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## Synthesis of *gem*-Difluoroalkenes by Copper-catalyzed Regioselective Hydrodefluorination of 1-Trifluoromethylalkenes

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A copper-catalyzed regioselective hydrodefluorination of 1-trifluoromethylalkenes with hydrosilanes has been developed. The copper catalysis is compatible with several functional groups, including alkyl chloride, ether, ester, nitrile, and imide moieties, to form the corresponding *gem*-difluoroalkenes in good yields. Additionally, the asymmetric induction is also possible by using the chiral DTBM-SEGPHOS ligand, and the *gem*-difluoroalkene with the point chirality at the allylic position is obtained with high enantioselectivity.

## **Keywords**: Copper, *gem*-Difluoroalkenes, Fluorine

Because of uniquely high electron-withdrawing nature, the gem-difluoroalkenes are isosteres of carbonyl group<sup>1</sup> and frequently observed in pharmaceuticals agrochemicals.<sup>2</sup> Additionally, they are good building blocks for the functionalized mono-fluoroalkenes, which can work as the biologically important amide isostere,3 via metal-catalyzed site-selective defluorofunctionalizations.<sup>4</sup> The gemdifluoroalkenes are generally prepared by the Wittig-type reaction of carbonyl compounds with difluoromethylenation reagents.<sup>5</sup> Moreover, some research groups have recently developed more convergent coupling-type processes with some CF<sub>2</sub> carbene precursors or CF<sub>3</sub>CH carbene equivalents.<sup>6</sup> However, despite the aforementioned certain advances, there still remains a need for efficient chemical synthesis of gemdifluoroalkenes owing to their ubiquity in fields of organic chemistry and medicinal chemistry.

Meanwhile, we recently developed the Cu/p-t-Bu-dppbzregioselective net-hydroamination<sup>7,8</sup> trifluoromethylalkene 1a with polymethylhydrosiloxane (PMHS) and the hydroxylamine derivative (Scheme 1a).9 Following our mechanistic considerations, 1a undergoes the insertion into in-situ generated copper hydride species A<sup>10</sup> to form the  $\alpha$ -CF<sub>3</sub> alkylcopper intermediate **B**. Subsequent electrophilic amination with the hydroxylamine delivers the corresponding hydroaminated product. In spite of the possibility of  $\beta$ -F elimination from  $\mathbf{B}$ ,<sup>4,11</sup> the *p-t*-Bu-dppbz ligand selectively promoted the net-hydroamination reaction During the optimization studies on this (Scheme 1b). chemistry, we serendipitously found the significant ligand effect of xantphos: even in the presence of hydroxylamine, the hydrodefluorinated product, namely gem-difluoroalkene 2a, was formed exclusively (Scheme 1c). The unique propensity with xantphos toward the β-F elimination prompted us to further investigate this reaction. Herein, we wish to report a Cu/xantphos-catalyzed hydrodefluorination of trifluoromethylalkenes with the hydrosilane to afford the gemdifluoroalkenes. The copper catalysis is tolerated with several functional groups, and the corresponding gem-difluoroalkenes are obtained in good yields with high regioselectivity. Additionally, by the judicious choice of supporting ligand, the asymmetric induction is also possible. The related boryl-, <sup>12</sup> silyl, <sup>13</sup> alkyl-, <sup>14</sup> and aryldefluorination <sup>15</sup> reactions of 1-trifluoromethylalkenes were reported by several groups, but the reductive defluorination process still remains underdeveloped. <sup>16</sup>

a) previous hydroamination with p-t-Bu-dppbz

b) mechanistic considerations

c) serendipity of hydrodefluorination with xantphos

$$t$$
-Bu  $t$ -Bu

Inspired by our initial finding (Scheme 1c), our optimization studies commenced with 1-trifluoromethylalkene 1a, PMHS, and CuCl/xantphos catalyst in 1,4-dioxane solvent, and some bases were initially screened (Table 1). As shown in Scheme 1c, CsOAc was a good base, giving the *gem*-difluoroalkene 2a in 75% <sup>1</sup>H NMR yield (entry 1). Consistent with our previous observation and recent mechanistic insight that smaller alkali metal cations can accelerate the β-F elimination process, lithium and sodium alkoxide bases also worked well, and the targeted 2a was obtained in yields comparable to that with CsOAc (entries 2–5). In particular,

NaO-t-Bu and NaOTMS showed better performance to deliver 2a in 87% and 92% isolated yields, respectively (entries 4 and 5). Other weaker non-alkoxide bases such as KOPiv resulted in no formation of 2a (data not shown), suggesting that relatively strong bases are necessary in the reaction. The effect of xantphos was critical: some other common phosphine ligands also promoted the reaction but with much lower efficiency (entries 6–11). We next examined other hydrosilanes. Among them, Ph<sub>2</sub>SiH<sub>2</sub> also gave an almost quantitative yield (entry 13), but from the viewpoints of stability and cost, we identified PMHS to be the best hydride source.

 $\textbf{Table 1.} \ \, \textbf{Optimization studies for regioselective hydrodefluorination of 1-trifluoromethylalkene } \textbf{1a.}^a$ 

Entry	H– <b>Si</b>	Ligand	Base	Yield <sup>b</sup> /%
1	PMHS	xantphos	CsOAc	75
2	PMHS	xantphos	LiO-t-Bu	72
3	PMHS	xantphos	LiOTMS	78
4	PMHS	xantphos	NaO-t-Bu	95 (87)
5	PMHS	xantphos	NaOTMS	>99 (92)
6	PMHS	dppbz	NaOTMS	13
7	PMHS	dppe	NaOTMS	21
8	PMHS	dppf	NaOTMS	45
9	PMHS	DPEPhos	NaOTMS	57
10	PMHS	rac-BINAP	NaOTMS	59
$11^c$	PMHS	PPh <sub>3</sub>	NaOTMS	43
12	(EtO) <sub>2</sub> MeSiH	xantphos	NaOTMS	38
13 <sup>d</sup>	$Ph_2SiH_2$	xantphos	NaOTMS	99

<sup>a</sup>Conditions: CuCl (0.025 mmol), ligand (0.025 mmol), **1a** (0.25 mmol), H–*Si* (0.75 mmol based on H–Si), base (0.50 mmol), 1,4-dioxane (1.5 mL), rt, 4 h, N<sub>2</sub>. <sup>b</sup>Estimated by <sup>1</sup>H NMR with CH<sub>2</sub>Br<sub>2</sub> as the internal standard. Isolated yields are in parentheses but somewhat lower than the <sup>1</sup>H NMR yield because of the volatility of **2a**. <sup>c</sup>With 0.050 mmol of PPh<sub>3</sub>. <sup>d</sup>With 0.50 mmol of Ph<sub>2</sub>SiH<sub>2</sub>.

With the optimal conditions in hand, we then investigated the scope of 1-trifluoromethylalkenes 1. The products 2 obtained are illustrated in Scheme 2, in which the best basic additive (NaO-t-Bu or NaOTMS) was dependent on the substrate structure and functional group. The simple aliphatic 1b was expectedly converted to the gemdifluoroalkene 2b in a good yield. The copper catalysis was compatible with alkyl chloride (2c), benzyl ether (2d), and silvl ether (2e) moieties. The carbonyl functions such as ester (2f) and nitrile (2g) were also tolerated under the standard conditions. Moreover, the substrates with protected nitrogen groups underwent the hydrodefluorination smoothly to form the corresponding functionalized gem-difluoroalkenes 2h and 2i in acceptable yields. Particularly notable is the successful conversion of trisubstituted alkene (2j) with the aid of TMSdppbz ligand instead of xantphos, thus giving a chance of asymmetric induction by an appropriate chiral ligand (vide infra).<sup>17</sup> On the other hand, the Ph-conjugated 1k resulted in a low yield of the corresponding gem-difluoroalkene 2k and competitive formation of the reduced byproduct 2k'. This is

probably because the regioselectivity in the insertion into the copper hydride (Scheme 1b) is not controlled. The Ph-vinyl conjugation increases the coefficient of LUMO at the carbon  $\alpha$  to CF3, thus competitively forming the regioisomeric  $\beta\text{-CF}_3$  alkylcopper intermediate. A similar trend was observed also in our previous work.  $^9$ 

**Scheme 2.** *gem*-Difluoroalkene products **2** by Cu-catalyzed hydrodefluorination of 1-trifluoromethylalkenes **1**. Conditions: CuCl (0.025 mmol), xantphos (0.025 mmol), **1** (0.25 mmol), PMHS (0.75

mmol based on H–Si), base (0.50 mmol), 1,4-dioxane (1.5 mL), rt, 4 h, N<sub>2</sub>. Isolated yields are shown. The base used (NaO-*t*-Bu or NaOTMS) is given in parentheses. <sup>a</sup>With PMHS (0.38 mmol based on H–Si). <sup>b</sup>At 50 °C. <sup>c</sup>With TMS-dppbz ligand instead of xantphos. <sup>d</sup>

TMS-dppbz

From a 9:1 *E/Z* mixture of **1k**. <sup>1</sup>H NMR yield. The reduced byproduct **2k'** was also formed in 3% <sup>1</sup>H NMR yield.

Ph Zk'

As a mechanistic probe, we subsequently performed the reaction of 1a with the Ph2SiD2 instead of PMHS under otherwise identical conditions (eq. 1). Expectedly, the deuterium atom was selectively incorporated at the allylic position, and the corresponding  $2a-d_1$  was obtained in a good yield with 96% D content. This result not only supports our initial hypothesis (Scheme 1b) but also provides access to deuterium-labeled gem-difluoroalkenes, which can find applications in biological labeling studies and drug discovery.18 Finally, we attempted to apply chiral bisphosphine ligands for the asymmetric synthesis of 2j (eq. 2). Gratifyingly, the (R)-DTBM-SEGPHOS ligand was found to be promising to produce the enantioenriched (S)-2i in 84% yield with >99:1 enantiomeric ratio (er). 19 The product could also be transformed to the borylalkene 3<sup>20</sup> under the reported conditions, 4c which should be a useful chiral building block for various chiral fluoroalkenes.

In conclusion, we have developed a copper/xantphoscatalyzed hydrodefluorination of 1-trifluoromethylalkenes with hydrosilanes to form the corresponding *gem*-difluoroalkenes in good yields with high regioselectivity. The copper catalysis can also provide an avenue to deuteriumlabeled as well as the enantioenriched *gem*-difluoroalkenes, which are of potent interest in medicinal and pharmaceutical chemistry. Ongoing work seeks to elucidate the origin of uniquely high  $\beta$ -F elimination performance with xantphos and develop related copper-catalyzed reductive transformations of 1-trifluoromethylalkenes.

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Supporting Information is available on http://dx.doi.org/xxxx.

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- 19 See the Supporting Information for optimization studies on asymmetric catalysis.
- 20 The absolute configuration of (S)-2j and 3 was determined by the specific rotation after the derivatization into the known product. See the Supporting Information for details.

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Me F F CuCI/(R)-DTBM-SEGPHOS Ph F PhHS  NaOTMS, 1,4-dioxane, rt 84%, >99:1 er					