

## Communication

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J. Am. Chem. Soc., 2008, 130 (4), 1128-1129 • DOI: 10.1021/ja077862I

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Published on Web 01/09/2008

#### Copper-Catalyzed Arylation and Alkenylation of Polyfluoroarene C-H Bonds

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Many directing group containing arenes can now be arylated or alkylated under palladium, rhodium, or ruthenium catalysis.<sup>1</sup> Even arenes lacking traditional directing groups can be functionalized representing the most atom-efficient method for creation of arylaryl bonds.<sup>2</sup> However, in this case, regioselectivity issues often are unsolved. An exception can be found in recent elegant work by Fagnou who showed that polyfluorobenzene C-H bonds can be arylated under palladium catalysis.3 The regioselectivity is imparted by the acidification of *ortho*-C-H bonds by fluorine substituents. While palladium catalysts can be replaced by copper in many coupling processes,<sup>4</sup> use of this cheaper and more convenient alternative in C-H activation reactions is not common.5 Fluorinated polyaryls are important in medicinal chemistry, and direct methods for synthesis of these compounds may allow the development of more efficient pathways to pharmaceuticals.<sup>6</sup> We report here a general method for the copper-catalyzed arylation and alkenylation of polyfluorobenzene C-H bonds.

We have recently shown that palladium catalyst can be replaced by copper in the arylation of electron-rich and electron-poor heterocycles.<sup>7</sup> The reaction mechanism presumably includes a deprotonation of an acidic heterocycle followed by lithium/copper transmetalation and reaction of the organocopper species with aryl iodide. It is reasonable to assume that relatively acidid robenzenes<sup>8</sup> can be arylated if a proper combination of ba and copper catalyst is used. A brief optimization of conditions was carried out. For pentafluorobenzene an aryl bromides, the best results were obtained in mixed D solvent (1:1) and by employing phenanthroline liga conversions were observed in DMF. Pentafluorobenze arylated by both aryl bromides and aryl iodides. For reactive substrates, the reaction with aryl iodide in DM results in higher yields. In most cases, potassium phos affords the best results. Less acidic fluorobenzenes the fewer than three fluorines in the molecule can be a employing lithium tert-butoxide base. The reaction respect to aryl halide is presented in Table 1. Both el (entries 1-5) and electron-poor (entries 6-9) aryl reactive. Functional groups, including ester (entry 8) (entry 9) are tolerated. Pyridyl (entries 10 and 11) and rich thienyl (entry 12) bromides can be employed for the If a 5.7/1 *E*/*Z* mixture of  $\beta$ -bromostyrene was used in all of pentafluorobenzene, a 5.7/1 E/Z ratio of pentafluorost obtained (analysis of crude reaction mixture; 77% E isolated, entry 13). Substantial steric hindrance is tolera aryl halide (entries 2 and 3). Mesitylation requires the iodide to obtain high yield. Mesityl bromide afforded isolated yield of the coupling product. Pentafluorol benzylated in fair yield (entry 14). A side reaction betw bromide and DMF solvent is responsible for the reduced yield. On a 10 mmol scale, yield is almost the same as on 1 mmol scale (88% vs 91%, entry 1).

c polyfluo-					
use, solvent,	7	$4\text{-}CF_3C_6H_4Br$	$C_6F_5 \longrightarrow CF_3$		
rylation by	8	$4\text{-}EtO_2CC_6H_4Br$			
MF/xylene nd. Lower	9	4-NCC <sub>6</sub> H <sub>4</sub> Br			
ene can be other, less	10	2-bromopyridine			
MF solvent			N		
sphate base	11	3-bromopyridine			
arylated by	12	2-bromothiophene			
scope with ectron-rich	13 <sup>d</sup>	β-bromostyrene	C <sub>6</sub> F <sub>5</sub> Ph		
and cyano d electron- e arylation.	14	benzyl bromide	C <sub>6</sub> F₅ Ph		
lkenylation tilbene was and 12% Z ated on the	<sup><i>a</i></sup> Substrisolated yie Z Mixture isomer.	rate (1.5 equiv), aryl hal elds. <sup>b</sup> Ten millimole sca (5.7/1) of bromoalkene	tide (1 equiv), base (2 equiv), base (2 equiv), base (2 equiv), $e$ Mesityl be used. $e$ Yield of $E$ iso		
use of aryl only 20%	The scope with respect to fluoroarene is preser Two of the three tetrafluorobenzenes can be arylate				
benzene is veen benzyl	(entries 1 and 2). The arylation of 1,2,3,4-tetrafluoro only a low yield of the cross-coupling product (				

	r 10 n phen	nol% Cul	
	F	DMF/xylene F R	
entry	F F halide	F F product	yield, %
1	4-MeC <sub>6</sub> H <sub>4</sub> Br	C <sub>6</sub> F <sub>5</sub> —	91 88 <sup>b</sup>
2	2-MeC <sub>6</sub> H <sub>4</sub> Br	C <sub>6</sub> F <sub>5</sub>	87
		Mé Me	
3	2,4,6-Me <sub>3</sub> C <sub>6</sub> H <sub>2</sub> I	C <sub>6</sub> F <sub>5</sub> ———————————————————————————————————	87 20 <sup>c</sup>
4	4-MeOC <sub>6</sub> H <sub>4</sub> Br		88
5	1- bromonaphthalene	C <sub>6</sub> F <sub>5</sub>	68
	oronionaphilatere		
6	4-FC <sub>6</sub> H <sub>4</sub> Br		92
7	$4\text{-}\mathrm{CF}_3\mathrm{C}_6\mathrm{H}_4\mathrm{Br}$	C <sub>6</sub> F <sub>5</sub> -CF <sub>3</sub>	88
8	4-EtO <sub>2</sub> CC <sub>6</sub> H <sub>4</sub> Br	C <sub>6</sub> F <sub>5</sub> -CO <sub>2</sub> Et	90
9	4-NCC <sub>6</sub> H <sub>4</sub> Br	C <sub>6</sub> F <sub>5</sub> -CN	95
10	2-bromopyridine		90
11	3-bromopyridine		86
12	2-bromothiophene		92
13 <sup>d</sup>	β-bromostyrene	C <sub>6</sub> F <sub>5</sub> Ph	77% <sup>e</sup> 12% <sup>f</sup>
14	benzyl bromide	C <sub>6</sub> F <sub>5</sub> Ph	31%

Table 1. Arylation Scope with Respect to Halides<sup>a</sup>

equiv). Yields are promide used. d E/ omer. f Yield of Z

nted in Table 2. ed in good yields obenzene affords (entry 3). 1,3,5-Trifluoro- and 1,3-difluorobenzenes are reactive; however, for the arylation of less acidic difluorobenzene, a stronger base, LiOtBu, is necessary. Fluorinated pyridines can also be arylated (entries 6



<sup>a</sup> Substrate (2-3 equiv), aryl halide (1 equiv), base (2-2.5 equiv). Yields are isolated yields. <sup>b</sup> Diarylation product (15%) also isolated. <sup>c</sup> Diarylation product (17%) also isolated. <sup>d</sup> Lithium t-butoxide base.

and 7). Again, for the less fluorinated 3-fluoropyridine, use of LiOtBu base is necessary. Fluorobenzene was unreactive under any conditions tried. As observed by Fagnou,<sup>3</sup> the most acidic C-H bonds, those flanked by two C-F bonds, are efficiently arylated.

It is known that pentafluorophenylcopper reacts with electrophiles, including aryl iodides, producing coupling products in high yield.9 It has also been reported that pentafluorobenzene reacts with aryl iodides in the presence of stoichiometric copper(I) oxide producing pentafluorobiaryls in moderate yields.<sup>10</sup> In light of this, it is reasonable to assume that the catalytic reaction also proceeds by base-promoted formation of fluoroarylcopper followed by the reaction of the copper species with aryl halide, affording the coupling product (Scheme 1).<sup>11</sup> If less acidic fluoroaryl is used, a stronger base is required to form the arylcopper species.

In conclusion, we have developed an efficient, copper-catalyzed method for the arylation, alkenylation, and benzylation of polyfluoroarenes. Arenes containing two or more fluorine substituents on the aromatic ring can be efficiently functionalized. The best Scheme 1. Mechanistic Considerations



results are obtained by using a combination of copper iodide catalyst, phenanthroline ligand, aryl bromide or aryl iodide coupling partner, and DMF or DMF/xylene mixed solvent.

Acknowledgment. We thank the Welch Foundation (Grant No. E-1571) and National Institute of General Medical Sciences (Grant No. R01GM077635) for supporting this research.

Supporting Information Available: Detailed experimental procedures and characterization data for new compounds. This material is available free of charge via the Internet at http://pubs.acs.org.

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- (11) Arylation of the mixture of pentafluorobenzene and 1,2,4,5-tetrafluorobenzene under usual reaction conditions resulted in preferential arylation of pentafluorobenzene (2/1). Additionally, it was shown that base-promoted H/D exchange in polyfluoroarenes occurs with the same efficiency both in the presence or absence of CuI. See Supporting Information for details.

JA077862L