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A practical cobalt-catalyzed cross-coupling of benzylic zinc reagents with aryl and heteroaryl bromides or chlorides†

Andreas D. Benischke, Irina Knoll, Alice Rérat, Corinne Gosmini and Paul Knochel*⁵

A catalytic system consisting of 5 mol% CoCl₂ and 10 mol% isoquinoline allows a convenient cross-coupling of benzylic zinc reagents with various aryl and heteroaryl bromides or chlorides leading to polyfunctionalized diaryl- and aryl-heteroaryl-methane derivatives.

Pd-Catalyzed cross-couplings between organozinc reagents and various organic halides constitute a major C-C bond formation methodology (Negishi cross-coupling). Due to the high price and toxicity of palladium, related transition metal-catalyzed cross-couplings involving zinc organometallics and Ni-, Fe- or Co-catalysts have been examined.²⁻⁴ Furthermore, the use of zinc organometallics is of special synthetic interest due to the high functional group compatibility of zinc reagents.⁵ Recently, we have reported several preparation methods of benzylic zinc halides and demonstrated that these reagents undergo smooth Negishi cross-couplings.⁶ Also Bedford reported that benzylic halides undergo useful Fe-catalyzed cross-couplings with arylzinc reagents.7 Gosmini has shown in one-pot procedures that arylzinc reagents generated in situ via a cobalt-catalyzed zinc insertion undergo cross-couplings with benzyl chlorides.8 Interestingly, Ingleson has described a transition metal free cross-coupling between relatively non-functionalized diarylzincs with benzylic bromides and chlorides performed in the

Herein, we report a practical cobalt-catalyzed cross-coupling promoted by 10 mol% of isoquinoline between various benzylic zinc reagents with aryl and heteroaryl bromides or chlorides resulting in the formation of valuable diaryl- and arylheteroarylmethane derivatives.¹⁰ Preliminary control experiments performed with benzylzinc chloride (1a; prepared via the oxidative insertion

of magnesium turnings into benzyl chloride (2a) in the presence of LiCl and ZnCl₂)¹¹ and 4-bromo-benzonitrile (3a) in a 2:1 THF: MTBE mixture¹² (MTBE = methyl tert-butyl ether) show that in the absence of transition catalysts no reaction is observed at 50 °C in 2 h (Table 1, entries 1 and 2). Also, Fe-catalysts such as Fe(acac)₃, Fe(acac)₂ or FeCl₂ were inefficient (Table 1, entries 3–5). ¹³ However, the use of 5 mol% CoBr2, Co(acac)2 and CoCl2 show the formation of the desired cross-coupling product (4a) in 47-76% GC-yield (Table 1, entries 6-8).14

Previously reported additives like 4-fluorostyrene, 15 TMEDA 16 or isoquinoline¹⁷ indicate a very positive effect of 10 mol% isoquinoline¹⁸ leading to an isolated yield of 82% for 4a (Table 1, entry 11; compared with entries 9 and 10). Decreasing the amount of isoquinoline to 5 mol% reduces somewhat the yield of 4a

Table 1 Screening of catalysts for the palladium-free cross-coupling of benzylzinc chloride (1a) with 4-bromobenzonitrile (3a)

Entry	Catalyst (mol%)	Additive (mol%)	Yield ^{a,b}
1	None	None	0
2	None	Isoquinoline (10)	0
3	$Fe(acac)_3$ (5)	None	0
4	$Fe(acac)_2$ (5)	None	Traces
5	FeCl ₂ (5)	None	Traces
6	$CoBr_2(5)$	None	47
7	$Co(acac)_2$ (5)	None	70
8	$CoCl_2$ (5)	None	76
9	$CoCl_2(5)$	4-Fluorostyrene (10)	66
10	$CoCl_2(5)$	TMEDA (10)	68
11	$CoCl_2(5)$	Isoquinoline (10)	$87 (82)^c (72)^d$
12	$CoCl_2(5)$	Isoquinoline (5)	75
13	CoCl ₂ ·2LiCl (5)	Isoquinoline (10)	69
14	CoCl ₂ ·2LiCl (5)	None	65

^a 1.1 equiv. of benzylzinc chloride (1a) was used. ^b Determined by GC-analysis with tetradecane as an internal standard. c Isolated yield of pure product. d CoCl2 with a purity of 99.999% was used.

absence of coordinating ethereal solvents.9

^a Department Chemie und Biochemie, Ludwig-Maximilians-Universität München, Butenandtstr. 5-13, D-81377 München, Germany. E-mail: knoch@cup.uni-muenchen.de; Fax: +49-89-2180-77680;

Tel: +49-89-2180-77681 ^b Laboratoire de Chimie Moléculaire, CNRS, Ecole Polytechnique, Université Paris Saclay, 91128 Palaiseau, France. E-mail: corinnne.gosmini@polytechnique.edu; Fax: +33-1-6933-4440: Tel: +33-1-6933-4412

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Communication ChemComm

(Table 1, entry 12). Also, we found that the use of CoCl₂·2LiCl¹⁹ was not advantageous (Table 1, entries 13 and 14). Additionally, we have examined the influence of the commercial origin of CoCl₂ as well as its purity. Thus, CoCl₂ having a purity of 99.999% provides under the same conditions (50 °C, 2 h) the diarylmethane 4a in 72% yield (compared to 82%; see Table 1, entry 11).^{20,21} The addition of MTBE as a cosolvent usually decreases the amount of homocoupling and therefore enhances the product yield. However, large amounts of MTBE reduce the reaction rate. We found the solvent mixture THF: MTBE 2:1 to be optimal.²² Concerning the need of isoquinoline as ligand, an extensive screening showed that N-heterocycles behave best, whereas various phosphines did not promote the cross-coupling.²³

With these optimized conditions in hand, we studied the reaction scope of the cross-coupling between various benzylic zinc chlorides (1a-i) with a broad range of aryl and heteroaryl bromides or chlorides. First, the treatment of benzylic zinc reagents (1a,b) in the presence of 5 mol% CoCl₂ and 10 mol% isoquinoline with 4-bromobenzonitrile (3a) at 50 °C within 2 to 4 h is leading to the diarylmethane derivatives 4a,b in 77-82% (Table 2, entries 1 and 2). Furthermore, the cross-coupling of an ortho-substituted benzylzinc chloride (1c) with 3a afforded the desired arene (4c) in 74% yield (Table 2, entry 3). Similarly, the two functionalized benzylic zinc reagents (1d,e) cross-coupled with 3a giving the products 4d,e in 70-79% (Table 2, entries 4 and 5). The ester-substituted benzylzinc chloride (1f) underwent a smooth cross-coupling with 3a leading to the functionalized diaryl-methane 4f in 62% yield (Table 2, entry 6). Additionally, the cross-couplings of the more electron-donating benzylic zinc reagents (1g,h) with 4-bromo-benzonitrile (3a) furnished the arenes 4g,h in 65-82% yield (Table 2, entries 7 and 8).

The reaction scope of this cross-coupling proved to be quite broad. Thus, 2-bromo-benzophenone (3b) underwent the cobaltcatalyzed cross-coupling with the benzylzinc chloride (1b) yielding to the corresponding ketone 5a in 64% yield (Table 3, entry 1). Similarly, the coupling of ethyl 4-bromo-benzoate (3c) with the two benzylic zinc reagents (1e,g) led to the functionalized diarylmethane derivatives (5b,c) in 54-70% yield (Table 3, entries 2 and 3). Remarkably, 2-chloropyridines react well with various benzylic zinc reagents (Table 3, entries 4-9). The cross-couplings of the benzylzinc chlorides (1b,e) with ethyl 2-chloronicotinate (3d) proceeded smoothly under these conditions affording the 2,3disubstituted pyridines (5d,e) in 60-95% yield (Table 3, entries 4 and 5). Also, 3-(ethoxycarbonyl)benzyl-zinc chloride (1f) underwent the coupling with the 2,3-di-substituted pyridine (3d) giving the functionalized aryl-hetero-arylmethane 5f in 68% yield (Table 3, entry 6). Furthermore, the cross-couplings of the benzylic zinc reagents (1d,g,i) with 2-chloro-nicotinonitrile (3e) led to the desired benzylated pyridines (5g-i) in 67-77% yield (Table 3, entries 7-9). Finally, the reaction of 3-fluorobenzylzinc chloride (1d) with ethyl 5-bromofuran-2-carboxylate (3f) afforded within 3 h the 2,5-disubstituted furan (5j) in 60% yield (Table 3, entry 10). The use of aryl bromides bearing electron-donating substituents led to low yields.24

Moreover, such benzylic zinc reagents undergo high yield cross-couplings with various chloro- or bromo-N-heterocycles.

Table 2 Isoquinoline-promoted Co-catalyzed cross-coupling of benzylic zinc reagents (1a-h) with 4-bromobenzonitrile (3a)

FG = 4-^tBu, 2-Cl, 3-F, 3-CF₃, 3-CO₂Et, 4-OMe, 4-SMe Entry Benzylic zinc reagent^a Electrophile Product, yield^{b,c} 4a: 82%, 2 h 1a За ZnCl 3a 1b 4b: 77%, 4 h ZnCl 3a **1c** 4c: 74%, 18 h ZnC 3a 4d: 79%, 1 h **1**d ZnCl 3a 4e: 70%, 2 h 1e ZnC 3a ĊO₂Et 1f 4f: 62%, 18 h 7 3a 1g 4g: 82%, 2 h ZnCl 3a

 a 1.3–1.5 equiv. of benzylic zinc reagent were used. b Isolated yield of pure product. c Less than 15% of homo-coupling of the zinc reagent was observed.

Thus, the reaction of 4-methoxybenzylzinc chloride (1g) with 2-bromopyrimidine (3g) and the two substituted pyridines, 2-chloro-5-(trifluoromethyl)pyridine (3h) and 2-chloro-6-fluoro-pyridine (3i),

4h: 65%, 18 h

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Table 3 Co-Catalyzed cross-coupling reactions of benzylic zinc reagents with aryl and heteroaryl halides

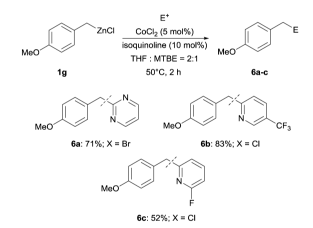
	FG = 4- ^t Bu, 3-F, 3-CF ₃ , 3-CO ₂ Et, 4-OMe, 4-Br				
Entry	Benzylic zinc reagent ^a	Electrophile	Product, yield ^{b,c}		
1	zno zno	Br O Ph	O Ph 'Bu 5a: 64%, 4 h		
2	ZnCl CF ₃ 1e	Br CO ₂ Er	CO ₂ Et 5b : 54%, 18 h		
3	MeO 1g	3c	MeO CO₂Et 5c: 70%, 1 h		
4	'Bu ZnC	CO ₂ Et	CO ₂ Et N 5d: 95%, 4 h		
5	ZnCl CF ₃	3đ	CO ₂ Et N CF ₃ 5e : 60%, 2 h		
6	ZnCl CO ₂ Et	3d	CO ₂ Et N CO ₂ Et 5f: 68%, 18 h		
7	ZnCl F 1d	CN CI N 3e	CN N F 5g: 67%, 3 h		

Table 3 (continued)

FG = 4-^tBu, 3-F, 3-CF₃, 3-CO₂Et, 4-OMe, 4-Br

Entry	Benzylic zinc reagent ^a	Electrophile	Product, yield b,c
9	ZnCl 1i	3e	5i: 68%, 18 h
10	ZnCl F 1d	Br O CO ₂ Et	O_CO ₂ Et F 5j : 60%, 3 h

^a 1.3-1.5 equiv. of benzylic zinc reagent were used. ^b Isolated yield of pure product. ^c Less than 15% of homo-coupling of the zinc reagent was observed.



Scheme 1 Isoquinoline-promoted cross-coupling of the benzylic zinc reagent 1g with selected N-heterocycles (3g-i).

led rapidly (within 2 h) to the functionalized aryl-heteroarylmethanes (6a-c) in 52-83% yield (Scheme 1).

In summary, we have reported a new practical Co-catalyzed, isoquinoline-promoted cross-coupling of various benzylic zinc chlorides with a range of aryl and heteroaryl bromides and chlorides, producing polyfunctionalized diaryl- or arylheteroaryl-methane derivatives. This method tolerates a variety of functional groups, such as esters, nitriles or ketones, and proceeds smoothly at 50 °C within 1-18 h. Remarkably, the combination of MTBE (MTBE = methyl tert-butyl ether) as co-solvent and isoquinoline as additive led only to small amounts of homo-coupling. In most cases, shorter reaction times and improved yields could be obtained. Further investigations towards the synthesis and applications of benzylic organo-metallics are underway in our laboratories.

5h: 77%, 2 h

1g

Communication ChemComm

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- 22 For a corresponding solvent screening, see: ESI,† Table S1.
- 23 For an extensive ligand screening, see: ESI,† Table S2.
- 24 Mechanistic studies are underway to explain these phenomena.