans.23

The reactivity and selectivity of the reaction can also be rationally optimized. For example, the coupling of benzyl diethyl phosphate ester24 was sluggish under our standard conditions and starting materials were not consumed (Table 2). Presumably, this was due to the reduced leaving group ability of OP(O)(OEt)2 compared to OMs.25 Increasing the amount of Co(Pc) from 1 mol% to 6 mol% as well as increasing the temperature from rt to 80 °C allowed the reaction to proceed in high yield to form diphenylmethane (3a').22 Notably, we observe no hydrodehalogenation under these conditions, perhaps because there is no acid (EtN(i-Pr)2·HOMs) present. We anticipate that a similar strategy could be employed to couple other less reactive electrophiles.

The likely intermediacy of a benzyl radical suggested to us that an enantioconvergent coupling of racemic secondary benzylic electrophile with an aryl bromide could be achieved.26,27 Although yields were low with secondary benzylic alcohol derivatives, α-chloroethylbenzene (7) coupled efficiently under slightly modified standard conditions to form diaryl methane 8(ref. 28) in 41% isolated yield and 43% ee (Scheme 3).29 This yield and enantioselectivity compares favorably to the results of Molander in the coupling of a racemic secondary benzylic reagent with an aryl halide using the same chiral
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18 Conditions: 1:1 PhI: BnBr, 7 mol% NiBr2·3H2O, 5 mol% dtbbpy, and 2 equiv. Zn dust in DMA (0.25 M) at 60 ºC for 10 min. BnBr is consumed, most of the iodobenzene remains.

19 Ligand-free nickel does not provide high yields (Table 2, entry 7) and excess nickel does not negatively impact reactions.†

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