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### **Nickel N-heterocyclic carbene-catalyzed cross-coupling reaction of aryl aldehydes with organozinc reagents to produce aryl ketones**



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# **Nickel N**‑**heterocyclic carbene-catalyzed cross-coupling reaction of aryl aldehydes with organozinc reagents to produce aryl ketones**

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**The transformation of aromatic aldehydes into aryl ketones by nickel-catalyzed cross-coupling has been developed. This transformation represents an efficient and attractive synthetic utilization of organozinc reagents. The reaction provides a mild, practical method toward aryl ketones, which are versatile intermediates and building blocks in organic synthesis.** 

Aryl ketones are multifaceted compounds that function as components of pharmacophores, photolabels and photosensitizers, organic electronics, and polymers.<sup>1</sup> Furthermore, manipulation of the carbonyl group provides a generic entry into molecules containing benzylic functionality. As a consequence, their preparation has received considerable attention in organic chemistry. Classical methods for the formation of these important structural motifs include the addition of arylmetal species to carbonyl compounds and Friedel-Crafts acylation reactions.<sup>2</sup> However, the presence of aromatic heterocyles can limit these strategies, as organometallic reagents can be incompatible with heteroaromatic nuclei and Friedel-Crafts acylations usually work best with *π*-rich arenes and sometimes result in isomeric mixtures. An alternative approach is the reaction of diazo compounds with aldehydes,<sup>3</sup> but use of structurally diverse diazo compounds is hampered by preparation and safety issues. In recent years, significant progress has been achieved in carbonylative Suzuki–Miyaura cross-coupling reaction. Various substituted aryl ketones have been successfully synthesized.<sup>4</sup> However, the high toxicity, and odorless and flammable character of CO gas means that transformations using CO gas must be operated with special care. More recently, metal-catalyzed C-H arylation

of aldehydes has provided new pathways for the construction of these molecules.<sup>5</sup> Related to this strategy, aryl ketones formation from aldehydes has been realized using NHC catalysis through reactions with aromatic electrophiles or with benzynes.<sup>6</sup> The carbon-carbon bond is the most widespread and fundamental bond existing in organic compounds. Very recently, preparation of aryl ketones by activation of C-C bonds has received much attention.<sup>7</sup> Despite these advances,<sup>8</sup> versatile and efficient methods for the direct construction of aryl ketones that are compatible with various functional groups and use readily available starting materials remain highly desirable.

Organozinc reagents have been known for more than 150 years. These reagents are known as unique nucleophiles compared to other common organometa-llic reagents, that is, the softer nucleophilicity of organozinc reagents shows higher functional-group tolerance.<sup>9</sup> They are increasingly commonly applied in bond construction reactions such as Negishi coupling,  $10$  Fukuyama coupling,  $11$ oxidative cross-coupling, $12$  and aldol reaction, etc.<sup>9</sup> Notably, the formation of alcohols from aldehydes and organozinc reagents has been particularly successful, whereas the synthesis of ketones from the above two substrates has still rarely been reported. As part of our ongoing efforts to develop transition-metal-catalyzed organic reactions, $^{13}$  we herein report an unprecedented nickel N‑heterocyclic carbene (NHC)-catalyzed crosscoupling reaction of aryl aldehydes with organozinc reagents that chemoselectively yields aryl ketones as the desired product. $14,15$  This reaction constitutes a novel transformation from organozinc reagents into aryl ketones. The method advantageously enriches and complements the existing toolbox used by synthetic chemists, and allows a straightforward access to a wide range of functionalized products (Scheme 1).

**The reported work**

 $Ar$   $-$  CHO + RZnBr  $\overline{\phantom{1}}$ OH R

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**Scheme 1** Reaction of aryl aldehydes and organozinc reagents.

Inspired by the pioneering works from the groups of Molander,  $^{16}$  Cheng,  $^{17}$  Matsubara<sup>18</sup> and Dong,  $^{15f}$  we initially chose benzaldehyde **1a** and PhZnBr **2a** as standard substrate to test the feasibility of cross-coupling reaction to form ketones. To our delight, when **1a** and **2a** were treated with a catalytic system consisting of **5a** (hydrogen acceptor, 1.5 equiv),  $MgCl<sub>2</sub>$  (1.0 equiv), Ni(cod)<sub>2</sub> (8 mol%), and IPr (ligands, 10 mol%) in THF at 30 °C, the desired benzophenone **3aa** was formed in 31 % yield (Table S1, entry 1). Tishchenko homodimer **4a** was obtained in 38% yield (Table S1, entry 1). Further explorations showed that hydrogen acceptor **5d** [2,2,2-trifluoro-1-(4-methoxyphenyl)ethanone] gave good results.<sup>19</sup> Next, different solvents were screened to enhance the efficiency of this reaction. It was found that a cosolvent system of 1,4 dioxane and THF was appropriate. Different nickel catalysts were screened, and  $Ni(cod)_2$  turned out to be the most effective catalyst (entries 6-9). Other ligands, including ItBu,  $PCy_3$ , and SIPr, were tested and found to be less effective than IPr (Table S1, entries 10-12). However, the absence of nickel catalyst resulted in no detectable amounts of 3aa (Table S1, entry 16). Without MgCl<sub>2</sub>, the reaction gave the cross-coupling product in 46% yield after 10 h (Table S1, entry 17). These results indicate that the structures of organozinc reagents are possibly not as simple as RZnX. $9b$  Coordination of THF and MgCl<sub>2</sub> to RZnX may significantly affect the transmetalation step.

With a set of optimized conditions in hand, we then investigated a series of aldehydes to extend the substrate scope. As summarized in Table 1, the standard reaction conditions were found to be compatible with a wide range of aldehydes **1**. As summarized in Table 1, both electrondonating and -withdrawing aryl aldehydes could be successfully converted to the corresponding aryl ketones in moderate to good yields. Several aromatic substituents, including Me, MeO, F and  $CF_3$ , were well-tolerated under the standard conditions (Table 1, entries 1-6); substituents at the *ortho*-, *meta*-, or *para*-positions had no distinct influence on the reaction (Table 1, entries 1, 3 and 4). It is noteworthy that trifluoromethyl-substituted aromatic aldehydes were tolerated well (Table 1, entry 6), thus leading to trifluoromethyl-substituted product, which has better membrane permeability and increased bioavailability.<sup>20</sup> Reaction of substrate **1h** with the polysubstituted aromatic aldehyde delivered the desired products **3ah** with a good yield (Table 1, entry 7). Aromatic aldehyde with a naphthyl group also participated in this Ni-catalyzed cross-coupling reaction, affording crosscoupling product in 58% yield (Table 1, entry 8). We also considered the reaction of **2a** with **1j** bearing furanyl moiety and obtained heteroaryl ketone **3aj** in 52 % yield (Table 1, entry 9), further extending the scope of this methodology. Application of aliphatic aldehyde did not deliver the corresponding ketone under the same reaction conditions (Table 1, entry 10). $41$ 

**Table 1** Scope of aldehydes 1

$R^{1} +$ 1	CHO PhZnBr 2a	<b>5d,</b> THF, $Ni(cod)_2$ IPr, 40 °C, MgCl <sub>2</sub> , Ar	о К Ph $R^{1 \frac{\sqrt{3}}{\sqrt{3}}}$ 3
entry	aldehydes 1	aryl ketones 3	yield <sup>b</sup> (%)
$\mathbf{1}$	CHO 1b	Ö Ph 3ab	74
$\overline{\mathbf{c}}$	CHO MeO 1c	ö 3ac MeC	74
3	CHO 1 <sub>d</sub>	Ph 3ad	56
$\overline{4}$	CHO 1e	Ph 3ae	54
5	CHO F 11	Ph 3af	61
6	CHO $F_3C$ 1g	Ο Ph 3ag $F_3C$	67
7	CHO 1h	Ph 3ah	69
8	CHO 11	Ö Ph 3ai	58
9	CHO 1j	U Ph $^{\circ}$ 3aj	52
10	CHO 1k	O Ph 3ak	$\boldsymbol{0}$

*<sup>a</sup>* Reaction conditions: **1** (0.3 mmol), **2a** (0.3 mmol), **5d** (1.5 equiv), MgCl2 (1.0 equiv), Ni(cod)2 (8 mol %), IPr (10 mol %), THF/1,4-dioxane (3 mL, 1:1), 30 ˚C, in Ar atmosphere for 8 h.*<sup>b</sup>* Isolated yield.

To further highlight the synthetic usefulness of our strategy we turned our attention to the scope of organozinc reagents (Table 2). Arylzinc reagents bearing electron-donating groups at the *para* position of the phenyl ring, such as methyl proceeded smoothly in the reaction to give the corresponding aryl ketones **3ab** in good yields (Table 2, entry 1). Substrates substituted with electron-withdrawing groups, such as Ph and ester, also

worked well and afforded the desired products **3ca**-**3da** in 49% and 73% yields, respectively (Table 2, entries 2-3). These results indicate that the electronic nature of the arylzinc reagents has little influence on the cross-coupling process. Heteroaryl-substituted substrates could also be converted into the desired products. For example, substrates **2e** and **2f** were transformed into products **3ea** and **3fa** in 70% and 47% yield, respectively (Table 2, entries 4-5). Efforts were made to apply this methodology for the synthesis of aryl alkyl ketones. It was found that reactions with **2f**-**2h** proceeded smoothly to give the aryl alkyl ketones in good yields (Table 2, entries 6-8).





*<sup>a</sup>* Reaction conditions: **1a** (0.3 mmol), **2** (0.3 mmol), **5d** (1.5 equiv), MgCl2 (1.0 equiv), Ni(cod)2 (8 mol %), IPr (10 mol %), THF/1,4-dioxane (3 mL, 1:1), 30 ˚C, in Ar atmosphere for 8 h. *<sup>b</sup>* Isolated yield.

To further probe the mechanism, control experiments with possible intermediates were designed and investigated. In the absence of organozinc reagents, we observed decarbonylation of *p*-tolualdehyde **1b** to produce toluene and reduction of the hydrogen acceptor **5a** (Scheme 2, 1).<sup>15f</sup> This result supports the intermediacy of an acyl nickel hydride species, which can undergo decarbonylation in the absence of a organozinc reagent. When a 1:1 mixture of 1a and  $1a-d_6$  was subjected to the Ni-catalyzed cross-coupling reaction conditions, we obtained the cross-coupling products  $3$ aa and  $3$ aa- $d_5$  in a ratio of  $5.9:1.^{22}$  The result suggests that C-H bond activation is rate-determining (Scheme 2, 2). Computational studies by Fu and co-workers support the concept of electron-deficient π-ligands on nickel promoting oxidative addition into aldehyde C-H bonds.<sup>23</sup>

**Scheme 2** Control Experiment



On the basis of these preliminary results and those of previous studies, a plausible mechanism for the Nicatalyzed cross-coupling reaction of aryl aldehydes with organozinc reagents yielding aryl ketones was proposed. Ketone **5d** binds to nickel to form complex **A**. Subsequently, complex **A** can coordinate to benzaldehyde **1a** to form intermediate **B**. Oxidative addition to the aldehyde C-H bond gives **C**. 15f Simultaneously, the hydrogen acceptor 5d is reduced by intermediate **C** to produce acyl nickel alkoxide **D**. Ligand exchange with organozinc reagent **2a** affords **E**, and reductive elimination provides the desired product 3aa. Coordination of ketone 5d may occur prior to or immediately after reductive elimination.<sup>24</sup> The coordination inhibits the formation of the undesired product 4a by disrupting the formation of a cycloisomerization intermediate between two aryl aldehydes.

**Scheme 3** Plausible Mechanism.



In summary, we have described a nickel N-heterocyclic carbene-catalyzed cross-coupling reaction of aryl aldehydes and organozinc reagents for the formation of complex aryl ketones. The process operates under mild conditions, involves simple experimental protocols, uses a commercially available N-heterocyclic carbene catalyst.

Further investigations of the mechanism of the reaction 13 (a) L. Gu, C. Jin, R. Wang and H. Ding, *ChemCatChem*., 2014, **6**, and its application are ongoing in our laboratory.

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The transformation of aromatic aldehydes into aryl ketones by nickel-catalyzed cross-coupling has been developed. This transformation represents an efficient and attractive synthetic utilization of organozinc reagents.