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## Schijnbare symmetrie bij esters van tetrathioorthokiezelzuur

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Document Version Publisher's PDF, also known as Version of record

Publication date: 1941

Link to publication in University of Groningen/UMCG research database

Citation for published version (APA):

Klasens, H. A. (1941). Schijnbare symmetrie bij esters van tetrathioorthokiezelzuur. Noordhoff Uitgevers.

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Download date: 28-10-2022

## XII. SUMMARY.

In the introduction we demonstrated theoretically that, by means of X rays, points of the crystal lattice may appear to be equivalent, though in reality they are not. It may also occur that the observed elements of symmetry do not exist, but are found with X rays in consequence of the cooperation of a great number of particles. The observed identical points are statistically identical and the observed elements of symmetry arise from a statistical distribution of particles over points possessing together those elements of symmetry.

Furthermore we pointed out that one has to reckon with such seeming symmetries, when deriving the symmetry of molecules from the X ray analysis of crystals. It is possible that these symmetries do not exist in reality. This is very obvious when the observed elements of symmetry are contradictory to the chemical image of the molecules. It appeared from the examples cited that isomorphism between related compounds of differing symmetry can point to the existence of such seeming symmetries.

Such an isomorphism has been found to exist between the compounds  $Si\{SC(CH_3)_3\}_4$  and  $Si\{SC(CH_3)_3\}_3SCH(CH_3)_2$ . The crystals of both esters are tetragonal.

The X ray analysis proves that the unit cell contains two molecules and that the space group  $D_{2d}^4$  (P  $\overline{42}_1c$ ) forms the foundation of the crystal lattice. Hence it follows that the molecular symmetry is  $S_4$  (or higher). This symmetry is possible for the tetra-tert. butyl ester but not for the tri-tert.butyl monoisopropyl ester.

In order to know to what extent the fourth alkyl group R in (BtS)<sub>3</sub>SiSR may differ from the three tertiary butyl groups Bt, without disturbing the X ray symmetry, we have introduced different groups. The new compounds were prepared according to the scheme:

$$(BtS)_3SiCl + NaSR = (BtS)_3SiSR + NaCl.$$

The following table gives

Tetragonal			
R	m.p.		
-C(CH <sub>3</sub> ) <sub>3</sub> *) -CH(CH <sub>3</sub> ) <sub>2</sub> *) -CH(CH <sub>3</sub> )(C <sub>2</sub> H <sub>5</sub> ) /CH <sub>2</sub> -CH <sub>2</sub>	160-161° 105° 79 <b>-</b> 80°		
−ĆH	104-105		
$CH_2$ — $CH_2$ — $C(CH_3)_2(C_2H_5)$	111-114		

Those esters which cry the tert. amyl ester) were e At the same time the te stannic and -orthogerman Si(SBt)<sub>4</sub>, were examined ments are brought togethe

To examine what influtert. butyl group by an icrystals, the following su

BtSSi{SCH(CH<sub>3</sub>)<sub>2</sub>}<sub>3</sub> (BtS)<sub>2</sub>Si{SCH(CH<sub>3</sub>)<sub>2</sub>}<sub>2</sub> (Measurable crystals of

The crystals of the di-itetragonal. We also examin the length of  $a_0$  and  $b_0$ 

<sup>1)</sup> The esters, indicated H. J. Backer and F. Stienstra

Tetragonal		Not tetragonal		
R	m.p.	R	m.p.	
-C(CH <sub>3</sub> ) <sub>3</sub> *) -CH(CH <sub>3</sub> ) <sub>2</sub> *) -CH(CH <sub>3</sub> )(C <sub>2</sub> H <sub>5</sub> ) -CH <sub>2</sub> -CH <sub>2</sub> -CH	160-161° 105° 79-80° 104-105°	$\begin{array}{c} -CH_3\ ^*) \\ -C_2H_5\ ^*) \\ -C_3H_7 \\ -C_4H_9 \\ -CH_2CH(CH_3)_2 \\ -CH(C_2H_5)_2 \\ /CH_2-CH_2 \end{array}$	43-44° 26-27° 62-62.5° b.p. 153-153.5°/1 mm 77-77.5° 27-29°	
$-C(CH_3)_2(C_2H_5)$	111-114°	$-CH$ $CH_2$ $CH_2$	64-65°	

ss (except the tert. amyl ester) were examined both optically and with X rays. At the same time the tetra-tert.butyl esters of tetrathio-orthostannic and -orthogermanic acids, which are isomorphous with Si(SBt)<sub>4</sub>, were examined with X rays. The results of the measurements are brought together in the following table.

	p.spec.	8	ω	$a_0$	Co
Sn(SBt) <sub>4</sub>	1.339	1.643 (Na-D)	1.640 (Na-D)	11.3	9.3
Ge(SBt) <sub>4</sub>	1.210	1.628 ( ,, )	1.622 ( ,, )	11.3	9.3
Si(SBt) <sub>4</sub>	1.087	1.615 (Hg <sub>578</sub> )	1.607 (Hg <sub>578</sub> )	11.2	9.2
(BtS) <sub>3</sub> SiCH(CH <sub>3</sub> ) <sub>2</sub>	1.068	1.600 (Na-D)	1.589 ( ,, )	11.2	9.3
$(BtS)_3SiCH(CH_3)(C_2H_5)$	1.080	1.600 ( ,, )	1.587 ( ,, )	11.4	9.4
(BtS) <sub>3</sub> SiSC <sub>5</sub> H <sub>9</sub>		1.614 ( ,, )	1.598 ( ,, )	11.3	9.4

To examine what influence the substitution of more than one tert. butyl group by an isopropyl group has on the form of the crystals, the following substances have been synthetised:

BtSSi{SCH(CH<sub>3</sub>)<sub>2</sub>}<sub>3</sub> (from NaSCH(CH<sub>3</sub>)<sub>2</sub> and BtSSiCl<sub>3</sub>) (BtS)<sub>2</sub>Si{SCH(CH<sub>3</sub>)<sub>2</sub>}<sub>2</sub> (out of NaSCH(CH<sub>3</sub>)<sub>2</sub> and (BtS)<sub>2</sub>SiCl<sub>2</sub>) Measurable crystals of the tri-isopropyl ester could not be made.

The crystals of the di-isopropyl ester are orthorhombic, pseudotetragonal. We also examined this ester with X rays. A difference in the length of  $a_0$  and  $b_0$  could not be observed. Both are 15.5 Å,

The following table gives a survey of these esters 1).

RY.	R	m.p.	R	m
theoretically that, by means ay appear to be equivalent, lso occur that the observed are found with X rays in at number of particles. The	$\begin{array}{c} -C(CH_3)_3 \ ^*) \\ -CH(CH_3)_2 \ ^*) \\ -CH(CH_3)(C_2H_5) \\ \hline \  \  \  \  \  \  \  \  \  \  \  \  \$	160-161° 105° 79-80° 104-105°	$ \begin{array}{c} -CH_3 \ ^*) \\ -C_2H_5 \ ^*) \\ -C_3H_7 \\ -C_4H_9 \\ -CH_2CH(CH_3)_2 \\ -CH(C_2H_5)_2 \\ /CH_2-CH_2 \end{array} $	43-4 26-2 62-6 b.p. 153-1 77-7 27-2
identical and the observed ical distribution of particles	$-C(CH_3)_2(C_2H_5)$	111-114°	-CH CH <sub>2</sub> -CH <sub>2</sub>	64-6
ements of symmetry. has to reckon with such			llise in the tetrago	

to exist between the com- $H_3$ )<sub>3</sub>}<sub>3</sub>SCH(CH<sub>3</sub>)<sub>2</sub>. The

ymmetry of molecules from

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to the chemical image of

oles cited that isomorphism symmetry can point to the

t cell contains two mole-(2,c) forms the foundation t the molecular symmetry le for the tetra-tert. butyl propyl ester.

fourth alkyl group R in ertiary butyl groups Bt, e have introduced diffeprepared according to

SiSR + NaCl.

<sup>1)</sup> The esters, indicated with \*), have already been synthetised by H. J. Backer and F. Stienstra, Rec. trav. chim. 54, 38 (1935).

 $c_0$  being 9.5 Å. The density is 1.070 and the unit cell contains two molecules.

The space group was found to be  $D_{2h}$  (Pnca). Hence it follows that the molecules possess a twofold axis of rotation.

To compare the esters of orthosilicic acid with those derived from tetrathioorthosilicic acid, some esters of the former acid have been synthetised according to the scheme:

 $SiCl_4 + 4 RONa = Si(OR)_4 + 4 NaCl$ 

In this way some new esters were obtained, namely the tetrasec.butyl, the tri-tert.butyl, the tri-tert.butyl monoethyl and the tri-tert.butyl monoisopropyl esters. Owing to steric hindrance, the tetra-tert.butyl ester could not be synthetised. An attempt to obtain this compound gave rise to the formation, among others, of the hexa-tert.butyl ester of disilicic acid, (BtO)<sub>3</sub>SiOSi(OBt)<sub>3</sub>). The tetraalkyl esters of orthosilicic acid are all liquid at room temperature.

STE

Het is mogelijk dat de sy Röntgenanalyse aan een mol is dan de werkelijke symmet

Haraszti en Széki hebben eugenolmethylaether een cyc J. Har

Het is niet waarschijnlijk,  $T_d^4$  ten grondslag ligt.

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De argumenten van Tay hydrolyse van alkylchlorid W. J

Uit de proeven van Ghebben verkregen.

M.