Bis(difluoromethyl)trimethylsilicate Anion: A Key Intermediate in Nucleophilic Difluoromethylation of Enolizable Ketones with Me₃SiCF₂H

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Dedicated to Professor K. Barry Sharpless on the occasion of his 75th birthday

Abstract: A pentacoordinate bis(difluoromethyl)silicate anion, [Me₅Si(CF₂H)₃]⁻, is observed for the first time by the activation of Me₃SiCF₂H with a nucleophilic alkali-metal salt and 18-crown-6. Further study on its reactivity by tuning the countercation effect led to the discovery and development of an efficient, catalytic nucleophilic difluoromethylation of enolizable ketones with Me₃SiCF₂H by using a combination of CsF and 18-crown-6 as the initiation system. Mechanistic investigations demonstrate that [(18-crown-6)Cs⁺][Me₅Si(CF₂H)₃]⁻ is a key intermediate in this catalytic reaction.

Hypercoordinate silicates are key intermediates in nucleophilic substitution on silicon and activation of organosilicon compounds. Many pentacoordinate silicates containing at least one electronegative heteroatom ligand such as F, Cl, O, and N have been prepared to study their structures and reactivities. However, pentaorganosilicate species containing five Si–C bonds are rare, and little is known about their carbon–ligand transfer reactivity. In 1999, the groups of Naumann and Röschenthaler independently reported the preparation and characterization of the pentaorganosilicate anion [Me₅Si(CF₂H)₃]⁻, which is derived from the interaction of a pentacoordinate [Me₅Si(F)]⁺ with the Ruppert–Prakash reagent (Me₅SiCF₃). The formation of such an acyclic pentaorganosilicate is attributed to the pronounced group electronegativity of the trifluoromethyl unit, which renders Me₅SiCF₃ of sufficient Lewis acidity to accept an incoming trifluoromethanide anion (CF₃⁻). In 2014, Prakash and co-workers elegantly demonstrated that the pentacoordinate [Me₅Si(CF₂H)₃]⁻ prepared from a bulky tert-butoxy anion with a [K(18-crown-6)]⁺ countercation and Me₅SiCF₃ can dissociate to give a CF₃⁻. They observed the CF₃⁻ species by NMR spectroscopy and ascertained its reactivity in nucleophilic trifluoromethylation.[5b]

Me₅SiCF₃H, as an analogue of Me₅SiCF₃, has recently emerged as a potentially useful difluoromethanide anion (HCF₂⁻) source for the nucleophilic introduction of a difluoromethyl group.[7–9] However, because of the weaker electron-withdrawing ability of the CF₂H group (compared with CF₃ group),[10] the reactivity of Me₅SiCF₂H is distinct from the well-developed Me₅SiCF₃ in nucleophilic fluoroalkylation reactions. The trifluoromethylation with Me₅SiCF₃ readily takes place under the activation of a wide range of Lewis bases,[11] whereas the similar difluoromethylation normally requires harsher reaction conditions, thus limiting the substrate scope.[7–9] To address these challenges in difluoromethylation with Me₅SiCF₂H, it is essential to investigate the hypervalent silicon intermediate in the nucleophilic activation of Me₅SiCF₂H and to probe its difluoromethyl-transfer reactivity. Herein, we describe the discovery and characterization of a unique pentacoordinate bis(difluoromethyl)silicate anion [Me₅Si(CF₂H)₃]⁻ derived from Me₅SiCF₂H and the tuning of its reactivity with the countercation [(18-crown-6)Cs⁺], as well as its application in the efficient difluoromethylation of enolizable ketones, a reaction which was previously difficult to achieve.[7–9]

We began our study with an investigation on the interaction between Me₅SiCF₂H and various nucleophilic activators in THF without adding the carbonyl substrate (Scheme 1a). Based on our previous report on the difluoromethylation of various aldehydes, diaryl ketones, and imines

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Scheme 1. Activation of Me₅SiCF₂H with various nucleophiles and observation of the pentacoordinate difluoromethylsilicate. THF = tetrahydrofuran.

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with Me$_3$SiCF$_3$H$_2$ stoichiometric amounts of CsF and tBuOK were initially tested as the activators. However, only the signals of CF$_3$H$_2$ and unreacted Me$_3$SiCF$_3$H were detected by $^{19}$F NMR spectroscopy at a wide range of temperatures ($-70^\circ$C to room temperature). The failure to observe any pentacoordinate difluoromethyl silicate species probably arises from the strong affinity of the HCF$_2$- to the alkali-metal cations, and thus leads to a spontaneous decomposition of the difluoromethyl silicates into HCF$_2$-, with subsequent protonation by either adventitious water or the solvent THF.

Considering that the countercation effect can significantly influence the stability of hypercoordinate silicate anions,[13,15] we turned our attention to the employment of a crown ether[12] as an additive to stabilize the silicate intermediate by minimizing the interaction between HCF$_2$- and the alkali metal cations (Scheme 1b). To our delight, a weak signal at around $\delta = \sim 130.0$ ppm was observed by $^{19}$F NMR spectroscopy at room temperature using the combination of CsF/18-crown-6 as the activator (see Figure S1 in the Supporting Information). Much stronger signals with the similar chemical shifts were observed at temperatures ranging from $-78^\circ$C to room temperature when the combinations of more soluble tBuOK/18-crown-6, tBuOCs/18-crown-6, and even tBuONa/18-crown-6 were used. Compared with the $^{19}$F NMR chemical shift of Me$_3$SiCF$_3$H ($\delta = \sim 140.7$ ppm), the downfield shift of the observed signal is likely to correspond to an anionic species.[13] To determine the structure of the observed species, we further carried out $^{29}$Si$^1$H NMR, $^{13}$C$^1$[H] NMR, $^1$H NMR, and heteronuclear multiple quantum coherence (HMQC) experiments of the reaction between Me$_3$SiCF$_3$H and tBuOCs/18-crown-6 in [D$_3$]THF (Figure 1; see Figure S2–S6 and Table S1).[14] In the $^{29}$Si$^1$H NMR spectrum, the signal is shifted upfield ($\delta = \sim 118.4$ ppm (quint, $J_{\text{Si-F}} = \sim 10.3$ Hz)) compared with that of Me$_3$SiCF$_3$H ($\delta = \sim 0.04$ ppm (t, $J_{\text{Si-F}} = \sim 28.8$ Hz)), and is in accordance with the formation of a pentacoordinated silicate species.[15,16] This chemical shift change is similar to that of the reported [Me$_3$Si(CF$_3$)$_2$] $^+$ ($\delta = \sim 112.1$ ppm) versus Me$_3$SiCF$_3$ $^+$ ($\delta = \sim 5.4$ ppm).[54] In the $^1$H NMR spectrum, both the signals of the CF$_3$H ($\delta = \sim 5.00$ ppm) and Me ($\delta = \sim 0.24$ ppm) groups are also detected to be upfield compared with those of Me$_3$SiCF$_3$H ($\delta = \sim 5.85$ ppm; $\delta$(Me) = 0.14 ppm). The $^1$J$_{\text{HF}}$ coupling constant is in good agreement with the corresponding coupling observed in $^{19}$F NMR ($\delta = \sim 130.0$ ppm (d, $J_{\text{HF}} = \sim 47.3$ Hz)). According to the ratio (2:9) of integrated area of the $^1$H NMR signals of CF$_3$H and Me, we inferred that the above silicon species is a pentacoordinated bis(difluoromethyl)silicate with five Si–C bonds, that is, $\left(18\text{-crown-6}\right)$[Me$_3$Si(CF$_3$)$_2$] $^+$.[16] Moreover, the $^{13}$C NMR signals of CF$_3$H ($\delta = 140.3$ ppm (t, $J_{\text{CF}} = \sim 285$ Hz)) and Me ($\delta = \sim 3.2$ ppm), which were assigned by $^{13}$C$^1$C HMQC analysis, appear downfield from those of Me$_3$SiCF$_3$H ($\delta$(CF$_3$H) = 123.7 ppm (t, $J_{\text{CF}} = \sim 253$ Hz); $\delta$(Me) = $\sim 6.5$ ppm). An increase in $^1$J$_{\text{CH}}$ coupling as well as a decrease in $^1$J$_{\text{CF}}$ coupling, which have been observed in the transformation of Me$_3$SiF$_3$ into [Me$_3$Si(CF$_3$)$_2$] $^+$,[55] are probably characters of bipyramidal [R$_3$Si(R$_3$)$_2$]$^+$ species with two R$_3$ groups at the axial positions.

It is noteworthy that these pentacoordinate silicates were relatively stable below $\sim 30^\circ$C and decomposed gradually to CF$_3$H$_2$ when slowly raising the temperature from $\sim 30^\circ$C to $20^\circ$C [for variable-temperature (VT) NMR study, see Figures S7 and S17]. Moreover, [Me$_3$Si(CF$_3$)$_2$] $^+$ was the only detectable difluoromethylated hypercoordinate silicon species regardless of the Me$_3$SiCF$_3$H/BuOM (either 1:1 or not). We also attempted to observe CF$_3$H$^-$ at a wide range of temperatures (from $\sim 78^\circ$C to $20^\circ$C), but no evidence supported the persistence of this species in our system, which is significantly different from the CF$_3$- anion derived from Me$_3$SiCF$_3$.[56] The generation of CF$_3$HD and CF$_3$H$_2$ as side products when using [D$_3$]THF as the solvent indicates that CF$_3$H$^-$ is kinetically unstable and has a high tendency to abstract a proton from both THF and 18-crown-6 (see Figures S3 and S13).

Having identified the relatively stable intermediate [Me$_3$Si(CF$_3$)$_2$] $^+$, we next sought to probe its reactivity in nucleophilic difluoromethylation reactions. The experiment was conducted by adding the electrophilic substrate to a THF solution of [Me$_3$Si(CF$_3$)$_2$] $^+$ pre-generated from stoichiometric amounts of tBuOM (M = Na, K, Cs), 18-crown-6, and Me$_3$SiCF$_3$H in a molar ratio of 1:1:2. To our surprise, enolizable ketones, which are challenging substrates under previously reported difluoromethylation conditions,[57,58] were readily difluoromethylated. Thus, the reaction between 1-(2-methoxyphenyl)ethanone (Ia) and [Me$_3$Si(CF$_3$)$_2$] $^+$, generated from tBuOCs/18-crown-6/Me$_3$SiCF$_3$H, gave the corresponding difluoromethylated alcohol in good yield (Scheme 2a; for details see Figure S8). However, when tBuONa was used instead of tBuOCs, the pre-generated [Me$_3$Si(CF$_3$)$_2$] $^+$ failed to undergo addition to Ia, only affording CF$_3$H$_2$ as the detectable side product. Based on these results, we concluded that the countercation effect not only influences the stabilization of [Me$_3$Si(CF$_3$)$_2$] $^+$, but it also dramat-
ically alters the reactivity of \([\text{Me}_3\text{Si(CF}_2\text{H)}_2]^@\) towards enolizable ketones.

Inspired by the observed excellent reactivity of \([\text{(18-crown-6)}\text{Cs}]+[\text{Me}_3\text{Si(CF}_2\text{H)}_2]^@\) towards the ketone \(1i\), we decided to develop a catalytic difluoromethylation method for synthesizing tertiary carbinols from \(\text{Me}_3\text{SiCF}_2\text{H}\) and the challenging enolizable ketones by employing the impressive countercation effect.\(^{[17]}\)

Thus, substoichiometric amounts of \(\text{tBuOM (M = Na, K, Cs)}/18\)-crown-6 were used as an initiator to investigate the reaction between \(1i\) and \(\text{Me}_3\text{SiCF}_2\text{H}\). As shown in Scheme 2b, a remarkable metal-ion effect was also found in this catalytic reaction and \([\text{(18-crown-6)}\text{Cs}]+\) again proved to be the most effective countercation.

To determine the role of \([\text{Me}_3\text{Si(CF}_2\text{H)}_2]^@\) in the catalytic cycle, we investigated the progress of the \((\text{tBuOCs}/18\)-crown-6\()-initiated reaction by VT \(^{19}\)F NMR experiments. The experiment was carried out by adding a mixture of \(1i\) (1 equiv) and \(\text{Me}_3\text{SiCF}_2\text{H}\) (2 equiv) to a THF solution of \(\text{tBuOCs}\) (20 mol%) and 18-crown-6 (20 mol%) at \(-70^\circ\text{C}\). As the reaction temperature gradually rose, the intermediate \([\text{Me}_3\text{Si(CF}_2\text{H)}_2]^@\) was observed first (\(-70^\circ\text{C}\) to \(-20^\circ\text{C}\)), followed by the observation of the difluoromethylation product (\(-20^\circ\text{C}\)). Interestingly, the silicate intermediate maintained a certain concentration even when the reaction temperature was elevated to \(5^\circ\text{C}\) (see Figures S10 and S11), thus indicating a continuous consumption and regeneration of \([\text{Me}_3\text{Si(CF}_2\text{H)}_2]^@\) during the reaction. In contrast, when \(\text{tBuONa}\) was used instead of \(\text{tBuOCs}\), although the formation of \([\text{Me}_3\text{Si(CF}_2\text{H)}_2]^@\) was observed first, it could not difluoromethylate \(1i\) to initiate the reaction. According to these results, we conclude that the cesium ion played an important role in \((\text{tBuOCs}/18\)-crown-6\()-initiated difluoromethylation of enolizable ketones.

Since \([\text{Me}_3\text{Si(CF}_2\text{H)}_2]^@\) can also be generated from the combination of CsF/18-crown-6, albeit in a low yield, we explored the difluoromethylation by employing the more readily available CsF instead of \(\text{tBuOCs}\) (Scheme 2c; for details see Table S1). We were pleased to find that reaction employing 10 mol% of CsF/18-crown-6 also provided a high yield of the difluoromethyl addition product. A screening of the solvent showed that the ether solvent dimethoxyethane (DME) is slightly superior to THF in improving the yield. Therefore, the combination of CsF/18-crown-6 was chosen as the optimal initiation system and DME was chosen as the optimal solvent for the reaction between \(\text{Me}_3\text{SiCF}_2\text{H}\) and enolizable ketones.

With the optimized reaction conditions in hand, we subsequently investigated the substrate scope. As shown in Scheme 3, most of the enolizable ketones examined provided good to excellent yields. In general, substituted aromatic ketones bearing electron-donating groups (such as Ph, \(\text{iPr}\), OMe, and NMe\(_2\)) showed higher reactivity (2b–f) than those

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**Scheme 2.** Nucleophilic difluoromethylation of enolizable ketones. (a) Yield was determined by \(^{19}\)F NMR spectroscopy. TBAF = tert-n-butylammonium fluoride.

**Scheme 3.** Direct nucleophilic difluoromethylation of various carbonyl compounds with \(\text{Me}_3\text{SiCF}_2\text{H}\). Reaction conditions (unless otherwise noted): \(\text{Me}_3\text{SiCF}_2\text{H}\) (1.2 mmol), \(1\) (0.6 mmol), CsF (10 mol%), 18-crown-6 (10 mol%), DME (3 mL). Yields of isolated products are reported. (a) CsF (100 mol%), 18-crown-6 (100 mol%). (b) CsF (20 mol%), 18-crown-6 (20 mol%). (c) Yield was determined by \(^{19}\)F NMR spectroscopy.
with electron-withdrawing groups (such as Br, Cl, COOME, and NO₂; 2g–k). 1-Phenylalkyl-1-ones containing alkyl groups of different chain lengths gave difluoromethylated products in good yields (2o–p). The CsF/18-crown-6/DME system was also suitable for difluoromethylation of aliphatic ketones (2s–v). Particularly, the readily enolizable 1,3-diphenylpropan-2-one and steroid could also react with Me₃SiCF₂H to afford the products 2t and 2v in 65 and 53% yields, respectively. When conjugated methyl ketone (E)-4-phenylbut-3-en-2-one was subjected to the reaction, the alcohol 2r resulting from the carbonyl addition was obtained as the sole product. To demonstrate both the potential pharmaceutical relevance and the functional-group tolerance of the present direct difluoromethylation protocol, we applied it in the synthesis of the compound (±)-2w, a potential antagonist of the orexin receptor. It was previously prepared using a two-step method: nucleophilic (phosphoryl) difluoromethylation of the corresponding ketone followed by removal of the phosphate group.[18] In addition to enolizable ketones, other carbonyl compounds including diaryl ketones, aromatic aldehyde, enolizable aliphatic aldehyde, phthalimide, and phthalide could also be difluoromethylated by Me₃SiCF₂H under the similar reaction conditions, thus affording the products 2x–ab in 37–90% yields.

Finally, based on our investigation, a mechanism involving [(18-crown-6)Cs][Me₃Si(CF₂H)]⁻ (A) as a key intermediate was proposed. As is shown in Scheme 4, the process commences with the initial generation of [Me₃Si(CF₂H)]⁻ from a catalytic amount CsF (or iBuOCSi)–18-crown-6 and Me₃SiCF₂H. The complexation of 18-crown-6 with Cs⁺ inhibits the formation of the strongly basic, free difluoromethaneide, thus stabilizing [Me₃Si(CF₂H)]⁻ and favoring the carbonyl addition rather than the enolation. The subsequent reaction between A and carbonyl substrate leads to the formation of the alcoholate B, which continues to attack the silicon atom of Me₃SiCF₂H to produce a new pentacoordinate silicate C. This step is followed by the transfer of a difluoromethanide anion to Me₃SiCF₂H to release the target product and to regenerate A. A will be constantly generated until all of the carbonyl substrate is consumed. In view of the fact that A is formed much faster than the carbonyl addition product, another pathway involving the difluoromethanide anion addition to the carbonyl group at the initiation stage (Scheme 4, dashed arrow), which is similar to the commonly assumed trifluoromethylation of carbonyl compounds with Me₃SiF,[11] is less likely to occur as a major pathway under our conditions.

In summary, [Me₃Si(CF₂H)]⁻, a pentacoordinate difluoromethylsilicate anion with five Si–C bonds, was observed for the first time through the activation of Me₃SiCF₂H with a nucleophilic alkali-metal salt and 18-crown-6. It is found that the countercation effect plays important roles in both stabilizing the [Me₃Si(CF₂H)]⁻ intermediate and improving its nucleophilic difluoromethylation potency. By employing the combination of a cesium salt and 18-crown-6 as the initiator, catalytic difluoromethylation of enolizable ketones was achieved in high yields because of the avoidance of the competitive enolation, which is usually encountered when using other initiators. During the whole reaction, [Me₃Si(CF₂H)]⁻ is not only the difluoromethaneide anion source, but also acts as a difluoromethaneide reservoir. The formation of the bis(difluoromethyl)trimethylsilicate intermediate with [(18-crown-6)Cs]⁺ as countercation alleviates the strong basicity of a difluoromethaneide, which is of great significance for the catalyzed difluoromethylation of enolizable ketones.

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During our preparation of this manuscript, a direct nucleophilic difluoromethylation of enolizable ketones with MeSiCF₂H/3BuONa/buBuONa/18-crown-6, see the Supporting Information.

For example, the 29Si NMR signal of lithium 2,2-biphenyldiyltrimethylsilicate appears at δ = −116.9 ppm. See Ref. [3c].


During our preparation of this manuscript, a direct nucleophilic difluoromethylation of enolizable ketones with MeSiCF₂H/3CF/HMPA was developed by Radchenko and co-workers. However, the yields are only moderate (see Ref. [7f]). In addition, both group and that of Radchenko (see Ref. [7f]) were not able to reproduce Tyutyunov's results of efficient difluoromethylation of enolizable ketones with MeSiCF₂H (see Ref. [7c]).