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# Conformations, Structures, and Vibrational Spectra of Triethylchloro- and Triethylbromosilane Using Theoretical Methods, Gas Phase Electron Diffraction, and IR and Raman Spectroscopy

Manuel Montejo,† Derek A. Wann,‡ Francisco Partal Ureña,† Fernando Márquez,† David W. H. Rankin,\*,‡ and Juan Jesús López González\*,†

Physical and Analytical Chemistry Department, University of Jaén, Campus Las Lagunillas, E-23071, Jaén, Spain, and School of Chemistry, University of Edinburgh, West Mains Road, Edinburgh, United Kingdom EH9 3JJ

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Conformational studies of triethylchlorosilane (TECS) and triethylbromosilane (TEBS) and the elucidation of their gas phase molecular structures have been accomplished by the combined use of theoretical (ab initio and density functional theory) calculations and experimental data from gas phase electron diffraction experiments. Additionally, analysis of the experimental features observed in the IR and Raman spectra recorded for both compounds has allowed us to propose a complete description of the vibrational spectra of both compounds, including an explanation of certain bands, which can only be correctly assigned when more than one conformer is considered to be present.

### Introduction

Organohalosilanes are an important family of compounds with a wide range of applications in organic and organometallic synthesis. 1 For instance, alkylhalosilanes are important precursors of alkylsilanols, which are well-known intermediates in industrially important sol-gel processes using organometallic compounds.<sup>2</sup> Following a series of previous studies dealing with the molecular structures and vibrational spectra of a number of trimethylsilane derivatives, $^{3-6}$  we carry out a joint theoretical and experimental study of the molecular structures and vibrational spectra of two related triethylsilylhalo derivatives, namely triethylchlorosilane (TECS) and triethylbromosilane (TEBS). Few works in the literature deal with the vibrational spectra of the title molecules.<sup>7,8</sup> In those that do, both TECS and TEBS are treated as symmetric top rotators belonging to the  $C_{3\nu}$ symmetry group. From a structural point of view, there is only one previous work in which Volkov et al. report the microwave spectrum of triethylchlorosilane.9

The presence of three rotating ethyl groups makes it likely that there will be more than one minimum on the potential energy surfaces of TECS and TEBS. As such, the first task of this work was to perform a conformational study on both systems based on the calculated populations obtained by the application of the Boltzmann distribution equation and the data from the gas phase electron diffraction (GED) experiments. Thereafter, the multiconformer molecular structures of both compounds have been determined using a combination of experimental data from the GED experiments and the results from theoretical ab initio (RHF, MP2) and density functional theory (DFT, B3LYP) calculations. Finally, the IR and Raman spectra of TECS and TEBS have been recorded. These data, and those from theoretical calculations using the B3LYP

method, have been employed to determine a complete vibrational assignment for both molecules.

### **Experimental Section**

Samples of TECS (purity >99%) and TEBS (purity  $\geq 97\%$ ) were purchased from Sigma-Aldrich and used without further purification.

Gas Phase Electron Diffraction. Data were collected for TECS and TEBS using the Edinburgh GED apparatus. <sup>10</sup> An accelerating voltage of around 40 kV was used, representing an electron wavelength of approximately 6.0 pm. Scattering intensities were recorded on Kodak electron image films at nozzle-to-film distances of 127.81 and 284.12 mm for TECS and 92.26 and 258.61 mm for TEBS. Sample and nozzle temperatures were held at 293 K in the case of TECS, whereas for TEBS the sample and nozzle temperatures were 335 and 342 K, respectively, for the shorter distance and 313 and 323 K for the longer distance. The differences between the nozzle-to-film distances used for TECS and TEBS correspond to the necessary use of different ports for the room-temperature and high-temperature nozzles.

The weighting points for the off-diagonal weight matrices, <sup>11</sup> correlation parameters, scaling factors, and nozzle-to-film distances for both TECS and TEBS are given in Table S1 (Supporting Information). Also included are the electron wavelengths as determined from the scattering patterns for benzene, which were recorded immediately before or after the patterns for the sample compounds. The scattering intensities were measured using an Epson Expression 1680 Pro flatbed scanner and converted to mean optical densities as a function of the scattering variable, *s*, using an established program. <sup>12</sup> The data reduction and least-squares refinement processes were carried out using the ed@ed program<sup>11</sup> employing the scattering factors of Ross et al. <sup>13</sup>

**IR and Raman Spectroscopy.** IR spectra were recorded in the 300-4000 cm<sup>-1</sup> range for both TECS and TEBS in the

<sup>\*</sup> Corresponding authors. E-mail: jjlopez@ujaen.es; d.w.h.rankin@ed.ac.uk.

<sup>†</sup> University of Jaén.

<sup>&</sup>lt;sup>‡</sup> University of Edinburgh.

liquid and gaseous phases using a Fourier transform (FT) IR Bruker Vector 22 spectrophotometer, equipped with a Globar source and a deuterated triglycine sulfate detector, using standard cells for liquid and gas (10 cm path length), both equipped with CsI windows. All spectra were obtained at room temperature with a resolution of 1.0 cm<sup>-1</sup> and 100 scans.

The Raman spectra in the liquid phase of the samples were recorded with a Bruker RF100/S FT-Raman spectrometer, equipped with a Nd:YAG laser (excitation line 1064 nm, 600 mW of laser power) and a cooled Ge detector at liquid nitrogen temperature, using a standard liquid cell. The spectra were again recorded with a resolution of 1.0 cm<sup>-1</sup> and 100 scans.

Computational Details. All molecular orbital and DFT calculations reported in this work were performed using the Gaussian 03 program package.<sup>14</sup> Geometry optimizations for TECS and TEBS were carried out using MP2 and B3LYP methods, each with three different basis sets, namely, the standard split-valence 6-31G\*,15 the so-called DZP+diff described elsewhere, 16 and the augmented correlation-consistent double-zeta basis set of Dunning (aug-cc-pVDZ).<sup>17</sup>

For each of the conformers identified for each compound, an RHF/6-31G\* force field was calculated, providing accurate amplitudes of vibration and vibrational correction terms for use in the GED refinement. The SHRINK program<sup>18</sup> was used to obtain corrections based on a curvilinear representation of the atomic motions.

For the vibrational study, frequency calculations were performed at the B3LYP level with 6-31G\* and DZP+diff basis sets. To achieve a better understanding of the PED matrices, and in order to implement the scaled quantum mechanics force field (SQMFF) methodology, 19-21 the Cartesian force fields were transformed into more appropriate sets of so-called natural coordinates, <sup>19</sup> listed in Table S2 (Supporting Information). Some examples of the internal coordinates used for the naturalcoordinate definitions are shown in Figure S1.

Following the SQMFF methodology, the quadratic force constants were scaled according to  $F'_{ii} = \lambda_i F_{ii}$  and  $F'_{ij} =$  $(\lambda_i \lambda_j)^{1/2} F_{ij}$ , where  $F_{ii}$  and  $F_{ij}$  are the quadratic quantum mechanically computed force constants, and  $F'_{ii}$  and  $F'_{ij}$  are the scaled equivalents. ( $\lambda_i$  and  $\lambda_i$  are the scaling factors.) The scaling factors (associated with the force constants expressed in natural coordinates) have been refined for the main conformer of each compound (taking 1.000 as the starting value) during the linear root-mean-square fitting of the theoretical frequencies to those observed experimentally. The scaling factors obtained for the main conformer were then transferred directly to the remaining conformers in order to predict their vibrational spectra. The force-field transformations, refinement of scaling factors, and normal-mode analyses were performed using the MOLVIB program.<sup>22</sup>

### **Results and Discussion**

Theoretical Conformational Analysis. Searches for conformers of TECS and TEBS were performed by rotating the three ethyl groups. These can adopt either gauche (g) or anti (a) orientations with respect to the Si-X (X = Cl, Br) bond. The presence of eclipsed conformations has been discounted as these would have much higher energies.

Five conformers were identified as real minima on the potential energy surfaces of both TECS and TEBS, with no imaginary frequencies for the calculations performed at the MP2/ 6-31G\* level. The five conformers can be defined as follows: conformer I  $(g^+g^+g^+)$  has  $C_3$  symmetry, with the three ethyl groups in gauche positions with respect to the Si-X bond, and

TABLE 1: Relative Energies (kJ mol<sup>-1</sup>) of the Five Conformers Defined for Each of TECS and TEBS, Calculated at All Levels Used in the Present Work

		$\Delta G$ , kJ mol $^{-1}$						
method	basis set	I	II	III	IV	V		
Triethylchlorosilane								
B3LYP	6-31G*	2.85	0.00	3.12	0.27	2.03		
	DZP+diff	2.65	0.00	3.06	0.41	2.27		
	aug-cc-pVDZ	2.70	0.00	3.60	0.21	1.80		
MP2	6-31G*	2.92	0.00	3.71	0.87	4.40		
	DZP+diff	3.55	0.00	3.90	0.89	4.82		
	aug-cc-pVDZ	3.44	0.00	3.99	0.91	4.14		
Triethylbromosilane								
B3LYP	6-31G*	2.59	0.00	3.84	0.44	2.20		
	DZP+diff	2.71	0.00	3.33	0.58	2.92		
	aug-cc-pVDZ	2.93	0.00	3.73	0.55	2.47		
MP2	6-31G*	2.82	0.00	4.05	0.75	3.67		
	DZP+diff	3.27	0.00	4.44	0.92	4.59		
	aug-cc-pVDZ	2.97	0.00	4.49	0.91	3.69		

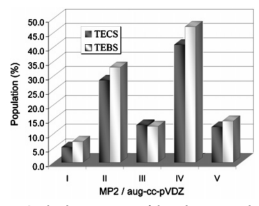
TABLE 2: Boltzmann Populations (%) for the Five Conformers of TECS and TEBS

		population (%)						
method	basis set	I	II	III	IV	V		
	Triethylchlorosilane							
B3LYP	6-31G*	5.0	21.7	13.7	39.2	20.4		
	DZP+diff	5.6	22.4	14.4	38.4	19.3		
	aug-cc-pVDZ	5.3	21.5	11.3	39.8	22.1		
MP2	6-31G*	6.3	28.0	14.2	40.6	11.0		
	DZP+diff	5.2	29.2	13.8	42.0	9.8		
	aug-cc-pVDZ	5.3	28.5	13.0	40.8	12.3		
Triethylbromosilane								
B3LYP	6-31G*	5.9	23.1	11.2	39.3	20.5		
	DZP+diff	5.9	24.3	14.1	39.2	16.5		
	aug-cc-pVDZ	5.4	24.0	12.1	39.2	19.3		
MP2	6-31G*	6.3	27.0	12.0	40.9	13.8		
	DZP+diff	5.9	29.6	11.5	42.2	10.8		
	aug-cc-pVDZ	6.3	28.3	10.7	40.3	14.4		

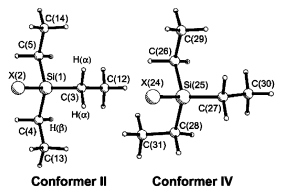
conformer II  $(ag^-g^+)$  has  $C_s$  symmetry. The remaining conformers have  $C_1$  symmetry, that is, conformer III  $(aag^+)$ , conformer IV  $(ag^+g^+)$ , and conformer V  $(g^+g^+g^-)$ . A  $C_{3\nu}$ symmetric (aaa) conformation was calculated to have one imaginary frequency, which described one of the three Si-C torsions. As such, the aaa conformation is not a real structure, rather a transition state on the way to conformer III.

The relative Gibbs free energies ( $\Delta G$ , kJ mol<sup>-1</sup>) calculated for each of the five conformers using MP2 and B3LYP methods with 6-31G\*, DZP+diff, and aug-cc-pVDZ basis sets are reported in Table 1 for both TECS and TEBS. Both the B3LYP and the MP2 methods calculate conformer II to be the global minimum.

The Boltzmann distribution equation has been used to take into account the relative multiplicities of the conformers based on the symmetry each possesses. When the Boltzmann equation distribution is implemented (using T = 293 K for TECS and T= 323 K for TEBS), both levels of calculation agree that conformer IV will have the highest population in the gas phase for both TECS and TEBS. As can be seen in Table 2, the calculated amounts of each conformer are quite similar when results from different methods are compared and also when TECS and TEBS are compared. Figure 1 compares the calculated (MP2/aug-cc-pVDZ) populations (in %) for the five conformers defined for both TECS and TEBS. Conformer IV (40.8% of TECS and 40.3% of TEBS) and conformer II (28.5% of TECS and 28.3% of TEBS) are calculated to be present in higher proportions than the remaining three conformers. The molecular structures and atom numbering for II and IV are



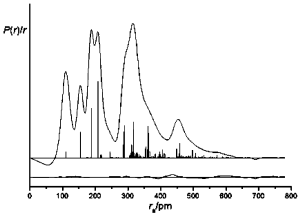
**Figure 1.** Graphical representation of the Boltzmann populations of the five conformers (MP2/aug-cc-pVDZ) for TECS and TEBS.



**Figure 2.** Molecular structures and atom numbering for the two main conformers of TECS (X = CI) and TEBS (X = Br).

presented in Figure 2. The sum of the amounts of conformers II and IV is approximately 70% of the total composition of each sample in the gas phase.

**GED Study.** A model, composed of geometrical parameters and containing the two conformers being studied, was written for TECS. An almost identical model was used in the refinement of TEBS, except that any parameters relating to chlorine were replaced by ones that substituted bromine.



**Figure 3.** Experimental and difference (experimental – theoretical) radial-distribution curves P(r)/r for TECS. Before Fourier inversion, the data were multiplied by  $s \exp(-0.00002s^2)/(Z_{Cl} - f_{Cl})(Z_C - f_C)$ .

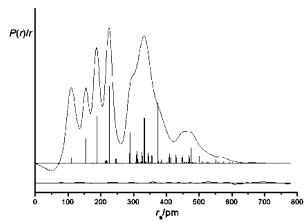
As can be seen in Tables S3 and S4 (Supporting Information), which list the calculated geometries of all the conformers at all levels of calculation, only small differences in parameters values are found depending on the method and on the basis set for both TECS and TEBS. Furthermore, the range of values for any parameter throughout the different conformers is quite small. This has allowed us to model the two conformers using the same set of 11 parameters. The only exception was the requirement for one extra torsional parameter  $(p_{12})$ , X(24)—Si-(25)-C(27)-C(30) (X = Cl, Br), for conformer IV. The 11 common parameters have as their starting points the average values calculated for the two conformers (MP2/aug-cc-pVDZ). Where differences exist in parameter values between conformers II and IV and the average values, these have been represented using fixed (i.e., non-refineable) differences, set at values determined from the MP2/aug-cc-pVDZ geometry optimiza-

Parameters 1–11 comprise four distances, namely, rSi-X ( $p_1$ ), the mean rC-H distance ( $p_2$ ), rC-C ( $p_3$ ), and rC-Si ( $p_4$ ). Four angle parameters were used:  $\angle \text{X}-\text{Si}-\text{C}$  ( $p_5$ ),  $\angle \text{Si}-\text{C}-\text{C}$  ( $p_6$ ), the average of the two ethyl  $\angle \text{H}-\text{C}-\text{Si}$  ( $p_7$ ), and the

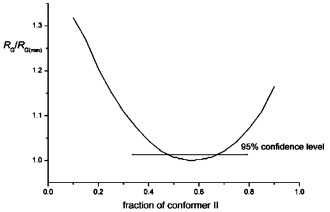
TABLE 3: Refined and Calculated Geometric Parameters with Ab Initio Values and Restraints for TECS (X = Cl) and TEBS (X = Br) from the GED Study (Distances in Picometers and Angles in Degrees)

parameter			triethylchlorosilane		triethylbromosilane		
descriptions		$\overline{\mathrm{GED}\left(r_{\mathrm{h1}}\right)}$	MP2/aug-cc-pVDZ <sup>a</sup>	restraint	$\overline{\mathrm{GED}\left(r_{\mathrm{h1}}\right)}$	MP2/aug-cc-pVDZ <sup>a</sup>	restraint
			Independent				
$p_1$	rSi $-$ X	208.6(1)	212.2		225.9(1)	227.9	
$p_2$	rC-H	110.6(1)	110.2		110.3(1)	110.2	
$p_3$	rC-C	154.5(1)	154.2		154.0(1)	154.1	
$p_4$	rSi-C	187.3(1)	188.8		187.2(1)	188.9	
$p_5$	$\angle X-Si-C$	106.7(1)	107.0		107.3(1)	106.9	
$p_6$	∠Si−C−C	114.0(1)	113.1		116.0(1)	113.1	
$p_7$	∠H−C−Si	109.2(4)	108.1	1.0	110.0(5)	108.1	
$p_8$	∠H−C−C	110.0(4)	111.3	0.5	111.5(3)	111.2	0.5
$p_9$	$\phi$ X-Si-C-C	62.7(8)	65.2	1.0	64.7(6)	65.1	1.0
$p_{10}$	$\phi$ X-Si-C-H( $\alpha$ )	58.6(19)	57.7	2.0	56.8(17)	56.9	2.0
$p_{11}$	$\phi$ X-Si-C-H( $\beta$ )	173.3(18)	172.3	2.0	171.9(16)	172.4	2.0
$p_{12}$	$\phi X - Si - C(27) - C(30)$	174.5(26)	174.5	3.0	177.7(25)	175.0	3.0
$p_{13}$	% conformer II	$57^{b}$	$41.1^{c}$		$33^{b}$	$41.3^{c}$	
			Dependent				
$p_{15}$	$\angle C(4)$ -Si-C(3)[II]	112.1(1)	-		111.5(1)		
$p_{16}$	$\angle C(4)$ -Si-C(5)[II]	112.0(1)			111.5(1)		
$p_{17}$	$\angle C(26)$ -Si- $C(27)[IV]$	112.2(1)	112.2		111.6(1)	112.3	
$p_{18}$	$\angle C(26)$ -Si- $C(28)[IV]$	112.2(1)			111.7(1)		
$p_{19}$	$\angle C(27)$ -Si- $C(28)[IV]$	112.0(1)			111.5(1)		

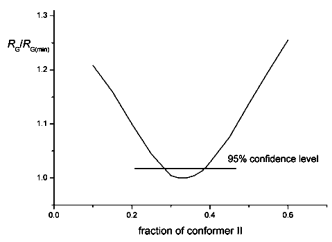
<sup>&</sup>lt;sup>a</sup> Average values for conformers II and IV; taken as starting point for the GED refinement. <sup>b</sup> See text and Figures 5 and 6 for uncertainties in the amounts of conformers. <sup>c</sup> Calculated from theoretical values considering the presence of only the two main conformers in the sample.



**Figure 4.** Experimental and difference (experimental – theoretical) radial-distribution curves P(r)/r for TEBS. Before Fourier inversion, the data were multiplied by  $s \cdot \exp(-0.00002s^2)/(Z_{Br} - f_{Br})(Z_C - f_C)$ .



**Figure 5.** Variation in  $R_G/R_{G(min)}$  with different amounts of conformer II for TECS.



**Figure 6.** Variation in  $R_G/R_{G(min)}$  with different amounts of conformer

average of three methyl  $\angle H-C-C$  (p<sub>8</sub>). The three dihedral angles common to the three conformers were  $\phi X-Si-C-C$  $(p_9), \phi X - Si - C - H(\alpha) (p_{10}), \text{ and } \phi X - Si - C - H(\beta) (p_{11}). (See$ Figure 2 for atom numbering and definitions of  $\alpha$  and  $\beta$ .) Parameter 12, as described in the previous paragraph, is only found in conformer IV. In addition to the 12 geometric parameters, a proportionality parameter has been included to allow the relative amounts of each conformer to be changed. Parameter 13 controls the amount of conformer II considered to be part of the gas mixture. Initially, this parameter remained fixed at the amount calculated using the Boltzmann equation,

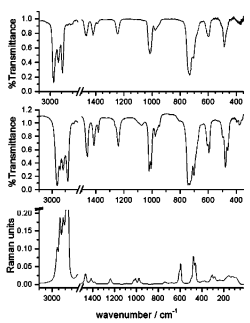


Figure 7. Experimental IR spectra of the gas phase (top) and the liquid phase (middle) and Raman spectrum of the liquid phase (bottom) for

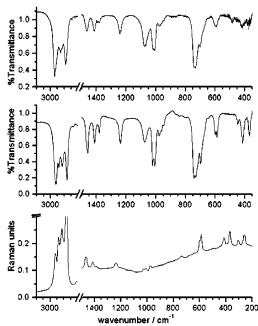


Figure 8. Experimental IR spectra of the gas phase (top) and the liquid phase (middle) and Raman spectrum of the liquid phase (bottom) for TEBS.

and only conformers II and IV (approximately 30% of conformer II and 40% of conformer IV, for both TECS and TEBS) were considered. Once the geometric parameters had been refined,  $p_{13}$  was varied to find the experimental composition of each of the gases.

Of the 12 geometric parameters used to define the twoconformer model of TECS, six were restrained using the SARACEN method.<sup>23</sup> For the refinement of TEBS, five were restrained. In Table 3, the refined parameters for TECS and TEBS are reported and compared with their calculated MP2/ aug-cc-pVDZ values. The C-Si-C angles are also included as dependent parameters. Additionally, eight groups of amplitudes of vibration were refined for both TECS and TEBS (no

TABLE 4: Experimental and Calculated Frequencies (in  $cm^{-1}$ ) of TECS<sup>a</sup>

		aled B3LYP					
conformer II		conformer IV			experimenta	$1^b$	
6-31G*	DZP+diff	6-31G*	DZP+diff	IR gas	IR liquid	Raman liquid	$P.E.D.^c$
44 A''	47 A"	47	51				$ au_{ m SiC}$
58 A'	64 A'	61	62				
80 A''	87 A"	71	73				
92 A'	92 A'	94	94				$\delta^{a}_{SiC3} + \rho_{SiC3} + sc_{Si}$
130 A'	129 A'	120	120			120 w,dp	$\rho_{\text{SiC3}}(\text{sc}_{\text{SiC3}})$
128 A"	135 A"	127	135			120 w, <b>a</b> p	$\delta^{\rm s}_{\rm SiC3}$
152 A"	154 A"	151	154			154 w,dp	$\rho_{\rm SiC3} + \delta^{\rm a}_{\rm SiC3}$
						134 w,up	$\rho_{\rm SiC3}$ 1 $\sigma_{\rm SiC3}$
162 A'	162 A'	155	157				
218 A"	219 A"	219	219			227 2	$ au_{\rm CC} ({ m sc}_{ m SiC3})$
234 A"	232 A"	233	232			235 w,p?	$ au_{ m CC}$
234 A'	234 A'	238	238				
		277	279			282 w,p	$ au_{ ext{CC}} + \delta^{ ext{s}}_{ ext{SiC3}} \  ext{sc}_{ ext{SiC3}} + \delta^{ ext{a}}_{ ext{SiC3}}$
299 A'	300 A'	277	217			305 w,p	Sesies 1 0 sies
303 A'	308 A'					, <b>r</b>	
			318				
360 A"	360 A"	201	200		200	250	
		381	380		380 m	378 vw <b>395 vw</b>	go (V)
				450 ab	461 a		$\operatorname{sc}_{\operatorname{SiC3}}(V)$
101 11	450 14	455	457.6	458 sh	461 s	461 m,p	$\nu_{\mathrm{SiCl}}\left(\mathbf{V}\right)$
481 A'	478 A'	477	476	486 s	478 s	476 m,p	$ u_{ m SiCl} $
				500 sh	497 m	500 sh	$\nu_{ m SiCl}$ (III)
		592	594	600 s	595 s	594 m,p	$ u_{ m SiC} $
605 A'	607 A'				606 s	608 m,p	
634 A''	634 A''	631	636				$ ho_{ m CH2} +  u_{ m SiC}$
		673	673		671 sh		
679 A''	680 A''				686 sh		
703 A'	706 A'	703	703	704 s	701 s		
724 A"	722 A"	700	, 00	724 sh	719 sh		$ ho_{\mathrm{CH2}}$
, 2	/22 11	730	730	732 vs	732 vs	733 w,dp	$\nu_{\rm SiC} + \rho_{\rm CH2}$
744 A'	744 A'	747	745	745 vs	743 vs	745 w,dp	VSiC   PCH2
				743 VS		743 w,up	. 1 4
937 A''	936 A"	939	942		947 w		$\rho_{\text{CH3}} + \text{tw}_{\text{CH2}}$
055 111	055 111	941	944		0.64 1		
955 A"	955 A"	961	961		964 sh		$ ho_{ m CH3} +  u_{ m CC}$
959 A'	960 A'	966	968				
971 A'	970 A'						
968 A"	971 A"	971	973	977 m	975 m	976 w,pp	$\nu_{\rm CC} + { m tw}_{ m CH2}$
974 A'	977 A'	975	979				
		1007	1005	1009 s	1006 s	1007 w,p	$\rho_{\text{CH3}} + \nu_{\text{CC}} + \text{wa}_{\text{CH}}$
013 A'	1011 A'	1018	1017	1013 s			$\nu_{\rm CC} + \rho_{ m CH3}$
014 A'	1011 A'						,
037 A"	1033 A"	1028	1025	1023 s	1021 s	1022 w,p	
238 A"	1233 A"	1242	1238	1244 m	1239 s	1236 w,p	$tw_{CH2}$
240 A"	1236 A"	1244	1240	1211111	1237 5	1230 W,p	CWCH2
245 A'	1244 A'	1248	1245				
256 A'	1244 A 1260 A'				1061 ab		
		1256	1258		1261 sh		wa <sub>CH2</sub>
259 A"	1261 A"	1257	1258				
262 A'	1265 A'	1262	1264				
385 A'	1383 A'	1379	1379	1385 w	1380 m	1381 w,p	$\delta^{ m s}_{ m CH3}$
386 A″	1383 A"	1380	1381				
387 A'	1384 A'	1381	1381				
415 A"	1412 A"	1401	1405	1419 m	1413 s	1412 w,dp	$sc_{CH2}$
421 A'	1419 A'	1418	1415				
425 A'	1422 A'	1423	1421				
456 A'	1455 A'	1455	1455	1465 m	1459 s	1464 w.dp	$\delta^{ m a}_{ m CH3}$
458 A"	1457 A"	1457	1458	1405 111	14373	1404 w,ар	O CH3
459 A'	1457 A'		1459				
		1458					
461 A"	1460 A"	1460	1461				
463 A'	1462 A'	1462	1462				
463 A''	1463 A"	1463	1463				
871 A''	2872 A"	2873	2874	2890 vs	2880 vs	2881 s,p	$ u^{ m s}_{ m CH2}$
875 A'	2875 A'	2875	2876				
881 A'	2878 A'	2884	2882				$ u^{ m s}_{ m CH3}$
885 A"	2881 A"	2887	2885				CIL
885 A'	2882 A'	2888	2886				
895 A'	2896 A'	2894	2894				v <sup>a</sup> arra
				2026 -	2015 -	2014	$ u^{ m a}_{ m CH2}$
912 A"	2914 A"	2910	2914	2926 s	2915 s	2914 s,p	
915 A'	2918 A'	2915	2919				
925 A''	2928 A"	2922	2925				$ u^{ m a}_{ m CH3}$
940 A'	2942 A'	2938	2942		2939 s	2939 s,p	

**TABLE 4: Continued** 

	theoretical/scaled B3LYP				$experimental^b$		
confo	ormer II	confo	ormer IV				
6-31G*	DZP+diff	6-31G*	DZP+diff	IR gas	IR liquid	Raman liquid	$P.E.D.^c$
2941 A"	2943 A"	2938	2943				
2941 A'	2944 A'	2940	2944				
2947 A"	2948 A''	2946	2948				
2956 A"	2955 A''	2955	2955	2968 vs	2961 vs	2960 vs,dp	
2957 A'	2956 A'	2955	2956			. 1	

<sup>a</sup> Calculations were performed at the B3LYP level using the 6-31G\* and DZP+diff basis sets for the three main conformers. The symmetry of each mode in conformer II is shown next to its calculated values. For conformer IV, all modes have A symmetry. The main terms of the PED for each mode are also shown. Bands that demonstrate the presence of different conformers are shown in bold. <sup>b</sup> Abbreviations used: vs = very strong, s = strong, m = medium, w = weak, vw = very weak, p = polarized, dp = depolarized, pp = partially polarized. <sup>c</sup> Abbreviations and Greek symbols (in order of appearance):  $\tau = \text{torsion}$ ,  $\delta = \text{deformation}$ ,  $\rho = \text{rocking}$ , sc = scissoring,  $\nu = \text{stretching}$ , tw = twisting, w = wagging. The superscripts a and s denote asymmetric and symmetric, respectively.

restraints were required). Tables S5 and S6 (Supporting Information) list the amplitudes of vibration for TECS and TEBS, respectively.

The success of the GED refinements can be gauged visually using Figures 3 and 4, where the radial-distribution curves and the theoretical minus experimental difference curves for TECS and TEBS are shown. The lines under the peaks in the radialdistribution curves represent individual interatomic distances, with the relative heights of the sticks proportional to the atomic numbers of the atoms involved and to the multiplicity of that distance and inversely proportional to the magnitude of the distance itself. The success of a refinement can also be quantified from the value of the goodness-of-fit function,  $R_G$ . For both TECS and TEBS, these values are low: 0.043 for TECS and 0.044 for TEBS. For TECS, the best fit relates to a composition of 57% conformer II and 43% conformer IV. For TEBS, these compositions were 33% conformer II and 67% of conformer IV. These results are in relatively good agreement with those obtained from the Boltzmann distributions at the experimental temperatures.

Figures 5 and 6 show the  $R_G/R_{G(min)}$  variations with respect to the composition of the sample as the amount of each conformer was changed. To quantify the uncertainties associated with these experimental compositions, lines have been drawn through the curves at  $R_{\rm G}/R_{\rm G(min)}=1.016$ , which, from Hamilton's tables,<sup>24</sup> represents a 95% confidence limit. The experimental mixtures for TECS and TEBS are therefore within two standard deviation of the calculated mixtures, albeit in different directions. Of course, there would also be small amounts of the other three conformers so the indicated uncertainties must be underestimates.

For further details of the refinement procedure, the leastsquares correlation matrices are given in Tables S7 and S8 (Supporting Information), and the GED atomic coordinates for each conformer of both TECS and TEBS are reported in Tables S9 and S10. Finally, the molecular scattering intensities curves are shown in Figures S2 and S3.

Vibrational Study. The IR spectra of the gas and liquid phases and the Raman spectrum of the liquid phase have been recorded for TECS and TEBS and are shown in Figures 7 and 8, respectively. As explained in previous sections, two different basis sets have been used for the vibrational analyses, in conjunction with the B3LYP functional. The calculated geometries at these levels (reported in Tables S3 and S4) are close to those determined by experiment for both compounds (see Table 3) and, therefore, no empirical corrections have been made to the calculated geometries.

The vibrational frequencies calculated theoretically at the B3LYP level with the 6-31G\* and DZP+diff basis sets have been fitted to the observed IR and Raman frequencies by means of a root-mean-squares procedure following the SQMFF methodology. 19-21 As explained in Computational Details, the scaling factors associated with the force constants, expressed in the set of natural coordinates defined for the TECS and TEBS molecules, were refined in order to reproduce the experimental frequencies, giving theoretical support to our proposal of vibrational assignment.

For the vibrational study, conformer IV has been used as a reference, since it is predicted to be the main conformer for both TECS and TEBS. The refined scaling factors obtained for this conformer were then transferred to the other four conformers, in order to predict their vibrational spectra. Using the scaled frequencies calculated for the two main conformers, we found it possible to explain almost all of the experimental features observed in the vibrational spectra. Some exceptions, which take into consideration conformers III and V, which are also predicted to be present in the sample in reasonable amounts, will be explained below. The scaled frequencies for the two main conformers at both levels of calculation, the experimental frequencies with their relative intensities, the proposed vibrational assignment, and the main terms of the PED are reported in Tables 4 and 5 for TECS and TEBS, respectively.

For clarity, and unless otherwise stated, throughout this discussion, we will refer to the B3LYP/DZP+diff results and the experimental bands observed in the IR spectra of the liquid phase. The latter can be justified by the small differences observed between the experimental gas and liquid frequencies and by the higher number of bands observed in the liquid phase, including those from Raman. Additionally, this strategy has given good results in previous studies with related systems.<sup>3-6</sup> The discussion below will concentrate on the parts of the vibrational spectra that cannot be explained if we consider only the presence of the main conformer. For the rest of the vibrational spectra, the scaled frequencies for the different conformers are almost identical. Complete vibrational assignments of the whole spectra have been accomplished for TECS and TEBS, as shown in Tables 4 and 5, respectively.

Triethylchlorosilane. The first evidence of the presence of different conformers of TECS, even in the liquid phase, is the weak polarized band observed at 305 cm<sup>-1</sup> in the Raman spectrum. That band, which cannot be assigned to conformer IV, is thus assigned to conformer II. This, in agreement with the results of the scaling, gives two normal modes at 300 and 308 cm<sup>-1</sup> (both belonging to the A' irreducible representation of the  $C_s$  symmetry group) describing the scissoring of the SiC<sub>3</sub> group. Moreover, a very weak band at 395 cm<sup>-1</sup> in the Raman spectrum cannot be assigned to any of the main conformers but can be assigned as the scissoring of the SiC<sub>3</sub> group in

TABLE 5: Experimental and Calculated Frequencies (in cm<sup>-1</sup>) of TEBS<sup>a</sup>

confo	conformer II conformer IV			experimental <sup>b</sup>			
6-31G*	DZP+diff	6-31G*	DZP+diff	IR gas	IR liquid	Raman liquid	$P.E.D.^c$
48 A"	47 A"	53	54	<u>U</u>	1	1	$ au_{ m SiC}$
57 A'	60 A'	59	61				₹SiC
80 A"	77 A"	75	72				
87 A'	85 A'	92	89				$\delta^{a}_{SiC3} + \rho_{SiC3} + sc_{SiC}$
119 A'	114 A'	110	108				$\rho_{\text{SiC3}}$ (sc <sub>SiC3</sub> )
122 A"	129 A"	127	132				$\delta^{\rm s}_{\rm SiC3}$
155 A'	151 A'	151	147				$\delta^{a}_{SiC3} + \rho_{SiC3}$
157 A"	155 A"	154	155				o SiC3   PSiC3
222 A"	219 A"	221	220				$\tau_{\rm CC} ({ m sc}_{ m SiC3})$
234 A'	231 A'	235	230				
238 A"	231 A"	240	237				$ au_{ m CC}$
262 A'	264 A'	252	253			259 w	$\nu_{\rm SiBr} + {\rm sc}_{\rm SiC3}$
300 A'	302 A'	301	303			305 w	
356 A"	359 A"	368	368		368 m	368 m	$sc_{SiC3} + \nu_{SiBr}$
							$sc_{SiC3} + \nu_{SiBr}$
413 A'	409 A'	418	418		415 s	413 m	$\nu_{\mathrm{SiBr}}$
		505	506	504 1	450 w	506	$\nu_{ ext{SiBr}}$ (III)
500 A/	500 A/	585	586	594 m,br	586 m	586 m	$ u_{ m SiC}$
599 A'	598 A'				599 m	596 m	
634 A"	634 A"	634	636				$ ho_{ m CH2} +  u_{ m SiC}$
681 A"	680 A"	674	672				
705 A'	704 A'	699	700	700 s	697 vs	699 vw	
724 A''	721 A"				713 sh		$\rho_{\mathrm{CH2}}$
		729	729	735 vs,br	730 vs	731 vw	$ u_{ m SiC} +  ho_{ m CH2}$
742 A'	741 A'	743	743		742 vs		
933 A"	937 A"	937	942	945 sh	946 w		$\rho_{\text{CH3}} + \text{tw}_{\text{CH2}}$
		940	944				
952 A"	955 A"	958	962		962 m		$\rho_{\text{CH3}}$
957 A'	960 A'	969	969				
965 A'	969 A'						
971 A"	972 A"	973	975	975 m	977 m	974 vw,p	$\nu_{\rm CC} + \rho_{\rm CH3}$
972 A'	978 A'	978	980			1	,
1011 A'	1011 A'	1007	1007	1008 s	1006 s	1005 vw,p	
1014 A''	1013 A"					·1	
1027 A'	1036 A'	1020	1020	1015 s	1022 s	1024 vw,p	
		1030	1028			· · · · · · · · · · · · · · · · ·	
1236 A"	1233 A"	1245	1237	1241 s	1238 s	1236 vw	tw <sub>CH2</sub>
1240 A"	1235 A"	1247	1239	12.15	12000	1200	*** C112
1242 A'	1243 A'	1247	1244				
1253 A'	1259 A'	1254	1257		1261 sh		wa <sub>CH2</sub>
1255 A"	1260 A"	1256	1258		1201 511		W CH2
259 A'	1264 A'	1261	1263				
1384 A'	1382 A'	1380	1379	1381 w	1380 w	1381 vw	$\delta^{ m s}_{ m CH3}$
1386 A"	1383 A"	1381	1380	1301 W	1300 W	1301 VW	O CH3
1386 A'	1384 A'	1382	1381				
1416 A"	1411 A"	1401	1405	1414 m	1412 m	1412 w,dp	CC CTTO
				1414 111	1412 111	1412 w,up	SCCH2
1422 A'	1418 A' 1421 A'	1416	1412 1419				
1427 A'		1423		1/65 h	1457 -	1462 4-	\$as
1455 A'	1454 A'	1454	1453	1465 br	1457 s	1462 w,dp	$\delta^{ m as}_{ m CH3}$
1458 A"	1456 A"	1457	1455				
1460 A'	1458 A'	1457	1456				
1460 A"	1459 A"	1460	1458				
1462 A'	1460 A'	1460	1459				
1464 A"	1461 A"	1462	1461	2000		***	
2871 A'	2868 A'	2871	2868	2889 vs	2878 vs	2879 vs,p	$ u^{\rm s}_{\rm CH2} $
2874 A'	2871 A'	2874	2871				
2881 A"	2876 A"	2884	2879				$ u^{\rm s}_{\rm CH3} $
2884 A'	2880 A'	2887	2882				
2885 A'	2881 A'	2888	2883				
2897 A''	2896 A"	2895	2893				$ u^{ m a}_{ m CH2}$
2914 A'	2913 A'	2911	2912	2925 s			
2917 A'	2917 A'	2916	2917				
2928 A"	2928 A"	2923	2924				
2939 A'	2941 A'	2937	2938		2938 vs	2937 s,p	$ u^{ m a}_{ m CH3}$
2941 A"	2941 A"	2939	2940		_,_,,,	»,P	. CID
2942 A'	2942 A'	2940	2941				
2942 A 2948 A''	2942 A 2947 A"	2947	2946				
2946 A 2955 A''	2947 A 2954 A"	2953	2952	2968 vs	2960 vs	2060 a n	
2955 A'	2954 A 2955 A'	2953 2954	2953	2700 VS	2700 VS	2960 s,p	

<sup>a</sup> Calculations were performed at the B3LYP level using the 6-31G\* and DZP+diff basis sets for the three main conformers. The symmetry of each mode in conformer II is shown next to its calculated values. For conformer IV, all modes have A symmetry. The main terms of the PED for each mode are also shown. Bands that demonstrate the presence of different conformers are shown in bold. <sup>b</sup> Abbreviations used: vs = very strong, s = strong, m = medium, w = weak, vw = very weak, p = polarized, dp = depolarized, pp = partially polarized. <sup>c</sup> Abbreviations and Greek symbols (in order of appearance):  $\tau$  = torsion,  $\delta$  = deformation,  $\rho$  = rocking, sc = scissoring,  $\nu$  = stretching, tw = twisting, wa = wagging. The superscripts a and s denote asymmetric and symmetric, respectively.

TABLE 6: Scaling Factors Obtained for Both Triethylsilyl **Derivatives** 

scaling	В3	LYP/6-3	31G*	B3I	YP/DZF	+diff
factor	TECS	TEBS	average	TECS	TEBS	average
SiC	1.031	1.022	1.026	1.019	1.016	1.017
CC	0.942	0.956	0.949	0.962	0.966	0.964
$CH (=CH_2)$	0.906	0.905	0.905	0.910	0.908	0.909
$CH$ ( $-CH_3$ )	0.897	0.897	0.897	0.903	0.901	0.902
CH <sub>2</sub> sc.	0.916	0.913	0.914	0.959	0.957	0.958
SiC <sub>3</sub> sc.	1.064	1.064	1.064	1.071	1.107	1.089
CH <sub>2</sub> rock	0.947	0.951	0.949	0.963	0.970	0.966
CH <sub>2</sub> wag	0.937	0.936	0.936	0.982	0.985	0.984
$CH_2$ tw.	0.925	0.951	0.938	0.982	0.975	0.978
sym. CH <sub>3</sub> def.	0.924	0.925	0.924	0.963	0.963	0.963
asym. CH3 def.	0.899	0.899	0.899	0.946	0.942	0.944
CH <sub>3</sub> rock	0.973	0.957	0.965	0.975	0.982	0.979
sym. SiC <sub>3</sub> def.	0.768	$0.768^{a}$	0.768	0.894	$0.894^{a}$	0.894
asym. SiC <sub>3</sub> def.	0.965	$0.965^{a}$	0.965	0.907	$0.907^{a}$	0.907
SiC <sub>3</sub> rock	0.736	$0.736^{a}$	0.736	0.714	$0.714^{a}$	0.714
SiC torsion	1.000	1.000	$1.000^{b}$	1.000	1.000	$1.000^{b}$
CC torsion	0.905	$0.905^{a}$	0.905	0.995	$0.995^{a}$	0.995

<sup>a</sup> These values were transferred from TECS, since there were no experimental bands for this region in the TEBS spectra. b Not refined because of the lack of experimental data.

conformer V, which agrees with the value calculated after the scaling of the force fields, that is,  $400 \text{ cm}^{-1}$ .

In the spectral region where the Si-Cl stretch appears, three bands are observed where only one would have been expected if there were just one conformer present. These bands appear at 461 cm<sup>-1</sup>, 478 cm<sup>-1</sup>, and 497 cm<sup>-1</sup>. The first is assigned as the SiCl stretching in conformer V (calculated at 457 cm<sup>-1</sup>). The second band is assigned to conformers II and IV, for which the calculated values are very close to one another and close to the observed value, 478 cm<sup>-1</sup> (A') in conformer II and 476 cm<sup>-1</sup> in conformer IV. The third band in that region is assigned to conformer III, for which the SiCl stretching normal mode is calculated to be at 502 cm<sup>-1</sup>.

At 606 cm<sup>-1</sup>, there is one strong band, which cannot be assigned to the main conformer. It is, therefore, assigned to conformer II in accordance with the calculated value for its SiC stretching normal mode, that is, 607 cm<sup>-1</sup> (A'). Similarly, the shoulder observed at 686 cm<sup>-1</sup> in the IR spectrum of the liquid is assigned to conformer II, which corresponds to a normal mode calculated at 680 cm<sup>-1</sup> with contributions from the CH<sub>2</sub> rocking and the SiC stretching vibrations.

Finally, another shoulder at 719 cm<sup>-1</sup> in the IR spectrum of the liquid (724 cm<sup>-1</sup> in the gas), which cannot be assigned to the main conformer, is assigned to the CH2 rocking mode of conformer II, in agreement with its calculated value of 722 cm<sup>-1</sup> (A"). As can be seen in Table 4, where the whole assignment is shown, the remaining bands can be assigned to conformer IV, and in the majority of the cases the scaled frequencies for the two main conformers are calculated to be very close.

Triethylbromosilane. In TEBS, the first evidence of the presence of more than one conformer is the weak band observed in the IR spectrum of the liquid phase at 450 cm<sup>-1</sup>, which is assigned to the SiBr stretching normal mode of conformer III, in agreement with its calculated value, 451 cm<sup>-1</sup>. This assignment is similar to that of the SiCl stretching mode in TECS, in which the band appearing in that region at a higher frequency was also assigned to conformer III. At higher frequencies, we observed two bands where only one was expected. The first, at 586 cm<sup>-1</sup>, is assigned to the SiC stretching normal mode of conformer IV (calculated at 586 cm<sup>-1</sup>), whereas the other band, observed at 599 cm<sup>-1</sup>, is assigned to the SiC stretch of conformer II, in accordance with its calculated value, 598 cm<sup>-1</sup> (A'). This assignment also agrees with that for TECS.

The last band that cannot be assigned to the main conformer is the shoulder at 713 cm<sup>-1</sup>. Once again, the proposed assignment is similar to that in TECS. The band has been assigned to conformer II, in accordance with the calculated value for the  $CH_2$  rocking in this conformer, that is, 721 cm<sup>-1</sup> (A"). In Table 5, the results for the whole vibrational assignment are shown and, as can be seen, the scaled frequencies for the two main conformers have very similar values and are in good agreement with the experimental bands.

In Table 6, the scaling factors obtained after the refinement procedure for TECS and TEBS, at the two levels employed, are reported. The values for TECS and TEBS are in good accordance, and so it is possible to average them, obtaining a set of parameters for each basis set.

#### **Conclusions**

It has been demonstrated that five different conformers represent minima on the PES of the title molecules. Theoretical analyses of the gas phase populations in terms of the Boltzmann distribution using  $\Delta G$  values calculated using the B3LYP and MP2 methods with the 6-31G\*, DZP+diff, and aug-cc-pVDZ basis sets indicated that conformer II  $(C_s)$  and conformer IV  $(C_1)$  account for around 70% of the sample composition.

The structures of TECS and TEBS have been determined experimentally by GED. For each, a model involving the two main conformers was used for the refinement of the experimental data. For TECS, the lowest  $R_G$  value was for a composition of 57% of conformer II and 43% of conformer IV, and for TEBS, the best fit represented 33% of conformer II and 67% of conformer IV, which are in good agreement with the results from the theoretical calculations.

The vibrational spectra of both TECS and TEBS have been completely assigned, following similar patterns in both compounds. The proposed assignment includes bands that could only be explained by considering the presence of more than one conformer. Although the majority of the bands could be explained by taking into account only the contributions from conformers II and IV, assignments of some bands used the scaled frequencies predicted for the remaining conformers, specifically, conformers III and V. Additionally, the use of the SQMFF methodology has allowed us to obtain two sets of scaling factors, one for each basis set used for the vibrational assignment. These parameters could, in the future, be transferred to other more reactive or unstable alkylsilane derivatives, such as silicon alkoxides or silanols, which take part in sol-gel processes of industrial interest, in order to predict their vibrational spectra. These data could be used in the monitoring of these processes, for which IR and Raman techniques have proved themselves to be very powerful tools.

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Supporting Information Available: Detailed information is available on the electron diffraction experiment (Table S1),

the natural-coordinate system employed (Table S2 and Figure S1), bond lengths, angles and torsions calculated at the MP2 and B3LYP levels (Tables S3 and S4), amplitudes of vibration from the GED experiment (Tables S5 and S6), correlation matrices from the least-squares refinements (Tables S7 and S8), coordinates from the GED structures (Tables S9 and S10), and molecular scattering intensity curves (Figures S2 and S3). This material is available free of charge via the Internet at http:// pubs.acs.org.

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