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## Catalytic metal-free Si-N cross-dehydrocoupling<sup>†</sup>

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The metal-free  $B(C_6F_5)_3$  catalyzed dehydrocoupling of hydrosilanes with anilines, carbazoles and indoles is reported. For anilines and carbazoles the reaction proceeds by the liberation of H<sub>2</sub> as the sole Si–N coupling byproduct. Indoles react with diphenyl(methyl) hydrosilane to give *N*-silyl indolines with high diastereoselectivity (d.r. 10:1) in excellent yields. A mechanism for this Si–N coupling/ hydrogenation sequence is proposed.

The cross-dehydrocoupling is an efficient methodology for the connection of two molecular entities.<sup>1</sup> Especially the dehydrocoupling of Si–H and N–H fragments provides an environmentally benign access to silyl-protected amines.<sup>2</sup> These ubiquitous structural motifs are usually obtained by the reaction of halosilanes with deprotonated amines, the generation of which often requires strong bases.<sup>3</sup> This is not only of concern for atom efficiency but also for functional group tolerance. In light of this, the Si–N dehydrocoupling proved very useful, *e.g.* for the protection of indoles using  $Zn(OTf)_2$  (10 mol%) in the presence of 0.5–1.0 equiv. of pyridine.<sup>4</sup> Oestreich's sulfur-bridged Ru–arene complex<sup>5</sup> is particularly effective in the base-free dehydrocoupling of silanes with other nitrogen-containing heterocycles, *e.g.* indole, carbazole and pyrrole derivatives using only 1 mol% of catalyst loading.<sup>6</sup> However, a metal-free variant has not yet been disclosed.<sup>7</sup>

We have shown earlier that the H<sub>2</sub>-activation product 1 of the frustrated Lewis pair (FLP) consisting of 2/3 is a transient species which readily releases H<sub>2</sub> at room temperature (Scheme 1, top).<sup>8</sup> Accordingly, the isostructural intermediate iso-1, generated through the silyl-transfer from the silane 4 to the aniline 5a, should readily liberate H<sub>2</sub> with concomitant release of the Si–N coupling product 6 (Scheme 1, bottom). As a potential silyl-transfer catalyst, borane 2 has attracted significant attention in hydrosilylation of aldehydes, ketones, imines and olefins.<sup>9</sup> An analogous mechanism was only recently proposed by Oestreich as a competing pathway in the borane-promoted imine reduction with hydrosilanes.<sup>9a</sup>



Indeed, when bis(4-toloyl)amine (5a) was reacted with diphenyl-(methyl) silane (4a) in the presence of 5 mol%  $B(C_6F_5)_3$  (2) at room temperature, the silylamine 6a was obtained in 95% yield accompanied with the evolution of H<sub>2</sub> (Table 1, entry 1). In the absence of the catalyst, the formation of 6a was not observed even when a mixture of 5a and 4a was heated to 90 °C for 12 h (Table 1, entry 2). The catalyst loading was reduced to 1 mol% with slight erosion in yield (73%, entry 3). Lower catalyst loadings of 0.1 mol% led to significantly reduced yields (entry 4). Further experiments were carried out with 1 mol% of 2 as catalyst.

The reaction displays a remarkable substrate scope. Besides diphenylamine derivatives (5a and 5b, entries 3 and 5), carbazole derivatives 5c-f also proved to be viable substrates and the products 6c-f were obtained in 83-97% yields (entries 6-9). The bibromo derivative required 70 °C to undergo Si-N cross-dehydrocoupling in 51% yield without the observation of dehalogenation (entry 8). The reduced yield was attributed to the very low solubility of 5e in toluene. Other silanes were also useful in the Si-N coupling reaction. Triethylsilane (4b) or 1,1,3,3-tetramethyldisiloxane (4c) readily reacted with carbazole (5c) or bis(4-tolyl)amine (5a) in high yields (entries 9 and 10). The silvlation of primary aniline derivatives proceeded at 60–70  $^{\circ}$ C in excellent yields (88-97%, entries 11-15).10 The electron-deficient anilines 5m and 5n were reactive even at room temperature and 6m and 6n were obtained in 88% and 97% yields (entries 16 and 17). Also the two diamines  $N_{N'}$ -(diphenyl)-1,4-phenylene diamine (50) and N,N'-(diphenyl)-ethylene diamine (5p) underwent silvlation with diphenylmethyl silane (4a) in high yields (entries 18 and 19).

Institute for Organic Chemistry, Karlsruhe Institute of Technology (KIT), Fritz-Haber-Weg 6, D-76131 Karlsruhe, Germany. E-mail: jan.paradies@kit.edu † Electronic supplementary information (ESI) available: Experimental procedures, analytical data. See DOI: 10.1039/c3cc49558b ± Both authors contributed evenly.

Table 1 Si–N cross-dehydrocoupling of aromatic amines with hydrosilanes<sup>a</sup>

1.0 equiv. R .	N <sup>,R'</sup> + <sup>·</sup> ∦5	1.0 equiv	cat. B(C <sub>6</sub> F <sub>5</sub> ) <sub>3</sub> ( <b>2</b> ) (1 mol%) CH <sub>2</sub> Cl <sub>2</sub> , temp.	R`N.R' I +H2 6 <i>SiR</i> 3		
	Entry	<i>t</i> [h]	$T [^{\circ}C]$	Product		Yield [%]
Diarylamines	1	1	25			95 <sup>b</sup>
	2	12	90	U,U	6a	$0^{c}$
	3	1	25	SiMePh <sub>2</sub>		73
	4	10	25	$\sim$		32"
	5	1	25	N SiMePh <sub>2</sub>	6b	91
	6	1	25	N SiMePh <sub>2</sub>	6c	97
	7	1	25	N SiMePh <sub>2</sub>	6d	83
	8	24	25	Br	6e	95 <sup>e</sup>
	9	1	25	N SiEt <sub>3</sub>	6f	95 <sup>e</sup>
	10	1	25	Me₂ Me₂ Tol <sub>N</sub> ,Si O Si N Tol Tol Tol	6g	97
Anilines	11	72	70	NH SiMePh <sub>2</sub>	6h	90 <sup>b</sup>
	12	48	70	NH SiMePh <sub>2</sub>	6i	90
	13	48	70	NH SiMePh <sub>2</sub>	6j	93
	14	36	60	CI-NH SiMePh <sub>2</sub>	6k	97
	15	24	60		61	91
	16	36	25		6m	88
	17	24	25	F <sub>3</sub> C NH SiMePh <sub>2</sub>	6n	97
Diamines	18	24	25	Ph N- Ph <sub>2</sub> MeSi <sup>'</sup> Ph	<sup>lePh</sup> 2 <b>60</b>	26 <sup><i>f</i></sup>
	19	24	70	$\stackrel{SiMePh_2}{\overset{Ph}{\underset{i}{\overset{N}{\overset{V}{\overset{V}}}}}}_{SiMePh_2}}$	6р	92 <sup>b</sup>
	20	24	60	Ph、H Ph~N <sup>Si</sup> N~Ph	6q	83
Indoles	21	144	70	SiMePh <sub>2</sub>	6r	50 <sup>e</sup>
	22	24	70	CI	6s	81
	23	24	70	N SiMePh <sub>2</sub>	6t	96
	24	24	70	Ph N SiMePh <sub>2</sub>	бu	97

1.0 (	equiv. $\stackrel{R}{}_{N} \stackrel{R'}{}_{H} + \frac{1}{5}$	1.0 equiv	: Н- <i>SIR<sub>3</sub></i> 4	cat. B(C <sub>6</sub> F <sub>5</sub> ) <sub>3</sub> ( <b>2</b> ) (1 mol%) CH <sub>2</sub> Cl <sub>2</sub> , temp.	R`N 6 <i>Si</i>	.R' + H <sub>2</sub> <b>R<sub>3</sub></b>
	Entry	<i>t</i> [h]	$T [^{\circ}C]$	Product		Yield [%]
	25	24	70	Me N N SiMePh <sub>2</sub>	бv	92

<sup>*a*</sup> Reactions were performed on a 1.0 mmol scale, 3 M in  $CH_2Cl_2$ . <sup>*b*</sup> 5 mol% 2. <sup>*c*</sup> Absence of 2. <sup>*d*</sup> 0.1 mol%  $B(C_6F_5)_3$ . <sup>*e*</sup> 10 mol% 2, 0.1 mmol scale, 3 M in  $CD_2Cl_2$ , yield determined by <sup>1</sup>H NMR. <sup>*f*</sup> 2 mol% 2.

Accordingly, the reaction of **5p** with phenylsilane (**4d**) provided the cyclic product **6q** in 83% yield (entry 20).

Finally, we investigated the potential of the Si–N dehydrocoupling for pyrrole and indole derivatives. While pyrrolederivatives were unreactive under our reaction conditions,<sup>11</sup> the indole-derivatives 5r-v displayed high reactivity. The indoles 5r-v were chemospecifically converted into the 1-silylated indoline derivatives 6r-v (entries 21–25) without the formation of unsaturated side products arising from N or C3-silylation.<sup>12</sup> Indole (5r) required prolonged reaction time (144 h, entry 21) for the domino silylation/ reduction sequence and indoline (6r) was obtained in 50% yield. The less electron-rich 6-chloroindole (5s) was transformed into 6s in excellent yield in only 24 h (95%, entry 22). Substituents in position 2 were well tolerated and the 2-methyl and 2-phenyl indolines 5t and 5u were obtained in quantitative yields (96% and 97%, entries 3–5). 2,3-dimethylindole (5v) was diastereoselectively reduced to *cis*-2,3dimethyl indoline (6v) in quantitative yield (98%, d.r. 10:1).<sup>13</sup>

The high chemospecificity and diastereoselectivity prompted us to investigate the Si–N cross coupling/hydrogenation reaction of **5v** with **4a** in detail (Scheme 2). Only resonances of the starting materials and the product **6v** were observed when the reaction was monitored by <sup>1</sup>H NMR (1 mol% 3,  $[D_8]$ -toluene). Neither the resonance of FLPactivated H<sub>2</sub> nor the resonance of dissolved H<sub>2</sub> was observed by <sup>1</sup>H NMR. Deuterium labeling experiments were conducted to investigate the fate of the hydridic and protic hydrogen atoms in silane **4a** 



**Scheme 2** Isotope labelling experiments with (a) 1-D-2,3-dimethylindole (1-(D)-4v), with (b) D-SiMePh<sub>2</sub> (D-4a) and (c) cross experiment.



Scheme 3 Proposed catalytic cycle for the Si–N coupling/hydrogenation domino reaction.

and indole 5v. The reaction of 1-D-2,3-dimethyl indole (1-D-5v, 95% D) with H-SiMePh<sub>2</sub> (4a) gave exclusively cis-3-D-2,3-dimethyl indoline (3-D-6v) in high yields (97%, 92% D-incorporation, Scheme 2a). The reaction of D-SiMePh<sub>2</sub> (D-2a, 95% D) with 5v provided exclusively cis-2-D-2,3-dimethyl indoline (2-D-6v) in 96% yield with 92% D-incorporation at position 2. Together the chemoselective deuteration and the absence of dissolved or FLPactivated H<sub>2</sub> or HD<sup>14</sup> strongly support a N-silvlation/rearrangement/ reduction mechanism (Scheme 3). The product of the  $B(C_6F_5)_3$ catalyzed silyl-transfer to 5v is 1-silyl-1-H-indol-1-ium 6, which rearranges to the more stable 1-silyl-3-H-indol-1-ium 7. Alternatively, an intermolecular proton-transfer might be conceivable. However, according to our cross experiment using 5t-u and 1-silyl-indole 8, the sigmatropic rearrangement mechanism is more likely (Scheme 2c). The indole derivatives 5t and 5u were equally reactive as 5v (96-98%, 24 h, see Table 1, entries 23-25) and should be readily protonated by transiently formed 6 (formed by the reaction of 5 and 4a, compare Scheme 3). However, the reaction of an equimolar mixture of 8, 5t-u, and 4a in the presence of 10 mol% 2 produced 6t or 6u as the product (6u/6v >95:5; 6t:6v >90:10). This is a strong indication that intermolecular proton-transfer is not operative in the silvlation/ hydrogenation reaction sequence. The final step in the catalytic cycle is the hydride transfer from  $[H-B(C_6F_5)_3]$  to the highly electrophilic iminium species 7 from the least hindered side liberating cis-6v and the catalyst 2.

In summary, we have developed the metal-free Si–N crossdehydrocoupling for primary and secondary aryl amines having solely molecular hydrogen as byproduct. Indole derivatives undergo *N*-silylation followed by a rearrangement/reduction sequence to furnish indolines in high yields and high diastereoselectivity (d.r. 10:1).

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