

Crucial role of the ligand of silyl Lewis acid in the Mukaiyama aldol reaction†

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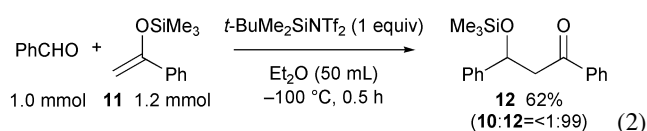
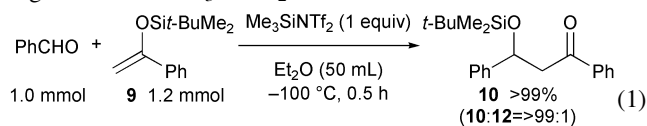
The Me_3SiX -induced Mukaiyama aldol reaction proceeds through each catalytic cycle under the influence of X^- : the silyl group of $\text{Me}_3\text{SiNTf}_2$ does not release from $-\text{NTf}_2$ and that of silyl enol ether intermolecularly transfers to the product, while the silyl group of Me_3SiOTf remains in the product and that of the silyl enol ether becomes the catalyst for the next catalytic cycle.

Ghosez^{1a} and Mikami^{1b} have independently introduced $\text{Me}_3\text{SiNTf}_2$ as a much more powerful carbonyl-activating reagent than Me_3SiOTf . Very recently, we demonstrated the extremely high activity of $\text{Me}_3\text{SiNTf}_2$ as a catalyst for the Mukaiyama aldol and Sakurai–Hosomi allylation reactions of not only aldehydes but also ketones.^{2,3} We report here that the Me_3SiX -induced Mukaiyama aldol reaction proceeds through each catalytic cycle under the influence of its ligand (X): specifically, there is a significant difference between $-\text{NTf}_2$ and $-\text{OTf}$, in that the silyl group of $\text{Me}_3\text{SiNTf}_2$ does not release from its ligand and that of silyl enol ether intermolecularly transfers to the product, while the silyl group of Me_3SiOTf remains in the product and that of the silyl enol ether becomes the catalyst for the next catalytic cycle.⁴

The three plausible reaction pathways for the Me_3SiX -induced Mukaiyama aldol reaction are outlined in Fig. 1.⁵ In the first stage of catalysis, the addition of a trialkylsilyl enol ether **2** to a Me_3Si -activated aldehyde **1** generates a siloxocarbenium ion intermediate **3**. In *path A*, R_3SiX is released into the reaction medium to give the trimethylsilyl aldolate **4** by the intramolecular transfer of X^- .⁶ Thus, R_3SiX becomes the catalyst for the next catalytic cycle.⁷ In *path B*, Me_3SiX is released into the reaction medium to give the trialkylsilyl aldolate **5** by the intramolecular transfer of R_3Si^+ .^{6,8} In *path C*, co-ordinated aldehyde as in **6** or **7**, which is activated by penta- or tetra-coordination^{4,9} of R_3Si^+ , reacts with **2** through the intermolecular silyl transfer of R_3Si^+ . In this catalytic process, **5** may be obtained as a major product.¹⁰

First, the mechanistic details of the $\text{Me}_3\text{SiNTf}_2$ ^{1a,11}-induced reaction were investigated. To clarify whether transformation from **2** to a silyl aldolate occurs through $-\text{NTf}_2$ transfer (*path A*) or $^+\text{SiR}_3$ transfer (*path B* or *C*), the reaction of *tert*-butyldimethylsilyl enol ether **9** derived from acetophenone with benzaldehyde was carried out in the presence of 1 equiv. of $\text{Me}_3\text{SiNTf}_2$ at -100°C for 0.5 h. Regardless of the order of the addition of substrates and $\text{Me}_3\text{SiNTf}_2$, only *tert*-butyldimethylsilyl aldolate **10** was obtained in high yield [eqn. (1)]. In contrast, the reaction of trimethylsilyl enol ether **11** with benzaldehyde in the presence of 1 equiv. of *t*- $\text{BuMe}_2\text{SiNTf}_2$ gave only trimethylsilyl aldolate **12** in moderate yield [eqn. (2)]. Control experiments for eqn. (1) showed that neither **9** and $\text{Me}_3\text{SiNTf}_2$ nor **12** and *t*- $\text{BuMe}_2\text{SiNTf}_2$ exchanged under similar conditions. These experimental results exclude *path A*. Although *path C* is also unlikely since **12** was not detected by GC analysis [eqn. (1)], we can not exclude *path C* because the equilibration between benzaldehyde and **1** may be quite fast, and the aldol reaction induced by *tert*-butyldimethylsilo-

ocarbenium ions **3** and **8** could be much faster than that induced by $\text{Me}_3\text{SiNTf}_2$. In fact, $\text{Me}_3\text{SiNTf}_2$ is an extraordinarily active catalyst,^{1,2} and its high turnover frequency can be reasonably explained by a $^+\text{SiR}_3$ -induced cascade process which avoids the regeneration of $\text{Me}_3\text{SiNTf}_2$.



For a crossover experiment,⁴ two silyl enol ethers **9** and **13** with comparable steric and electronic properties were combined with benzaldehyde [eqn. (3)]. Benzaldehyde was added to a solution of a 1 : 1 mixture of **9** and **13** (0.6 equiv. of each) in the presence of 1 mol% of $\text{Me}_3\text{SiNTf}_2$ at -78°C . The product composition was determined by GC analysis. A mixture of

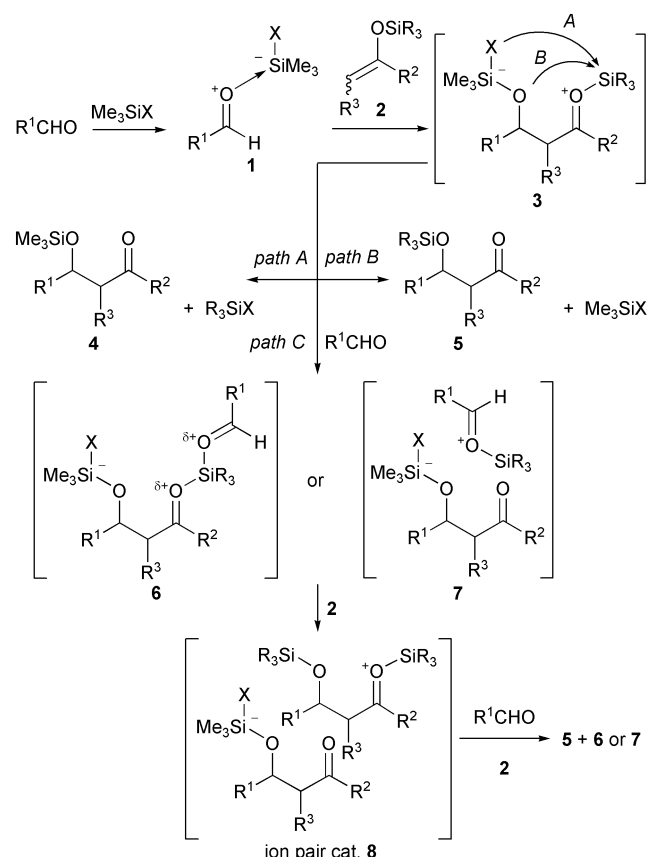
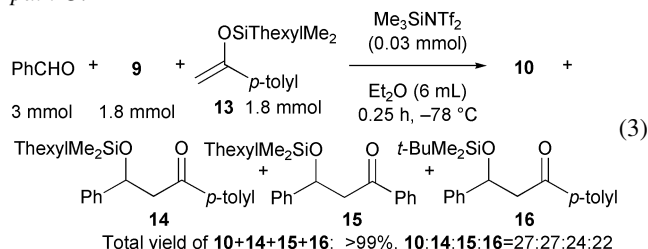


Fig. 1 An outline of the three possible mechanisms for the Mukaiyama aldol reaction induced by Me_3SiX .

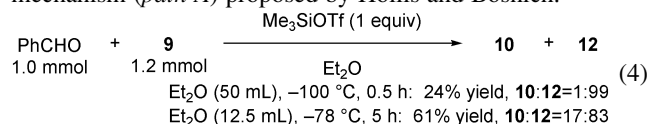
† Electronic supplementary information (ESI) available: experimental section. See <http://www.rsc.org/suppdata/cc/b2/b203838b/>

scrambled silyl ethers was isolated in a ratio of 27:27:24:22 (**10**:**14**:**15**:**16**). Control experiments showed that neither the starting silyl enol ethers nor the silylated aldol products exchanged under similar reaction conditions. Based on these observations, *path B* was unambiguously precluded. Therefore, it is likely that the Me₃SiNTf₂-induced reaction occurs through *path C*.



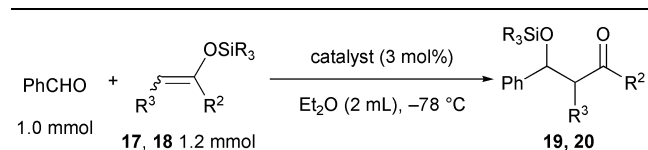
The *syn/anti* selectivities in the aldol reactions of (*E*)- and (*Z*)-silyl enol ethers induced by silyltriflylimides were investigated (Table 1). Similar *syn/anti* ratios of silyl aldolates, **19** and **20**, were obtained independent of the trialkylsilyl group of the silyltriflylimides. On the other hand, the ratios were correlative to the trialkylsilyl group of silyl enol ethers. These results also support *path C*.

Next, we performed a mechanistic study of the Me_3SiOTf -induced Mukaiyama aldol reaction.¹² Reactions analogous to eqn. (1) were conducted using Me_3SiOTf [eqn. (4)]. A mixture of **10** and **12** was obtained in 24% yield at a molar ratio of 1:99 under the same conditions with eqn. (1). The starting materials were not completely consumed upon prolonged stirring at -78°C with a concentration three-fold higher than that in eqn. (1), and a mixture of **10** and **12** was obtained in 61% yield at a molar ratio of 17:83 after 5 h at -78°C . The reaction of **9** with benzaldehyde did not proceed at -100°C in the presence of *t*-BuMe₂SiOTf in a control experiment for eqn. (4). Further control experiments for eqn. (4) showed that neither **9** and Me_3SiOTf nor **10** and Me_3SiOTf exchanged under similar conditions. These experimental results strongly support the mechanism (*path A*) proposed by Hollis and Bosnich.^{4d}

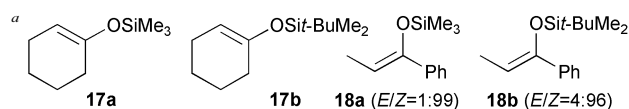


The present observations indicate that the ligand of silyl Lewis acid plays a crucial role in the aldol reaction. The transfer

Table 1 *Syn/anti* selectivity in the Mukaiyama aldol reaction



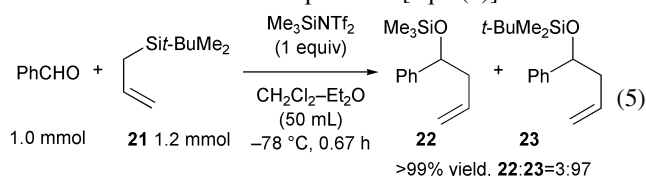
		Silyl aldolates	
Silyl enol ether ^a	Catalyst	Yield (%)	<i>Syn:Anti</i>
17a	Me ₃ SiNTf ₂	19a , 97	60:40
17a	<i>t</i> -BuMe ₂ SiNTf ₂	19a , > 99	62:38
17b	Me ₃ SiNTf ₂	19b , > 99	52:48
17b	<i>t</i> -BuMe ₂ SiNTf ₂	19b , > 99	51:49
18a	Me ₃ SiNTf ₂	20a , 90 (1) ^b [9] ^c	61:39 (48:52) ^b
18a	<i>t</i> -BuMe ₂ SiNTf ₂	20a , 89 (2) ^b [12] ^c	58:42 (45:55) ^b
18b	Me ₃ SiNTf ₂	20b , > 99	87:13
18b	<i>t</i> -BuMe ₂ SiNTf ₂	20b , > 99	87:13



^b Data of alcohol adducts are given in parentheses. ^c Yields of dimeric ethers of alcohols are given in brackets.²

of X- from **3** is expected to occur by electrophilic attack of the 'R₃Si-O⁺ silicon' of **3**. In the Me₃SiOTf-induced reaction, R₃SiOTf would be generated by electrophilic attack of the 'R₃Si-O⁺ silicon' to the 'S=O oxygens' or the 'S-O oxygen' of -OTf (*path A*). In the Me₃SiNTf₂-induced reaction, on the contrary, less nucleophilicity¹⁴ and/or more bulkiness of -NTf₂ may suppress the electrophilic attack of the 'R₃Si-O⁺ silicon' to the nitrogen¹¹ or oxygen atoms¹³ of -NTf₂, and may increase Lewis acidity of siloxocarbenium ions **3** and **8** (*path C*).

Lastly, Sakurai-Hosomi allylation reaction induced by $\text{Me}_3\text{SiNTf}_2$ also occurs through $+\text{SiR}_3$ transfer, according to results of a stoichiometric experiment [eqn. (5)].²



These findings may provide a basis for the future development of not only chiral silyl Lewis acid catalysts but also other chiral metal catalysts for carbon–carbon bond-forming reactions of silyl nucleophiles with carbonyl compounds.¹⁵

Notes and references

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