Nickel-on-Charcoal-Catalyzed Reductions of Aryl Chlorides

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Warmly dedicated to Professor Ryoji Noyori in recognition of his outstanding contributions to synthetic organic chemistry

Abstract: Exposure of (functionalized) aryl chlorides to catalytic quantities of nickel-on-charcoal in the presence of stoichiometric amounts of Me₂NH•BH₃/K₂CO₃ in refluxing acetonitrile leads to high yields of reduced arenes. PCBs are also reduced under these conditions. The method is highly tolerant of moisture.

Key words: aryl chlorides, dechlorination, heterogeneous catalysis, nickel-on-charcoal

Reduction of an aryl chloride to the corresponding arene is usually accomplished via a process involving palladium catalyzed transfer hydrogenation using formic acid or its salts as the source of hydrogen.¹ While several other methods have appeared over the years which, likewise, effect a net aryl C-Cl to C-H conversion {e.g., Raney Ni,² "KCuH2",³ cat Ni(II)/Zn(0)/EtOH,⁴ cat Pd(0)/NaOCH3,⁵ cat NiCl₂/MgH₂,⁶ cat [Ru]/s-BuOH,⁷ cat [Rh]/HSiEt₃,⁸ cat Cp₃La/NaH,⁹ and cat Cp₂TiCl₂/RMgBr},¹⁰ functional group compatibility is rarely addressed. Many reflect development driven by environmental issues, specifically the goals of modifying dioxins and PCBs,¹¹ where only reduction of the carbon-chlorine bond(s) is of paramount concern. In fine chemicals synthesis, however, clean and chemoselective reductive methods which involve neutral conditions and thus tolerate a variety of functionality, especially electrophilic centers, could prove to be quite valuable. For example, nature provides a plethora of chlorinated, physiologically active compounds,¹² including such topical examples as the antitumor agent cryptophycin 1¹³ and the clinically critical antibiotic vancomycin (cf. aglycon below),¹⁴ the des-chloro derivatives of which might be of interest from the perspective of structure-activity relationships.

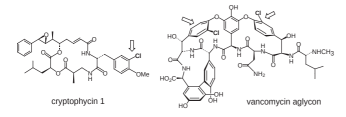
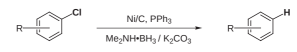


Figure 1

Moreover, a relatively strong C-Cl bond could be viewed as a blocking group for an aryl site, and/or it might also function stereoelectronically to direct electrophilic attack onto an aryl ring. We now describe an operationally simple, environmentally friendly, and inexpensive method for reducing aryl chlorides using nickel on charcoal as catalyst (Scheme 1).



Scheme 1

Previously, we have demonstrated that readily prepared Ni/C is highly effective at catalyzing C-C bond constructions of the Negishi,^{15a} Kumada,^{15b} and Suzuki^{15c} types. More recently, Ni/C has been shown to catalyze aminations of aryl chlorides.¹⁶ In turning our attention to a reductive process which could take advantage of the ability of Ni(0) to oxidatively insert into an aryl-chlorine bond, it seemed reasonable that a suitable source of hydride could be found which would participate in the catalytic cycle. Our initial attempts utilizing H₂ at atmospheric pressure or above (up to 27 psi in a Parr hydrogenator)¹⁷ were fruitless (i.e., no reaction occurred), as were reactions employing either HCO₂H or HCO₂NH₄.¹ Products of reduction could be obtained in simple systems using Red-Al, although functional group compatibility is likely to be compromised with this potent hydride source on more highly derivatized cases. Excellent results were eventually noted using a commercially available, inexpensive amineborane¹⁸ in the presence of K_2CO_3 . Thus, treatment of an aryl chloride with 1.1 equivalents of both Me₂NH•BH₃ and K₂CO₃ under the influence of 5% Ni/C and 20% triphenylphosphine in refluxing acetonitrile required only 5-10 hours to effect complete reduction. Further streamlining this methodology is the fact that prior reduction of the Ni(II)/C to Ni(0)/C with two equivalents (i.e., 10 mol%) of *n*-BuLi in an inert solvent (e.g., THF)¹⁵ is not necessary. Thus, the slight excess of Me₂NH•BH₃/K₂CO₃ (i.e., 10 mol%) being used is presumably responsible for in situ generation of the catalyst in active zero valent form.

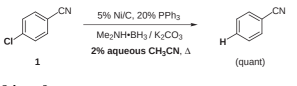
Table 1 illustrates several representative aryl chlorides examined, including those bearing electrophilic centers (entries 7, 8 and 10). Considering that a nitrile is used as solvent, clearly this residue is inert as well (entries 1 and 2). Based on the observation that relatively acidic protons on nitrogen in amide 2 and indole 3 did not interfere, an experiment was conducted using moist acetonitrile as the

Table 1 Representative Reductions of Aryl Chlorides Using 5%Ni/C+20%PPh3 as Catalyst

Entry	Aryl chloride	Time (h)	Yield(%) ^a
1		5	100
2	CN	9	99
3	CL	7.5	99
4	MeO	8	91
5	CF3 CI	7	98
6	F	5	97
7	Eto	8	100
8	Ph 2	7	96 ^b
9		6	95
10		6 Et	96 ^b

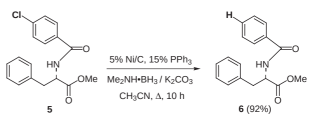
^aBy quantitative GC. ^bIsolated yield.

reaction medium. Treatment of **1** in 2% aqueous CH_3CN under otherwise identical conditions (*cf.* Table 1, entry 1), afforded the desired reduced arene quantitatively (Scheme 2).



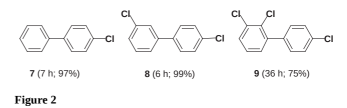


Substrates bearing *ortho*-substituents were smoothly reduced (entries 2, 6, and 10), while both electron-rich and electron-poor cases (entries 4 and 7, respectively) afforded similar results at surprisingly similar rates. Several of the above observations are embodied in the efficient conversion of a chlorinated racemic biaryl ether to its dechlorinated derivative **4** (entry 10). Phenylalanine methyl ester **5** { $[\alpha]_D^{25} = -63.7^\circ$ (c = 1.1, 95% EtOH; lit -64° }¹⁹ could be smoothly reduced to **6** in high isolated yield essentially without loss of optical purity { $[\alpha]_D^{25} = -44.4^\circ$ (c = 1.0, 95% EtOH; lit. -45.3° ;¹⁹ 98% ee; Scheme 3}. This method is limited, however, to non-aldehyde- and -ketone-containing educts, as *p*-chlorobenzophenone led mainly to diphenylcarbinol.



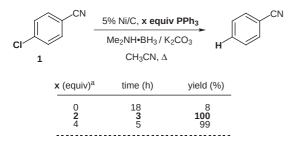
Scheme 3

Application of this Ni/C-amine-borane combination to reductions of (poly)chlorinated biphenyls (PCBs)¹¹ was also possible. Mono-, di-, and trichlorobiphenyls, (**7-9**, respectively) were all successfully converted to biphenyl (Figure 2).²⁰ In the case of **9**, however, incomplete reduction was observed even after 36 hours, the remaining material consisting of *ortho*-chlorobiphenyl.



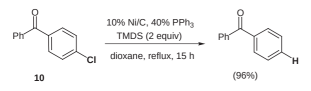
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Prior experience with Ni(0)/C suggests that a phosphine ligand (usually PPh₃) is required for a synthetically useful level of conversion, and ultimately, isolated yield.¹⁵ Typically, 3-4 equivalents are needed relative to the percentage of nickel loaded onto the solid support, an assumption used in carrying out the examples illustrated in Table 1. Although seemingly the norm for Ni(0)/C-mediated C-C bond constructions, it is not obvious that a reductive process will have identical requirements.¹⁶ Indeed, in the test case of *p*-chlorobenzonitrile (1, Table 2), only two equivalents of PPh₃ relative to nickel were needed to obtain the desired product quantitatively. Moreover, the rate of this reduction was actually somewhat faster than that noted under conditions where twice the amount of phosphine (i.e., 4 equivalents) was present (cf. Table 1, entry 1). In the complete absence of phosphine, the level of conversion was dramatically reduced.



^arelative to nickel

In an attempt to broaden the scope of this process, a silane was envisioned as a particularly mild donor of hydride. Tetramethyldisiloxane (TMDS)²¹ was therefore selected to serve in this capacity. Although reduction of chloroketone 10 to benzophenone required higher temperatures (i.e., refluxing dioxane), it took place very efficiently and without competitive 1,2-carbonyl addition (Scheme 4). Unfortunately, it was found that a considerable amount of nickel had bled off the solid support. Thus, whereas analyses via inductively coupled plasma (ICP)²² indicated that reductions involving an amine-borane led to at most a 3% loss of nickel from the 5% Ni/C being used, 80% of the Ni(II) which had been mounted on charcoal and used as catalyst in the reaction of 10 could be detected in the aqueous sample prepared from the filtrate of the crude reaction mixture involving TMDS.

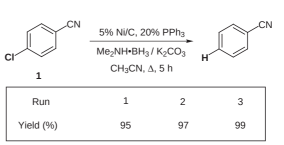


Scheme 4

This catalyst bleeding appears to be an unusual and unexpected phenomenon characteristic (thus far) exclusively of silanes, as similar observations were made when TMDS was replaced by polymethylhydrosiloxane (PM-HS)²³ or Et₃SiH.

Finally, the question of catalyst-recycling has been addressed using this procedure for aryl chloride reductions. Thus, upon completion of the reduction of **1**, filtration of the reaction mixture through a sintered glass frit allowed for recovery of the spent Ni/C. This marterial was subjected to two additional reaction cycles using fresh substrate and amine-borane to afford essentially identical results (Scheme 5).

In summary, a new method has been developed which allows for high yield reductions of aryl chlorides to the corresponding arenes mediated by the heterogeneous catalyst Ni/C.²⁴ The procedure involved is straightforward and especially forgiving with respect to levels of moisture present in the medium. Reaction work-up is particularly





simple (i.e., filtration and solvent evaporation), as is characteristic of this type of catalysis. Several of the more common electrophilic functional groups of interest are tolerated,²⁵ only stoichiometric amounts of hydride are needed (1.0 equivalent relative to substrate), and given the likely in situ generation of the kaliated form of Me_2NH •BH₃, the reaction conditions are essentially neutral.

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- (25) We have recently observed that isolated olefins, unfortunately, are reduced under these conditions.

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