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Synthesis of multi-substituted vinylsilanes *via* copper(ı)-catalyzed hydrosilylation reactions of allenes and propiolate derivatives with silylboronates†

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An efficient and general copper(i)-catalyzed method for the synthesis of multi-substituted vinylsilanes is reported. Multi-substituted allenes with electron-withdrawing groups and propiolate derivatives reacted well with (dimethylphenylsilyl)boronic acid pinacol ester to afford silyl-substituted butenoate derivatives and β -silyl-substituted acrylate derivatives, respectively. The corresponding products could be obtained in moderate to high yields and with good to excellent stereoselectivities.

The development of new and efficient methods for the synthesis of versatile vinylsilanes has attracted the interest of many organic chemists.¹ Accordingly, various noble transition-metal catalyzed hydrosilylation of alkynes using silanes has been reported to furnish this class of compounds in good yields.² Recently, silylboronate reagents developed by Suginome's group have been widely used for the silylation reactions to forge new C-Si bonds.³ The research groups of Ito,4 Tanaka,5 Suginome with Ohmura,6 Oestreich, Moberg, Sato, Cheng, Hayashi, Hoveyda, Riant, Riant, and Sawamura¹⁴ et al. have contributed greatly in this field. Among the metals, copper salts have been recognized to be a good choice as catalysts because of their low cost and environmental friendly merits.¹⁵ However, not much attention has been focused on the copper-catalyzed Si-B activation and transformations. The pioneering work on silylcupration reaction of alkynes with Me₂PhSi-BEt₃Li has been reported by Oshima and co-workers. 16 In this work, for most of the cases, a mixture of vinylsilane regioisomers was obtained. It was not until about 25 years later that Loh and co-workers developed a CuCl/Johnphos system catalyzed silylcupration reaction of terminal alkynes using Suginome's reagent ((dimethylphenylsilyl)boronic acid pinacol ester 1) in the presence of MeOH to afford the corresponding branched vinylsilanes in good yields and

Initially, 1 and 2a were chosen as the model substrates to optimize this reaction. In the presence of a strong base KO^tBu, only 32% yield of the desired product was obtained (Table 1, entry 1). To clarify what leads to the relatively low yield of the product, a few control experiments were carried out. We found that when 2a, CuCl (10 mol%) and KO^tBu (11 mol%) in THF were stirred for 24 hours in the absence of 1, the starting material 2a was decomposed. Without the use of KO^tBu, only a trace amount of the desired product was obtained but most of 2a could be recovered (Table 1, entry 2). On this basis, we attempted to use less basic Et₃N as the additive. To our delight, 14% product yield could be obtained with the recovery of starting material 2a (Table 1, entry 3). Diverse solvents were then screened and it was found that the yield of the desired product could be improved to 69% by using a protic solvent such as methanol (Table 1, entry 6). The use of tert-butanol gave the best results in terms of yield (Table 1, entries 7-12 and 16). We observed that other phosphine ligands were efficient but not necessary to afford the desired product 3a in this reaction (Table 1, entries 10-12 and 15). Further control experiments were carried out and it was shown that both the copper catalyst and Et₃N were crucial in this catalytic reaction (Table 1, entries 13 and 14). Ultimately, the optimized reaction conditions were established as follows: 10 mol% CuBr as a catalyst and 11 mol% dppe as a ligand using 11 mol% Et₃N as an additive in tert-butanol, heated at 40 °C.

high selectivities.¹⁷ Subsequently, Tsuji and co-workers elegantly described a Cu(i)-catalyzed silacarboxylation reaction of internal alkynes with carbon dioxide and silylboranes.¹⁸ In 2013, Hoveyda's group reported an efficient NHC–Cu-catalyzed hydrosilylation of terminal alkynes also using **1** to generate linear (*E*)-β-vinylsilanes.¹⁹ So far, there has been no report on the copper-catalyzed hydrosilylation of electron-deficient allenes or propiolate derivatives.²⁰ The need to use a strong base NaO'Bu to initiate the copper(i) catalytic cycle in the previously reported works by using silylboronate reagent **1** limits the use of this method for base sensitive substrates.^{17–19} To circumvent this issue in order to generate multi-substituted vinylsilanes, we report a copper(i)-catalyzed protosilylation reaction of allenes and propargyl ester derivatives to produce multi-substituted vinylsilanes with very high stereoselectivities under very mild reaction conditions.

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Table 1 Optimization of the reaction conditions^a

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PhMe ₂ Si-B(pin)	+ = -\\POPh₂	cat. (10 mol%) L (11 mol%) additive (11 mol%) alcohol (2 equiv.)	∑Ph Si—
4	20	colvent (1 ml.)	_ POPh₂

PI	nMe ₂ Si-B(p	in) + ==	—∖ POPh₂	additive	1 mol%) e (11 mol%) ol (2 equiv.)	- =	ši—
	1	2	2a		ent (1 mL)	3a	POPh ₂
Entry	Catalyst	Ligand	Additive	Alcohol	Solvent	T (°C)	Yield ^b (%)
1	CuCl	dppe	KO^tBu	МеОН	THF	30	32
2	CuCl	dppe	_	MeOH	THF	30	Trace
3	CuCl	dppe	NEt_3	MeOH	THF	30	14
4	CuCl	dppe	NEt_3	MeOH	Toluene	30	No reaction
5	CuCl	dppe	NEt_3	MeOH	DMSO	30	38
6	CuCl	dppe	NEt_3	_	MeOH	30	69
7	CuCl	dppe	NEt_3	_	^t BuOH	30	75
8	CuBr	dppe	NEt_3	_	^t BuOH	30	75
9	CuI	dppe	NEt_3	_	^t BuOH	30	74
10	CuBr	dppe	NEt_3	_ _ _	^t BuOH	30	70
11	CuBr	Johnphos	NEt_3	_	^t BuOH	30	68
12	CuBr	Xantphos	NEt_3	_	^t BuOH	30	67
13	CuBr	dppe	_	_	^t BuOH	30	No reaction
14	_	dppe	NEt_3	_	^t BuOH	30	No reaction
15	CuBr	_	NEt_3	_	^t BuOH	30	46
16	CuBr	dppe	NEt_3		^t BuOH	40	81

^a All reactions unless otherwise stated were carried out with 2a (0.3 mmol), 1 (0.33 mmol), 10 mol% of the copper catalyst (0.03 mmol), 11 mol% ligand, 11 mol% additive and 2 equiv. of alcohol in the given solvent (1.0 mL). b Isolated yield.

With the optimized conditions in hand, we next tested the scope of allenes for this Cu(1)-catalyzed hydrosilylation reaction using (dimethylphenylsilyl)boronic acid pinacol ester 1. The results are shown in Table 2. Firstly, we extended this reaction to the monosubstituted allenic ester and a sulfone. The desired products were obtained in good yields, respectively (Table 2, entries 3b and 3c). In all the cases using 1,1-disubstituted allenes, very high yields of the

Table 2 Synthesis of multi-substituted vinylsilanes using internal alkynes^a

β-silvlation products were obtained (Table 2, entries 3d-3g). 1,3-Disubstituted allenes were also treated as substrates in our reaction and the corresponding trisubstituted vinylsilanes could be generated in reasonable yields (Table 2, entries 3h, 3i and 3j). Furthermore, we noticed that the multi-substituted allenes (Table 2, entries 3k, 31 and 3m) are good substrates for this reaction. In all cases, only the β-substituted products were obtained in high yields.

After the successful synthesis of dimethylphenylsilyl-substituted butenoate derivatives, next we turn our attention to study the hydrosilylation of propiolate derivatives which would provide very useful Si-substituted acrylate derivatives. 21 We chose methyl non-2ynoate and (dimethylphenylsilyl)boronic acid pinacol ester 1 to optimize the reaction conditions (see the details in the ESI†). To our surprise, we found that in the absence of any phosphine ligand, the exclusive (E)-isomer 5a could be generated in a very high yield (92%) when the reaction was carried out using 10 mol% CuBr as a catalyst combined with 11 mol% Et₂N in MeOH.²²

Next, we examined the scope of the reaction using various propiolate derivatives. It was found that all the substrates possessing aliphatic side-chains could provide the desired products in very high yields (Table 3, entries 5a-5g). Functional groups

Table 3 Synthesis of multi-substituted vinylsilanes using internal alkynes^a

^a Unless noted otherwise, the reaction conditions are as follows: allene (0.3 mmol), 1 (0.33 mmol), CuBr (10 mol%), dppe (11 mol%) and Et₃N (11 mol%) in t BuOH (1.0 mL) heated at 40 ${}^{\circ}$ C for 24 h. b Isolated yields.

^a Unless noted otherwise, the reaction conditions are as follows: alkyne (0.3 mmol), 1 (0.6 mmol), CuBr (10 mol%) and Et $_3$ N (11 mol%) in MeOH (1.0 mL) stirred at 28 °C for 24 h. b Isolated yields.

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Scheme 1 Proposed mechanism for the synthesis of vinylsilanes from allenes.

including chloride (Table 3, entry 5c), protected hydroxy (Table 3, 5e) and bulky cyclopropane groups (Table 3, entry 5f) are well tolerated in this reaction. β -Aromatic ring substituted propiolates also worked well in this reaction (Table 3, entries 5h-5m). The electronic properties of the substituents on the aromatic ring have no apparent effect on the yield of the products. Heterocycles such as a thiophene group containing substrate also furnished the product 5p in 87% yield (Table 3, entry 5n). When the substrates were changed to butynone derivatives, the desired products could also be obtained in moderate to good yields (Table 3, entries 5o and 5p).

Based on the observed results, a possible mechanism was proposed as shown in Scheme 1. Firstly, the Et₃N could help to activate the B–Si bond *via* possible coordination between nitrogen and boron atoms.²³ In the presence of a copper catalyst, species **A** could be generated. Due to the electron density bias of allenes with an electron-withdrawing group, an addition of Si–Cu into the double bond will afford an allylic copper species **B** which could then undergo protonolysis in the presence of ^tBuOH to give the desired product 3.²⁴ The reactive L–Cu–O^tBu complex is then regenerated and will be involved in the next catalytic cycle. As for the hydrosilylation of propiolate derivatives, a similar mechanism as depicted in Scheme 1 could also be proposed. The higher reactivity of propiolate derivatives permits an easy transformation of the silyl group into the triple bond.

In conclusion, we have found that copper salts can catalyze the silylation reactions of allenes and propiolate derivatives with silylboronates under mild reaction conditions. These methods provide facile access to versatile silyl-substituted homoallylic esters and β -silyl-substituted acrylates in moderate to excellent E/Z selectivities. This study also reveals that weak bases such as triethylamine could activate the Si–B bond to initiate the catalytic cycle for the transformation of a silyl group into unsaturated bonds. The use of this strategy to carry out other organic transformations is in progress.

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