Comparing -Carbanion-Stabilizing Ability of Substituents Using the Brook Rearrangement

Kei Takeda,^{*} Haruka Ubayama, Ayako Sano, Eiichi Yoshii, Toru Koizumi

Faculty of Pharmaceutical Sciences, Toyama Medical and Pharmaceutical University, 2630 Sugitani, Toyama 930-0194, Japan

Abstract

The -carbanion-stabilizing ability of the phenylthio and trimethylsilyl groups was compared based on the relative rate of the base-catalyzed Brook rearrangement of the -substituted -silylallylalcohol.

Keywords: Carbanions; Rearrangements; Silicon and Compounds; Substituent effects

In connection with our studies of the mechanism of the Brook rearrangement-mediated $[3 + 2]^{[1]}$ and $[3 + 4]^{[2]}$ annulations, we needed to determine the relative stability of -heteroatom-substituted carbanions in solution, particularly -trimethylsilyl and -phenylthio carbanions. Although both silicon and sulfur atoms are well recognized to stabilize -carbanions by negative hyperconjugation and/or (p-d) bonding, the difference in the degree of the stabilization by both atoms, especially in solution, has been much less known.^{[3],[4]} We envisaged that the relative stability of the -heteroatom-substituted carbanions would be roughly evaluated on the basis of the ratio of 1,2-adduct **2** to Brook rearrangement product **3** and Brook rearrangement/allylic rearrangement (B-A) product **4** in the reaction of -heteroatom-substituted acryloylsilane **1** with an appropriate nucleophile.



This stems from the fact that the Brook rearrangement,^[5] a 1,2-anionic shift of the silyl group from the carbon atom to the oxygen atom, is facilitated by electron-withdrawing

substituents. Thus, the ratio of **3** and **4** to **2** would increase as the -carbanion-stabilizing ability of the substituent X in **5**, a B-A intermediate, is more increased.

As the nucleophile, we selected the lithium enolate of *t*-butyl acetate **7** based on the preliminary experiments using the reaction of benzoyltrimethylsilane $6^{[6]}$ with the enolate in which the 1,2-adduct **8**, Brook rearrangement product **9**, and silanol elimination product of the rearrangement product **10** were obtained in an appropriate ratio.^[7]



The reaction was carried out using crotonoylsilane $1d^{[8]}$ and -phenylacryloylsilane $1c^{[8]}$ in addition to -trimethylsilyl and -phenylthio derivatives $1a, b^{[1],[9]}$ in THF (0.02 M) at -80 °C for 1 h and then quenched with acetic acid (1 equiv) to give both the 1,2-addition product and the B-A product except for 1d (Table 1). This observation that the reaction of the -phenyl derivative 1c affords both the 1,2-adduct and the B-A product in contrast to the reaction of the -methyl derivatives with only the formation of the 1,2-adduct 11d suggests that the system may be used for comparing -carbanion stabilization of the substituents. The phenylthio derivative 1a affords more of the rearrangement product 12 relative to the trimethylsilyl derivative 1b, indicating that the phenylthio group stabilizes the -carbanion more strongly than does the trimethylsilyl group. A troubling aspect of this system, however, was the lack of reproducibility; the ratio of 11 to 12 varied over a considerable range, particularly in the -phenylthio derivative, presumably because even subtle changes in experimental conditions affect the ratio.

Table 1



Next we turned our attention to comparison of the relative rate of base-catalyzed rearrangement of the 1,2-adduct **11** to **12**. Besides **11a-c**, the -sulfinyl, -stannyl, -chloro and -bromo derivatives were prepared by quenching of the reaction of **1** with **7** at low temperature by addition of acetic acid (Table 2). Table 2

x	O SiMe	e₂Bu ^t +	OLi OBu ^t	THF -85 °C	SiMe ₂ Bu ^t
		Х	time	yield (%)	
	11a	SPh	30 min	31	
	11b	SiMe ₃	60 min	33	
	11c	Ph	40 min	56	
	11e	S(O)Ph	10 min	4^b	
	11f	$SnBu_3$	60 min	50	
	11g	Cl^{a}	30 min	44	
	11h	Br	30 min	52	

a Z-isomer was used.

b Major product was the Brook rearrangement product.

Effective reaction conditions for the rearrangement were examined on **11a** and **11b** using several amine bases including diethylamine, diisopropylethylamine, pyridine, and DBU in $CDCl_3$ and d_6 -DMSO on monitoring by ¹H NMR.^[10] The reaction proceeded at a reasonable rate when using 0.2 equiv of DBU in d_6 -DMSO (50 mM) at 23 °C. Under these conditions, the half-lives of the reactions for **11** were measured (Table 3). Unfortunately, for the tributylstannyl derivative **11d**, the half-life could not be measured because of its low solubility in d_6 -DMSO under the conditions.

Table 3

HO X	OF SiMe ₂	Bu ^t DB d ₆ -DM Bu ^t	5U (0.2 eq) ▲ 1SO (50 mM) 23 °C	H OSiMe ₂ X O OB 13	₂ Bu ^t + H u ^t X	OSiMe ₂ Bu ¹ OOBu ¹
			product	t_{1}	/2 (min)	
	11a	SPh	12		3.2	
	11b	SiMe ₃	12 , 13 (ca	2:1) 2	27.5	
	11c	Ph	12		5.5	
	11e	S(O)Ph	12	<	<1.0	
	11g	Cl	13	11 : 13 = 2	2.2 : 1 (25	h)
	11h	Br	13	11:13=4	4.5 : 1 (43	h)

These results show that the -carbanion-stabilizing ability of the phenylthio group is considerably greater than that of the trimethylsilyl group, even than that for the phenyl group. Particularly noteworthy is the fact that, in the case of the trimethylsilyl derivative, the Brook rearrangement product 13 was formed in addition to the B-A product 12, sug-

gesting that -carbanion stabilization by the trimethylsilyl group is not much greater than that by the siloxy group, although in the latter case, stabilization by the -carbonyl group must be considered. In the case of phenyl sulfinyl derivative **11c**, the rearrangement occurred spontaneously. The reaction of -halogen derivatives reached equilibration with the B-A product **13**, not unexpectedly, indicating that the stabilization by the halogen groups was quite small.

In summary, we have demonstrated that comparison of the rate of Brook rearrangement in -substituted -silylallylalcohol **11** has the potential of becoming a tool for the assessment of the -carbanion-stabilizing ability of the -substituent in solution.

References and Notes

- (a) Takeda K, Fujisawa M, Makino T, Yoshii E, Yamaguchi K. J. Am. Chem. Soc. 1993;115:9351-9352.
 (b) Takeda K, Sakurama K, Yoshii E. Tetrahedron Lett. 1997;38:3257-3260.
- [2] Takeda K, Takeda M, Nakajima A, Yoshii E. J. Am. Chem. Soc. 1995;117:6400-6401. (b) Takeda K, Nakajima A, Yoshii E. Synlett;1996:753-754. (c) Takeda K, Nakajima A, Takeda M, Okamoto Y, Sato T, Yoshii E, Koizumi T, Shiro M. J. Am. Chem. Soc. 1998;120:in press.
- [3] For the silicon atom, see : (a) Bassindale AR, Taylor PG. Activating and directive effects of silicon. In: Patai S, Rappoport Z, editors. The Chemistry of Organic Silicon Compounds. New York: John Wiley & Sons, 1989:893-963. (b) Wetzl DM, Brauman JI. J. Am. Chem. Soc. 1988;110:8333-8336 and references cited therein. For the sulfur atom, see: (c) Wiberg KB, Catejon H. J. Am. Chem. Soc. 1994;116:10489-10497 and references cited therein. (d) Bernasconi CF, Kittredge KW. J. Org. Chem. 1998;63:1944-1953. Also, see: (e) Römer B, Gatev GG, Ahong M, Brauman JI. J. Am. Chem. Soc. 1988;120:2919-2924.
- [4] For pKa values for many types of acids, including one containing -heteroatoms, see: Bordwell FG. Acc. Chem. Res. 1988;21:456-463.
- [5] (a) For reviews of the Brook rearrangement, see: (a) Brook AG, Bassindale AR. Molecular rearrangements of organosilicon compounds. In: de Mayo P, editor. Rearrangements in Ground and Excited States. New York: Academic Press, 1980;Vol.2:149-221. (b) Brook AG. Acc. Chem. Res. 1974;7:77-84.
- [6] Picard JP, Calas R, Dunogues J, Duffaut N, Gerval J, Lapouyade P. J. Org. Chem. 1979;44:420-424.
- [7] Reaction of 6 with ketone enolates afforded 1,2-cyclopropanediol derivative ii by way of Brook rearrangement of the initial 1,2-adduct i and subsequent internal attack on the -carbonyl group. No 1,2-addition product was obtained, which could be explainable by acceleration of the Brook rearrangement by the more electrophilic ketone carbonyl group relative to the ester carbonyl group. see: Takeda K, Nakatani J, Nakamura H, Sako K, Yoshii E, Yamaguchi K. Synlett;1993:841-843.





- [8] Nowick JS, Danheiser RL. J. Org. Chem. 1989;54:2798-2802.
- [9] Reich HJ, Kelly MJ, Olson RE, Holtan RC. Tetrahedron 1983;39:949-960.
- [10] Brook and coworkers reported the rate of amine base-catalyzed rearrangement of a wide variety of -silylcarbinols. Brook AG, LeGrow GE, MacRae DM. Can. J. Chem. 1967;45:239-253.