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Gas Phase Formation of c-SiC₃ Molecules in the Circumstellar Envelope of Carbon Stars

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Abstract

Complex organosilicon molecules are ubiquitous in the circumstellar envelope of the Asymptotic Giant Branch (AGB) star IRC+10216, but their formation mechanisms have remained largely elusive until now. These processes are of fundamental importance in initiating a chain of chemical reactions leading eventually to the formation of organosilicon molecules - among them key precursors to silicon carbide grains – in the circumstellar shell contributing critically to the galactic carbon and silicon budgets with up to 80% of the ejected materials infused into the interstellar medium. Here we demonstrate via a combined experimental, computational, and modeling study that distinct chemistries in the inner and outer envelope of a carbon star can lead to the synthesis of circumstellar silicon tricarbide (c-SiC₃) as observed in the circumstellar envelope of IRC+10216. Bimolecular reactions of electronically excited silicon atoms $(Si(^{1}D))$ with allene (H₂CCCH₂) and methylacetylene (CH₃CCH) initiate the formation of SiC_3H_2 molecules in the inner envelope. Driven by the stellar wind to the outer envelope, subsequent photodissociation of the SiC₃H₂ parent operates the synthesis of the c-SiC₃ daughter species via dehydrogenation. The facile route to silicon tricarbide via a single neutral-neutral reaction to a hydrogenated parent molecule followed by photochemical processing of this transient to a bare silicon-carbon molecule provides evidence for a shift in currently accepted views of the circumstellar organosilicon chemistry and provides an explanation for the previously elusive origin of circumstellar organosilicon molecules, eventually constraining the level of molecular complexity, which can be synthesized in carbon-rich, circumstellar environments.

Significance Statement

Since the detection of methylidyne (CH) in deep space over 80 years ago, about 200 molecules and molecular ions from molecular hydrogen to fullerenes have been observed in interstellar and circumstellar environments, but the synthesis of organosilicon molecules has remained largely elusive. Exploiting silicon tricarbide (c-SiC₃) as a benchmark, we provide evidence from laboratory experiments, electronic structure calculations, and astrochemical modeling that organosilicon species can be synthesized via distinct chemistries in the inner and outer envelope of carbon stars like IRC+10216 via bimolecular reactions followed by photochemical dehydrogenation to bare silicon carbon molecules. These mechanisms are of fundamental significance to facilitate an understanding how carbon and silicon chemistries can be coupled to synthesize organosilicon molecules in the universe.

Keywords: Reaction dynamics; Circumstellar chemistry; Organosilicon molecules; Astrochemistry

\body

For the last decades, the carbon-rich Asymptotic Giant Branch (AGB) star IRC+10216 (CW Leo) has been widely recognized as a rich, natural laboratory for advancing our fundamental understanding of the chemical evolution of the carbon-rich circumstellar envelope (CSE) through astronomical observations combined with astrochemical modeling exploiting complex gas phase reaction networks involving ion-molecule¹⁻⁶ and neutral-neutral reactions.^{4,5,7-11} However, with about 80 molecules detected in circumstellar environments¹² – primarily hydrogen-deficient carbon chains and molecules with exotic chemical bonding containing metals and silicon - these models have, as yet, failed to account for the synthesis of ubiquitous silicon carbide molecules and forecasting their molecular fractional abundances, such as cyclic silicon dicarbide (c-SiC₂),¹³ bicyclic silicon tricarbide $(c-SiC_3)^{14}$ and linear silicon tetracarbide (SiC_4) ,¹⁵ which diverge by up to two orders of magnitudes compared with astronomical observations.^{16,17} Hence, although circumstellar molecules - among them precursors to silicon carbide grains - contribute critically to the galactic carbon and silicon budgets with up to 80 % of the ejected material infused into the interstellar medium,^{18,19} the formation routes of organosilicon molecules in circumstellar envelopes and the fundamental processes involved in coupling the carbon and silicon chemistries are largely unknown to date.^{17,20,21}

Modern astrochemical models propose that the very first silicon-carbon bonds are formed in the inner envelope of the carbon star, which is undergoing mass loss at the rates of several 10⁻⁵ solar masses per year.^{22,23} Pulsations from the central star may initiate shocks, which (photo) fragment the circumstellar material.^{24,25} These non-equilibrium conditions cause temperatures of 3,500 K or above¹⁹ and lead to highly reactive metastable fragments such as electronically excited silicon atoms, which can attack hydrocarbon molecules to initiate the formation of silicon-carbon bonds.^{24,26} The newly synthesized molecules are proposed to be driven by the stellar wind to the outer envelope, where they are photolyzed by the interstellar ultraviolet field.²⁴ Considering the strength of silicon-carbon double (450-550 kJ mol⁻¹) versus carbonhydrogen bonds of about 400 kJ mol⁻¹, the photochemistry is expected to preferentially cleave carbon-hydrogen bonds eventually synthesizing bare silicon-carbon molecules. However, the validity of these processes has not been confirmed, although these pathways are fundamental to an intimate understanding of the evolution of circumstellar envelopes at the molecular level, eventually constraining the pathways to bare silicon-carbon molecules in our galaxy.

To fill this gap, we present a combined crossed molecular beam, high-level *ab initio*, and astrochemical modeling investigation on the formation of (hydrogenated) circumstellar silicon tricarbide (c-SiC₃) via distinct chemistries in the inner and outer envelope of the carbon star IRC +10216. The reactions of electronically excited silicon atoms (Si(¹D)) with allene (H₂CCCH₂) and methylacetylene (CH₃CCH) (reaction (1)) were explored under single collision conditions, revealing first the formation of distinct SiC₃H₂ isomers in the inner envelope of IRC+10216. Ejected by the stellar wind to the outer envelope, photochemical processing of these SiC₃H₂ parent molecules leads to the synthesis of the bicyclic silicon tricarbide daughter species as observed in the envelope of IRC+10216 via dehydrogenation (reaction (2)).

(1) $\operatorname{Si}(^{1}\mathrm{D}) + \mathrm{C}_{3}\mathrm{H}_{4} \rightarrow \mathrm{Si}\mathrm{C}_{3}\mathrm{H}_{2} + \mathrm{H}_{2}$

(2) $SiC_3H_2 + h\nu \rightarrow c-SiC_3 + H_2$

Results

Crossed Molecular Beams Studies - Laboratory Frame

The gas phase reactions of electronically excited silicon atoms (Si(¹D)) with allene (H₂CCCH₂) and methylacetylene (CH₃CCH) were explored experimentally under single collision conditions in a crossed molecular beam machine by intersecting supersonic beams of electronically excited silicon atoms with the hydrocarbon beams perpendicularly at collision energies of 30 ± 2 kJ mol⁻¹ (Materials and Methods). The primary beam, generated via photolysis of 0.5 % disilane seeded in helium, contains ground and excited state silicon atoms (Si(³P)/Si(¹D)) along with the silylidyne radical (SiH(X²Π)). The reaction dynamics of the silylidyne radical with allene and methylacetylene have been explored previously,^{27,28} while ground state silicon atoms do not react with allene or methylacetylene (SI Appendix). Therefore, the present study allows us to discriminate scattering signal obtained from the reaction of electronically excited silicon atoms from signal originating from the silylidyne radical with allene and methylacetylene, respectively. These neutral reaction products were ionized via electron impact at 80 eV within a triply differentially pumped quadrupole mass spectrometric detector, and then mass- and velocity-analyzed to record time-of-flight (TOF) spectra of the ionized products.²⁹ First, in both systems, reactive scattering signal with allene and methylacetylene was observed at mass-to-

charge ratio (m/z) of 68 (SiC₃H₄⁺), 67 (SiC₃H₃⁺), and 66 (SiC₃H₂⁺). Signal at m/z of 68 (SiC₃H₄⁺) was found to originate from the reaction of the silvlidyne radical (SiH($X^2\Pi$)) with allene and methylacetylene leading to the formation of 2-methyl-1-silacycloprop-2-enylidene (c-SiC₃H₄) plus atomic hydrogen.^{25,26} Second, in both reactions, the TOF spectra at m/z of 68 (SiC₃H₄⁺) and 67 (SiC₃H₃⁺) were indistinguishable after scaling, revealing that signal at m/z of 67 originates from the dissociative electron impact ionization of the 2-methyl-1-silacycloprop-2-enylidene parent molecules in the electron impact ionizer. Third, TOF spectra taken at m/z of 66 (SiC₃H₂⁺) for both systems are distinct compared to the TOFs taken at m/z of 68 (SiC₃H₄⁺) and 67 $(SiC_3H_3^+)$. Therefore, signal at m/z of 66 does not solely arise from dissociative ionization of the SiC₃H₄ parent products formed in the reaction of the silvlidyne radical with allene and methylacetylene, but also from a molecular hydrogen loss channel in the reaction of the excited silicon with allene and methylacetylene leading to SiC₃H₂ isomer(s) (reaction (1)), provided that the ground silicon does not react with these two C₃H₄ isomers under our experimental conditions. Fig. 1 depicts the laboratory angular distributions and selected TOF spectra recorded at m/z of 66 (SiC₃H₂⁺), which were found to be fitted in two separated reaction channels. The first channel arises from the dissociative ionization of the SiC₃H₄ product from the silvlidyne radical reaction with allene and methylacetylene, whereas the second channel originates from reactive scattering within the Si(¹D)-C₃H₄ systems leading to SiC₃H₂ product(s) via molecular hydrogen losses (reaction (1)). Quantitatively, the contributions of the SiH- C_3H_4 system presents a minor channel contributing about 10 % of the ion signal at m/z of 66. As revealed in Fig. 1, the laboratory angular distributions spread over 45° and are nearly forward-backward symmetric around the center-of-mass (CM) angle of 33° for the Si(¹D) reactions with both C₃H₄ isomers. These data suggest that both Si(¹D) reactions with allene and methylacetylene proceed by indirect reaction dynamics involving the formation of SiC₃H₄ complex(es).

Having established that in the reactions of excited state atomic silicon with allene and methylacetylene, the molecular hydrogen loss leads to the formation of SiC_3H_2 isomer(s) (reaction (1)), we are attempting now to elucidate the position of the molecular hydrogen loss. This can be conducted for the methylacetylene reactant since the methyl and acetylenic hydrogen atoms are chemically non-equivalent, and its partially deuterated D3-methylacetylene reactant is commercially available. Reaction with Si(¹D) can proceed via HD loss to the formation of

SiC₃D₂ (68 amu) (reaction (3)), whereas molecular deuterium (D₂) loss should yield signal at m/z of 67 (SiC₃DH) (reaction (4)). Note that products formed via the HD loss (SiC₃D₂) cannot undergo dissociative electron impact ionization to signal at m/z of 67. Therefore, ions detected at m/z of 68 and 67 originate – with the exception of 3.3 % ¹³C and 5.1 % ²⁹Si substituted products – from SiC₃D_{2⁺} and SiC₃DH⁺, respectively. Fig. 2 shows the TOF data collected for the HD and D₂ losses at m/z of 68 (SiC₃D_{2⁺}) and 67 (SiC₃DH⁺) for the reaction of Si(¹D) with D3-methylacetylene. Due to the high cost of D3-methylacetylene, data were collected only at the CM angle. As evident from Fig. 2, we detected signal at m/z = 68 (SiC₃D_{2⁺}) and 67 (SiC₃DH⁺) related to the HD and D₂ loss with branching ratios of 58 ± 14% and 42 ± 14%, respectively. Overall, we conclude that in the reaction of electronically excited silicon atoms with D3-methylacetylene, both the molecular deuterium and hydrogendeuteride are open leading to distinct isomers.

(3)
$$Si(^{1}D) + CD_{3}CCH \rightarrow SiC_{3}D_{2} + HD$$

(4)
$$Si(^{1}D) + CD_{3}CCH \rightarrow SiC_{3}DH + D_{2}$$

Crossed Molecular Beams Studies – Center-of-Mass Frame

The interpretation of the raw data provided conclusive evidence that in the reaction of excited silicon atoms with allene and methylacetylene, an organosilicon molecule with the molecular formula SiC₃H₂ is synthesized via molecular hydrogen elimination. In the case of excited silicon atoms reaction with methylacetylene, the hydrogen atoms are ejected from distinct carbon atoms of the methyl and acetylenic group. We attempt now to identify the product isomer(s) formed. This requires an extraction of the underlying reaction dynamics by converting the laboratory data into the CM reference frame. Fig. 3 depicts the CM translational energy distributions $P(E_T)$ s and CM angular distributions $T(\theta)$ s for the molecular hydrogen loss channel leading to SiC₃H₂ isomer(s). Both the Si(¹D)–allene and Si(¹D)–methylacetylene systems could be fit with similar $P(E_T)$ s, i.e. 218 ± 28 kJ mol⁻¹ and 216 ± 26 kJ mol⁻¹ respectively. Therefore, the $P(E_T)$ s can extend up to 217 ± 29 kJ mol⁻¹ for the Si(¹D)-C₃H₄ systems, which presents the sum of reaction energy plus the collision energy for those products born without internal (ro-vibrational) excitation. By subtracting the collision energy from the high-energy cutoff, the reaction is determined to be exoergic by 187 ± 31 kJ mol⁻¹. Also, the $P(E_T)$ s peak away from zero

translational energy depicting a broad maximum around 50-80 kJ mol⁻¹; this finding indicates the existence of a tight exit transition state for the decomposition of the SiC₃H₄ complex(es) and a significant electron density rearrangement upon molecular hydrogen loss.³⁰ Finally, the $T(\theta)$ s are similar in both systems and provide additional information on the reaction dynamics. The $T(\theta)$ s cover the complete angular range from 0° to 180° with a nearly forward-backward symmetry. This suggests that the reaction involve indirect scattering dynamics via long-lived SiC₃H₄ intermediates whose lifetimes are longer than or at least competitive with their rotation periods.³¹ Also, the distribution maxima around 90° ('sideways' scattering) are clearly visible; this feature proposes geometrical constraints with the molecular hydrogen leaving parallel to the total angular momentum vector almost perpendicular to the rotational plane of the decomposing SiC₃H₄ intermediate(s).³¹

Electronic Structure Calculations – Bimolecular Reactions

Molecular Hydrogen Loss Channel: Having identified SiC₃H₂ isomers as the product of the bimolecular gas phase reaction of electronically excited silicon atoms (Si(¹D)) with allene and methylacetylene under single collision conditions, we next combine these findings with the computational results to untangle the underlying chemical dynamics and reaction mechanism(s). Reactants, products, intermediates, and transition state structures relevant to the reaction of electronically excited atomic silicon with methylacetylene, allene and D3-methylacetylene have been characterized in the SI Appendix (Tables S1-S3). A summary of these results is given shown in Fig. 4, in the form of zero-point corrected energies, which are slightly simplified by eliminating all transition states that are above the experimental collision energy of 30 ± 2 kJ mol⁻¹ and hence cannot be overcome; see SI Appendix for a full version of the surface). For both reaction systems, the formation of isomers **p1** – **p4** correlates well with the experimentally determined reaction exoergicity of 187 ± 31 kJ mol⁻¹ (Fig. 4).

In detail, the computations predict that both reactions are initiated by a barrierless addition of the Si(¹D) atom to the π electron density of the hydrocarbon reactant. For methylacetylene, Si(¹D) adds to the carbon-carbon triple bond, yielding intermediate [i1], which is the most stable of the SiC₃H₄ isomers. From [i1], the system can isomerize to [i2], [i4], [i5], and [i6] with barriers of 305, 188, 239, and 256 kJ mol⁻¹, respectively. Despite the lower barrier to [i4], rapid unimolecular decomposition to **p2** via [i4] \rightarrow [i13] \rightarrow **p2** is stifled by a barrier of 260 kJ mol⁻¹

for the two-step process. Isomer [i5] can readily isomerize to [i2] with a barrier of 19 kJ mol⁻¹ or to [i9] with a barrier of 99 kJ mol⁻¹. From [i6], there is very facile isomerization to [i9] and [i12], with barriers of 42 and 60 kJ mol⁻¹, respectively, and also access to [i10] through a barrier of 143 kJ mol⁻¹. Most of the intermediates are thereby capable of interconverting, if products are not too rapidly formed.

Qualitatively this is confirmed by examining the exit barriers for molecular hydrogen loss to form SiC₃H₂. The most favorable exit transition states are the [i6] \rightarrow p4 transition state at -114 kJ mol⁻¹, i.e. a barrier of 188 kJ mol⁻¹, the [i9] \rightarrow p1 transition state at -89 kJ mol⁻¹ with a barrier of 190 kJ mol⁻¹, and two [i12] \rightarrow **p3** transition states at -76 and -70 kJ mol⁻¹ with barriers of 172 and 178 kJ mol⁻¹, but from a high energy intermediate. Exit transition states to **p2** are much higher in energy, with energies of 7 kJ mol⁻¹, 46 kJ mol⁻¹, and 46 kJ mol⁻¹ for the [i1] \rightarrow **p2**, [i6] \rightarrow p2, and [i13] \rightarrow p2 transition states, respectively. Detailed prediction of branching ratios requires solving the coupled kinetic equations for all of these pathways, after applying RRKM theory to obtain the individual rate constants (SI). The outcome for this network is a predicted product distribution of 38.2% p1, 0.6% p2, 11.1% p3, and 45.8% p4. The experimental results from the silicon-D3-methylacetylene system, which reveal a branching ratio of $58 \pm 14\%$ HD loss and $42 \pm 14\%$ D₂ loss, allows for validation of our computed reaction network. Comparing the moderate exit barriers to the comparatively much smaller barriers to isomerization, we expect significant isomerization. Including all possible constitutional isomers in our reaction network, our RRKM calculations predict a product distribution of 56.8 % HD loss and 43.2 % D₂ loss (SI Appendix). This result is in good agreement with experiment, and suggests that the energy landscape shown in Fig. 4 is faithfully capturing the critical aspects of the experimental dynamics observed for $Si(^{1}D)$ with methylacetylene.

Turning next to the computational results for the reaction of Si(¹D) with allene, it is evident from Fig. 4 that the silicon atom can add barrierlessly to a C=C bond, yielding a 3-membered ring structure, [i7], or attack the central carbon atom to form the ring-opened isomer [i10]. There is virtually no barrier to ring-opening [i7] to yield [i10]. Intermediate [i10] provides access to the main intermediates already seen in silicon/methylacetylene system. For instance, the sequences $[i10] \rightarrow [i2] \rightarrow [i1]$ and $[i10] \rightarrow [i6] \rightarrow [i1]$ both access intermediate [i1], the initial encounter product for Si(¹D) + methylacetylene. Therefore the initial collision complexes of the siliconmethylacetylene ([i1]) and silicon–allene ([i7]/[i10]) surfaces are effectively coupled via intermediates [i2] and [i6] with reaction pathways eventually leading to **p1-p4** via H₂ loss involving tight exit transition states located 40 to 167 kJ mol⁻¹ above the energy of the separated products. The predicted product distribution for silicon/allene system is therefore very similar to that discussed above for the silicon/methylacetylene system: 38.7% **p1**, 0.5% **p2**, 11.3% **p3**, and 46.5% **p4**.

Atomic Hydrogen and Methyl Loss Channels: Having established that the molecular hydrogen loss channel leads predominantly to the formation of the 1-sila-1,2,3-butatrienylidene molecule $(H_2C=C=C=Si; X^1A_1; p1)$, and the less stable 4-membered ring structure (c-SiCH=C=CH; p4) we also investigated computationally the possibility of atomic hydrogen (H) and methyl (CH₃) loss channels; recall that these channels were not detected experimentally. For the atomic hydrogen loss channel, we located 22 SiC₃H₃ isomers (SI Appendix, Table S4). Six isomers – one acyclic (n1) and five cyclic molecules (n2-n6) – are energetically accessible under our experimental conditions at collision energies of 30 ± 2 kJ mol⁻¹ holding overall reaction energies from -10 to 24 kJ mol⁻¹. For the methyl loss channels, four SiC₂H isomers could be identified (SI Appendix, Table S5), two of which were permitted under experimental conditions with overall reaction energies of -30 and -5 kJ mol⁻¹ for m1 and m2, respectively. Statistical (RRKM) calculations were conducted to examine the branching ratios for the energetically accessible hydrogen atom and methyl loss channels (SI Appendix, Table S9) versus molecular hydrogen loss. These radical product channels were found to contribute only about 4 % in total to the overall product stream for silicon/methylacetylene, and only about 3 % of the product stream for the silicon/allene system (SI Appendix). These results bolster our experimental findings that the molecular hydrogen loss is the primary dissociation channel and also support the experimental non-detection of atomic hydrogen and methyl group loss channels.

Electronic Structure Calculations – Photodissociation of SiC₃H₂ to yield SiC₃

We have so far concluded that the bimolecular reactions of electronically excited atomic silicon with allene and methylacetylene lead primarily to the formation of 1-sila-1,2,3-butatrienylidene (H₂C=C=C=Si; X¹A₁) (**p1**) and other SiC₃H₂ isomers (particularly the 4-membered ring, **p4**) under experimental conditions mimicking the conditions in the inner envelope of carbon stars. Ejected by the stellar wind to the outer envelope, the SiC₃H₂ isomers

may be dehydrogenated subsequently via photodissociation to silicon tricarbide isomers (SiC₃) such as to the astronomically observed silicon tricarbide (c-SiC₃) by, e.g., interstellar Lyman- α photons (10.199 eV or 984 kJ mol⁻¹ for hydrogen) Our computations on the interconversion and molecular hydrogen loss pathways of SiC₃H₂ isomers are summarized in Fig. 5; see also SI Appendix (Table S6). From Fig. 5, it is evident that H₂C=C=C=Si (**p1**) can readily isomerize via ring closure to **p2**; likewise c-SiCH=C=CH (**p4**) can readily isomerize to **p3**.

Considering that the total system energy under experimental conditions is 220 kJ mol⁻¹ above **p1**, it is clear that molecular hydrogen loss and other isomer interchange pathways such as **p1** \leftrightarrow **p3** are closed unless the SiC₃H₂ isomers are further activated. This is most likely to occur by absorbing Lyman- α photons (10.2 eV or 984 kJ mol⁻¹) from the interstellar radiation field. As shown in the SI Appendix (Table S7), all five product isomers have excited states that carry significant oscillator strength in the region of 10.2 eV, with the strongest absorbers being **p2** (10.2 eV, f = 0.20) and **p4** (10.2 eV, f = 0.25). Upon photon absorption, and rapid non-radiative relaxation to an extremely vibrationally hot electronic ground state, this very large excess internal energy is sufficient to interconvert all SiC₃H₂ isomers and to readily surmount the barriers to molecular hydrogen loss and formation of three isomers of SiC₃, [1], [2] and [3] as shown on Fig. 5.

In agreement with our results, previous calculations revealed that c-SiC₃(X¹A₁) [1] represents the thermodynamically most stable silicon tricarbide isomer comparing with c-SiC₃(X¹A₁) [2], which has a transannular silicon-carbon bond, being 8 to 52 kJ mol⁻¹ less stable than [1].³³⁻⁴⁰ A second linear isomer (CSiCC) has been predicted to be about 318 kJ mol⁻¹ higher in energy compared to (l-SiC₃; X³Σ⁻).⁴¹ The rhomboidal C_{2v} symmetric isomer c-SiC₃ (X¹A₁) [1] that contains a transannular carbon-carbon bond,^{42,43} has been observed in the circumstellar envelope of IRC +10216. We also computationally investigated alternative photodissociation pathways from p1, p2, and p5 via atomic hydrogen loss (SI Appendix, Table S8). These channels are endoergic between 387 and 359 kJ mol⁻¹. While these reaction energies are close to the transition state energies for the molecular hydrogen loss, the molecular hydrogen loss channels are thermodynamically preferred and would be enhanced via tunneling leading us to suggest that molecular hydrogen loss is the main photodissociation pathway, and will produce c-SiC₃ (X¹A₁) [1].

Astrochemical Modeling

Having established that the cyclic silicon tricarbide molecule (c-SiC₃) can be formed via photodissociation of 1-sila-1,2,3-butatrienylidene (SiC₃H₂; H₂C=C=C=Si; X¹A₁), which is generated via bimolecular reactions of electronically excited atomic silicon atom with allene and methylacetylene, we explore how our findings transfer to the circumstellar envelope of IRC+10216 through astrochemical modeling. This is central since crossed molecular beam experiments along with computations cannot adequately simulate conditions in circumstellar environments, such as the presence of multiple reactants and where the photodissociation of newly formed molecules plays a fundamental role.¹⁷ We performed simulations using a network of gas phase reactions in the circumstellar envelope of the carbon rich star IRC+10216 via the chemical kinetic data of the RATE12 release from the UMIST Database for Astrochemistry (UDfA).¹⁷ Physical parameters for the simulations of the inner to outer envelopes were adopted from Li et al..⁴⁴ This reaction network was updated by the additional reactive and photodissociation chemistries of the SiC₃H₂ molecule as elucidated in the present work with initial fractional abundances of the reactive species relative to molecular hydrogen taken from the models of Refs. 17 and 44.

These modeling studies lead to fascinating conclusions. First, the astrochemical modeling reveals multiple sources of electronically excited silicon atoms in the inner envelope: photodissociation of silane (SiH₄)⁴⁵ and (photo) fragmentation of two silicon-bearing diatomic molecules silicon monoxide (SiO) and silicon monosulfide (SiS), which in turn can be generated from shock-induced sputtering of silicon-bearing grains in the inner envelope of the carbon star.^{19,46} Silane, silicon monoxide, and silicon monosulfide were detected in the circumstellar envelope at fractional abundances of 2.2×10^{-7} , 1.8×10^{-7} , and 1.3×10^{-6} with respect to molecular hydrogen.^{44,47} Note that Suto and Lee examined the photodissociation of silane in the range 106-160 nm revealing that electronically excited silicon atoms contribute up to 1.2 % of the photodissociation yield.⁴⁵ Based on that the overall silicon budget reaches peak fractional abundance above 1.0×10^{-6} , electronically excited silicon atoms constitute up to 1% due to photodissociation. Second, the coupling of the silicon with the carbon chemistries leads to

present-day fractional abundances of allene (H₂CCCH₂), methylacetylene (CH₃CCH),⁴⁸ and silicon tricarbide (c-SiC₃) between 7.0×10^{-9} and 8.0×10^{-9} at a radius of about 4×10^{16} cm, while silicon tetracarbide (SiC₄) shows a much lower peak abundance of 2.0×10^{-10} (Fig. 6).

The modeling suggests that our proposed mechanism contributes up to an 80% increase in the column density of the total silicon tricarbide budget. Our new channels constitute an experimentally and computationally validated reaction mechanism based on neutral-neutral reactions followed by photo processing, which complements previously proposed networks of ion-molecule reactions that are less well characterized (SI Appendix). It should be highlighted that the fractional abundance of silicon tricarbide (c-SiC₃) likely represents a lower limit since only Lyman- α photons were considered in our model to photodissociate SiC₃H₂ (SI Appendix, Table S7). Further, the modeled fractional abundances of SiC₃ and of SiC₄ clearly indicate that SiC₄ cannot represent a precursor of SiC₃. Note that in an early study, Apponi et al. commented that SiC₃ and SiC₄ held column densities of 4.3×10^{12} cm⁻² and 2.7×10^{12} cm⁻² assuming that both have the same 40" diameter shell as silicon dicarbide (SiC₂); however, up-to-date models predict that 20" and 25" diameter shells have to be considered for SiC₃ and SiC₄, respectively,^{44,47} thus suggesting that conclusions drawn from older models have to be adjusted.

Finally, it is important to note that fractional abundances of SiC_3H_2 of about 10^{-10} are still predicted to be present today and are hence observable in prospective searches in the circumstellar shell. Therefore, the astrochemical modeling reveals the potential for bimolecular reactions of electronically excited silicon atoms (Si(¹D)) with allene (H₂CCCH₂) and methylacetylene (CH₃CCH) to initiate the formation of SiC₃H₂ molecules in the inner envelope. Then, driven by the stellar wind to the outer envelope, subsequent photodissociation of the SiC₃H₂ parent is efficient in the formation of the bicyclic silicon tricarbide (c-SiC₃) daughter species via dehydrogenation.

Conclusions

On the bimolecular reactions of electronically excited silicon atoms (Si(¹D)) with allene and methylacetylene, our combined experimental and computational investigation unraveled indirect scattering dynamics initiated through barrierless addition of silicon to the π electronic systems of the hydrocarbon reactants leading to distinct SiC₃H₄ collision complexes. These intermediates are formed with 250-400 kJ mol⁻¹ of internal energy enabling them to interconvert, and

ultimately undergo unimolecular decomposition by molecular hydrogen loss to form at least 1sila-1,2,3-butatrienylidene (H₂C=C=C=Si; X¹A₁) (**p1**) among other isomers. The product isomer distribution is then predicted by a kinetic network developed computationally that is validated by excellent agreement with the H₂/HD ratio observed experimentally for the silicon-D3methylacetylene system. The experiment was conducted at a collision energy of 30 ± 2 kJ mol⁻¹, which is equivalent to a temperature of about 3,500 K. 3500 K is comparable to conditions in circumstellar envelopes of carbon-rich stars reached by pulsations and inherent shocks in the ejected stellar material.¹⁹ The newly formed sila-1,2,3-butatrienylidene (SiC₃H₂) formed in the inner envelope via equation (1), can then be transported by the stellar wind to the thinner, outer envelope. In these regions, as demonstrated computationally and via astrochemical modeling, the interstellar ultraviolet field can photolyze sila-1,2,3-butatrienylidene (SiC₃H₂) to the bicyclic silicon tricarbide isomer c-SiC₃(X¹A₁) **[1]** as observed in the envelope of IRC+10216 via dehydrogenation (equation (2)).

This system illustrates the concept of an excited state induced synthesis of exotic organosilicon molecules (SiC₃H₂) in the inner circumstellar envelope via the reaction of electronically excited silicon atoms with ubiquitous hydrocarbons with the parent species undergoing photochemical processing in the outer envelope to cyclic silicon tricarbide ($c-SiC_3$). Previous models proposed that silicon tricarbide is formed by photodissociation of silicon tetracarbide (SiC₄),^{4,15,49} but the predicted column density of SiC₄ is two orders of magnitude less than those being observed astronomically in the circumstellar envelope of IRC+10216.^{17,44} Therefore, although some c-SiC₃(X^1A_1) may originate from photodissociation of SiC₄(X^1A_1), the computations along with astrochemical models study identify hydrogenated silicon-carbon species – the isomers of SiC_3H_2 – as a key source of c-SiC₃(X¹A₁).⁵⁰ This facile route to silicon tricarbide via a single, bimolecular neutral-neutral reaction to a hydrogenated parent molecule followed by photochemical dehydrogenation to a bare silicon-carbon molecule represents a shift away from currently accepted views. It provides an explanation for the previously elusive origin of circumstellar organosilicon molecules eventually constraining the level of molecular complexity, which can be expected in carbon-rich, circumstellar environments. Although the interstellar photon field is more complex than considered in our photodissociation study, which was restricted to Lyman- α , this proof-of-concept study opens up a hitherto overlooked possibility

of distinct chemistries in the inner and outer circumstellar envelop of IRC+10216 eventually leading to silicon tricarbide c-SiC₃(X¹A₁). This result challenges the conventional wisdom that circumstellar silicon carbide molecules can only be formed at elevated temperatures via complex ion-molecule reactions or photodegradation of higher molecular weight silicon carbon molecules. With the commission of the Atacama Large Millimeter/ Submillimeter Array (ALMA), the detection of unusual silicon-bearing molecules will continue to grow, and an understanding of these data will rely on advances in experimental and computational laboratory astrophysics as proposed here, thus eliminating the gap between observational and laboratory data on the circumstellar organosilicon chemistry that has existed for decades.

Materials and Methods

Experimental. The crossed beam reactions of ground and excited state silicon atoms $(Si(^{3}P)/Si(^{1}D))$ with allene $(H_{2}CCCH_{2}; X^{1}A_{1})$ and methylacetylene $(CH_{3}CCH; X^{1}A_{1})$ were conducted in a universal crossed molecular beams machine under single collision conditions.^{29,51-} ⁵³ A pulsed supersonic beam of silicon atoms (Si(³P)/Si(¹D)) along with silvlidyne radicals (SiH(X²Π)) was generated via photolysis of 0.5 % disilane (Si₂H₆; 99.998 %; Voltaix) seeded in helium (He; 99.9999 %; Gaspro). This gas mixture was fed into a pulsed valve equipped with a piezo crystal (Piezo Disk Translator P-286.23; Physik Instrumente) operating at a repetition rate of 120 Hz, pulse width of 80 µs, peak voltage of -350 V, and backing pressure of 1,520 Torr in the primary source chamber. Since disilane is an extremely flammable gas with a flammability limit as low as 0.2 % in air (Disilane MSDS, Linde Electronics and Specialty Gases), a liquid nitrogen cold trap (FTN-6-1502-NWB, Nor-Cal products) was incorporated between the turbo molecular pump and the roots scroll backing pump system to freeze out the disilane. To generate the reactive silicon species, the output of an excimer laser (ArF, 193 nm, 30 mJ per pulse, Coherent) was focused with a lens (1.5-meter focal length; CVI Melles Griot) downstream of the pulse valve nozzle to an area of $1 \text{ mm} \times 4 \text{ mm}$. The pulsed beam of the silvlidyne radicals and of the silicon atoms passed through a skimmer; a four-slit chopper wheel rotating at 120 Hz selected a part of the supersonic beam with a well-defined peak velocity (v_p) and speed ratio (S) of 1735 ± 20 ms⁻¹ and 19 ± 2 , respectively. In the interaction region, this section of the pulse intercepted the most intense sections of pulsed beams of allene (98 %; Organic Technologies)

and methylacetylene (99 %; Organic Technologies) at a backing pressure of 550 Torr perpendicularly. Peak velocities and speed ratios of the hydrocarbons of the pulse were determined to be $790 \pm 10 \text{ ms}^{-1}$ and 12 ± 1 , for both allene and methylacetylene. This resulted in nominal collision energies of $30 \pm 2 \text{ kJ}$ mol⁻¹ as well as a center-of-mass (CM) angle of $33 \pm 1^{\circ}$. To elucidate the position of the molecular hydrogen loss, experiments also were carried out with partially deuterated D3-methylacetylene (CD₃CCH; 99 atom % D; CDN Isotopes), but only at the CM angle for economic reasons. The primary pulsed valve was triggered after 1886 µs with respect to the time zero defined by the infrared diode pulse mounted on the chopper wheel; since the secondary beam was slower than the primary beam, the secondary pulsed valve had to be triggered 35 µs prior to the primary one. Finally, the excimer laser was triggered 155 µs later than the primary pulsed valve to allow a maximum overlap of the gas pulse with the laser beam. To allow a 'laser-off' minus 'laser-on' background subtraction, both pulsed valves were triggered at 120 Hz, but the laser at half of the repetition rate at 60 Hz.

The reactively scattered products were mass filtered after ionization utilizing a quadrupole mass spectrometer (QMS) operated in the time-of-flight (TOF) mode; the ions were detected by a Daly-type detector housed in a rotatable, triply-differentially pumped ultrahigh vacuum chamber $(1 \times 10^{-11} \text{ Torr})$ after electron-impact ionization of the neutral products at an electron energy of 80 eV and an emission current of 2 mA. This detector can be rotated within the plane defined by the primary and the secondary reactant beams to record angular-resolved TOF spectra. At each angle, up to 6×10^5 TOF spectra were accumulated to obtain good signal-to-noise ratios. The recorded TOF spectra were then integrated and normalized to the intensity of the TOF at the CM angle to extract the product angular distribution in the laboratory frame. In order to acquire information on the scattering dynamics, the laboratory data were then transformed from the laboratory into the CM reference frame utilizing a forward-convolution routine.⁵⁴⁻⁵⁶ This iterative method employs a parametrized or point-form angular flux distribution, $T(\theta)$, and translational energy flux distribution, $P(E_T)$, in the CM system. Laboratory TOF spectra and the laboratory angular distributions are calculated from the $P(E_{\rm T})$ and $T(\theta)$ functions, and are averaged over a grid of Newton diagrams accounting for the apparatus functions, beam divergences, and velocity spreads. For the fitting, we considered a reactive scattering cross section of an $E_{\rm T}$ -^{1/3} energy dependence, where $E_{\rm T}$ stands for the translational energy. This method

is usually adopted within the line-of-center model for exoergic and entrance barrierless reactions dominated by long-range attractive forces.^{57,58}

Theoretical. Structures for the reactants, intermediates, and products were obtained via geometry optimizations and frequency calculations using the ω B97X-V⁵⁹ density functional and the cc-pVTZ basis set.⁶⁰ This functional is known to be amongst the most accurate density functionals for thermochemistry and reaction barrier heights.⁶¹ Transition state structures were calculated using the freezing string method (FSM)⁶² to generate an initial structure and Hessian which were then refined by a transition state search using the partitioned-rational function optimization (P-RFO) eigenvector following method⁶³ and followed by a frequency calculation. These calculations were also carried out at the $\omega B97X-V/cc-pVTZ$ level of theory. The vibrational analysis was used to confirm that the minima have no imaginary frequencies and the transition states have only one imaginary frequency each, as well as to calculate harmonic zeropoint energy corrections for the structures. Density functional theory (DFT) calculations were all carried out using an ultra-fine integration grid consisting of 99 radial points and 590 angular points. To further improve the accuracy of the results for relative energies and barrier heights, single point energy calculations were performed at the minima and transition states. The objective was to approach the complete basis set (CBS) limit using coupled cluster with single, double, and perturbative triple excitations (CCSD(T)).⁶⁴ To this end, CCSD(T)/cc-pVTZ calculations using a frozen core approximation were combined with second-order Møller-Plesset perturbation theory using the resolution of the identity approximation (RI-MP2),^{65,66} in larger basis sets. The working expression used for the energy of a given structure was:

E(CCSD(T)/CBS)

$$= E(HF/cc - pV5Z) + E^{corr}(RI - MP2/CBS_{3,4,5}) + E^{corr}(CCSD(T)/cc - pVTZ) - E^{corr}(RI - MP2/cc - pVTZ) + ZPE(\omega B97X - V/cc - pVTZ).$$

 $E^{corr}(RI-MP2/CBS_{3,4,5})$ is the extrapolated RI-MP2 correlation energy using the cc-pVTZ, cc-pVQZ, and cc-pV5Z basis sets to fit

$$E^{corr}(RI - MP2/cc - pVNZ) = E^{corr}(RI - MP2/CBS_{3,4,5}) + AN^{-3}$$

where *N* denotes the cardinal number for the cc-pVNZ basis sets.⁶⁷ These CCSD(T)/CBS energies are typically accurate to about 3-4 kJ mol⁻¹. The energy of the silicon atom in the ¹D state was calculated by computing the energy of the atom on its triplet ground state and corrected

with an experimentally derived triplet-singlet gap (75.4 kJ mol⁻¹) taken from the NIST Atomic Spectra Database.⁶⁸ Time-dependent density functional theory⁶⁹ calculations were also carried out on select SiC₃H₂ species to search for electronic excited states corresponding to the absorption of a Lyman-α photon. All calculations were performed using the Q-Chem suite of electronic structure programs.⁷⁰ For further analysis, the Rice-Ramsperger-Kassel-Marcus (RRKM)⁷¹ rate constants for select dehydrogenation pathways of SiC₃H₄ isomers were approximated using a Beyer-Swinehart direct state counting algorithm⁷² modified to include vibrational tunneling as suggested by Miller.⁷³

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Fig. 1. Laboratory angular distributions and selected TOF spectra. Laboratory angular distributions of the products recorded at m/z of 66 (SiC₃H₂⁺) for the reactions of excited state silicon atoms (Si(¹D)) (blue) along with the silylidyne radical (SiH(X²Π)) (green) with allene (a) and methylacetylene (b). The solid circles represent the experimental data, C. M. designates the center-of-mass angle, the error bars represent the 1 σ standard deviation and red solid lines represent the overall fit. Selected TOF spectra in subpanels (c) and (d) represent the reactions of excited state silicon atoms (Si(¹D)) (blue) along with the silylidyne radical (SiH(X²Π)) (green) with allene (c) and methylacetylene (d) with the overall fit defined by the red lines, while the open circles depict the experimental data points.



Fig. 2. TOF spectra for the reaction of excited silicon atoms with D3-methylacetylene (CD₃CCH). Recorded at the respective CM angle, the two TOF spectra depict the signal for m/z of 68 (red line, HD loss) and 67 (blue line, D₂ loss), corresponding to singly ionized SiC₃D₂ and SiC₃DH molecules, respectively.



Fig. 3. CM translational energy distributions $P(E_T)$ s and CM angular distributions $T(\theta)$ s. Subpanels (a) and (b) are responsible for $P(E_T)$ s while subpanels (c) and (d) depict $T(\theta)$ s, in the reactions of excited silicon atoms with allene ((a) & (c)) and with methylacetylene ((b) & (d)), leading to the products of SiC₃H₂ isomers and molecular hydrogen. The shaded areas represent the acceptable fits accounting for the 1 σ error limits of the laboratory distributions.



Fig. 4. Pruned potential energy surface for the reactions of silicon (Si(¹D)) with allene and methylacetylene. Relative energies are given in units of kJ mol⁻¹. The potential energy surface is pruned by removing barriers above the 30 kJ mol⁻¹ collision energy (refer to the SI Appendix (Fig. S1) for the full version). Colors of the atoms: silicon (green), carbon (black) and hydrogen (light grey).



Fig. 5. Potential energy surface for interconversion of SiC₃H₂ isomers and H₂ elimination to yield SiC₃. Relative energies are given in units of kJ mol⁻¹; the zero of energy is taken as the ZPE-corrected energy of H2C=C=C=Si. The total internal energy of the system under experimental conditions is +220 kJ/mol. Colors of the atoms: silicon (green), carbon (black), and hydrogen (light grey).



Fig. 6. Modeled present-day fractional abundances of key species related to the formation of silicon tricarbide (c-SiC₃) relative to molecular hydrogen. The relative fractional abundances of methylacetylene (CH₃CCH), allene (H₂CCCH₂), 1-sila-1,2,3-butatrienylidene (SiC₃H₂), silicon tricarbide (SiC₃), and silicon tetracarbide (SiC₄) are plotted as a function of radius for the circumstellar envelope of the carbon star IRC +10216.