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Nickel-catalyzed cyanation of aryl halides and triflates using acetonitrile *via* C-CN bond cleavage assisted by 1,4-bis(trimethylsilyl)-2,3,5,6-tetramethyl-1,4-dihydropyrazine†

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We developed a non-toxic cyanation reaction of various aryl halides and triflates in acetonitrile using a catalyst system of $[Ni(MeCN)_6](BF_4)_2$, 1,10-phenanthroline, and 1,4-bis(trimethylsilyl)-2,3,5,6-tetramethyl-1,4-dihydropyrazine ($Si-Me_4$ -DHP). $Si-Me_4$ -DHP was found to function as a reductant for generating nickel(0) species and a silylation reagent to achieve the catalytic cyanation via C-CN bond cleavage.

Nitriles are highly versatile functional compounds often used to prepare the corresponding amines, amides, and carboxylic acids.1 Several useful synthetic protocols have been established as not only stoichiometric reactions, such as the Sandmayer reaction and Rosenmund-von Braun reaction using CuCN,2 but also metal-catalyzed cyanation reactions of aryl halides or aryl triflates using toxic cyanide reagents, 1f,3 such as NaCN,4 KCN,5 CuCN,6 Zn(CN)2,7 and Me3SiCN.8 Recent efforts have been aimed at developing non-toxic cyanation reactions using alkylnitriles,5h,9 despite the relatively high bond dissociation energy of the C-CN bond (e.g. 133 kcal mol⁻¹ for the C-CN bond of acetonitrile) compared with that of the typical C(sp³)-C(sp³) bond (ca. 83 kcal mol⁻¹).¹⁰ In fact, pivotal studies using acetonitrile as a cyano source were recently achieved at high temperature (120–160 °C): Cheng et al. used phosphine complexes of palladium and nickel for the catalytic cyanation of ortho-mono and ortho-disubstituted aryl halides with acetonitrile in the presence of zinc powder at elevated temperature (160 °C) (Fig. 1a), 9d and Shen et al. applied a catalyst system of copper(II) nitrate with 2,2,6,6-tetramethylpiperidine-1-oxyl (TEMPO) for the cyanation of aryl iodides in acetonitrile at high temperature (150 °C) (Fig. 1b). 9e,f Quite recently, Morandi et al. demonstrated that a combination of bis(acetylacetonato) nickel(II) with Xantphos became a catalyst for the cyanation of aryl chlorides and aryl triflates with *n*-butyronitrile as a cyano

source in the presence of highly reactive Lewis acids such as tri(isobutyl)aluminum at high temperature (120 °C) (Fig. 1c). 9g

In this context, we have continuously focused our efforts on finding a catalyst system active for such cyanation using acetonitrile under milder reaction conditions. Herein, we report a nickel-catalyzed cyanation of aryl halides and aryl triflates with acetonitrile at 80 °C or 100 °C in the presence of an organosilicon compound, 1,4-bis(trimethylsilyl)-2,3,5,6-tetramethyl-1,4-dihydropyrazine (Si-Me₄-DHP), which served as both an effective reductant and a silylating reagent for nickel species (Fig. 1d).¹¹ It is noteworthy that C–CN bond cleavage of acetonitrile was achieved by α -methyl elimination from an [N-(trimethylsilyl)iminoacyl]nickel intermediate, which was generated by insertion of acetonitrile into Ni–SiMe₃ species.

We started by searching for the best nickel catalyst system for the cyanation of 4-bromoanisole (1a) in acetonitrile, and the results are summarized in Table 1. The reaction mixture of 1a and Si-Me₄-DHP (2.5 equiv.) in the presence of catalytic amounts of [Ni(MeCN)₆](BF₄)₂ (2, 5 mol%) and 2,2'-bipyridine (L1, 5 mol%) in acetonitrile was heated at 80 °C for 24 h to give p-anisonitrile (3a) in 33% yield (entry 1). We further examined other nitrogen-based bidentate ligands: 4,4'-disubstituted-2,2'bipyridine L2-L4 afforded 3a in low yields (entries 2-4), whereas 1,10-phenanthroline (L5) drastically improved the yield of 3a up to 91% (entry 5). Other 1,10-phenanthroline-based ligands bearing substituents at 4,7-positions, such as methyl (L6), methoxy (L7), and phenyl (L8), were less effective than L5, giving 3a in 58%, 30%, and 59% yields, respectively (entries 6-8). Accordingly, we selected L5 as the best ligand. Furthermore, we found that the catalytic activity was quite sensitive to the amounts of L5: 2 equiv. of L5 slightly decreased the yield (86%) of 3a and 3 equiv. of L5 significantly suppressed the activity to 54% yield (entries 9 and 10), the latter of which was consistent with the observation that an isolated dicationic tris(1,10-

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Recent catalytic cyanation reactions using alkylnitriles.

Optimization for the cyanation of 1a with acetonitrile^a

Entry	Ligand (x mol%)	Reductant	Yield ^b (%)
1	L1 (5)	Si-Me ₄ -DHP	33
2	L2 (5)	Si-Me ₄ -DHP	19
3	L3 (5)	Si-Me ₄ -DHP	17
4	L4 (5)	Si-Me ₄ -DHP	14
5	L5 (5)	Si-Me ₄ -DHP	91
6	L6 (5)	Si-Me ₄ -DHP	58
7	L7 (5)	Si-Me ₄ -DHP	30
8	L8 (5)	Si-Me ₄ -DHP	59
9	L5 (10)	Si-Me ₄ -DHP	86
10	L5 (15)	Si-Me ₄ -DHP	54
11 ^c	_ ` ´	Si-Me ₄ -DHP	61
12^d	L5 (5)	Si-Me ₄ -DHP	75
13^e	L5 (5)	Si-Me ₄ -DHP	86
14	_	Si-Me ₄ -DHP	10
15	L5 (5)	Zn	0
16	L5 (5)	Mn	0
17	L5 (5)	TDAE	0
18	L5 (5)	_	0

 $[^]a$ Reaction conditions: 1a, 0.10 mmol (0.025 M). b 1 H NMR yield using 1,3,5-trimethoxybenzene as an internal standard. c [Ni(L5)₃](BF₄)₂ was used instead of 2. d 60 °C. e 100 °C.

phenanthroline)nickel complex, [Ni(L5)₃](BF₄)₂, afforded 3a in 61% yield (entry 11). Upon varying the reaction temperature at 60 °C and 100 °C, we obtained 3a in lower yields (75% and 86% yields, respectively) compared with the reaction conducted at 80 °C, giving 3a in 91% yield (entries 12 and 13 vs. entry 5). The addition of L5 was essential: the catalyst system without L5 gave 3a in a low yield (10%) (entry 14). The selection of the reducing reagents is the most significant factor for the cyanation reaction: no cyanation was observed when applying typical metal-based reducing reagents, such as zinc and manganese powder, as well as a typical organic reductant, tetrakis(dimethylamino)ethylene (TDAE) (entries 15-17), and the reaction without Si-Me₄-DHP did not produce 3a (entry 18), indicating that Si-Me₄-DHP was

indispensable for the catalytic cyanation reaction (vide infra). As a result, we selected the mixture of catalytic amounts of 2 (5 mol%) and L5 (5 mol%) in combination with an excess amount of Si-Me₄-DHP (2.5 equiv.) as the best catalyst system. 12

With the best catalyst system in hand, we examined the substrate scope of this nickel-catalyzed cyanation reaction (Table 2). Aryl bromides with electron-donating groups such as methoxy (1a), dialkylamino (1b-1d), and alkyl groups (1e and 1f) at the para-position gave the corresponding aryl nitriles 3a-3f in excellent to good yields (70-88%). Additionally, aryl bromides with electronically neutral para-substituents such as trimethylsilyl (1g), phenyl (1h), and p-methoxyphenyl (1i) were efficiently transformed into the corresponding aryl nitriles 3g-3i in good yields (61-77%). On the other hand, aryl bromides with electron-withdrawing groups significantly decreased the yield of the corresponding aryl nitriles though the aryl bromides were completely consumed; 4-fluorobromobenzene (1j) afforded 4fluorobenzonitrile (3i) in 36% yield together with unidentified byproducts, while aryl bromides with cyano, methoxycarbonyl, nitro, and trifluoromethyl groups resulted in a complex mixture. When the amount of Si-Me₄-DHP was reduced to 1.1 equiv. for the cyanation of 1j, 3j was obtained in 50% yield together with 4-fluoroacetophenone (14% yield) as the main byproduct after acidic work-up. Typical protecting groups such as benzyloxycarbonyl (Cbz) and benzyl (Bn) groups were well tolerated to give the corresponding aryl nitriles in 54-86% yields (3k-3m). Notably, this nickel catalyst system was less sensitive to the steric hindrance around the bromide group: 4-methoxybromoarenes **1n-1p**, which respectively contained 3-Me, 2-Me, and 3,5-Me2 substituents, afforded the corresponding arylnitriles 3n-3p in 65-88% yields. In addition, the cyanation of a polycyclic compound, 1-bromo-4-methoxynaphthalene (1q), gave 3q in 83% yield. We further investigated the catalytic cyanation of several aryl triflates (4a, 4h, 4r, and 4s), which were effectively converted to the corresponding aryl nitriles 3a, 3h, 3r, and 3s in 59-97% yields by increasing the catalytic loadings

Table 2 Substrate scope of substituted aryl halides and triflates^a

(15 mol%) and temperature (100 °C). Moreover, the cyanation of aryl chlorides (5a and 5t–5v) afforded 3a and 3t–3v in 59–77% yields, whereas the cyanation of 4-iodoanisole resulted in complex mixtures.

This nickel catalyst system was applicable to a wide range of heterocyclic compounds (Table 3): excellent to moderate yields were obtained for the cyanation of five- and six-membered heterocycles such as indoles (3w-3z) (53-99%), carbazole (3aa) (86%), indoline (3ab) (70%), dibenzofuran (3ac) (88%), benzothiophene (3ad) (48%), benzoxazole (3ae) (61%), pyrazole (3af) (52%), and pyrimidine (3ag) (62%). In all cases, no decomposition of the heterocycle motif was observed.

Regarding the reaction mechanism, we conducted some controlled experiments. The reaction of $Ni(cod)_2$ with 4-methoxy-2-methylbromobenzene (**10**) in the presence of 1 equiv. of **L5** afforded a nickel(π) σ -aryl complex, $Ni(C_6H_3$ -4-OMe-2-Me)Br(**L5**) (**6**), in 76% yield, π which was characterized by spectral data as well as X-ray diffraction study of its single crystal. We then monitored the solution of **6** in acetonitrile with or without Si-Me₄-DHP at 80 °C for 24 h: the reaction of **6** with acetonitrile in the presence of Si-Me₄-DHP proceeded to give **3n** in a substantial yield (24%), whereas no reaction of **6** with

acetonitrile was observed in the absence of Si-Me₄-DHP or in the presence of other reducing reagents (eqn (1)), clearly indicating that complex **6** was a key intermediate and Si-Me₄-DHP played a key role for giving **30**. In fact, cyanation of **1a** in the presence of 10 mol% of **6** produced **3a** in 80% yield with stoichiometric amounts of **3o** from **6** (eqn (2)). It is noteworthy that Si-Me₄-DHP was consumed in the presence of Ni(cod)₂ (10 mol%) at room temperature to produce (Me₃Si)₂ and 2,3,5,6-tetramethylpyrazine (Me₄-pyrazine).¹⁴ We thus assume that Si-Me₄-DHP acts as a silylation reagent of nickel species.

^a Isolated yields. ^b Reaction conditions for aryl bromides **1a–1s** and aryl chloride **5a: 1** or **5**, 0.40 mmol (0.027 M); **2** (5 mol%); **L5** (5 mol%); **80** °C. ^c Reaction conditions for aryl triflates **4a**, **4h**, **4r**, and **4s** and aryl chlorides **5t–5v**: **4** or **5**, 0.40 mmol (0.027 M); **2** (15 mol%); **L5** (15 mol%); **100** °C. ^d ¹H NMR yield. ^e Si–Me₄-DHP (1.1 equiv.) was used.

Table 3 Substrate scope of heterocyclic aryl bromides^a

 a Reaction conditions: 1, 0.40 mmol (0.027 M). Isolated yield for all the products.

Scheme 1 shows a plausible catalytic cycle for the nickelcatalyzed cyanation reaction with acetonitrile as the sole cyano source based on the results of the controlled experiments shown in eqn (1) and (2). The initial step is the reduction of 2 by Si-Me₄-DHP in the presence of L5, producing a 1,10phenanthroline-coordinated Ni(0) species A.116 Subsequently, oxidative addition of aryl halides or aryl triflates (ArX) to A gives a Ni(II) species, Ni(Ar)X(L5) (B), X of which is silylated by the reaction with Si-Me₄-DHP to afford Ni(Ar) (SiMe₃) (L5) (C) with a release of Me₃SiX and Me₄-pyrazine. Insertion of acetonitrile into the Ni-Si bond of species C forms N-(trimethylsilyl)iminoacyl species D.15 α-Methyl elimination from D generates isonitrile species E, and the subsequent 1,1-insertion of CNSiMe₃ into the nickel-aryl moiety results in the formation of F. β-Silyl elimination of F spontaneously induces a liberation of aryl nitriles accompanied by the generation of Ni(Me) (SiMe₃) (L5) (G).16 Finally, the second insertion of acetonitrile into the Ni-Si bond of G followed by reductive elimination of N-(trimethylsilyl)propan-2-imine from H regenerates A. Notably, this is consistent with the formation of 4-fluoroacetophenone after acidic work-up when aryl bromide 1j with an electronwithdrawing substituent is used as a substrate; reductive elimination from species D or F is competitive with the formation of G and 3j.

Scheme 1 Plausible reaction mechanism.

Furthermore, we conducted the catalytic reaction in benzylnitrile under the same catalytic conditions, and we obtained a mixture of **3a** (46%), 1-benzyl-4-methoxybenzene (33%), and dibenzylketone (12%) (eqn (3)).¹⁷ Thus, the formation of 1-benzyl-4-methoxybenzene corresponded to reductive elimination from a similar intermediate, Ni(Ar) (Bn) (CNSiMe₃) (L5) (E' in SI), and the formation of dibenzylketone corresponded to the formation of acetone after hydrolysis of *N*-(trimethylsilyl)propan-2-imine derived during the reaction course.

Conclusions

We developed a toxic cyanide-free cyanation of aryl halides and aryl triflates using a nickel catalyst system of $[Ni(MeCN)_6](BF_4)_2$ and 1,10-phenanthroline along with a stoichiometric amount of an organosilicon reductant, $Si-Me_4$ -DHP, in acetonitrile. Noteworthy was that the mechanism of our catalytic system differs from the well-established direct C-CN bond cleavage involving Lewis acid-assisted oxidative addition to electron-rich low-valent group 10 metal complexes. 9g,18,19 In this catalytic system, $Si-Me_4$ -DHP had dual functions to reduce the nickel(π)

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catalyst precursors but also to serve as a silylation reagent for nickel(II) σ -arvl species. Further application of Si-Me₄-DHP as a silylating reagent to transition metals is ongoing in our laboratory.

Conflicts of interest

The authors declare no conflict of interest.

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