Electron Diffraction Investigation of the Molecular Structure of Trifluoromethanesulphonic acid (triflic acid)

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The molecular geometry of triflic acid is characterized by the following bond lengths (r_g) and bond angles from an electron diffraction study: S-C 183.3 \pm 0.5, F-C 133.2 \pm 0.2, S=O 141.8 \pm 0.2, S-O 155.8 \pm 0.3 pm, S-C-F 110.3 \pm 0.3, F-C-F 108.6 \pm 0.3, C-S=O 105.4 \pm 1.1, C-S-O 102.3 \pm 2.3, O-S=O 109.9 \pm 0.7, and O=S=O 122.0 \pm 1.3°. The heavy-atom-skeleton is staggered with respect to the rotation about the S-C bond with an estimated barrier of rotation of 15 kJ mol⁻¹.

The molecular structure of triflic acid has been investigated by electron diffraction as a continuation of our studies on the sulphone series [1-3]. The electron diffraction patterns were taken at a nozzle temperature of about 80 °C with the Oslo KD-G2 apparatus (for further details and references, cf. [4]). The reduced molecular intensities and radial distributions are shown in Figs. 1 and 2.

The rotation-dependent part of the radial distribution confirmed the expectation for a staggered form with respect to rotation around the C-S bond. Refinement of the rotation angle F-C-S-O showed some slight torsion similarly to CCl_3SO_2Cl [3]. On the other hand, the position of the O-H bond relative to the S-O bond could not be determined. The C-S and O-H bonds were assumed to be in anti position, and the O-H bond length and S-O-H bond angle were fixed at 96 pm and 115° in the final calculations. The CF₃ group had threefold symmetry, the C-S bond coincided with the symmetry axis, and the C-S-O plane bisected the O=S=O bond angle in our model. The results

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of the least-squares refinement are given in Table 1. Mean amplitudes of vibration have been coupled in groups as shown in Table 1.

The bond lengths (r_g) and bond angles characterising the triffic acid molecule together with the estimated total errors [5] are collected in the Abstract.

The sulphur bond configuration and the geometry of the CF₃ group are normal and consistent with earlier observations for analogous molecules. The C-S bond is rather long and is closer to that in CF₃SO₂Cl, 185.7 ± 0.6 pm [2] than to that in CH₃SO₂Cl, 176.3 ± 0.5 pm [6]. The lengthening of C-S bonds may be related to the electron-withdrawing ability of the CF₃ group [7].

Table 1. Results from the least-squares refinement of the structural parameters of triflic acid (the standard deviations are parenthesized as units in the last digit). The bond lengths (r_g) and bond angles with estimated total errors (see [5]) are given in the Abstract.

	r a (pm)/∢(°)	l (pm)	Key to the coupling scheme for the <i>l</i> -values
Independent parameters			
s-c	183.2 (3)	4.9 (4)	i
F-C	133.0 (1)	4.9 (2)	ii
S=0	141.7 (1)	3.9	ii
S-0	155.7 (2)	4.5	ii
=00=	248.0 (11)	6.4(2)	iii
S-C-F	110.3 (2)		
C-S=O	105.3 (8)		
C-S-O	102.3 (16)		
F-C-S-O	10.5 (13)		
Dependent parameters			
S.F	261.1 (2)	7.3	iii
=0C	259.6 (14)	9.5	iii
-0C	264.4 (30)	9.5	iii
=0F	373.4 (10)	6.8 (9)	iv
=0F	284.8 (16)	14.2 (10)	v
=0F	292.8 (20)	13.8	v
=0F	311.5 (22)	13.8	v
=0F	302.2 (20)	13.8	v
=0F	371.9 (9)	6.8	iv
-0F	308.1 (34)	16.2	v
-0F	380.6 (23)	7.8	iv
-0F	288.0 (39)	16.2	v
=00-	243.6 (7)	7.8	iii
$\mathbf{F} \dots \mathbf{F}$	216.0 (1)	5.9 (2)	vi
0 - 8 = 0	109.9 (5)		
F-C-F	108.6 (2)		
0=S=0	122.0 (9)		

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Fig. 1. Experimental (E) and theoretical (T) molecular intensities and the difference curves (Δ). The theoretical distributions were calculated from the parameters of Table 1.

The S=0 bond length is consistent with the empirical relationships established for XSO₂Y sulphones [8] between r(S=0) and the group electronegativities χ_X and χ_Y as well as between r(S=0) and the bond stretching frequencies (for frequency data, see [9, 10]).

The F...F distance of the CF₃ group $(r_g =$ 216.1 ± 0.3 pm) is the same as the one observed to

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- [5] Here $r_g \approx r_a + l^2/r_a$. The estimated total errors for distances $\sigma_t = [(0.001 r)^2 + 2 \sigma^2]^{1/2}$, for angles $\sigma_t =$ $\sqrt{2}\sigma$, where σ is the standard deviation from the least-squares results (see Table 1). For error estimation, cf. K. Hedberg and M. Iwasaki, J. Chem. Phys. 36, 589 (1962).
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Fig. 2. Experimental (E) and theoretical (T) radial distributions.

be strikingly constant in a large series of trifluoromethyl derivatives [11]. The O... O distances are considerably larger than twice the postulated nonbond radius of oxygen (viz. 113 pm [12]). It has been suggested that the O ... O distances in various OSO moieties are determined by a fine balance of non-bonded interactions and electron pair repulsions [13, 14].

The slight deviation from C_s symmetry of the CF₃SO₂O-skeleton as determined from electron diffraction may be a consequence of the torsional motion around the C-S bond. Accordingly, the average angle of the deviation, 10.5°, would suggest a barrier to internal rotation of about 15 kJ mol⁻¹ (cf. [15]). Similar barriers to rotation have been estimated for CCl₃SO₂Cl [3] and CF₃SO₂Cl [2].

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