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Trimethylsulfonium Methanesulfonate

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Trimethylsulfonium methanesulfonate

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Key indicators

Single-crystal X-ray study T = 120 KMean $\sigma(S-O) = 0.002 \text{ Å}$ R factor = 0.039 wR factor = 0.089Data-to-parameter ratio = 17.7

For details of how these key indicators were automatically derived from the article, see http://journals.iucr.org/e.

In the title compound, $C_3H_9S^+.CH_3O_3S^-$, a thermal decomposition product of dimethyl sulfoxide, both cation and anion lie on mirror planes. In the cation, the S atom lies 0.792 (2) Å out of the plane defined by the three C atoms, with S—C distances of 1.781 (2) and 1.786 (3) Å. In the anion, the S—O distances are 1.4556 (14) and 1.4646 (19) Å, and the S—C distance is 1.759 (3) Å.

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Comment

We have been studying the structure and mechanism of formation of colored products derived from resorcinarene macrocycles upon heating in dimethyl sulfoxide (DMSO) (Lewis et al., 1997, 2000; Davis et al., 1999). We isolated the title compound, (I), a decomposition product formed by prolonged heating of a solution of macrocycle in DMSO, and determined its structure to ascertain its identity. While 38 salts of the trimethylsulfonium ion and 62 salts of the methylsulfonate anion are present in the Cambridge Structural Database (December 2000, 224400 entries; Allen & Kennard, 1993), the structure of the title compound has not been previously reported. Decomposition of DMSO or its complexes to form trimethylsulfonium methanesulfonate has been previously reported as a result of heating (Banci, 1967; Arsenin et al., 1988) and γ irradiation (Gutierrez et al., 1977).

Both cation and anion lie across crystallographic mirrors. In the cation, the S atom lies 0.792 (2) Å out of the plane defined by the three C atoms. Geometric parameters (Table 1) are normal. Most of the H atoms are involved in $C-H\cdots O$ hydrogen bonding (Table 2)

Experimental

For the preparation of (I), the tetramethylresorc[4]arene 2,8,14,20-tetramethylpentacyclo[19.3.1.13,7.19,13.115,19]octacosa-1(25),3,5,-7(28),9,11,13(27),15,17,19(26),21,23-dodecaene-4,6,10,12,16,18,22,24-octol (250 mg, 0.46 mmol) was placed in a sealed tube along with 12 ml of DMSO and 3 ml of water. The mixture was heated at 493 K for 36 h. An aliquot was removed, and the solvent evaporated, yielding colorless crystals of the title compound.

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Crystal data

C3H9S+CH3O3S Mo $K\alpha$ radiation $M_r = 172.26$ Cell parameters from 7642 Orthorhombic, Pnma reflections a = 12.6157 (4) Å $\theta = 2.5 - 30.0^{\circ}$ b = 8.2419 (4) Å $\mu = 0.62 \text{ mm}^{-1}$ c = 7.5397 (8) Å $T=120~\mathrm{K}$ $V = 783.96 (9) \text{ Å}^3$ Plate, colorless $0.12 \times 0.10 \times 0.02 \text{ mm}$ 7 = 4 $D_x = 1.459 \text{ Mg m}^{-3}$

Data collection

KappaCCD diffractometer (with 7642 measured reflections Oxford Cryosystems Cryostream 1222 independent reflections cooler) 838 reflections with $I > 2\sigma(I)$ $R_{\rm int}=0.047$ ω scans with κ offsets $\theta_{\rm max} = 30.1^{\circ}$ Absorption correction: multi-scan (HKL SCALEPACK; Otwi $h=-17\to17$ $k = -11 \rightarrow 11$ nowski & Minor 1997) $T_{\min} = 0.936, T_{\max} = 0.988$ $l = -10 \rightarrow 10$

Refinement

Refinement on F^2 $w = 1/[\sigma^2(F_o^2) + (0.0326P)^2]$ $R[F^2 > 2\sigma(F^2)] = 0.039$ + 0.2622P] $wR(F^2) = 0.089$ where $P = (F_o^2 + 2F_c^2)/3$ S = 1.02 $(\Delta/\sigma)_{\text{max}} < 0.001$ $\Delta \rho_{\text{max}} = 0.40 \text{ e Å}$ 1222 reflections $\Delta \rho_{\min} = -0.37 \text{ e Å}^{-3}$ 69 parameters Only coordinates of H atoms Extinction correction: SHELXL97 Extinction coefficient: 0.0067 (18) refined

Table 1 Selected geometric parameters (Å, °).

S1-O1	1.4646 (19)	S2-C2	1.786 (3)
S1-O2	1.4556 (14)	S2-C3	1.781(2)
S1-C1	1.759 (3)		
O2-S1-O1	112.20 (7)	O2-S1-C1	106.11 (8)
$O2^{i}-S1-O2$	113.45 (12)	$C3-S2-C3^{ii}$	101.31 (15)
O1-S1-C1	106.11 (13)	C3-S2-C2	102.00 (10)

Symmetry codes: (i) $x, \frac{1}{2} - y, z$; (ii) $x, \frac{3}{2} - y, z$.

Table 2 Hydrogen-bonding geometry (Å, °).

$D-\mathrm{H}\cdots A$	D-H	$H \cdot \cdot \cdot A$	$D \cdot \cdot \cdot A$	$D - H \cdot \cdot \cdot A$
$C1-H1B\cdots O2^{i}$ $C2-H2A\cdots O1^{ii}$ $C3-H3A\cdots O1^{ii}$ $C3-H3B\cdots O2^{iii}$ $C3-H3C\cdots O1$	0.95 (2)	2.54 (2)	3.463 (2)	163.5 (18)
	1.06 (3)	2.41 (3)	3.382 (4)	152 (2)
	0.97 (2)	2.48 (2)	3.388 (3)	156.0 (19)
	0.95 (2)	2.44 (2)	3.279 (3)	147.3 (19)
	0.92 (3)	2.41 (3)	3.280 (2)	157.1 (19)

Symmetry codes: (i) $\frac{3}{2} - x$, -y, $z - \frac{1}{2}$; (ii) 1 - x, $\frac{1}{2} + y$, -z; (iii) 1 - x, $\frac{1}{2} + y$, 1 - z.

The coordinates of H atoms were refined, while their isotropic displacement parameters were assigned as $U_{\rm iso}=1.5U_{\rm eq}$ of the attached atom.

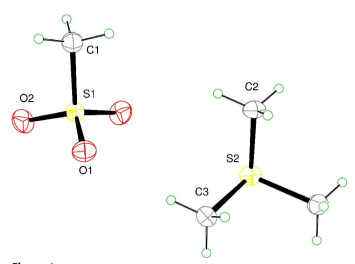


Figure 1
The atom-numbering scheme for (I) with ellipsoids at the 50% probability level.

Data collection: *COLLECT* (Nonius, 2000); cell refinement: *DENZO* and *SCALEPACK*; data reduction: *DENZO* and *SCALEPACK* (Otwinowski & Minor, 1997); program(s) used to solve structure: *SIR* (Altomare *et al.*, 1994); program(s) used to refine structure: *SHELXL*97 (Sheldrick, 1997); molecular graphics: *ORTEP*-3 (Farrugia, 1997); software used to prepare material for publication: *SHELXL*97.

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