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Identifying a cobalt catalyst for highly selective hydrosilylation of allenes†

Zheng Yang, $\ddagger^{a,b}$ [D](http://orcid.org/0000-0002-2866-2431)ongjie Peng, $\ddagger^{a,b}$ Xiaoyong Du,^{a,b} Zheng Huang*^{a,b} and Shengming Ma **D** *a,b,c

An efficient method of cobalt-catalyzed allene-hydrosilylation is developed. The reaction enjoys an excellent regio- and stereoselectivity and a broad scope affording Z-allylic silanes. Many synthetically useful functional groups can be tolerated. A Co(I)-species involved mechanism is proposed.

Featuring reasonable stability, low-toxicity, and ease of handling, silane reagents are valuable intermediates for synthetic transformations, such as Hiyama cross-couplings¹ and Tamao oxidation reactions to form the carbonyl group.² The hydrosilylation of unsaturated C–C bonds such as alkenes,³ alkynes, 4 and dienes, 5 which is considered to be one of the most synthetically efficient approaches to organosilane compounds, has been achieved with various catalysts. However, reports on the hydrosilylation of allenes are relatively rare, partially due to the difficulty in controlling the regio- and stereoselectivity. Most of the reported allene hydrosilylations occurred at the non-terminal $C=C$ bond, affording vinylsilanes using Pd,⁶ Ni,⁷ Au,⁸ Ru⁹ or Al¹⁰ catalysis (Scheme 1a) or branched allylsilanes using $Pd^{6,7}$ catalysis (Scheme 1b). In 2016, Asako and Takai reported a molybdenum-catalyzed hydrosilylation at the terminal $C=C$ bonds of allenes that yielded linear allylic silanes (Scheme $1c$).¹¹ However, the stereoselectivity of this reaction is rather poor. To the best of our knowledge, the allene hydrosilylation catalyzed by earthabundant base metal iron or cobalt has not been established yet. Here we report a cobalt-catalyzed hydrosilylation of allenes to form linear (Z)-allylsilanes, enjoying an excellent regio- and stereoselectivity. **RESEARCH ARTICLE**

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Non-precious metal iron and cobalt catalysts in general offer lower activity than Pd and other precious metal catalysts, thus identifying that a suitable ligand is the key to achieve an efficient and selective hydrosilylation of allenes. We employed

iron and cobalt complexes of phosphinite-iminopyridine¹² $(P^{O}NN)$ and phosphine-iminopyridine¹³ ($P^{C}NN$) ligands developed in one of our laboratories (Chart 1), which exhibited an excellent catalytic activity in the hydrosilylation of alkenes.

We commenced the study by examining an iron catalyst generated from $(^{tBu}P^ONN^{iPr})FeCl_2$ (1) and NaBHEt₃ for the hydrosilylation of phenyl allene (3a) with phenylsilane. The initial run in toluene using only 0.5 mol% of the catalyst yielded (Z)-phenyl(3-phenylallyl)silane Z-4a in 93% yield as determined by ¹H NMR analysis. Neither the regioisomers 4a' and 4a″ nor the stereoisomer E-4a was detected. Solvent variations have a large effect on the cobalt-mediated hydrosilyl-

Chart 1 Fe and Co complexes with $P^{O}NN$ and $P^{C}NN$ ligands.

^aState Key Laboratory of Organometallic Chemistry, Shanghai Institute of Organic Chemistry, Chinese Academy of Sciences, 345 Lingling Lu, Shanghai 200032, P. R. China. E-mail: masm@sioc.ac.cn, huangzh@sioc.ac.cn

^bUniversity of Chinese Academy of Sciences, Beijing 100049, P. R. China

c Department of Chemistry, Fudan University, 220 Handan Lu, Shanghai 200433, P.R. China

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[‡]These authors contributed equally.

Table 1 Optimization of the reaction conditions^a

Entry	Precatalyst	Solvent	TICIA OI (Z) -4a ^b (%)	RECOVETY of $3a^b$ (%)
$\mathbf 1$	1	Toluene	93 (91)	
$\overline{2}$	1	THF	70	
3	1	Hexane	81	
4	1	MeCN	9	62
5		DCM		76
6	1	DMF	Ω	64
7	2a	Toluene	34	44
8	2 _b	Toluene	94 (92)	
9	2c	Toluene	54	25
10	2d	Toluene	13	39
11		Toluene	Ω	94
12^c	2 _b	Toluene	Ω	82
13	CoCl ₂	Toluene	0	89^d

^a Reaction conditions: 3a (0.5 mmol), PhSiH₃ (0.5 mmol), 2.5 µmol of precatalyst, 5.0 µmol of NaBHEt₃, in solvent (0.5 mL). ^b Determined by ¹H NMR analysis with CH₃NO₂ as the internal standard. Values in parentheses are yields of the isolated products. CNo NaBHEt₃ was added. $\frac{d}{d}$ Isolated recovery of 3a.

ation. The yield dropped when the reaction was conducted in THF, hexane or acetonitrile (Table 1, entries 2–4), while the reaction did not occur in dichloromethane or DMF (Table 1, entries 5 and 6). Next, we screened a series of complexes of Fe and Co ligated by P^C NN ligands (Table 1, entries 7-10) which revealed that $(^{tBu}P^CNN^{iPr})CoCl_2$ (2b) gave the best result (Table 1, entry 8) and the steric effect of the complexes greatly affects the reaction. Lower yields were observed in the runs using the complexes with less bulky substituents on the ligands (Table 1, entries 9 and 10). No conversion was observed when running the reaction in the absence of the metal complex or N aBHE t_3 , or replacing cobalt complexes with $CoCl₂$, indicating the important effect of the ligand on the hydrosilylation (Table 1, entries 11–13). Notably, a minor impurity was observed in the isolated product when using complex 1 as the precatalyst (entry 1), while the reaction with the precatalyst 2b afforded the product in a very high purity (entry 8). Therefore, the parameters used in entry 8 have been chosen as the optimized reaction conditions for further study.

Next, we investigated the scope of the cobalt-catalyzed hydrosilylation with respect to the allene substrates. All the reactions of 3-aryl or alkyl substituted 1,2-dienes employed 0.5 mol% 2**b** as the precatalyst, furnishing linear (Z) -allylsilanes in high isolated yields with an excellent (Z)-selectivity (Schemes 2 and 3). Aryl-substituted allenes bearing either electron-donating (4b and 4g) or electron-withdrawing groups (4c) were hydrosilylated with high regio- and stereoselectivity, and

Scheme 2 Highly regio- and stereoselective hydrosilylation of aryl substituted allenes. The reaction was carried out with 1.0 mmol of 3, 1.0 mmol of PhSiH₃, 5.0 µmol of 2b, 10.0 µmol of NaBHEt₃, in 1 mL toluene at room temperature for 5 h. Yields of the isolated products are given

Scheme 3 Highly regio- and stereoselective hydrosilylation of alkyl substituted allenes. The reaction was carried out with 1.0 mmol of 3, 1.0 mmol of PhSiH₃, 5.0 µmol of 2b, 10.0 µmol of NaBHEt₃, in 1 mL toluene at room temperature for 5 h. Yields of the isolated products are given. a ^a The reaction was carried out with 5 mmol of 3i to afford 1.2 g of Z-4i.

substituents in the *ortho*, meta, and para positions of the phenyl ring are compatible with the reaction conditions (4d–f). The naphthyl substituent is also tolerated, giving a high yield of 4h and no isomeric product was observed (Scheme 2).

Alkyl substituted allenes also reacted with phenylsilane smoothly under the standard reaction conditions. Synthetically useful functional groups such as acetoxy (4j), cyano (4k), benzyl (4l), and ketal (4m) could be tolerated. Notably, even the reactive acetyl (4n) could be accommodated in this hydrosilylation reaction. Furthermore, the allene showed a higher reactivity than the internal alkyne as demonstrated by the isolation of 4o in a high yield with an exclusive chemoselectivity towards the allene unit. No side-products resulting from the hydrosilylation of the acetyl and internal alkyne groups were observed in these reactions. Moreover, these Organic Chemistry Frontiers **Research Article** Chemistry Frontiers **Research Article** Chemistry Frontiers **Research Article**

reactions could be carried out on a one-gram scale, affording 4i in 93% yield.

The reaction of unsymmetric 1,1-disubstituted allene 3p also proceeded smoothly to afford linear allylsilanes Z-4p in a decent yield (Scheme 4). Significantly, only the Z-isomer was formed, demonstrating the capability of the cobalt catalyst to discriminate between Me and a larger substituent in the 1,1-disubstituted allene substrate.

Scheme 4 Highly regio- and stereoselective hydrosilylation of 1,1-disubstituted allenes.

Other silanes such as Ph_2SiH_2 , Et_2SiH_2 , and Et_3SiH were also tried for this reaction but only complicated mixtures were obtained.

We propose a rationale for the cobalt-catalyzed hydrosilylation of allenes on the basis of the precedents of the relevant $(P^{C}NN)$ Co-catalyzed alkene and alkyne hydrosilylations^{13,14} (Scheme 5). The hydrosilylation process starts with the activation of $2b$ by using NaBHEt₃, followed by the reaction with PhSiH₃ to form a cobalt(1) silyl intermediate **Int-1.**¹⁵ Most likely due to the steric repulsion between the P^{C} NN ligand and the substituent groups of the allenes, the terminal $C=C$ double bond of substrate 3 coordinates to the metal center of Int-1 from the less hindered side, resulting in the generation of the allene adduct Int-2. The subsequent insertion of the $C=C$ double bond into the Co–Si bond would afford the vinyl

Scheme 5 Proposed mechanisms

cobalt intermediate Int-3 with the Co center located at the cis position relative to the small substituent due to the steric effect. The vinyl complex Int-3 then reacts with $PhSiH₃$, via either sigma-bond metathesis or a silane oxidative addition/ reductive elimination pathway, to deliver the hydrosilylation product and regenerate Int-1. Alternatively, the catalytic process may involve a cobalt (i) hydride intermediate Int-1'.¹⁶ The corresponding allene adduct **Int-2'** then undergoes insertion to form the allyl cobalt intermediate Int-3′, which further reacts with $PhSiH₃$ to form the desired product and regenerate the cobalt(I) hydride. Further detailed mechanistic studies are ongoing to establish unambiguously the real mechanistic nature of the reaction. Organic Chemistry Frontiers

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In conclusion, we have developed a highly regio- and stereoselective cobalt-catalyzed allene-hydrosilylation method for the synthesis of linear (Z)-allylsilanes. Both mono and 1,1-disubstituted allenes are applicable for this transformation and a variety of synthetically useful functional groups could be tolerated. Further investigations including mechanistic studies and synthetic applications of the allylsilane products have been pursued in this laboratory.

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