1,4-Carbosilylation of 1,3-Dienes via Palladium Catalyzed Three-Component Coupling Reaction

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Abstract: Three-component coupling reaction of acid chlorides, organodisilanes, and 1,3-dienes achieves 1,4carbosilylation of the 1,3-dienes to afford allylic silanes as the product. Bis(dibenzylideneacetone)palladium, a naked Pd(0) complex without donating ligand, showed high catalytic activity. A carbon and a silicon substituent are introduced at 1- and 4-positions of the 1,3-dienes regio- and stereoselectively with concomitant decarbonylation of the acid chlorides. A wide variety of allylic silanes are synthesized in high yields from these easily accessible substrates. On the other hand, a bulky acid chloride such as adamantane-1-carboxylic acid chloride did not undergo the decarbonylation reaction but afforded allylic silanes containing acyl functionality. In all these reactions, transmetalation of the disilanes with η^3 -allylchloropalladium intermediates might be a critical step in the catalytic cycle. As a model reaction for the transmetalation, reaction of di- μ -chlorobis[(1,2,3- η)-4-phenyl-2-butenyl]dipalladium with disilanes was carried out. Although intermediate η^3 -allylsilylpalladium species could not be detected, the corresponding allylic silanes, silyl chlorides, and Pd(0) metal were formed during the reaction. Furthermore, a similar three-component coupling reaction using aryl iodides, organosilylstannanes, and dienes also proceeded. However, the selectivity and the yield decreased considerably.

Introduction

Addition of organosilanes (H-Si) to unsaturated compounds $(hydrosilylation)^{1}$ is a most useful reaction for synthesis of a wide variety of organosilicon compounds [Scheme 1(i)]. On the other hand, if a carbon and a silicon unit, instead of a hydrogen and a silicon, are introduced simultaneously into unsaturates (carbosilylation), the reaction might be far more beneficial as synthetic method. However, while some carbonmetal bonds² such as C-AI and C-Cu are reactive enough to undergo so-called carbometalation reaction, most C-Si bonds are inert under usual reaction conditions. Therefore, it seems extremely difficult to activate C-Si bonds directly³ toward insertion of unsaturates into these bonds [Scheme 1(ii)]. In order to resolve this dilemma, we explored three-component coupling reaction,⁴ in which a carbon and a silicon substituents are introduced from different sources into unsaturated substrates [Scheme 1(iii)]. We adopted 1,3-dienes as the unsaturated substrates, since selective introduction of the carbon and the silicon units at 1- and 4-positions of the 1,3-dienes gives

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Scheme 1

Hydrosilylation
H-Si
$$\lesssim$$
 + $X=Y \lesssim$ ----- H-X-Y-Si \lesssim (i)

Carbosilylation (via C-Si cleavage)

Carbosilylation (via three-component coupling)

synthetically important allylic silanes. Allylic silanes⁵ are highly versatile synthetic intermediates due to their regioselective reactions with various electrophiles.⁶ Therefore, much attention has been paid to preparation method⁷ of allylic silanes including allylic Grignard reactions,^{7a,b} hydrosilylation of 1,3-dienes,^{7c-e} and Wittig reactions with β -silylethylidenephosphorane.^{7f,g}

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1,4-Carbosilylation of 1,3-Dienes

In this paper, we describe a palladium catalyzed threecomponent coupling reaction of organodisilanes (Si-Si),⁸ 1,3dienes, and acid chlorides to realize the overall carbosilylation reaction. The reaction is highly regio- and stereoselective providing a useful synthetic method of allylic silanes from these easily accessible starting materials. We also developed similar three-component coupling reaction with 1,3-dienes, where organosilylstannanes $(Si-Sn)^9$ are employed as the silicon source and aryl iodides as the carbon source.

Results and Discussion

Carbosilylation of 1,3-Dienes Using Acid Chlorides and Organodisilanes. It is well-known that acid chlorides react with low valent transition metal complexes by oxidative addition¹⁰ and generate carbon-transition metal bonds. On the other hand, organodisilanes⁸ are very useful silylating reagents in the presence of a transition metal catalyst. First we employed acid chlorides as a carbon source and organodisilanes as a silicon source in the three-component coupling reaction with 1,3-dienes (eq 1).¹¹ The results are summarized in Table 1. Benzovl chloride (1a) reacts with hexamethyldisilane (2a) and 1,3butadiene (3a) in the presence of a catalytic amount (5 mol %) of $Pd(DBA)_2^{12,13}$ (DBA = dibenzylideneacetone) to afford the coupling product (4a) in high yield (entry 1). The phenyl and the silvl substituents are introduced regio- and stereoselectively at the 1- and 4-positions of 3a providing only the (E) isomer. Decarbonylation from the acid chloride¹⁴ proceeded completely and coupling products containing carbonyl functionalities were not detected.



The reaction also proceeds with substituted benzoyl chlorides in good to high yields. Thus, chloro (1c), bromo (1d), nitro (1e), and keto (1f and 1g) functionalities were tolerated in the reaction (entries 3-7). Usually direct synthesis of these functionalized allylic silanes are highly difficult by the conventional procedures.⁷ 2- and 1-Naphthoyl chlorides (1h and 1i) also afforded the corresponding products in excellent yields (entries 8 and 9). 2-Furoyl chloride (1j), 5-bromo-2-furoyl chloride (1k), and 2-thiophenecarbonyl chloride (1l) also afforded the (E)-1,4-adducts in good yields (entry 10-12). Terephthaloyl chloride (1m) afforded the corresponding compound having two allylic silane side chains (entry 13). However, o-phthaloyl chloride and 2-(p-toluoyl)benzoyl chloride were not consumed in the reaction, suggesting steric congestion with these substrates affects the reaction. Various (E)-alkenovl chlorides (1n-r) can be employed in the present coupling reaction and provided the corresponding (E)-1,4 adducts regioand stereoselectively (entries 14-18). Alkynoyl chlorides (1s and 1t) also afforded the corresponding products in high yields (entries 19 and 20). As 1,3-dienes, isoprene (3b) and 2,3dimethyl-1,3-diene (3c) can be employed. The reaction gave the products (4u-w) regioselectively (entries 21-23); with isoprene (3b) the trimethylsilyl group is introduced on the methylene carbon closer to the methyl substituent. In these cases, however, stereoselectivity was often modest. Symmetrically substituted disilanes such as 1,1'-diphenyl- (2b), 1,1'di-(4-fluorophenyl)- (2c), 1,1'-dichloro- (2d), and 1,1'-difluorotetramethyldisilanes (2e) afforded the corresponding carbosilvlation products (4x- α) with high regio- and stereoselectivity (entries 24-27). As for monosubstituted unsymmetrical disilanes, phenylpentamethyldisilane (2f) afforded the dimethylphenylsilyl derivative (4x) exclusively (entry 28), whereas fluoropentamethyldisilane (2g) gave fluorodimethyl isomer (4 α) as a major product (entry 29).

Effects of some reaction conditions and selected catalyst precursors were examined in the coupling reaction using **1a**, **2a**, and **3a**. As a solvent, aromatic hydrocarbon such as toluene gave the best results (entry 1); the reaction in tetrahydrofuran (THF) and dimethylformamide (DMF) lowered the yields considerably (15% and 5%, respectively). The reaction proceeds smoothly at 80 °C, while lower (50 °C) or higher (130 °C) reaction temperature reduced the yield considerably (19% and 36%, respectively). As a catalyst precursor, Pd(DBA)₂,^{12,13} a naked Pd(0) complex without donating ligand, is most effective. An addition of AsPh₃ (As/Pd = 4) or P(OEt)₃ (P/Pd = 4) to Pd(DBA)₂ reduced the yield of **4a** to 41% or 13%, respectively.

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Table 1. 1,4-Carbosilylation of 1,3-Dienes Using Acid Chlorides and Organodisilanes^a

entry	acid chloride	disilane	diene	product	yield / % ^b	entry	acid chloride	disilane	diene	product	yield / % ^b
1		2a	3a	Q4a	86(93)	15	Q 10	2a	3a	Q 40	78
2	we [°] c⊨ 1b	2a	38	the Carrow silde, 4b	77	16		2a	3a	Sillie ₃	72
3	ci Ci ic	2a	3a	CL Bilden 4c	80	17	Land a	2a	38	Silver 4q	57
4	Brown Ci 1d	22	3a	Br 4d	90	18		22	3a	June 141	81
5	NO2 Ci le	2a	3a	NO2 SIMO2 40	51	19	o is	2a	3a	\$1Me; 45	85
6	Martin St. 1f	2a	38	He Silles 41	92	20		2a	3a	Silves 4t	75
7		29	3e	4g	74	21		2a	3b	Company 44d	82(91)
8		2a	3a	4h	95	22	J r	2a	3b	Silles due	80
9		2a	3a		92	23		2a	3c	Sildes 4wr	47
	\mathbf{U}					24	C ¹ ci 1a	2b	3a	Q→→ silder	68
10	ر بر الرقم ال	2a	3a	Sille, 4j	61	25	C in	2c	3a	Q	71
11	Br-Co-Ci 1k	2a	3a	Br Silles 4k	63	26	Constant in	2d	3a	Sillez ^{CI} 4z	61
12	5 6 1	22	3 a	S Silves 41	64	27	° Ci la	2e	3a	Silder 4a	40
13	cr√ ^ℓ ci 1m	2ª°	3a	Mea Si Am	70	28	ب م ا	21	3a	Q 4x	89
14		2a	3 a	Constitutes 4n	94	29	C ² ci is	2g	3a	4a. + 70 %	4a 9%

^{*a*} Conditions: acid chloride (1: 0.50 mmol), disilane (2: 0.50 mmol), 1,3-diene (3: 1.5 mmol), Pd(DBA)₂ (0.025 mmol; 5 mol %), and toluene (2.0 mL) at 80 °C for 4 h. ^{*b*} Isolated yields. Numbers in parentheses show GC yields determined by the internal standard method. ^{*c*} The disilane (1.0 mmol). ^{*d*} E/Z = 75/25. ^{*c*} E/Z = 91/9. ^{*f*} E/Z = 65/35.

Furthermore, the addition of PBu₃ (P/Pd = 4) or bidentate ligands such as DPPE [1,2-bis(diphenylphosphino)ethane], DPPP [1,2-bis(diphenylphosphino)propane], and DPPF [1,1'bis(diphenylphosphino)ferrocene] (for these bidentate phosphines, ligand/Pd = 2) totally suppressed the conversion of **1a**. Other selected transition metal precursors (5 mol %) such as Pd(PPh₃)₄, PdCl₂(PPh₃)₂, Pt(DBA)₂, and Pt(CO)₂(PPh₃)₂ were not active as the catalyst and the acid chlorides remained intact. Thus, palladium(0) complex without coordinating ligands is essential as the catalyst precursor for the present threecomponent coupling reaction.

In the reactions listed in Table 1, the decarbonylation^{14,15} from the acid chlorides (1) took place completely. Usually, *catalytic* decarbonylation of acid chlorides or aldehydes are sluggish,¹⁶ especially at lower reaction temperatures. In the present coupling reaction, highly unsaturated nature of the palladium catalyst center might facilitate such fast and complete decarbonylation of the acid chlorides. After the reaction, comparable amount of carbon monoxide was detected in the gas phase by GC analysis (on Molecular Sieve 13X-S, oven temperature at 50 °C). When the reaction was carried out under carbon monoxide pressure (10 kg/cm²), consumption of the acid chlorides was low (<10%), and **4** was not obtained at all. Yamamoto and co-workers reported that palladium catalyzed reaction of acid chlorides (1) with Me₃SiSiMe₃ (2a) affords acyl silanes (5) by suppressing the decarbonylation of 1 (eq 2).¹⁷ The key to the suppression was added triethyl phosphite $[P(OEt)_3]$ ligand to $[PdCl(\eta^3-C_3H_5)]_2$. We attempted the same catalyst system under the reaction conditions of entry 1. However, conversion of the acid chloride (1a) was quite low (<10%), and the yield of the coupling product (4a) was only 7% with concomitant formation of a trace of PhCOSiMe₃ (5a) (eq 3). Allylic silanes having carbonyl functionality such as 6a were not obtained at all. Other ligands such as P(OPh)₃, P(OPr')₃, and P(OCH₂)₃CEt added to Pd(DBA)₂ or [PdCl($\eta^{3-}C_3H_5$)]₂ could not suppress the decarbonylation; usually these added ligands lowered the conversion of the acid chlorides considerably.



The acid chlorides (1) provide the coupling products (4) via the decarbonylation. Other substrates which can react with a

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low valent metal center by oxidative addition might work as the carbon source in an analogous three-component coupling reaction. Thus, several candidates for the carbon source are examined. Benzoyl bromide^{14b} and benzoyl fluoride¹⁸ did not afforded the coupling product (4a) at all; with benzoyl bromide the reaction was complicated, while benzoyl fluoride did not convert. Heptanoyl chloride did not afford the carbosilylation products at all presumably due to β -hydrogen elimination of hexyl palladium intermediate generated by the oxidative addition of the acid chloride followed by the decarbonylation. As for an aryl halide, bromobenzene afforded 4a in 40% yield. However, with iodobenzene, its conversion was low (40%), and the yield of 4a was only 8%. Chlorobenzene did not convert at all. β -Bromostyrene afforded the corresponding coupling product (4n) in lower yield (32%) than the corresponding cinnamoyl chloride (entry 14). With 4-bromobenzoyl chloride (1d) as mentioned above (entry 4), the reaction occurred at the acid chloride site rather than the aromatic bromide site, suggesting the reaction at the former site seems to be faster than the latter. This rate difference was confirmed in a competitive reaction between 4-methylbenzoyl chloride (1b; 1.0 equiv) and bromobenzene (1.0 equiv) in the presence of Me₃SiSiMe₃ (2a; 0.50 equiv) and 1,3-butadiene (3a; 3.0 equiv) under standard reaction conditions [at 80 °C, for 2 h, with 5 mol % of $Pd(DBA)_2$ based on 2a]. In the reaction, most of the bromobenzene was recovered and the coupling product came almost from the acid chloride: yield of 4b, 78%; 4a, trace. The same result was observed with benzoyl chloride (1a; 1.0 equiv), 4-bromotoluene (1.0 equiv), 2a (0.50 equiv), and 3a (3.0 equiv) indicating the *p*-methyl substituents have virtually no effects on the reactivity: yield of 4a, 89%; 4b, trace. Moreover, esters having good leaving groups such as phenyl trifluoromethanesulfonate (C_6H_5OTf) and diethyl phenyl phosphate [C_6H_5OP - $(O)(OC_2H_5)_2$ have no reactivities in the reaction. We have recently reported dimerization-double silvlation of 1,3-dienes using 2a catalyzed by Pd(DBA)₂, in which the reaction smoothly proceeds at room temperature and provides the product (7) in high yield regio- and stereoselectively (eq 4).8ª When the substrates of the carbon source were apparently inert [with C₆H₅-COF, C_6H_5Cl , C_6H_5OTf , and $C_6H_5OP(O)(OC_2H_5)_2$], 7a was formed in high yields (>80%).



In the present reaction, one of the silvl moieties of the disilanes (2) is incorporated in the allylic silane (4) as the product. Fate of the other silyl moiety of 2 was examined in the reaction using Me₃SiSiMe₃ (2a) by measuring ²⁹Si-NMR spectra of the resulting reaction mixture. After the reaction was carried out under the same reaction conditions as entry 1, the expected amount of Me₃SiCl (30.31 ppm; lit.^{19a} 30.27 ppm) was found along with excess 2a (-20.51 ppm; lit.^{19b} -20.51 ppm), indicating that one of the silvl moieties of the disilane (2) is trapped as the silyl chloride. Strong Si-Cl bond (Si-Cl bond energy of Me₃SiCl:²⁰ 113 kcal mol⁻¹) may be a driving force of the reaction.

Scheme 2



A possible catalytic cycle for the present reaction via the decarbonylation (Table 1) is shown in Scheme 2. Oxidative addition of acid chloride (1) to the Pd(0) active catalyst species (8) initiates the catalytic cycle and gives acyl palladium species (9). After the decarbonylation to 10, 1,3-diene (3) inserts into the resulting C-Pd bond providing allylchloropalladium intermediate (11). Transmetalation of disilane (2) with 11 may provide allylsilyl species (12) with a concomitant formation of the silvl chloride. Finally, reductive elimination of 12 affords the allylic silane (4) as the product and regenerates the Pd(0)active catalyst (8).

4x: C.H.

4β:CH₂CH=CH₂

6

0.001 94

66

4.46

0.54

With regard to the catalytic cycle, oxidative addition of acid chlorides¹⁰ (1) to a low valent metal complex $(8 + 1 \rightarrow 9)$ and decarbonylation of acyl metal complexes¹⁴ ($9 \rightarrow CO + 10$) are well-known. Furthermore, insertion of 1,3-diene (3) into carbon-metal bond $(10 + 3 \rightarrow 11)$ has been also observed in several transition metal reactions.²¹ However, there is no precedent for explicit transmetalation of disilanes to form silylmetal species $(11 + 2 \rightarrow 12)$. Thus, we examined a model reaction of the transmetalation step. In the carbosilylation reaction of 1,3-butadiene (3a) using benzoyl chloride (1a) and Me₃SiSiMe₃ (2a) (entry 1), chloro[$(1,2,3-\eta)$ -4-phenyl-2-butenyl]palladium intermediate (11a: $R = C_6H_5$) might be involved in the catalytic cycle. As a model complex for **11a**, di- μ -chlorobis- $[(1,2,3-\eta)-4$ -phenyl-2-butenyl]dipalladium (13) was prepared from the corresponding chloride and Na₂PdCl₄.^{33c,35} Then, a stoichiometric reaction of 13 with Me₃SiSiMe₃ (2a; 2 equiv) was carried out at room temperature in toluene- d_8 (Scheme 3). In the reaction, a clear yellow solution gradually turned brown, and finally palladium mirror and a small amount of palladium powder appeared in 6 h. ²⁹Si-NMR spectrum of the colorless

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filtrate showed that Me₃SiCl (30.06 ppm) and 4a (7.17 ppm) were formed in equal amounts; yield of the latter was 64% which was determined with ¹H-NMR spectrum using mesitylene as an internal standard. The transmetalation would proceed via intermediate allylsilyl metal species (14) with a concomitant formation of silyl chloride. We attempted to isolate the intermediate 14 by adding PPh3 or PMe3 (2 equiv) to the reaction mixture. However, the added ligand apparently hindered the transmetalation, and any silvl species such as 14 could not be detected with ²⁹Si-, ¹H-, and ¹³C-NMR spectra. 1,1'-Diphenyltetramethyldisilane (2b) also reacted with 13 under the same reaction conditions. The reaction was faster than that of 2a; in 1 h, the palladium mirror appeared. With 1,1'-diallyltetramethyldisilane (2h), the reaction was much faster; within 5 s black powders precipitated. In these reaction mixtures, comparable amount of the corresponding silvl chloride [C₆H₅-Me₂SiCl²² or (CH₂=CHCH₂)Me₂SiCl] and allylic silane (4x or 4 β) were detected by ²⁹Si-NMR. The yields of 4x and 4 β were determined with ¹H-NMR spectra using mesitylene as the internal standard. Precoordination through the unsaturated substituents of 2b and 2h might accelerate the transmetalation reaction. In these cases, the postulated intermediate 14 could not be detected, either. As soon as 14 is formed, fast reductive elimination of these highly unsaturated (14e) species may afford 4 and Pd(0) very quickly.

Coupling Reaction without Decarbonylation of Acid Chlorides. The acid chlorides such as listed in Table 1 undergo the complete decarbonylation reaction (vide supra). In contrast, adamantane-1-carboxylic acid chloride (1u) did not undergo the decarbonylation, but selectively afforded **6b** containing the acyl functionality in 70% yield (eq 5). Notably, the coupling product



via the decarbonylation (4γ) was not afforded at all. Similar result was obtained with *tert*-butylacetyl chloride (1v), affording **6c** in 74% yield without formation of 4δ (eq 6). With these



sterically demanding acid chlorides, equilibrium between acyl and alkylcarbonyl species might lie far toward the former.²³ On the other hand, pivaloyl chloride (1w) afforded a mixture of two allylic silanes formed with and without the decarbonylation (6d and 4ϵ in 58% yield; $6d/4\epsilon = 1/1$) (eq 7). Thus, difference



between the reaction with and without the decarbonylation seems to be very subtle. A competitive reaction between 1u and 1a was carried out (eq 8). In the reaction, only 4a derived from



1a via the decarbonylation was afforded exclusively; 6b, 6a, and 4γ were not obtained at all. The similar reactivity was observed in the competitive reaction between 1u and 1o, affording only 4o derived from 1o; 6b, 6e, and 4γ were not obtained (eq 9). Insertion of the diene into aryl- or alkenyl-



Pd bond $(10 + 3 \rightarrow 11)$: in Scheme 2) might be much faster than the insertion into acyl-Pd bonds $(9 + 3 \rightarrow 15)$: in Scheme 4, cycle A). From benzoylformyl chloride (1x), the main product is the double decarbonylation product (4a), while 6awas obtained only in low yield (eq 10). Moreover, phenylacetyl



chloride (1y) did not afford the corresponding allylic silane, but afforded only acylsilane (5b) in high yield (eq 11). In this



case, the oxidative addition of **1y** to the Pd(0) affords a phenylacetylpalladium intermediate. This particular metal species is well-known to be easily enolized: $C_6H_5CH_2(C=O)M \leftrightarrow C_6H_5CH=CH(OH)M.^{24}$ The enolization might affect the reaction and prevent the insertion of the 1,3-diene (Scheme 4, cycle B).

Carbosilylation of 1,3-Dienes Using Aryl Iodides and Organosilylstannanes. In the carbosilylation of 1,3-dienes (3) using disilanes (2) (eq 1), the transmetalation of 2 is a key step in the catalytic cycle (Scheme 2). On the other hand, organo-

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Scheme 4



stannanes are well-known to undergo the transmetalation efficiently.²⁵ Thus, we tried to utilize organosilylstannanes (18) in place of 2 as the silicon source. When 18 was used in place of 2 in eq 1, conversion of 18 was low and the corresponding carbosilylation products (4) were obtained only in low yields (<5%). After examination of several possible carbon source, we found that aryl iodides (19) are suitable in the carbosilylation reaction using 18 (eq 12). The results are listed in Table 2. Iodobenzene (19a) reacted with n-Bu₃SnSiMe₃ (18a) and 1,3butadiene (3a) to afford the corresponding carbosilylation products (entry 30). Unlike the reaction using disilanes (2) as the silicon source, the reaction provided only a mixture of 1,4-(4a: E/Z mixture) and 1,2-adduct (20a) in 50% yield. Similar selectivity and yields were obtained when 19b, 19c, or 1-iodonaphthalene (19d) was used as the aryl iodide (entries 31-33). Aryl bromides and chlorides did not provide the corresponding carbosilylation products at all. Isoprene (3b; entries 34-36), 2,3-dimethyl-1,3-butadiene (3c; entry 37), and 2-phenyl-1,3butadiene (3d; entries 38 and 39) also provided the corresponding coupling products. However, selectivity of the reaction was also moderate; a mixture of 1,4- and 4,1-adducts were obtained when the unsymmetrical dienes (3b and 3d) were employed.



Effect of the catalyst precursor was examined in the reaction using **18a**, **3b**, and **19a**. Among the complexes examined, Pd(DBA)₂ shows the best result (entry 34); the yield is 39% with PdMe₂(COD), 40% with Pd(η^3 -C₃H₅)(η^5 -C₅H₅), 40% with PdBr₂(COD)₂, 32% with PdCl₂(COD)₂, 20% with Pd(PPh₃)₄, and 16% with Pd(OAc)₂. Naked palladium complex without donating ligands seems most favorable, which is reminiscent of the carbosilylation using the organodisilanes (**2**) and the acid chlorides (**1**) (eq 1). Other selected catalyst precursors such as RhCl(PPh₃)₃, [Rh(COD)₂]BF₄, and Pt(CO)₂(PPh₃)₂ did not show any catalytic activity. As for the organosilylstannane, *n*-Bu₃SnSiMe₃ (**18a**) was the most suitable. Other silylstannanes

Table 2. Carbosilylation Using n-Bu₃SnSiMe₃ (18a)^a

		product					
entry	diene	Ar-I	yield, ^b %	distribution ^c			
30	3a	19a	(50) 42	4a:20a:21a = 84(88):16:-			
31	3a	19b	40	4b:20b:21b = 85(88):15:-			
32	3a	19c	56	4c:20c:21c = 88(87):12:-			
33	3a	19d	61	4i:20i:21i = 85(87):15:-			
34	3b	19a	(57) 48	4u:20u:21u = 75(55):13:12			
35 ^d	3b	19a	(39)	4u:20u:21u = 71(55):13:16			
36	3b	19d	62	4ζ :20 ζ :21 ζ = 75(64):13:12			
37	3c	19a	44	4w:20w:21w = 100(68):0:-			
38	3d	19a	67	$4\eta: 20\eta: 21\eta = 88(67): 6:6$			
39	3d	19d	85	$4\dot{\theta}:20\dot{\theta}:21\dot{\theta}=84(70):8:8$			

^{*a*} n-Bu₃SnSiMe₃ (18a: 0.50 mmol), 3 (1.5 mmol), 19 (0.50 mmol), Pd(DBA)₂ (0.025 mmol), toluene (2.0 mL), at 100 °C for 4 h. ^{*b*} Isolated yield. Numbers in parentheses show GC yields determined by the internal standard method. ^{*c*} Percentages of the *E*-isomers of 4 are shown in the parentheses. ^{*d*} PdMe₂(COD) (0.025 mmol) is used in place of Pd(DBA)₂.

such as $Me_3SnSiMe_3$, $Me_3SnSiMe_2(t-Bu)$, $Me_3SnSiMe_2Ph$, $Ph_3SnSiMe_3$, and $n-Bu_3SnSiMe_2SiMe_3$ were totally ineffective and did not afford any carbosilylation products.

Alkenyl iodides such as 1-iodo-1-octene and 1-iodo-1cyclohexene reduced the yields appreciably (<20%), and the selectivities of the reaction were still moderate. The reaction path changed dramatically, when phenyl trifluoromethanesulfonate (Ph-OTf), diethyl phenyl phosphate (PhOP(=O)-(OEt)₂), or phenyl *p*-toluenesulfonate (Ph-OTs) was employed as the carbon source (eq 13). The reaction proceeded even at room temperature. However, no carbosilylation products were obtained, but an equimolar mixture of all four stereoisomers of **22** were isolated in 75–85% yields. The phenyl group was not incorporated in the products. As for the fate of the phenyl functionality, comparable formation of benzene was confirmed by GC analysis (on Chromosorb 103 at 200 °C). The mechanism of this reaction is not clear at the present stage.

We have recently reported 1,4-silylstannation of 1,3-dienes (3) using 18 catalyzed by $Pt(CO)_2(PPh_3)_2$, which affords 23 selectively.^{9a} There might be some possibility that the 1,4silylstannation of the 1,3-dienes occurred first, and its product (23) was an intermediate for following Stille coupling reaction²⁵ to give 4a and/or 20a. However, in the reaction of 23 with iodobenzene (eq 14), no reactions occurred, and both 23 and 19a were recovered unchanged.²⁶ Therefore, intermediacy of the 1,4-silylstannation products (23) is unlikely.

A possible catalytic cycle for the carbosilylation using **18a** is illustrated in Scheme 5. Oxidative addition of aryl iodide (**19**) to palladium(0) species (step a) initiates the catalytic cycle.

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Scheme 5



Insertion of 1,3-diene (3) into the resulting Pd-C bond (step b) affords η^3 -allyl palladium intermediate. Then, transmetalation with 18a (step c) provides allylsilyl palladium species and tributyltin iodide. Formation of the latter was confirmed by GC analysis (OV-17). Finally, reductive elimination of allylic silanes (step d) affords the products and regenerates the active catalyst species. The regiochemistry of the reaction is determined by steps b and d, while the stereochemistry would be governed by relative stability between syn and anti η^3 allyliodopalladium intermediates.

Conclusion

Carbosilylation of 1,3-dienes has been achieved by a threecomponent coupling reaction of the 1,3-dienes, organodisilanes, and acid chlorides. The reaction proceeds highly regio- and stereoselectively to afford various allylic silanes as the product. In the catalytic cycle, transmetalation of the disilanes with η^3 allylchloropalladium intermediate might be a critical step. Similar three-component coupling reaction proceeds using organosilylstannanes, aryl iodides, and 1,3-dienes, although the selectivity and the yield were considerably low.

Experimental Section

Materials. The reagents and the solvents were dried and purified before use by usual procedures.²⁷ Hexamethyldisilane (2a) was purchased from Aldrich. 1,2-Difluorotetramethyldisilane,^{28a} fluoropentamethyldisilane,^{28a} 1,2-dichlorotetramethyldisilane,^{28b} and chloropentamethyldisilane^{28b} were prepared by the methods reported by Kumada. Other disilanes were prepared from chloropentamethyldisilane or dichlorotetramethyldisilane by the reactions with corresponding organolithium (for 2b, 2c, and 2f) or organomagnesium (2h) reagents. 2-Phenyl-1,3-butadiene was prepared by the known method.²⁹ Some acid chlorides were prepared from the corresponding carboxylic acids with thionyl chloride^{30a} (for 1i-j, 1p-q) or with oxalyl chloride^{30b} (for 1f). Phenyl trifluoromethanesulfonate^{31a} and diethyl phenyl phosphate^{31b} were prepared by the reported procedures. Organosilylstannanes (18) were prepared by the methods by Chenard.³² The following catalyst precursors were prepared by the published methods: $Pd(PPh_3)_{4,33a} PdCl_2(PhCN)_{2,33b} [PdCl(\eta^3-C_3H_5)]_{2,33c} PdCl_2$ (PPh₃)₂,^{33d} PdMe₂(COD),^{33e} Pd(η^3 -C₃H₅)(η^5 -C₅H₅),^{33c} PdBr₂(COD),^{33f}

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PdCl₂(COD),^{33f} Pt(DBA)₂,^{33g} Pt(CO)₂(PPh₃)₂,^{33h} Ru(COD)(COT),³³ⁱ [Rh(COD)2]BF4 33j

Analytical Procedure. The NMR spectra (in CDCl₃) were recorded with JEOL α -400 and GX-270 spectrometers. The mass spectra were measured on Shimadzu QP-1000 (EI) and Shimadzu 9020-DF (HRMS) equipped with a PAC 1100S computer system. The GC analysis was made on a Shimadzu GC-8APF equipped with an integrater (C-R6A) with a column (3 mm i.d. \times 3m) packed with Silicon OV-17 (2% on Uniport HP, 60/80 mesh) or Apiezon Grease L (5% on Uniport HP, 60/80 mesh). Elemental analysis was performed at the Microanalytical Center of Kyoto University.

Coupling of Acid Chlorides (1), Organodisilanes (2), and 1,3-Dienes (3) (Eq 1). A typical procedure is described for the synthesis of 4a. A mixture of 1,3-butadiene (3a; 1.5 mmol, 0.94 mL of 1.6 M stock solution in toluene), Pd (DBA)₂ (14 mg, 0.025 mmol), benzoyl chloride (1a; 70 mg, 0.50 mmol), hexamethyldisilane (2a; 73 mg, 0.50 mmol), toluene (2.0 mL), and a magnetic stirring bar were placed under argon flow in a 30 mL stainless steel autoclave containing an inserted glass liner. An air purge was confirmed by three pressurization (20 atm)-depressurization sequences with argon. The autoclave was heated to 80 °C in 10 min and held at this temperature for 4 h. The reaction was terminated by rapid cooling, and the autoclave was discharged. The resulting reaction mixture was passed through a short Florisil column (8 mm i.d. × 50 mm). GC analysis (OV-17) with phenanthrene as an internal standard showed the product (4a) was formed in 93% yield. The product (4a) was isolated in 86% yield (88 mg) by mediumpressure column chromatography (silica gel: Wakogel 300, 45-75 mm; hexane as an eluent) followed by Kugelrohr distillation (Büchi; pot temperature 80 °C/0.3 mmHg).

Coupling of Aryl Iodides (19), Organosilylstannanes (18), and 1,3-Dienes (3) (Eq 12). A typical reaction procedure is described for the synthesis of 4a. A mixture of 1,3-butadiene (3a; 1.5 mmol, 0.94 mL of 1.6 M stock solution in toluene), Pd(DBA)₂ (14 mg, 0.025 mmol), 19a (102 mg, 0.5 mmol), 18a (181 mg, 0.5 mmol), toluene (2.0 mL), and a magnetic stirring bar were placed under argon flow in a 30 mL stainless autoclave containing an inserted glass liner. An air purge was confirmed by three pressurization (20-atm)-depressurization sequences with argon. After the reaction, the reaction mixture was treated with saturated ammonium fluoride solution to remove tributyltin iodide as insoluble tributyltin fluoride. The product was isolated in 42% yield (43 mg) by medium-pressure chromatography (silica gel/ hexane) followed by Kugelrohr distillation.

Preparation of Di-µ-chlorobis[(1,2,3-η-4-phenyl-2-butenyl]dipalladium (13). 1-Chloro-4-phenyl-2-butene was prepared by the reaction of 1,4-dichloro-2-butene with phenyllithium in the presence of nickel catalyst.³⁴ Then, according to the published procedure,^{33c,35} di-µ-chlorobis[(1,2,3-η-4-phenyl-2-butenyl]dipalladium (13) was prepared as a yellow solid (1.06 g, 78%) from 1-chloro-4-phenyl-2-butene and Na₂PdCl₄ with bubbling carbon monoxide: ¹H-NMR (CDCl₃) δ 3.04 (d, J = 12 Hz, 2H), 3.12-3.22 (m, 4H), 4.04 (d, J = 6 Hz, 2H),4.19 (m, 2H), 5.35 (dt, J = 6 Hz, 12 Hz, 2H), 7.22–7.35 (m, 10H); $^{13}\text{C-NMR}$ δ 38.05 (t), 59.32 (d), 84.87 (d), 110.20 (t), 126.66 (d), 128.67 (d), 128.85 (d), 137.82 (s).

Reaction of Di-µ-chlorobis[(1,2,3-η)-4-phenyl-2-butenyl]dipalladium (13) with Disilanes (2a, 2f, 2h) (Scheme 3). The reaction using 2a is typical. In a 5 mm ϕ NMR tube, 13 (15 mg, 0.03 mmol) was dissolved in an argon-degassed toluene-d₈ (0.5 mL). Hexamethyldi-

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silane (2a) (9 mg 0.06 mmol) was added into the solution at 0 °C. Then, the mixture was left at room temperature. In 6 h, palladium mirror appeared on the wall of the NMR tube. The liquid part was transferred through a short Celite plug to another NMR tube. The formation of 4a was confirmed by ¹H- and ¹³C-NMR spectra. On the ¹H-NMR spectrum, the yield of 4a was determined (64%) using mesitylene as an internal standard. Furthermore, the ²⁹Si-NMR spectrum is most diagnostic of comparable formation of trimethylsilyl chloride (30.06 ppm; lit.^{19a} 30.27 ppm); also two other resonances appeared at -19.86 ppm (for excess 2a; lit.^{19b} -20.51 ppm) and 7.17 ppm (for 4a).

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Supporting Information Available: Spectral (¹H and ¹³C NMR and MS) and some elemental analysis data for $4\mathbf{a}-\mathbf{z}$, $4\alpha,\beta, 4\epsilon-\theta, 5\mathbf{b}, 6\mathbf{b}-\mathbf{d}, 20\mathbf{a}-\mathbf{c}, 20\mathbf{i}, 20\mathbf{u}, 20\xi-\theta, 21\mathbf{u}, 21\xi-\theta$, and 22 (9 pages). This material is contained in many libraries on microfiche, immediately follows this article in the microfilm version of the journal, can be ordered from the ACS, and can be downloaded from the Internet; see any current masthead page for ordering information and Internet access instructions.

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