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A facile access to a novel NHC-stabilized silyliumylidene ion and C–H activation of phenylacetylene

A facile one-pot synthetic route for the synthesis of donor stabilized silyliumylidene ions has been devised. Additionally, an unusual C–H insertion product was isolated from the reaction of NHC-stabilized arenylsilyliumylidene ion with three equivalents of phenylacetylene.

A facile access to a novel NHC-stabilized silyliumylidene ion and C–H activation of phenylacetylene†

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Taking advantage of two N-heterocyclic carbenes (NHCs), novel silyliumylidene ions 1a and 1b are prepared by a facile one-pot reaction of the corresponding dichlorosilanes with three equivalents of NHCs. For the first time, a C–H insertion reaction of phenylacetylene by a novel silyliumylidene ion is reported. The treatment of m-terphenyl substituted silyliumylidene ion 1a with three equivalents of phenylacetylene results in the formation of m-terphenyl substituted 1-alkeny1-1,1-dialkynylsilane 2.

Silylium ions [R₃Si⁺], heavier analogues of carbenium ions, are among the strongest Lewis acids. It took a deliberate effort of over half a century towards their successful isolation.1 In general, the factors like specially designed non-coordinating counter anions, donor free solvents and kinetically stabilized bulky substituents were crucial towards the isolation of free silylium ions.2 The trivalent silicon centre in silylium ions is highly electrophilic and now Lewis acid catalysis as well as C–F bond activation are the most prominent applications of silylium ions.3 On the other hand, silylenes [R₂Si:], heavier analogues of carbenes, have attracted much attention in the past 20 years and show interesting reactivities and potential applications in transition metal catalysis.5

Meanwhile, silyliumylidene ions [RSi:+] bearing the best combined character of both silylium ions and silylenes. For example, the electrophilicity is more pronounced since the silicon centre possesses four valence electrons, two vacant orbitals and a lone pair of electrons. Consequently the isolation of silyliumylidene ions gets even more challenging. Several silyliumylidene cations, however, have been reported in the past ten years (Chart 1).6–11 This has been achieved either by employing well-designed ligands or multi-step synthetic methods. For example, the seminal work in this field is reported by Jutzi for silyliumylidene ion I, thanks to the stabilization effect of the pentamethylcyclopentadienyl ligand.6 Driess and co-workers utilized the intramolecular stabilization effect for the isolation of the silicon(II) cation II7 which also shows aromatic stabilization. The same group later on, reported on the synthesis of III8 and IV9 by the incorporation of especially designed bisiminophosphorane and bis N-heterocyclic carbene ligands, respectively. Moreover, So and coworkers reported on the synthesis of silyliumylidene ion V stabilized by DMAP and the amidinate ligand.10 It should also be mentioned that Filippou and co-workers reported on the striking example of silyliumylidene VI stabilized by two NHCs in a three step synthetic methodology starting from SiI₄.11a

Despite their interesting character, the reactivity study of silyliumylidenes is still in its infancy.12 For example, the catalytic behavior of I in the controlled degradation of ether,13 the remarkable synthesis of a silylone from IV9 and activation of elemental sulfur from III10 and V10 remain as the highlights of the reactivity of silyliumylide ions. Herein we report the C–H insertion in phenylacetylene using a novel silyliumylidene ion synthesized in a facile single step methodology.

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The computational studies performed by Müller suggested a stable silylimidylenide ion substituted by the kinetically stabilizing terphenyl group at the silicon-center. This prompted us to employ the terphenyl substituents for the synthesis of a novel silylimidylenide ion. On the other hand, N-heterocyclic carbenes (NHCs) are employed as external donors to provide the desired stabilization for the low-valent silicon compounds. Thus, we employed two NHCs parallel to the methodology used by Filippou towards the isolation of NHC-stabilized chlorosilylene. Interestingly, we have found that the target m-terphenyl substituted silylimidylenide ion \( \text{1a} \) stabilized by two NHCs can be obtained in a very facile one step experimental procedure (Scheme 1). In addition, we have also employed the trisopropylphenyl substituent at the silicon centre in order to generalize this convenient synthesis (Scheme 1). The synthetic methodology involves the addition of a solution of three equivalents of Me_4NHC (Me_4NHC = 1,3,4,5-tetramethylimidazol-2-ylidene) to a solution of the corresponding dichlorosilane \( \text{1a} \) for \( m \)-Ter = 2,6-Mes_2C_6H_3, Mes = 2,4,6-trimethylphenyl, Tipp = 2,4,6-trisopropylphenyl). Compound \( \text{1a} \) is obtained as a bright yellow crystalline product in 45% yield. Compound \( \text{1b} \), however, is synthesized by slow addition of NHC to a heated solution of TippSiHCl_2 in benzene at 50 °C. Consequently, the solution is separated from the imidazolium salt and the solvent is reduced in volume and allowed to stand overnight at room temperature. Compound \( \text{1b} \) is obtained as a white yellow crystalline product in 45% yield.

The \( ^1 \)H NMR spectrum of \( \text{1a} \) and \( \text{1b} \) at room temperature displays one set of signals for the respective \( m \)-terphenyl and Tipp groups as well as the two coordinated NHCs. Additionally, for \( \text{1a} \), four broad singlets (corresponding to the N-Me and C-Me protons of NHCs, the 3,5-Mes protons and the ortho-Me protons of the mesityl group) observed at room temperature split up into two singlets each at –20 °C. On the other hand, the \( ^1 \)H NMR spectrum of \( \text{1b} \) does not show a similar broadening of the chemical shifts as seen for \( \text{1a} \). The \( ^{13} \)C NMR resonances for the carbene-carbons of \( \text{1a} \) and \( \text{1b} \) are observed at 160.3 ppm and 159.7 ppm as singlets, in the \( ^{13} \)C NMR spectrum, respectively. One sharp signal was observed at –68.85 ppm for \( \text{1a} \) in the \( ^{29} \)Si NMR spectrum, whereas the \( ^{29} \)Si NMR spectrum of \( \text{1b} \) displayed a chemical shift at –69.50 ppm as a singlet. These are downfield shifted compared to \( \text{V} \) (\( \delta = –82.3 \) ppm), but upfield shifted compared to \( \text{IV} \) (\( \delta = –58.4 \) ppm). The calculated values of \( ^{29} \)Si NMR resonances of \( \text{1a} \) and \( \text{1b} \) (\( \delta = –67.32 \) ppm and –68.55 ppm, respectively, B3LYP/6-31G(d)/[C,N,H]/6-31G(3d)/[Si]) are in good agreement with the experimental values.

Compound \( \text{1a} \) crystallizes in the monoclinic space group \( P2_1/c \) as separated ion pairs (the shortest Si-Cl distance is 6.234 Å). The molecular structure of \( \text{1a} \) is depicted in Fig. 1. The silicon centre is three fold coordinated to the two N-heterocyclic carbenes and ipso carbon (C1) of the \( m \)-terphenyl group. The sum of the bond angles around the Si1 atom is 310.2°. The Si-C bond lengths [Å] and angles [°]: Si(1)–C(1) 1.9355(19), Si(1)–C(32) 1.9481(20), Si(1)–C(25) 1.9665(19), C(1)–Si(1)–C(32) 105.06(8), C(1)–Si(1)–C(25) 111.36(8), and C(32)–Si(1)–C(25) 93.78(8).

Scheme 2 presents the possible mesomeric structures of compounds \( \text{1a} \) and \( \text{1b} \). A donor–acceptor stabilized silicon(II) cation is the mesomeric structure based on the high degree of pyramidalization observed for \( \text{1a} \) (310.21). The other mesomeric form is the zwitterionic structure, where the positive charge is dispersed over the two NHC backbones.

Furthermore, DFT calculations for the cationic part of \( \text{1a} \) and \( \text{1b} \) were carried out at B3LYP/6-31G(d) level of theory. The HOMO of \( \text{1a} \) shows mainly the lone pair orbital at the silicon centre, whereas the LUMO of \( \text{1a} \) is dispersed over the NHC skeletons (Fig. 2). Molecular orbitals of \( \text{1b} \) are similar to that of...
The NBO charge clearly shows that the silicon centre bears positive net charge (+0.798 for 1a and +0.804 for 1b).

Silylenes undergo cycloaddition reaction with internal alkynes and terminal alkynes as well as activate C-H bonds in terminal alkynes. It is of note that Müller has predicted the potential of silyliumylidene ions towards C-H activation through the generation of a silylum ion by proton abstraction from solvent by an intermediate silyliumylide. We have therefore embarked on reactivity investigation of the silyliumylidene 1a with the terminal alkyne. Interestingly, the unprecedented reactivity of the silyliumylidene 1a towards phenylacetylene was observed. The reaction of 1a with three equivalents of phenylacetylene in acetonitrile yields the m-terphenyl substituted 1-alkenyl-1,1-dialkynylsilane 2 in 68% yield, solely as the Z-isomer (Scheme 3).

Compound 2 was fully characterized by multinuclear NMR spectroscopy, ESI-HRMS as well as single crystal X-ray analysis. The 1H-NMR of 2 displays one set of signals for the m-terphenyl group, the alkynyl substituents and the alkenyl substituent. The ethylene (Si–CH = CH–Ph) protons are observed as doublets at 5.16 and 6.97 ppm. In the 13C NMR spectrum of 2, the ethylene carbon resonances (Si–C≡C–Ph) appear at 90.9 and 106.7 ppm, whereas the ethylene carbon chemical shifts (Si–CH = CH–Ph) are observed at 126.5 and 145.3 ppm. The 29Si-NMR chemical shift of 2 is observed at −62.28 ppm as a sharp singlet, which is downfield shifted in comparison to that of precursor 1a (δ = −68.85 ppm). In addition, this value fits well with the calculated value (δ = −59.02 ppm, B3LYP/6-31G(d)[C,N,H]/6-311G(3d)[Si]). Furthermore, DFT calculations [RI-B97-D/cc-pVTZ(SMD = acetonitrile)/RI-B97-D/6-31G* level of theory] were performed to suggest a plausible mechanism for the formation of 2 as a Z-isomer as the sole product (see ESI†). According to the DFT calculations, the formation of the E-isomer requires high activation barriers and therefore the formation of the E-isomer is kinetically not favored (for details see also the ESI†).

Compound 2 crystallizes in the monoclinic space group C2/c and the molecular structure is shown in Fig. 3. The central silicon possesses a distorted tetrahedral geometry with two alkynyl groups (C33–C41 and C41–C50) and one alkylene group (C25–C26) terminally coordinated to the silicon, whereas the maximum steric room is occupied by the umbrella shaped m-terphenyl substituent on the silicon. The broadest angle at the Si centre is displayed between C(terphenyl)–Si–C(alkylene) as 115.03°.

The alkylene substituent (C25–C26) reveals the Si–C(alkenyl) bond (1.87(2) Å) is slightly longer than that of 1-alkenyl-1-alkynylsilole (1.83(2) Å), however similar to those of the 1-alkenyl-1-alkynylsilole reported by Cui and co-workers.17 Another feature to be compared with the 1-alkenyl-1-alkynylsilole is the Si–C bond lengths. While the Si–C(alkenyl) bonds (1.82(15) Å and 1.83(15) Å) of 2 are similar to those of the 1-alkenyl-1-alkynylsilole (1.83(2) Å), however the Si–C(alkenyl) bond (1.87(25) Å) is slightly longer than that of 1-alkenyl-1-alkynylsilole (1.83(2) Å) which leads back to electronic interaction of silicon and butadiene in the silole ring.17

In conclusion, we report on the synthesis of novel m-terphenyl and Tipp substituted silyliumylidene ions 1a and 1b stabilized by two NHCs through a facile synthetic route. This striking one pot reaction will be further generalized by employing various substituents at the silicon center in our laboratories. In addition, the promising reactivity of 1a is evident from its reaction with phenylacetylene leading to the C–H insertion product 2. Both compounds 1a and 1b are characterized by multinuclear NMR as well as single-crystal X-ray diffraction analysis. Moreover, DFT calculations are performed to establish the energy pathway for the formation of 2 from 1a, which rationalizes the formation of the Z-isomer in this reaction. Further reactivity studies on 1a and 1b are ongoing and will be reported in due course.

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**Notes and references**


