See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/6876256

Gold-Catalyzed Intramolecular Allylation of Silyl Alkynes Induced by Silane Alcoholysis

ARTICLE IN JOURNAL OF THE AMERICAN	CHEMICAL SOCIETY	· SEPTEMBER 2006
Impact Factor: 12.11 · DOI: 10.1021/ja062560p · Source: Pul	bMed	

CITATIONS	READS
50	10

2 AUTHORS, INCLUDING:



Daesung Lee

University of Illinois at Chicago

123 PUBLICATIONS 2,572 CITATIONS

SEE PROFILE

1 7

Published in final edited form as:

J Am Chem Soc. 2006 August 23; 128(33): 10664–10665. doi:10.1021/ja062560p.

Gold-Catalyzed Intramolecular Allylation of Silyl Alkynes Induced by Silane Alcoholysis

Sangho Park and Daesung Lee*

Department of Chemistry, University of Wisconsin, Madison, WI 53706

Abstract

The activation of alkynyl allyl silanes with cationic gold catalyst in the presence of alcohols provides a vinyl silanes that contain the allyl group at the β -position and the alkoxysilyl group in *cis*-orientation. The bond reorganization processe is most consistent with the involvement of a carbocationic intermediate, which undergoes a nucleophilic attack by an alcohol selectively at the silicon center. The cis vinyl silyl ether products can be further elaborated by ring-closing and cross metathesis to form more substituted 1,4-dienyl silanes.

The pursuit of synthetic efficiency has promoted constant development of new concepts and innovated synthetic arsenals. ¹ One of the most effective ways of achieving synthetic efficiency is to implement tandem reactions ² into a synthetic sequence whereby many bond-forming and cleaving events can occur in one synthetic operation. Following from our interest in silyl etherbased metathesis chemistry, ³ we envisioned a tandem reaction to form C–H, C–C, and Si–O bonds in one step to generate alkenyl and alkynyl silyl ethers of stereochemically defined vinyl silanes (Eq 1). ⁴ Overall, this is a net addition of H–OR to the alkynyl allyl silanes accompanied by an allyl transfer from silicon to carbon. ⁵ To achieve this tandem bond formation efficiently, we propose to use carbophilic metal catalysts instead of strong mineral acid catalysts to promote the allyl transfer and the addition of H–OR to carbon and silicon centers. Herein we report a stereoselective intramolecular allylation of silyl alkynes to generate alkoxy vinyl silanes via a gold-catalyzed alcoholysis of alkynyl allyl silanes. ⁶

$$R_1$$
 R_2 R_2 R_2 R_3 R_4 R_5 R_5 R_6 R_1 R_1 R_2 R_2 R_2 R_3 R_4 R_5 R_5 R_6 R_1 R_1 R_2 R_2 R_2 R_3 R_4 R_5 R_5 R_5 R_5 R_6 R_7 R_1 R_2 R_2 R_2 R_3 R_4 R_5 R_5 R_5 R_5 R_5 R_6 R_7 R_7

(1)

The general reactivity feature of allyl alkynyl silanes was examined with 1-octynyl allyl dimethylsilane ${\bf 1a}$ by using several metal catalysts under different conditions (Table 1). ⁸ The reaction of ${\bf 1a}$ with Ph₃PAuCl/AgSbF₆ in dry CH₂Cl₂ at room temperature provided a trace amount of ${\bf 2a}$ and ${\bf 3}$ (entry 1). ⁷ However, under otherwise identical conditions in undistilled CH₂Cl₂, desilylated product ${\bf 3}$ was obtained in 93% yield with small amount of ${\bf 2a}$ (entry 2). Reaction with Ph₃PAuCl/AgOTf gave no conversion (entry 3) whereas PtCl₂ (toluene, 90 °C) gave product ${\bf 2a}$ and ${\bf 3}$ in 20% yield (entry 5). ⁹ The extent of desilylation was reduced by replacing the dimethyl silyl with diphenyl silyl group in ${\bf 1b}$, which however, was recovered unchanged (entry 6) under the conditions (Ph₃PAuCl/AgSbF₆ in undistilled CH₂Cl₂) where ${\bf 1a}$ gave high conversion. On the other hand, the same reaction with added ⁱPrOH (1 equiv) provided silyl ether ${\bf 2b}$ in good yield (75%) (entry 7). This clearly indicates that the nucleophilic assistance is crucial for an efficient transfer of allyl group.

Having established optimized conditions for an intramolecular allyl transfer assisted by oxygen-based nucleophiles, we next examined the scope of this reaction by employing allyl silane 1c and a variety of alcohols (Table 2). Treatment of 1c with $Ph_3PAuCl/AgSbF_6$ in dry CH_2Cl_2 with 1° , 2° , and 3° alcohols gave good yield of products 4a-i as inseparable mixtures of Z/E- isomers in the range of 1:1.7 to 10:1 ratio. A salient feature of these reactions is that the alkene and alkyne functionalities in the alcohol counterpart do not interfere with the reaction.

To broaden the substrate scope, substituents on the alkyne and allyl moieties were introduced (Table 3). Reaction of **1b** with 4-penten-2-ol under the optimized conditions gave **5** in 71% yield with a 10:1 Z/E ratio (entry 1). Substrate 1d with terminal alkyne provided 6 in 85% yield, however, the stereochemistry of the double bond was scrambled, giving 1:1.7 mixture of Z/E- isomers (entry 2). This is, probably, the consequence of isomerization of the initially formed Z-isomer catalyzed by either the gold catalyst or a proton. ¹⁰ Substrates **1e**–**g** with methyl, phenyl, and benzyloxymethyl substituent on the alkyne behave uneventfully, giving 7-10 in good yields and Z/E-selectivity (entries 3-6). On the other hand, substrate 1h generated more 12 than the expected product 11 (81%, 12:11 = 3:1). Presumably, this is the consequence of more favorable activation of the allyl group by the gold catalyst due to the presence of sterically hindered t-butyl group on the alkyne. Substrates with alkyl substituent on the allyl segment showed variable reactivity depending on the position of the substituent compared to that of the parent system. 11 Thus, 1i bearing a crotyl group gave excellent yields and Z/Eselectivity of 13 and 14 (entries 8 and 9), whereas the reaction of methallyl-bearing substrate 1j provided 1:5.5 mixture of 15 and the methallyl-displaced product 16 in 89% overall yield (entry 10).

From a mechanistic standpoint, we surmised that the activation of the alkynyl moiety of $\bf 1$ by a carbophilic catalyst would induce 6-endo mode attack 12 over that of 5-exo by the pendant allyl silyl moiety to generate intermediate $\bf 17$ due to the β -silyl effect on the alkyne moiety (Scheme 1). In the subsequent step, carbocation $\bf 17$ would undergo a nucleophilic attack at the silicon center by an alcohol to give the final product $\bf 4$ – $\bf 11$ and $\bf 13$ – $\bf 15$ after protonolysis of the C-Au bond. Despite the sterically hindered environment around the silicon center, presumably, the formation of a strong Si–O bond is the driving force to form the observed products. Although the formation of a putative carbenoid $\bf 18$ followed by its alcoholysis is conceivable, products $\bf 19$ or $\bf 20$ were not observed. $\bf 13$ A direct alcoholysis of the allyl moiety of $\bf 1$ was observed when the allyl becomes a methallyl group, which is the consequence of preferential activation of the more electron rich methallyl group over the alkyne by the catalyst, thereby giving product $\bf 21$.

The utility of this tandem bond-forming technology was further expanded by the ring-closing metathesis 14 of alkoxy hydroallylation products (Scheme 2). Silyl ethers 4c and 4f/4g could

be cyclized by Grubbs complex 22^{15} to form 10- and 8-membered siloxanes 23-25 in good yields. Also, the cross metathesis 16 of 4i with 4-penten-1-ol provided the cross metathesis product 26 in 42% yield as a mixture of Z/E isomers.

In conclusion, we have developed a gold-catalyzed tandem intramolecular allyl transfer reactions induced by an alcoholysis of alkynyl allyl silanes, which generate alkoxy vinyl silanes ¹⁷ in high yield and *Z/E*-selectivity. Synthetic application of this tandem bond-forming process will be reported in due course.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgment

We thank NIH (RO1 CA106673) and the Sloan Foundation for financial support of this work as well as the NSF and NIH for NMR and Mass Spectrometry instrumentation.

References

- 1. Atom-economy: (a) Trost BM. Science 1991;254:1471. [PubMed: 1962206]. Step-economy: (b) Wender PA, Bi FC, Gamber GG, Gosselin F, Hubbard RD, Scanio MJC, Sun R, Williams TJ, Zhang L. Pure Appl. Chem 2002;74:25.
- 2. Ho, T-L. Tandem organic reactions. Wiley; New York: 1992. (b) Tietze LF. Chem. Rev 1996;96:115. [PubMed: 11848746]
- 3. (a) Miller RL, Maifeld SV, Lee D. Org. Lett 2004;6:2773. [PubMed: 15281766] (b) Park S, Kim M, Lee D. J. Am. Chem. Soc 2005;127:9410. [PubMed: 15984868]
- 4. For an intermolecular allyl silylation: Yoshikawa E, Gevorgyan V, Asao N, Yamamoto Y. J. Am. Chem. Soc 1997;119:6781.
- Silane alcoholysis during enyne cyclization: Fernández-Rivas C, Méndez M, Echavarren AM. J. Am. Chem. Soc 2005;127:12468. [PubMed: 16144376]
- Alcoholysis of C-Si bond to form silyl ethers: (a) Grimm JB, Lee D. J. Org. Chem 2004;69:8967.
 [PubMed: 15575786] Acid catalyzed alcoholysis of allylsilane: (b) Morita T, Okamoto Y, Sakurai H. Tetrahedron Lett 1980;21:835.
- 7. Gold catalyzed alkyne activation, see recent reviews: (a) Ma S, Yu S, Gu Z. Angew. Chem., Int. Ed 2006;45:200. (b) Hashimi ASK. Angew. Chem., Int. Ed 2005;44:6990. (c) Höffmann-Röder A, Krause N. Org. Biomol. Chem 2005;3:387. [PubMed: 15678171] (d) Hashimi ASK. Gold Bull 2004;37:51. (e) Dyker G. Angew. Chem., Int. Ed 2000;39:4237.
- 8. For the preparation of alkynyl allyl silanes, see Supporting Information
- Selected examples of most recent PtCl₂-catalyzed reaction of enynes: Fürstner A, Davies PW, Gress T. J. Am. Chem. Soc 2005;127:8244. [PubMed: 15941232] and references therein.
- 10. Another possibility of the scrambling is based on the ipso protonation followed by demetallation of the vinyl metal species.
- 11. The relative reactivity of allyl and substituted allyl groups in Lewis acid-catalyzed allyl silylation, see Ref 4.
- 12. A theoretical study for a 6-endo preference for 1,6-enyne possessing a terminal alkyne, see: Soriano E, Ballesteros P, Marco-Contelles J. Organometallics 2005;24:3182.
- 13. Reactions with all carbon-based system following 18 to 19 and 20, see: Nieto-Oberhuber C, Paz Munoz M, López S, Jiménez-Núnez E, Nevado C, Herrero-Gómez E, Raducan M, Echavarren AM. Chem. Eur. J 2006;12:1677. and references therein.
- 14. Reviews: (a) Grubbs RH, Chang S. Tetrahedron 1998;54:4413. (b) Fürstner A. Angew. Chem., Int. Ed 2000;39:3012. (c) Schrock RR, Hoveyda AH. Angew. Chem., Int. Ed 2003;42:4592. (d) GrubbsRHHandbook of Metathesis20032Wiley-VCHWeinheim
- 15. Scholl M, Ding S, Lee CW, Grubbs RH. Org. Lett 1999;1:953. [PubMed: 10823227]

16. For a review, see: Connon SJ, Blechert S. Angew. Chem., Int. Ed 2003;42:1900.

17. The use of alkoxy vinyl silanes in C–C bond formation, see: Denmark SE, Sweis RF. Acc. Chem. Res 2002;35:835. [PubMed: 12379136]

Scheme 1. Gold-Catalyzed Reorganization of Alkynyl Allyl Silane

Scheme 2. Ring-Closing and Cross Metathesis of Silyl Ethers

Table 1

I a Optimization of Catalyst and Reaction Conditions^a

try	enyne	R	catalyst/conditions	yield(2+3,%)	$2:3^c$
	1a	Me	Ph ₃ PAuCl/AgSbF ₆ , rt, dry CH ₂ Cl ₂	traces	2:1
			$Ph_3PAuCI/AgSbF_6$, rt, wet $CH_2CI_2^2d$	63°	1:20
			$Ph_3PAuCI/AgOTf$ rt, wet $CH_2CI_2^d$	traces	I
			AuCl ₂ rt, wet $CH_2CI_2^d$	traces	I
				20	2:1
	116	Ph	$Ph_2PAuCI/AgSbF_6$, rt, wet $CH_2CI_2^d$	traces	I
			Ph,PAuCl/AgSbF6, rt, CH,Cl, + PrOH	25	9:1

 a Reactions with 1 mol % catalyst for 5–10 h.

b Isolated yields.

 c The ratios were determined by 1 H NMR.

 $^d {\it Undistilled CH}_2{\it Cl}_2.$

 $\label{eq:Table 2} \mbox{Intramolecular Allylation-Alcoholysis Catalyzed by $\operatorname{Gold}^{a,b}$}$

 $[^]a$ Reactions with 1 mol % catalyst for 10 min.

 $b_{
m Isolated\ yields.}$

 $^{^{\}it c}$ The ratios of major:minor were determined by $^{\it 1}$ H NMR.

1 1 1b R = Hex (C ₆ H ₁₃) 2 1d R = H 3 1e R = Me 4 m m 1 m	ε-< ²	$\begin{array}{c} \begin{array}{c} \begin{array}{c} \mathbb{R}^{\mathbb{R}^n} \\ \end{array} \end{array}$ 5 R = Hex (C_6H_{13}) 6 R = H 7 R = Me	71 85 80 80 55	10:1 1:1.7 10:1 8:1
	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	5 $R = Hex (C_6H_{13})$ 6 $R = H$ 7 $R = Me$	71 85 80 55	10:1 1:1.7 10:1 8:1
	₹⊸	$\begin{array}{c} 6 \ \mathbf{R} = \mathbf{H} \\ 7 \ \mathbf{R} = \mathbf{M} \mathbf{e} \\ 7 \ \mathbf{e} \\ 9$	85 80 55	1:1.7
	₹⊸	7 R = Me	80 55	10:1 8:1
	₽	SI-O-ISI	55	8:1
	~			10.1
	w.	Ph C C C C C C C C C C C C C C C C C C C	89	10:1
	i	***************************************		
	5-<	Con Constitution of the Co	73	15:1
Leave the user	5_<	18 C 1 = 27.13) 10 C 1 = 27.13		
žő.	Ю	«√ £ø	86	11:1
9 olde 4	R A	13 R = Me 14 R = Abyl	85	10:1
	H_	8 8		
10 OMe 1	\	OM4 16 (16:16 = 1:5.5)		

 $^{\rm \it d}_{\rm \it Reactions}$ with 1 mol % catalyst for 10 min.

bIsolated yields.

 c The $Z\!/E$ ratios of were determined by 1 H NMR.

 d The initially formed isomers with the silyl and allyl groups in cis-orientation are defined Z-isomer.