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Structures of Tetrasilylmethane Derivatives (XMe₂Si)₂C(SiMe₃)₂ (X = H, Cl, Br) in the Gas Phase, and their Dynamic Structures in Solution

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ABSTRACT: The structures of the molecules $(XMe_2Si)_2C(SiMe_3)_2$, where X = H, Cl, Br, have been determined by gas electron diffraction (GED) using the SARACEN method of restraints, with all analogues existing in the gas phase as mixtures of C_1 -and C_2 -symmetric conformers. Variable temperature ¹H and ²⁹Si solution-phase NMR studies, as well as ¹³C NMR and ¹H/²⁹Si NMR shift correlation and ¹H NMR saturation transfer experiments for the chlorine and bromine analogues, are reported. At low temperatures in solution there appear to be two C_1 conformers and two C_2 conformers, agreeing with the isolated-molecule calculations used to guide the electron diffraction refinements. For (HMe₂Si)₂C(SiMe₃)₂ the calculations indicated six conformers close in energy, and these were modeled in the GED refinement.

KEYWORDS: gas electron diffraction, molecular structure, conformational analysis, tetrasilylmethane derivatives, variable-temperature NMR

■ INTRODUCTION

The chemistry of tetrasilylmethane derivatives has been the subject of numerous studies, and many novel structures and unusual reactivities have been attributed to having four silicon centers in a sterically crowded environment.^{1–4} The most widely studied tetrasilylmethane derivatives have the general structures $(XMe_2Si)_4C$, $(Me_3Si)_3CSiRR'X$, $(PhMe_2Si)_3CSiRR'X$, and $(Me_3Si)_2C(SiMe_2X)(SiR_2Y)$ (where R and R' = Me, Et, Ph *etc.*, and X and Y = H, halide, OAc *etc.*).^{1–5}

A range of simple *bis*-functionalized tetrasilylmethanes $(XMe_2Si)_2C(SiMe_3)_2$ (for example, X = H,⁶⁻⁹ F,⁹⁻¹¹ Cl,^{7,9-14} Br,^{6,7,9,15,16} I,^{7,9,11,12,17} OH,^{7,9,10,18,19} OMe,^{10,20-22} OAc,^{7,10,20} O₂CCF₃,^{10-12,16,19} OClO₃,⁹ OSO₂CF₃,⁹ OSO₂-*p*-C₆H₄Me,⁹ and vinyl^{12,15}) are known, but apart from the diol (HOMe₂Si)₂C(SiMe₃)₂,¹⁸ little structural information is available for them. The structure of the permethyl species, (Me₃Si)₄C, has, however, been studied by NMR spectroscopy,^{23,27} X-ray diffraction,²⁸⁻³⁰ gas electron diffraction (GED),^{31,32} computational methods^{33,34} and vibrational spectroscopy.³⁴

Solution-phase NMR spectroscopy has been used to probe dynamic processes in bulky tetrasilylmethane derivatives previously, for example, in $C(SiMe_3)_2(SiMePh_2)(SiMe_2ONO_2)$,³⁵ $C(SiMe_3)_2(SiClPh_2)(SiMe_2OMe)$,³⁶ (Me_3Si)_3CSiX_3 (X = Cl or Br)³⁷ and (PhMe_2Si)_3SiCl_3.³⁷ GED studies have also been carried out on the tetrasilylmethane (Me_3Si)_3CSiCl_3³⁸ and on (HMe_2Si)_3CSiH_3,³⁹ the latter of which showed the presence of eleven distinct conformers.

The work presented here comprises two related studies. First, the structures of $(XMe_2Si)_2C(SiMe_3)_2$ [X = H (1), Cl (2), Br (3)] are described in the gas phase as determined by the combination of GED experiments and *ab initio* calculations, and

secondly, NMR studies of the dynamic processes occurring for the same species in solution are presented.

EXPERIMENTAL SECTION

Syntheses. The syntheses of $(HMe_2Si)_2C(SiMe_3)_2$,⁹ $(CIMe_2Si)_2C(SiMe_3)_2$,¹³ and $(BrMe_2Si)_2C(SiMe_3)_2$ ⁹ were carried out according to the literature methods shown in Scheme 1, and the compounds were purified for structural studies by sublimation. Thus, treatment of $(Me_3Si)_3CSiMeCII$ with ICl leads to a rearrangement of the type often seen in tetrasilylmethane derivatives¹ to give $(CIMe_2Si)_2C(SiMe_3)_2$ (**2**), which is readily reduced by LiAlH₄ to give $(HMe_2Si)_2C(SiMe_3)_2$ (**1**), which then affords $(BrMe_2Si)_2C(SiMe_3)_2$ (**3**) in high yield upon treatment with bromine.^{*}

Scheme 1 Synthetic routes to $(XMe_2Si)_2C(SiMe_3)_2$ (X = H, Cl, Br)



NMR measurements. ¹H, ¹³C and ²⁹Si NMR spectra were recorded in CDCl₃/CD₂Cl₂ or CDCl₃/acetone- d_6 solutions using a Bruker AMX 500 spectrometer at 500, 126, and 99 MHz, respectively, unless otherwise stated. The ²⁹Si{¹H} NMR INEPT spectra were

^{*} The fluorine analogue $(FMe_2Si)_2C(SiMe_3)_2$ can be prepared by reaction between $(AcOMe_2Si)_2C(SiMe_3)_2$ and CsF,¹⁰ while the iodine analogue $(IMe_2Si)_2C(SiMe_3)_2$ can be prepared by reaction of $(HMe_2Si)_2C(SiMe_3)_2$ and I_2 ,⁹ though neither compound was readily available for use in the current study.

recorded on a Bruker AMX 500 NMR spectrometer at 99 MHz, and ²⁹Si{¹H} inverse-gated NMR spectra were recorded on a Bruker Avance 600 spectrometer at 119.23 MHz. Chemical shifts of all NMR spectra are reported in ppm relative to TMS.

All solid-state ¹³C{¹H} and ²⁹Si{¹H} MAS NMR spectra were recorded on a Bruker DSX 200 WB NMR spectrometer. Samples were spun at 3–5 kHz and simple Bloch decay techniques (standard single-pulse excitation method) were used. Approximately 1000 scans per sample were collected. The operating frequencies for ¹³C and ²⁹Si NMR experiments were 50.28 and 39.7 MHz, respectively.

X-ray Crystallography. The attempted single-crystal X-ray study for $(BrMe_2Si)_2C(SiMe_3)_2$ was carried out using an OD Xcalibur 3 diffractometer, using X-rays of wavelength 0.71073 Å, at a temperature of 100 K.

Computational Methods. Previous studies for similar molecules³⁹ suggested that $(XMe_2Si)_2C(SiMe_3)_2$ compounds would have a series of potential-energy minima leading to a number of conformational isomers, dependent on the relative rotations of the two XMe_2Si groups. Figure 1 shows one possible conformation of $(XMe_2Si)_2C(SiMe_3)_2$ as an illustration.

Figure 1. Structure, with atom numbering, of one conformer of $(XMe_2Si)_2C(SiMe_3)_2$. Hydrogen atoms have been removed for clarity. Atoms in subsequent conformers are numbered by adding multiples of 47 to these.



These minima occurred at three approximate X(14/15)–Si(2/3)–C(1)–Si(3/2) dihedral angles: 80, –40 and –160°. By independently setting the two XMe₂Si to all possible combinations of these angles, it can be seen that there are nine (=3²) possible minimumenergy conformers for each of **1–3**. Experience of studying a similar set of species (XMe₂Si)₄C, where X = H, F, Cl, Br, has shown that the opposite sense of each angle (*i.e.* – 80, 40 and 160°) should also be considered when looking to identify all possible conformers.⁴⁰

All calculations used Gaussian 09⁴¹ on either the University of Edinburgh's ECDF cluster⁴² or the UK's National Service for Computational Chemistry Software clusters.⁴³ Geometry optimizations and frequency calculations were carried out to determine which ground-state conformers had the lowest energies. For comparison, both the B3LYP⁴⁴⁻⁴⁶ and M06-2X⁴⁷ methods with the 6-31G(d)^{48,49} basis set were used for these calculations.

Further geometry optimizations and frequency calculations were carried out on conformers deemed to have low lying energies. The B3LYP hybrid method with the aug-cc-pVDZ basis set^{50,51} was used for **1** and **2**, whilst the pseudopotential basis set aug-cc-pVDZ-PP^{52,53} was used for **3**. These basis sets will be denoted as aug-cc-pVDZ(-PP) from hereon in. As a comparison, geometry optimizations were performed for every conformer using the M06-2X method and the aug-cc-pVDZ(-PP) basis set, as well as calculations using the MP2 method⁵⁴ with the 6-31G(d) and aug-cc-pVDZ(-PP) basis sets. The relative amounts of each conformer that would be present in the GED samples at the temperature of each experiment were calculated using the Gibbs free energy for each conformer (obtained from quantum calculations carried out at 0 K) and the Boltzmann distribution equation:

$$\frac{N_i}{N} = \frac{g_i \exp\left(-\frac{\Delta G_i}{RT}\right)}{\sum_i g_i \exp\left(-\frac{\Delta G_i}{RT}\right)},$$

where *N* is the total number of molecules, and N_i is the number of molecules in a given state *i*, at temperature *T*. *R* is the gas constant, whilst ΔG_i and g_i are the Gibbs free energy difference (with respect to the lowest energy conformer) and degeneracy, respectively, of state *i*, where g_i is equal to 1 for C_1 symmetric, and 2 for C_2 symmetric molecules.

Gas Electron Diffraction (GED). Data for 1, 2, and 3 were collected using the GED apparatus that was used in Edinburgh until 2010.⁵⁵ An accelerating potential of 40 keV was applied, producing electrons with an approximate wavelength of 6.0 pm. Each molecule was analyzed with two different nozzle-to-camera distances, increasing the range of data collected. Exact nozzle-to-camera distances were calibrated by analyzing the results of benzene diffraction experiments that were carried out immediately after collecting data for the molecules of interest. The scattering intensities were recorded on Kodak Electron Image films, and measured with the use of an Epson Expression 1680 Pro flat-bed scanner and converted to mean optical densities using a method described elsewhere.⁵⁶ A full list of experimental parameters, including the measured nozzle and sample temperatures for each experiment, can be found in Table S1.

The data were analyzed using the ed@ed least-squares refinement program v3.0,⁵⁷ incorporating the scattering factors of Ross *et al.*⁵⁸ Weighting points for the off-diagonal weight matrices, and scale factors can be found in Table S1, while Tables S2–S4 show the correlation matrices.

RESULTS AND DISCUSSION

Gas-Phase Static Structures. By starting geometry optimization for structures with all possible combinations of minimum-energy dihedral angles, six unique conformers were identified. Frequency calculations, carried out using M06-2X/6-31G(d) and B3LYP/6-31G(d), suggested that all of the unique conformers of **1** had similar ground-state energies, and hence all could be present in the gas phase at the temperature of the experiments. Three of these conformers have C_1 symmetry (**1a–c**), and three have C_2 symmetry (**1d–f**). The calculations also suggested that for **2** and **3** four of these six conformers were likely to be observable in the gas electron diffraction experiments. For each of these molecules, two conformers have C_1 symmetry (**2a/b** and **3a/b**) and two have C_2 symmetry (**2c/d** and **3c/d**). Tables 1–3 show the zero-point-corrected ground-state Gibbs free energies for all conformers of **1**, **2**, and **3**, as obtained from the B3LYP/aug-cc-pVDZ(-PP) calculations, and relate these to the relative abundance of each conformer at the temperature of the experiment, which for each species **1–3** is an average of the recorded temperatures for both nozzle and sample (seen in Table S1) at both camera distances.

	indicative	point-group	relative			
conformer	dihedral angle ^b	symmetry	energy ^c	proportion ^d		
1a	-160 / -40	C_1	1.47	0.204		
1b	-160 / 80	C_1	0.00	0.311		
1c	80 /40	C_1	0.31	0.285		
1d	-160/-160	C_2	4.13	0.048		
1e	80 / 80	C_2	2.18	0.083		
1f	-40 / -40	C_2	2.87	0.069		

 Table 1. Indicative Dihedral Angle, Symmetry, Relative Energy, and Proportion for

 Each Conformer of 1^a

^{*a*} Calculations performed using B3LYP/aug-cc-pVDZ. ^{*b*} These are the starting values of the H(14)-Si(2)-C(1)-Si(3) / H(15)-Si(3)-C(1)-Si(2) dihedral angles in degrees; no interconversion was observed upon optimization. See Figure 1 for atom numbering. ^{*c*} Gibbs free energy in kJ mol⁻¹ (ZPE corrected). ^{*d*} Calculated at 431 K.

	indicative	point-group	relative	
conformer	dihedral angle ^b	symmetry	energy ^c	proportion ^d
2a	-160 / -40	C_1	0.00	0.720
2b	-160 / 80	C_1	7.56	0.106
2c	-160/-160	C_2	6.19	0.075
2d	80 / 80	C_2	5.07	0.099

Table 2. Indicative Dihedral Angle, Symmetry, Relative Energy, and Proportion for Each Conformer of 2^a

^{*a*} Calculations performed using B3LYP/aug-cc-pVDZ. ^{*b*} These are the starting values of the Cl(14)-Si(2)-C(1)-Si(3) / Cl(15)-Si(3)-C(1)-Si(2) dihedral angles in degrees; no interconversion was observed upon optimization. See Figure 1 for atom numbering. ^{*c*} Gibbs free energy in kJ mol⁻¹ (ZPE corrected). ^{*d*} Calculated at 485 K.

 Table 3. Indicative Dihedral Angle, Symmetry, Relative Energy, and Proportion for

 Each Conformer of 3^a

	indicative	point-group	relative	
conformer	dihedral angle ^b	symmetry	energy ^c	proportion ^d
3a	-160 / -40	C_1	0.00	0.785
3b	-160 / 80	C_1	8.76	0.086
3c	-160 / -160	C_2	8.63	0.044
3d	80 / 80	C_2	6.07	0.085

^{*a*} Calculations performed using B3LYP/aug-cc-pVDZ-PP. ^{*b*} These are the starting values of the Br(14)–Si(2)–C(1)–Si(3) / Br(15)–Si(3)–C(1)–Si(2) dihedral angles in degrees; no interconversion was observed upon optimization. See Figure 1 for atom numbering. ^{*c*} Gibbs free energy in kJ mol⁻¹(ZPE corrected). ^{*d*} Calculated at 486 K.

The geometry optimizations showed that the four silyl branches surrounding the central carbon atom were arranged in a near-tetrahedral formation. In order to refine the experimental GED data, parameterized models were written in FORTRAN for each of **1–3**, describing all conformers of each species that were likely to appear in the sample. The parameters used in the models were based on the bond lengths and angles of the most abundant conformer of each species, due to the small difference (less than 0.5 pm) as suggested by the MP2/aug-cc-pVDZ(-PP) geometry optimizations, for the equivalent atomic distances between the conformers. Slight deviations in bond lengths and angles between different conformers were accounted for by applying fixed (non-refinable)

differences to the parameters. For **1**, **2**, and **3**, thirty-two, twenty-six and twenty-six parameters were used to describe six, four, and four conformers, respectively. A full and complete description of the models used to describe the molecules can be found in the Supporting Information, with full atomic coordinates for each conformer **1–3** can be found in Tables S5–S7.

Refinements of the experimental data were carried out using the SARACEN method,^{59–61} with adjustments made for the effects of vibrational motions using data from SHRINK.⁶² SARACEN restraint values were based on the MP2/aug-cc-pVDZ(-PP) calculations, while the ranges of values from a series of geometry optimizations were used to estimate the uncertainties in these values.

Of the parameters, twenty-seven, twenty-two, and twenty parameters were restrained, for **1–3**, respectively, while the rest refined freely. In each model parameters $p_{1}-p_{6}$ describe distances between pairs of atoms in the molecule, parameters $p_{7}-p_{14}$ are bond angles used to position atoms relative to one another, while parameters $p_{15}-p_{26}$ (and additionally $p_{27}-p_{32}$ for **1**) are sets of dihedral angles to position the four main branches in each molecule relative to each other. Tables S8–S10 contain full lists of parameters and values for each of **1–3**, respectively.

To refine the amplitudes of vibration, the individual atomic distance that produced the largest scattering effect under a particular peak was selected. All other atomic distances under that same peak (not including distances related to hydrogens on a methyl group) had their amplitudes of vibration tied to the selected amplitude at the calculated ratio, with the single amplitude was refined. For 1, 2 and 3, eleven, nine, and eleven amplitudes, respectively, were refined, five, one, and five of these restrained.

The refinements were initially carried out with the amount of each conformer fixed to the calculated proportions reported in Tables 1–3 for 1–3, respectively. Once the optimal refinement was obtained with these conformer amounts, some of the values were varied in order to determine the optimal amounts of conformer.

Full lists of interatomic distances, amplitudes of vibration, distance corrections, and SARACEN restraints for **1**, **2**, and **3** can be found in Tables S11–S13.

Figures 2(a)–(c) shows the experimentally obtained radial distribution curves for **1**, **2**, and **3**, respectively. Difference curves can also be seen underneath each radial distribution curve, showing how good a fit was obtained to the experimental data. The related molecular scattering curves for each molecule can be seen in Figure S1 a–c. The R_G factors obtained for the least-squares refinements of **1**, **2**, and **3** were 6.1%, 8.7%, and 10.9%, respectively, with R_D factors (which ignore off-diagonal elements of the weight matrix) of 3.1%, 7.4%, 7.2%, respectively. Ref. 63, and other references therein, gives a full explanation of the differences between R_G and R_D . The refined coordinates of each atom for all conformers of all three species studied can be found in Tables S14–S16.

Figure 2. Radial distribution curves and difference curves between theoretical and experimental data for molecules 1 (a), 2 (b) and 3 (c).



Tables 4 and 5 contain selected parameters that demonstrate the typical bond lengths, bond angles, and dihedral angles observed for each of **1–3**. As the models for the GED refinement were based on the most abundant conformer of each species, with fixed differences to allow for slight deviations between that and other conformers, the bond lengths and angles relating to the main conformer are shown. However, X–Si–C–Si

dihedral angles for all conformers are shown as these differ considerably between conformers of the same species. Experimental geometric parameters are presented as r_{h1} values, which are formally derived from the vibrationally averaged r_a values that are yielded by the electron diffraction experiments (and which are listed for each pair of atoms in Supplementary Information Tables S11–S13). Vibrational corrections are applied to the r_a distances, first accounting for the amplitude of vibrations, u_{h1} , which act along the vectors between atom pairs, and then by applying the perpendicular vibrational correction, k_{h1} , which is calculated using the SHRINK program. In total this means that for any given atom pair $r_{h1} \approx r_a + u_{h1}^2/r_a - k_{h1}$. The r_e values quoted are determined from the theoretical equilibrium distances obtained from the various quantum chemical calculations.

parameter	r _{h1}	r _e B3LYP	r _e MP2	<i>r</i> _e M06-2X
rC(48)–Si(49)	190.1(6)	193.0	191.6	190.2
<i>r</i> C(48)–Si(51)	191.0(6)	193.6	191.8	190.5
rSi(49)–C(59)	188.7(1)	190.2	190.1	190.2
<i>r</i> Si(49)–H(61)	153.3(21)	150.1	150.3	149.8
rSi(51)–C(57)	189.0(1)	190.2	190.3	189.4
$\angle C(48) - Si(49) - H(61)$	108.3(6)	107.2	107.6	107.6
$\angle C(48) - Si(49) - C(59)$	115.2(9)	115.0	114.0	112.9
$\angle C(48) - Si(51) - C(57)$	113.3(6)	112.7	112.2	112.1
$\angle C(57) - Si(51) - C(58)$	107.0(7)	106.0	106.4	106.4
$\angle C(59) - Si(49) - C(60)$	102.2(20)	105.0	105.8	106.6
<i>ϕ</i> H(14)–Si(2)–C(1)–Si(3)	-160.2(35)	-163.7	-160.6	-159.2
<i>ϕ</i> H(15)–Si(3)–C(1)–Si(2)	-41.2(15)	-42.5	-41.6	-42.1
<i>ϕ</i> H(61)–Si(49)–C(48)–Si(50)	-163.9(12)	-160.0	-159.5	-159.6
<i>φ</i> H(62)–Si(50)–C(48)–Si(49)	79.3(30)	75.9	76.2	78.9
<i>ϕ</i> H(108)–Si(96)–C(95)–Si(97)	81.8(16)	78.8	78.8	78.4
<i>ϕ</i> H(109)–Si(97)–C(95)–Si(96)	-43.3(13)	-40.0	-39.7	-40.3
<i>φ</i> H(155)–Si(143)–C(142)–Si(144)	-161.5(7)	-162.7	-161.7	-162.3
<i>φ</i> H(202)–Si(190)–C(189)–Si(191)	79.7(8)	77.7	78.4	78.2
<i>φ</i> H(249)–Si(237)–C(236)–Si(238)	-42.6(13)	-46.8	-45.3	-45.7

Table 4. Selected Experimental (r_{h1}) and Theoretical (r_e) Geometric Parameters for 1^a

^{*a*} Distances (*r*) are in pm, angles (\angle) and dihedral angles (ϕ) are in degrees. Atom numbering as described in Figure 1. *r*_e values were calculated using the aug-cc-pVDZ basis set for each respective theory.

parameter	2				3			
	$r_{\rm h1}$	r _e B3LYP	r _e MP2	<i>r</i> _e M06-2X	$r_{\rm h1}$	r _e B3LYP	r _e MP2	<i>r</i> _e M06-2X
rC(1)-Si(2)	190.1(3)	192.7	190.8	189.7	189.6(13)	193.0	190.7	190.1
rC(1)-Si(4)	193.8(3)	196.2	193.8	192.9	195.0(13)	196.5	193.8	193.2
rSi(2)-C(12)	187.7(7)	188.7	188.6	187.6	187.4(2)	188.9	188.6	187.7
rSi(2)–X(14)	208.3(2)	215.1	213.8	213.3	227.7(2)	232.3	229.1	230.3
rSi(4)-C(10)	188.7(5)	189.9	189.7	188.8	188.7(2)	189.8	189.7	188.7
$\angle C(1) - Si(2) - X(14)$	109.6(6)	109.5	108.2	108.6	110.6(7)	111.1	108.9	110.1
$\angle C(1)$ -Si(2)-C(12)	116.1(2)	115.6	115.4	115.3	116.3(7)	115.3	115.6	115.1
$\angle C(1) - Si(4) - C(10)$	111.9(3)	112.8	112.1	112.1	112.0(6)	112.8	112.2	112.2
$\angle C(10) - Si(4) - C(11)$	105.9(7)	105.5	106.1	106.0	106.0(7)	105.5	105.9	105.9
$\angle C(12) - Si(2) - C(13)$	107.9(13)	107.4	107.9	107.7	106.8(21)	107.8	108.5	108.4
$\phi X(14) - Si(2) - C(1) - Si(3)$	-156.7(9)	-159.2	-158.6	-158.9	-158.7(12)	-158.9	-157.9	-158.8
$\phi X(15) - Si(3) - C(1) - Si(2)$	-43.2(7)	-41.8	-41.2	-41.9	-41.9(13)	-42.3	-41.1	-42.2
<i>φ</i> X(61)–Si(49)–C(48)–Si(50)	-160.8(5)	-165.8	-165.6	-166.0	-163.1(11)	-166.4	-166.5	-166.8
<i>φ</i> X(62)–Si(50)–C(48)–Si(49)	74.7(7)	77.6	78.0	77.6	75.7(11)	78.04	78.4	78.2
<i>φ</i> X(108)–Si(96)–C(95)–Si(97)	-161.5(6)	-161.8	-161.1	-161.6	-161.3(13)	-162.0	-161.0	-161.7
<i>φ</i> X(155)–Si(143)–C(142)–Si(144)	76.9(4)	75.5	75.7	75.4	76.5(8)	75.2	75.4	75.0

Table 5. Selected Experimental (r_{h1}) and Theoretical (r_e) Geometric Parameters for 2 (X = Cl) and 3 (X = Br)^a

^{*a*} Distances (*r*) are in pm, angles (\angle) and dihedral angles (ϕ) are in degrees. Atom numbering as described in Figure 1. *r*_e values were calculated using the aug-cc-pVDZ basis set for **2**, and aug-cc-pVDZ-PP for **3**, for each respective theory, and are based on the most abundant conformer for each molecule.

For 1, it can be seen from Table 4 that the distances to the central carbon atom, C(48) for the most abundant conformer, have a range of only around 1 pm. This is true for both the experimental and computational results. There is generally good agreement between the GED-derived distances and those from quantum chemical calculations, with the largest deviation observed for the Si–H distance. It is possible that this is due to the poor scattering ability of the lighter H atoms, but it is also likely to be a product of the anharmonicity observed in the vibration between the relatively heavy Si and light H atoms. Comparing the calculations themselves, which were all performed using the aug-cc-pVDZ basis set, both the MP2 and M06-2X theories give values that match the GED values well. The largest deviation observed relates to $\angle C(59)$ –Si(49)–C(60), which differs by just under 3 degrees from the theoretical value. However, the parameters relating to this angle have been restrained according to the SARACEN method, so we should accept this value.

For 2 and 3, it can be seen from Table 5 that there is also reasonable agreement between calculated and experimental values. Perhaps the most striking difference between the structure of 1, and those for 2 and 3, is the effect of the electronegative Cl and Br atoms in the latter. For 2 and 3, the electron withdrawing properties of atom X, cause C–Si distances to the central atom to have a range of around 3 pm, and this is observed for both experimental and calculated values. Bonds to SiMe₃ groups [e.g. C(1)–Si(4)] are longer than those to SiMe₂X groups [*e.g.* C(1)–Si(2)]; this was not observed for 1. Again MP2 and M06-2X theories produced calculated values that are closest to the experimental values. The most significant deviations were for the Si–X distances [X = Cl (2), Br (3)], and this is likely due to the small size of the basis sets used (necessary because of computational restrictions).

As one would expect when replacing the H of **1** with the larger Cl and Br atoms in **2** and **3**, the experimentally-defined angles for C(1)–Si(2)–C(12) and C(12)–Si(2)–C(13) are larger in each case than the equivalent values for **1**. Such trends are also observed from the computational results.

As mentioned before, to find all possible conformers of each of 1–3, calculations were started with each XMe₂Si group set to one of three dihedral angles (–40, –160, and 80°), and the majority of optimized dihedral angles fell within 5° of the expected angles. While most of the refined dihedral angles were close to the computationally predicted values, the dihedral angle ϕ Cl(61)–Si(49)–C(48)–Si(50) for 2, deviated from the predicted computational range by 5°. However, we might expect more freedom in the range of dihedral angles.

All three theoretical methods (B3LYP, M06-2X, and MP2) gave similar dihedral angles for the same sets of atoms, with the largest discrepancies being 0.7 and 1.2° between predicted values for **2** and **3**, respectively. The three methods also predicted similar dihedral angles for the same set of atoms for **1**, although there were some larger discrepancies between methods, with the largest being 4.5° [relating to ϕ H(14)–Si(2)–C(1)–Si(3)]. The refined dihedral angles, which were all restrained using SARACEN,^{59–61} fell within 5° of the predicted calculations, tending also towards the expected dihedral angles.

In terms of the amount of each conformer present for each molecular species, it was found that for **1** there was little change in the R_G value of the refinement as the conformer ratio was adjusted. This is to be expected of **1** due to the nature of the hydrogen atoms on the silicon group, allowing for free rotation of the SiHMe₂ groups. Therefore the refinement for this species was performed with the conformers fixed at the proportions predicted in Table 1. For 2 and 3, a noticeable change in the R_G value was observed as the relative amounts of the two lowest energy conformers **a** and **b** (as predicted in Tables 2 and 3) were adjusted. The amounts of **c** and **d** remained fixed. How the R_G values vary for each of 2 and 3 is illustrated in Figure 3, which also shows 95% confidence level (represented by a horizontal bar). For 2, a relatively shallow minimum is observed around the proportion predicted in Table 2. Because of this, the conformer ratio for 2 was kept at the values seen in Table 2 for the final refinement. For 3, a more pronounced minimum is observed in Figure 3, with the final refinement performed where the proportion of conformers **a**:**b**:**c**:**d** was 0.764:0.106:0.045:0.085. The results for 2 and 3 show that the theory was accurate in predicting the relative amounts of the most abundant conformers.

Figure 3. Variation in $R_G/R_{G(min.)}$ for (a) **2** and (b) **3** as the proportions of conformers **a** and **b** are varied relative to each other. The proportion of conformers **c** and **d** remained fixed. The horizontal bar represents the 95% confidence limit for the data.



The refinements performed here for **1** can be compared to those for $(HMe_2Si)_3CSiH_3$, eleven conformers of which are reported in Ref. 39. The average bond lengths observed for the various C–Si distances for $(HMe_2Si)_3CSiH_3$ (equivalent to rC(48)–Si(49) and rSi(49)– C(59) in Table 4) were 189.8 and 188.8 pm, respectively. These differ only by 0.3 and 0.1 pm, respectively, from values seen for similar bonds in **1**. Slight deviations from the angles observed for $(HMe_2Si)_3CSiH_3$ are noted for $\angle C(48)$ –Si(49)–C(59), with that reported in Ref. 39 being 113.8(4)°, while for **1** the value was 115.2(9)°. The larger angle observed in **1**, is most likely due to added strain on the branch due to larger groups around the central carbon (two SiMe₃ groups and an SiMe₂H group), compared to less bulky groups (two SiMe₂H and one SiH₃) for (HMe₂Si)₃CSiH₃.

In the case of **2**, comparisons can be made with $(Me_3Si)_3CSiCl_3$, as seen in Ref. 38. Whilst the structures are quite similar, some structural differences are observed. In general, the bonds in $(Me_3Si)_3CSiCl_3$ are shorter than those in **2** by 1 to 5 pm. For example, the average distance from the central carbon atom to silicon [i.e. the mean of C(1)–Si(2/3/4/5)] for $(Me_3Si)_3CSiCl_3$ is 190.9(8) pm, compared to 192.0 pm for **2**. The average Si–C distance for an SiMe₃ branch is also shorter for $(Me_3Si)_3CSiCl_3$, at 187.8(6) pm compared to 188.7(5) pm for **2**. The largest observed difference in bond lengths occurs with the Si–Cl distance: 203.3(6) pm for $(Me_3Si)_3CSiCl_3$, and 208.3(2) pm for **2**. This is not surprising as the chlorine-containing moiety is quite different; $(Me_3Si)_3CSiCl_3$ exhibits stronger Si–Cl bonds than those in **2**, which is most likely due to that region being highly electronegative and drawing electrons towards it.

Differences are also observed between the two in relation to similar bond angles, with angles generally being wider for $(Me_3Si)_3CSiCl_3$ than for **2**. The C–Si–C angle in $(Me_3Si)_3CSiCl_3$ [which is equivalent to $\angle C(10)$ –Si(4)–C(11) in **2**] is 107.0(11)°, compared to 105.9(7)° in **2**, though this difference is not significant. The biggest difference is once again for a parameter relating to the chlorine atoms. The C–Si–Cl angle [$\angle C(1)$ –Si(2)– Cl(14)] in (Me_3Si)_3CSiCl_3 is 114.6(11)°, while it is only 109.6(6)° in **2**. This may be due to the added steric hindrance of three chlorine atoms in close proximity.

Solution-Phase Dynamic Structures. Extensive NMR experiments were performed for **2** and **3**, with full details given in the Supporting Information.

 $(BrMe_2Si)_2C(SiMe_3)_2$. The 400 MHz ¹H NMR spectrum of $(BrMe_2Si)_2C(SiMe_3)_2$ (**3**) shows, as would be expected, two resonances at room temperature: a slightly broadened singlet for the SiMe₂Br protons and a sharp singlet for the SiMe₃ signal (see Figure S2 in the Supporting Information). However, on lowering the temperature a much more complicated spectrum emerges and, at 213 K, the spectrum shows numerous signals in both the SiMe₂Br and SiMe₃ regions (Figure S2). The ¹H NMR spectrum at 201 K recorded at higher field (500 MHz, Figure S3) shows the SiMe₂Br region to have six large signals and at least six smaller signals, while the SiMe₃ region has eight larger signals and at least six smaller signals together with several unidentified signals thought to belong to impurities (Figure S3). The ²⁹Si NMR spectrum recorded at 300 K shows a signal at -0.35 ppm, corresponding to the SiMe₃ groups and a broad signal due to the SiMe₂Br region, which has begun to split out into several signals, extending from 24.38 to 22.11 ppm. These two main signals again split into numerous signals at 201 K (Figure S4) and, together with the ¹H spectra, this indicates the presence of more than one conformer at low temperature.

A 2D ¹H/²⁹Si NMR shift correlation spectrum of (BrMe₂Si)₂C(SiMe₃)₂ was recorded at 201 K in order to correlate ¹H NMR signals with ²⁹Si NMR signals. Each ²⁹Si NMR signal in the SiMe₂Br region of the spectrum (shown in Figure 4 and Figure S5) is expected to be associated with two different proton signals. This spectrum shows that proton signals at 0.94 and 0.87 ppm, labelled A and B, correlate with the ²⁹Si NMR resonance at 24.60 ppm (labelled I). The ²⁹Si NMR signal at 23.40 ppm (II) is associated with proton signals C and F at 0.86 and 0.779 ppm. The third, large silicon signal III at 21.85 ppm is linked to proton signals at 0.81 ppm (D) and 0.784 ppm (E). Proton signals α and γ concealed under a large peak at 0.86 ppm and at 0.767 ppm, correlate with the small silicon signal 1 at 24.00 ppm.

Silicon signal 2 at 23.29 ppm is linked with the ¹H NMR signals a and b at 0.96 ppm and hidden under a large peak at 0.779 ppm. Proton signals β and δ at 0.788 and 0.74 ppm, are associated with the ²⁹Si NMR signal at 20.92 ppm labelled 3.



Figure 4 2D 1 H/ 29 Si NMR shift correlation spectrum of the SiMe₂Br region of (BrMe₂Si)₂C(SiMe₃)₂ in CDCl₃/CD₂Cl₂ at 201 K.

As proton signals a and b are assumed to be due to a minor C_2 conformer of $(BrMe_2Si)_2C(SiMe_3)_2$, silicon signal 2 must also be associated with this conformer. Likewise, proton signals α , β , γ and δ assigned to minor conformer C_1 are linked to silicon signals 1 and 3, which must therefore be due to the same conformer.

For the SiMe₃ region of the ²⁹Si NMR spectrum (see Figure 5 and Figure S6) each signal is expected to be associated with three ¹H NMR signals. Interpretation of the shift correlation spectrum in a manner similar to that used for the SiMe₂Br region yields the assignments summarized in Table 6. Several small signals labelled by asterisks do not seem to correlate in a similar way to the ¹H NMR signals and are assumed to be due to impurities which can also be seen at low intensity in the room-temperature spectra.

Figure 5 2D 1 H/ 29 Si NMR shift correlation spectrum of the SiMe₃ region of (BrMe₂Si)₂C(SiMe₃)₂ in CDCl₃/CD₂Cl₂ at 201 K. A star denotes a peak assigned to an impurity.



Table 6 Summary of the 2D 1 H/ 29 Si NMR shift correlation assignments in (BrMe₂Si)₂C(SiMe₃)₂.

Major Conformers (C_1 and C_2)		Minor Conformers (C_1 and C_2)		
²⁹ Si	$^{1}\mathrm{H}$	²⁹ Si	$^{1}\mathrm{H}$	
Ι	A and B	1	α and γ	
II	C and F	2	a and b	
III	D and E	3	β and δ	
IV	K, L and N	4	g, h and k	
V	H, J and M	5	i, m and n	
VI	G, I and O	6	j, l and o	

Several ¹H NMR saturation transfer experiments on (BrMe₂Si)₂C(SiMe₃)₂ were recorded at 201 K to probe the exchange processes between different conformers at low temperature. Both large and small proton signals in the SiMe₂Br region of the proton spectrum were irradiated and it was clear that exchange between the major and minor conformers was occurring, but a full assignment of the enhanced signals is ambiguous in some cases due to

overlap between the signals (Figure S7). Similar experiments were also carried out for the SiMe₃ region signals and again while exchange processes could be observed, a full assignment could not be made (Figure S8).

A series of 126 MHz ¹³C{¹H} NMR spectra of $(BrMe_2Si)_2C(SiMe_3)_2$ was recorded from 293 to 213 K (Figure S9). At 293 K signals corresponding to the SiMe_2Br and SiMe_3 groups, are seen at 9.89 and 4.97 ppm respectively. As was seen for the ¹H and ²⁹Si spectra, the signals in the ¹³C spectrum split into a complicated pattern as the temperature is lowered (Figure S10). The signals are again consistent with the presence of major C_1 and a major C_2 conformers but a full analysis is hampered by the complexity and overlapping of several signals.

 $(ClMe_2Si)_2C(SiMe_3)_2$. The degree of steric crowding in $(ClMe_2Si)_2C(SiMe_3)_2$ is between that of $(HMe_2Si)_2C(SiMe_3)_2$ and $(BrMe_2Si)_2C(SiMe_3)_2$ but, at readily accessible temperatures, its NMR spectra are much more similar to those of the bromide described above. Variable-temperature ¹H NMR spectra are shown in Figures S11 and S12, and are reminiscent of those for $(BrMe_2Si)_2C(SiMe_3)_2$, showing two signals at room temperature and many at low temperature. Again this is consistent with the presence of more than one conformation being present at low temperature. The ²⁹Si{¹H} inverse-gated NMR spectrum of $(ClMe_2Si)_2C(SiMe_3)_2$ in CDCl₃ at 300 K shows two broad resonances at 25.66 and -1.09 ppm, corresponding to the SiMe_2Cl and SiMe_3 groups, respectively. On lowering the temperature, these signals split into several new peaks (Figure S13), leading to numerous signals with an overall chemical shift pattern similar to that seen for $(BrMe_2Si)_2C(SiMe_3)_2$ at 201 K (see Figure S4). 2D ¹H/²⁹Si NMR shift correlation spectra of $(ClMe_2Si)_2C(SiMe_3)_2$ in a CDCl₃/acetone-*d*₆ solvent mixture were recorded at 203 K (Figures S14 and S15) and

have a similar appearance to the analogous spectra for the bromide (Figures 3 and 4). However, the spectra for (ClMe₂Si)₂C(SiMe₃)₂ are less well resolved than for the bromide analogue and although they are consistent with the presence of a major C_1 and a major C_2 conformer together with minor conformers a detailed analysis has not been possible (See Supporting Information for a more detailed discussion). Several ¹H NMR saturation transfer experiments (Figures S16 and S17) at 203 K were carried out in a similar manner to those described above for (BrMe₂Si)₂C(SiMe₃)₂. The results of these saturation transfer experiments again appear to confirm that energy exchange processes at 203 K occur between small and large population conformers as well as between different low abundance conformers. It is not known what the symmetries of these conformers are. Comparison of all ¹H NMR saturation transfer experiments of (ClMe₂Si)₂C(SiMe₃)₂ with those of the analogous compound (BrMe₂Si)₂C(SiMe₃)₂ show that the energy exchange processes occur, as might be expected, in similar ways in both compounds in solution at low temperature, although exchange processes between minor conformers of (ClMe₂Si)₂C(SiMe₃)₂ cannot be confirmed. A series of 126 MHz ¹³C {¹H}NMR spectra of (ClMe₂Si)₂C(SiMe₃)₂ was also recorded from 293 to 203 K, (Figure S18 and S19). Unfortunately, the low-temperature spectra were complicated, and while they are generally consistent with the presence of the conformers described above several peaks are not observed, presumably due to accidental signal overlap. Thus, a full analysis cannot be given. The Supporting Information provides further data and a more detailed discussion.

To summarize, at low temperatures it was possible to assign peaks in the multinuclear NMR spectra to conformers with different point-group symmetries. For example, a C_1 and a C_2 conformer of **2** or **3** would be expected to give rise to six and three different proton

signals in the Me₃Si region, respectively. Therefore, the presence of nine large and nine small proton signals in the ¹H NMR spectra recorded for **2** and **3** suggests that are two C_1 and two C_2 conformers present. This is in close agreement with the results of the quantum-chemical calculations.

 $(HMe_2Si)_2C(SiMe_3)_2$. Similar ¹H and ²⁹Si NMR spectra were recorded for $(HMe_2Si)_2C(SiMe_3)_2$ at 213 K. These, however, showed no significant changes when compared to the corresponding spectra recorded at ambient temperature. The ¹H NMR spectrum at 213 K showed a singlet at 0.15 ppm (SiMe_3), a doublet (SiMe_2H) at 0.24 ppm and a septet (SiMe_2H) at 4.04 ppm. Two signals were seen in the ²⁹Si{¹H} INEPT NMR spectrum of (HMe_2Si)_2C(SiMe_3)_2, one at -16.49 ppm due to the SiMe_2H groups and one at -0.46 ppm due to the SiMe_3 groups. The proton coupled ²⁹Si NMR spectrum shows complicated multiplets. The signal at -16.49 ppm splits into two multiplets which selective decoupling ²⁹Si DEPT NMR experiments show are due to the expected large doublet ¹J_{Si-H} (182.3 Hz), a septet ²J_{Si-H} (6.4 Hz, coupling to two Me groups) and smaller doublet ³J_{Si-H} (3.2 Hz, coupling to Si-H on remote Si). The lack of dynamic processes being observed at low temperatures is presumably due to the relatively small size of H compared to the halides.

X-ray Crystallographic Study. Several unsuccessful attempts were made to carry out single-crystal X-ray diffraction structural analysis of $(Me_3Si)_2C(SiMe_2Br)_2$ at 100 K in an attempt to freeze out any dynamic disorder present. $(Me_3Si)_2C(SiMe_2Br)_2$ was determined to belong to the cubic space group $Pa\overline{3}$ with unit cell lengths of 12.58 Å. This space group requires complete disorder of bromine positions along with at least two different sets of silicon positions. The disorder present precluded the identification of any specific

conformer and no model structures could be obtained. A similar problem was noted previously for C(SiMe₂I)₄, which also gave a cubic cell with a = 12.982(1) Å.⁶⁴

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ASSOCIATED CONTENT

Supporting Information

Additional details relating to the GED experiments (Table S1); least-squares correlation matrices (Tables S2–S4); calculated coordinated and energies (Tables S5–S7); details from the GED models and refinements, including amplitudes of vibration and curvilinear distance corrections (Tables S8–S13); final GED coordinate (Tables S14–S16); plots of molecular-scattering intensity curves (Figure S1); details of the NMR spectroscopic studies

(Tables S17–S21, Figures S2–S19). This material is available free of charge via the Internet at http://pubs.acs.org.

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GRAPHICAL ABSTRACT

