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Introduction

Nitriles are important synthesis intermediates in transformation¹ and are key components in various natural products, medicinal pharmacophores and drugs,² and in organic synthesis, the cyano group is equivalent to an amine or carbonyl group. The catalytic addition of the cyano group to the $C=C$ bond has been established as one of the most direct pathways for the synthesis of nitriles.^{3,4} Among the important nitriles, β cyano-ketones are commonly utilized in organic synthesis.⁵ One of the classical approaches to deliver these compounds is the catalyzed conjugate addition of cyanide to α , β -unsaturated carbonyl compounds (hydrocyanation procedure, Scheme 1a),⁶ which utilizes the highly toxic and explosive HCN gas as the cyano source. An alternative strategy is transfer hydrocyanation, which involves the commercially available, less-toxic and lessexplosive cyanohydrin to deliver nitriles (Scheme 1b), but with low atom economy.⁷ Recently, Morandi developed a nickelcatalyzed transfer hydrocyanation reaction between alkyl nitriles and alkenes or aryl chlorides, which utilizes non-toxic alkyl nitriles as the cyanide source.⁸ Although the catalyzed hydrocyanation and transfer hydrocyanation reactions have been well developed,^{6,7} it remains an important challenge to bypass the usage of toxic HCN gas as the cyano source and overcome the issue of atom-economy. To mitigate these concerns and inspired by the atom- and step-economical procedure of β -alkylation of secondary alcohols with aldehydes via borrowing hydrogen reactions,^{9,10} we postulated that the cyano group could be tolerated in the reaction with a mechanism analogous to the borrowing hydrogen reactions. **EDGE ARTICLE**
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Cyano-borrowing reaction: nickel-catalyzed direct conversion of cyanohydrins and aldehydes/ketones to *β***-cyano ketone**†

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A direct nickel-catalyzed, high atom- and step-economical reaction of cyanohydrins with aldehydes or ketones via an unprecedented "cyano-borrowing reaction" has been developed. Cleavage of the C–CN bond of cyanohydrins followed by aldol condensation and conjugate addition of cyanide to α , β unsaturated ketones proceeded to deliver a range of racemic β -cyano ketones with good to high yields. The practical procedure with the use of a commercial and less-toxic CN source bodes well for wide application of this protocol.

$$
R \xrightarrow{\begin{array}{c}\nO \\
R\n\end{array}} \n\begin{array}{ccc}\nO & \text{Catalyst} \\
H \text{V} & \text{CNO} \\
\end{array}
$$

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d) This work: Nickel-Catalyzed Cyano-Borrowing Reaction

High atom-economy: H₂O as the sole side product C-CN bond cleavage and formation C Commercial, non-toxic CN source

Scheme 1 Development of cyano-borrowing

[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/c9sc00640k

Results and discussion

To test our hypothesis, we initiated the cyano-borrowing reaction using commercially available acetophenone cyanohydrin 1a which could be prepared from acetophenone and TMSCN and benzaldehyde 2a for the optimization of the reaction conditions. After screening an array of transition metal catalysts, we found that nickel complexes showed better performance. To our delight, a cocktail consisting of $NiBr₂$, $PPh₃$, and LiOH as the base in dioxane at 100 $^{\circ}$ C could deliver the desired racemic b-cyano ketone 3aa in 52% yield (Table 1, entry 1). Notably, determined by the crude ${}^{1}\mathrm{H}$ NMR of the reaction mixture, not even a trace of 1,2-addition products was observed.¹⁴ As shown in Table 1, we then examined an extensive array of parameters. By varying the anion of the nickel salt $(i.e.,$ $NiCl₂, Ni(OAc)₂, Ni(OTf)₂ and Ni(acac)₂ are precatalysts, NiCl₂$ exhibited the best yield (Table 1, entries 3–5 vs. 2). The screening of different phosphines revealed that this transformation requires a bulky, electron rich ligand, and ${}^t\mathrm{BuPAd}_2$ showed the highest reactivity with 82% yield (Table 1, entry 11 vs. 2 and 6–10). Stronger bases such as NaOH and $\mathrm{KO}^t\mathrm{Bu}$ show low reactivity. Meanwhile, Cs_2CO_3 and organic bases (*i.e.*, DMAP and DBU) failed to provide the target products (Table 1, entries 14–16). Optimization of the solvent led to no improvement (Table 1, entries 17-19). Lowering the temperature to 80 \degree C gives a lower yield (Table 1, entry 20). Control experiments verified that the presence of a Ni-complex was necessary to achieve high yield in the cyano-borrowing reaction (Table 1, entry 21). No reaction occurred in the absence of LiOH (Table 1, entry 22). After the screening of the reaction parameters, we found that NiCl₂ and ⁿ⁻BuPAd₂ as the precatalyst and LiOH as the base in dioxane at 100 °C for 20 h (82% yield) were the optimal conditions.

Having identified the optimized reaction conditions, we further explored the substrate scope of this reaction. Various commercially available ketone cyanohydrins were examined with benzaldehyde (2a) and the results are summarized in Scheme 2. The electron-deficient substrate with varying substituent patterns at the *p*-position did not dramatically influence yields (3ba–3da). Substituting the cyanohydrin with p methyl also shows high reactivity and 72% of the desired product was obtained (Scheme 2, 3ea). However, substituting

Table 1 Screening studies of β -alkylation of cyanohydrin 1a with benzaldehyde 2a^c

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Guided by these considerations, we envisioned that, as shown in Scheme 1c, under the catalysis of a transition metal, there is		Table 1 Screening studies of β -alkylation of cyanohydrin 1a with benzaldehyde 2a ^a				
cleavage of the C-CN bond ^{11,12} of the cyanohydrin and delivery of the corresponding ketones and metal-cyano intermediate $([M]^n$ -CN), followed by aldol condensation of ketones with aldehydes, and subsequent conjugate addition of cyanide to chalcones and utilization of the $[M]^{n+1}$ -CN as the cyano donor to deliver the desired products (Scheme 1c). This hypothesis is		HQ CN 1a	CHO 2a	[Ni] (5 mol%) L (10 mol%) Base (3.0 equiv.) Solvent, 4A MS 100 °C		CN Заа
notable in that cyanohydrin plays a dual role both as the source	Entry	[Ni]	L	Base	Solvent	Yield b (%)
of ketone and the cyanide donor with high atom- and step- economy. Herein, we report the first catalytic process of the direct transformation of cyanohydrins with aldehydes to deliver β-cyano ketones as the sole product via nickel catalyzed cyano- borrowing reactions. ¹³	1 2 3 4 5 6	NiBr ₂ NiCl ₂ $Ni(OAc)_{2}$ $Ni(OTf)_2$ $Ni (acac)_2$ NiCl ₂	PPh ₃ PPh ₃ PPh ₃ PPh ₃ PPh ₃ PCy_3	LiOH LiOH LiOH LiOH LiOH LiOH	Dioxane Dioxane Dioxane Dioxane Dioxane Dioxane	52 76 66 62 71 76
Results and discussion	7 8	NiCl ₂ NiCl ₂	dppe dppp	LiOH LiOH	Dioxane Dioxane	72 78
Open Access Article. Published on 06 May 2019. Downloaded on 5/21/2025 8:30:06 AM. This article is licensed under a Creative Commons Attribution 3.0 Unported Licence. To test our hypothesis, we initiated the cyano-borrowing reac- tion using commercially available acetophenone cyanohydrin 1a which could be prepared from acetophenone and TMSCN and benzaldehyde 2a for the optimization of the reaction conditions. After screening an array of transition metal cata- lysts, we found that nickel complexes showed better perfor- mance. To our delight, a cocktail consisting of NiBr ₂ , PPh ₃ , and LiOH as the base in dioxane at 100 °C could deliver the desired racemic β-cyano ketone 3aa in 52% yield (Table 1, entry 1). Notably, determined by the crude ${}^{1}H$ NMR of the reaction	9 10 11 12 13 14 15 16 17 18 19 20 ^c 21	NiCl ₂ NiCl ₂	dppb BINAP n -BuPAd ₂ n -BuPA d_2 n -BuPAd ₂ n -BuPAd ₂	LiOH LiOH LiOH NaOH KOt Bu Cs_2CO_3 DMAP DBU LiOH LiOH LiOH LiOH LiOH	Dioxane Dioxane Dioxane Dioxane Dioxane Dioxane Dioxane Dioxane Toluene TBME THF Dioxane Dioxane	37 44 82 28 Trace $\boldsymbol{0}$ $\boldsymbol{0}$ 0 22 23 72 59 < 10
mixture, not even a trace of 1,2-addition products was observed. ¹⁴ As shown in Table 1, we then examined an extensive array of parameters. By varying the anion of the nickel salt $(i.e.,$ NiCl ₂ , Ni(OAc) ₂ , Ni(OTf) ₂ and Ni(acac) ₂) as precatalysts, NiCl ₂ exhibited the best yield (Table 1, entries $3-5$ vs. 2). The screening of different phosphines revealed that this trans- formation requires a bulky, electron rich ligand, and t BuPAd ₂ showed the highest reactivity with 82% yield (Table 1 entry 11	n -BuPAd ₂ 22 NiCl ₂ $\overline{0}$ Dioxane a The reaction was carried out with 0.4 mmol of 1a, 0.4 mmol of 2a, 5 mol% [Ni], 10 mol% ligand (L) and 300 mol% base in 0.5 mL of solvent at 100 °C for 18 h. $\frac{b}{c}$ Isolated yield. $\frac{c}{c}$ The reaction was carried out at 80 °C. dppm = bis(diphenylphosphino)methane; dppe = bis (diphenylphosphino)ethane; dppp = bis(diphenylphosphino)- propane; dppb = bis(diphenylphosphino)butane; "BuPAd ₂ = di(1- adamantyl)- n butylphosphine; TBME = methyl tert-butyl ether.					

the *para* position with a more electron-rich functionality $(i.e., p-)$ MeO) led to diminished yields, but the desired product could be delivered with 86% yield while increasing the reaction temperature to 120 °C (Scheme 2, 3fa). Reactions of cyanohydrin with ortho- and meta-substitution on aryl groups gave excellent yields (Scheme 2, 3ga–3ja). The reaction proceeds smoothly in the case of the cyanohydrin bearing a thienyl group, affording the corresponding product 3la in 82% isolated yield. In terms of the alkyl substituents, cyanohydrin derived from butanone proceeded smoothly to afford the desired product with 40% yield (Scheme 2, 3ma). The substrate bearing a cyclopropyl group was well tolerated, leading to the cyclopropyl substituted product with 57% yield (Scheme 2, 3na). Importantly, the cyclopropyl group remains untouched, which indicates that this nickelcatalyzed protocol does not proceed via a radical pathway.

Next, various aldehydes were investigated with acetophenone cyanohydrin (1a) using the optimized reaction conditions, and representative results are summarized in Scheme 3. Benzaldehydes bearing various electron-deficient (3ab-3ad and 3ag-3ah), electron-neutral (3al and 3am) and electron-rich (3ae–3af

Scheme 2 Reaction scope of ketone cyanohydrins. ^aThe reaction was carried out at 120 °C

Scheme 3 Reaction scope of aldehydes. ^aThe reaction was carried out at 120 \degree C

and 3ai–3ak) substituents reacted with 1a to deliver the desired products in moderate to excellent yields (58–90%). Furthermore, the reaction is also compatible with heteroaryl rings, such as 2-furanyl (3an), 2-thiophenyl (3ao) and unprotected 3-indolyl (3ap), providing diverse β -cyano ketones in 51–77% yield. Remarkably, compared with the aryl aldehydes, the alkyl substituted aldehydes (e.g., 2-phenylacetaldehyde and cyclohexanecarbaldehyde) reacted with 1a to afford the corresponding products 3aq and 3ar in 64% and 80% yields, respectively. The aldehyde containing a sulfur atom is also tolerated under the nickel-catalyzed protocol and delivers the desired product 3as with a slightly lower yield. To demonstrate the practicality and scalability of our protocol, we proceeded to carry out a gram-scale reaction with 5.0 mmol 2i reacted with 10.0 mmol 1a catalyzed by 5 mmol% of NiCl₂, affording 1.12 g 3ai in 85% yield, suggesting that this procedure is quite reliable and practically applicable.

Inspired by the success with the nickel-catalyzed cyanoborrowing reaction of cyanohydrins derived from ketones 1 with aldehydes, we further demonstrate the cyano-borrowing protocol with more challenging substrates such as aldehyde cyanohydrin 5. To our delight, benzaldehyde cyanohydrin 5a reacted with acetophenone 4a smoothly under standard conditions, yielding the corresponding product 3aa with full conversion and 83% isolated yield. To indicate the generality of this protocol, we then examined the scope of benzaldehyde cyanohydrins and ketones. In addition to the phenyl group, it was found that substrates bearing electron-rich or electrondeficient substituents on the benzene ring were well tolerated to give β -cyano ketones in moderate to good yields (3ba, 3ea, 3ad and 3af in Scheme 4). Moreover, heteroaryl and alkyl substituents also participated in this protocol very well (Scheme 4, 3la, 3ma, 3ao and 3ar).

To illustrate the scope and limitations of the new transformation, challenging substrates beyond methyl ketones were further examined, shown in Scheme 5. The propiophenone cyanohydrin 6 was selected to react with benzaldehyde 2a under standard reaction conditions, and the desired product 8 was obtained in 31% isolated yield and with high diastereoselectivity $(dr > 10 : 1)$ (Scheme 5a, left). Furthermore, we also examined benzaldehyde cyanohydrin 5a and propiophenone 7, with a nickel catalyst, and the corresponding product 8 was obtained in 25% yield (Scheme 5a, right). Notably, the bioactive

Scheme 4 Examples of aldehyde cyanohydrin with ketone. ^aThe reaction was carried out at 120 \degree C.

Scheme 5 Cyano-borrowing beyond methyl ketones.

ketone, epiandrosterone was tested in this reaction with the partner of benzaldehyde cyanohydrin 5a, and the corresponding product 10 was delivered in 57% yield with 5 : 1 dr (Scheme 5b), which shows the potential of this nickel-catalyzed cyano borrowing process for the selective modification of bioactive ketones.

To shed light on the mechanism for the nickel-catalyzed cyano-borrowing protocol, a series of control experiments were conducted. As shown in Scheme 6, acetophenone cyanohydrin 1a reacted smoothly with chalcone 11 and give the corresponding product 3aa in excellent yield, which indicated that cyanohydrin is the source of the cyano group in transfer hydrocyanation under standard reaction conditions. Benzaldehyde cyanohydrin 5a could also react with chalcone 11 efficiently, delivering 3aa with 89% yield, and the hydrogenborrowing product 12 was not observed, showing that cleavage of the C–CN bond is easier than that of the C–H bond in cyanohydrins. Meanwhile, in the crossover reaction of 1a, 2e and 11 under standard conditions, we got the corresponding products 3aa and 3ae with the ratio of 1.05 : 1, which shows that the cyano group from the cleavage of the C–CN bond of cyanohydrin was a free anion in this nickel-catalyzed protocol

Scheme 6 Mechanism studies.

and has the same opportunity to conjugate to each chalcone. Together, these experimental results support our hypothesis on nickel-catalyzed step- and atom-economical cyano-borrowing reaction of cyanohydrin with aldehydes or ketones (Scheme 1c) (for more details of the mechanism studies, please see the ESI†).

Experimental

General procedure

Method A. To a vial equipped with a dried stir bar was added aldehydes (0.2 mmol), ketone cyanohydrins (0.4 mmol), $NiCl₂$ (5 mol%), ⁿ⁻BuPAd₂ (5 mol%), LiOH (0.6 mmol), 100 mg 4Å MS and anhydrous dioxane (1 mL) in a glovebox. The reaction mixture was taken outside the glovebox and allowed to stir at room temperature for 30 min. After that, the reaction mixture was allowed to stir at 100 $^{\circ}$ C for 18 hours. The crude reaction mixture was concentrated under reduced pressure and directly purified by silica gel chromatography to give pure products.

Method B. To a vial equipped with a dried stir bar was added ketones (0.2 mmol), aldehyde cyanohydrins (0.4 mmol), $NiCl₂$ (5 mol%), ⁿ⁻BuPAd₂ (5 mol%), LiOH (0.6 mmol), 100 mg 4Å MS and anhydrous dioxane (1 mL) in a glovebox. The procedure was the same as Method A.

Conclusions

In conclusion, we have developed an unprecedented nickelcatalyzed protocol for the direct conversion of cyanohydrins and aldehydes or ketones into racemic β -cyano ketones via a nickel-catalyzed cyano-borrowing reaction. A range of cyanohydrins derived from aldehydes or ketones could be tolerated and delivered products with high regioselectivity and good to excellent yields. To the best of our knowledge, catalytic conversion of cyanohydrins into β -cyano ketones by reaction with aldehydes/ketones has not been reported. Further studies of catalytic cyano-borrowing reaction of cyanohydrins are in progress in our research lab and will be reported in due course.

Conflicts of interest

There are no conflicts to declare.

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