A New Supramolecular Assembly Obtained by Reaction Between Thiosaccharin and Hexamethylenediamine

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The crystal structure of hexamethylenediammonium bis(thiosaccharinate) dihydrate, $[H_3N-(CH_2)_6-NH_3](tsac)_2 \cdot 2H_2O(tsac = C_7H_4NO_2S_2)$, the anion of thiosaccharin), was determined by single-crystal X-ray diffraction. It crystallizes in the monoclinic space group $P2_1/a$ with Z = 4. The thiosaccharinate moiety is planar and shows small but significant modifications in the bonding of the thioamide functional group as compared with the protonated neutral molecule. The ionic crystal is further stabilized by an extensive H-bonding network, which links the anions and cations into an infinite three-dimensional supramolecular assembly. The FTIR spectrum of the compound is briefly discussed in comparison with those of the neutral constituent molecules.

Key words: Hexamethylenediammonium Bis(thiosaccharinate) Dihydrate, Crystallographic Data, Supramolecular Adduct, IR Spectra

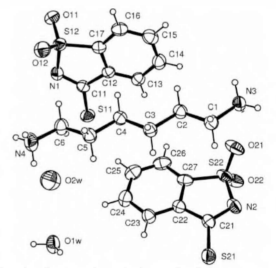
Introduction

In the last years we have synthesized and characterized a number of metallic complexes of thiosaccharin (for a recent review cf. [1]), and have now initiated a search to verify the possible formation of supramolecular adducts between thiosaccharin and some simple organic amines. In this paper, we present the first compound of this type, hexamethylenediammonium bis (thiosaccharinate) dihydrate, generated by interaction of thiosaccharin with hexamethylenediamine (1,6-hexanediamine).

Results and Discussion

Crystal and molecular structure

An ORTEP [2] drawing of the structure is presented in Fig. 1. Selected bond lengths and angles are shown in Table 1. As expected, the thiosaccharinate $C_7H_4NO_2S_2^-$ molecular skeletons are nearly planar (r.m.s. distance of atoms from least-squares planes less than 0.0194 Å). A comparison with the structure of the neutral thiosaccharin (tsac) molecule [3] shows that the major bonding changes in the charged moi-



1. ORTEP plot of the molecular structure of Fig. $[H_3N-(CH_2)_6-NH_3](tsac)_2 \cdot 2 H_2O$ showing the labeling of the non-H atoms and their displacement ellipsoids at the 30 % probability level.

ety occur at the thioamide functional group, as also predicted by a previous quantum chemical study [4]. In fact, upon deprotonation, the excess nitrogen elec-

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Table 1. Selected interatomic bond lengths (Å) and angles (°) in $[NH_3-(CH_2)_6-NH_3](tsac)_2 \cdot 2 H_2O$.

S(11)-C(11)	1.681(4)	N(1)–C(11)	1.326(5)
S(12)–O(11)	1.430(3)	N(3)–C(1)	1.488(5)
S(12)–O(12)	1.435(3)	N(4)–C(6)	1.478(6)
S(12)–N(1)	1.632(4)	C(1)–C(2)	1.500(5)
S(12)-C(17)	1.752(4)	C(3)–C(4)	1.505(5)
S(21)-C(21)	1.676(4)	C(5)–C(6)	1.506(5)
S(22)-C(27)	1.742(5)	C(11)–C(12)	1.479(5)
C(12)–C(17)	1.394(5)	C(22)–C(27)	1.394(5)
O(11)–S(12)–O(12)	115.6(2)	N(4)-C(6)-C(5)	111.6(4)
O(11)–S(12)–N(1)	110.0(2)	N(1)-C(11)-C(12)	114.0(4)
O(12)–S(12)–N(1)	110.1(2)	N(1)-C(11)-S(11)	123.8(3)
O(11)–S(12)–C(17)	111.9(2)	C(12)-C(11)-S(11)	122.1(3)
O(12)-S(12)-C(17)	111.0(2)	N(3)-C(1)-C(2)	110.3(3)
N(1)-S(12)-C(17)	96.6(2)	C(1)–C(2)–C(3)	113.2(3)
C(11)-N(1)-S(12)	111.4(3)	C(4)–C(3)–C(2)	112.7(3)

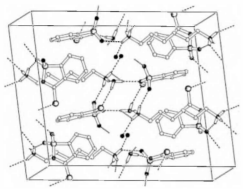


Fig. 2. Crystal packing of $[H_3N-(CH_2)_6-NH_3](tsac)_2$ · 2 H_2O as viewed down the *b* axis (PLATON [9]). The crystal *a* axis is along the vertical. For clarity, only hydrogen atoms involved in H-bonding are shown. H bonds are indicated by dashed lines. Sulfur atoms are denoted by larger shaded circles, while oxygen and nitrogen atoms are indicated by the smaller black discs.

tron mainly contributes to a delocalized N–C–S bonding. This is shown by the shortening observed in both N–C bonds (from 1.384 Å in neutral tsac to 1.326(5) and 1.327(5) Å in the two anions present in the investigated adduct) and the lengthening of the C–S bonds (from 1.622 Å in tsac to 1.681(4) and 1.676(4) Å in the present compound). In contrast, no significant changes are observed for the C(12)–C(17) and C(22)–C(27) and the sulfonyl S–O bond lengths comparing the neutral and charged molecules. On the other hand, the bond lengths and angles in the title compound agree well with those reported for the sodium, potassium, ammonium and bis(triphenylphosphinium) thiosaccharinate salts [5–8].

The crystal is further stabilized by the extensive 3-D net of hydrogen bonds shown in the PLA- TON [9] plot (Fig. 2). The H-bonding structure involves both hexamethylenediammonium NH₃ groups as donors, and as acceptors the exocyclic sulfur atom of one thiosaccharinate $[d(N3 \cdots S21) = 3.210 \text{ Å} \text{ and}$ \angle (N3–H···S21) = 161.4°], the sulforyl oxygen atoms of neighboring thiosaccharinate ions [the strongest with $d(N3\cdots O22) = 2.937$ Å and $\angle (N3\cdots O22) =$ 159.0° and the water oxygen atoms [the strongest] with $d(N4 \cdots O1w) = 2.813$ Å and $\angle (N4 - H \cdots O1w) =$ 161.2°]. This latter water molecule, in turn, acts as donor in a pair of H bonds, one with the other exocyclic S atom $[d(O1w \cdots S11) = 3.230 \text{ A and } \angle (O1w H \cdots S11$ = 158.1°], the other with the imino N atom (N1) of one thiosaccharinate anion $[d(O1w \cdots N1) =$ 2.835 Å and \angle (O1w–H···N1) = 168.4°]. Further details of the crystal H-bonding are presented in Table 2.

In the case of the recently described adduct of saccharin, $[H_3N-(CH_2)_6-NH_3](sac)_2$ (space group $P2_1/c$, Z = 4), the extended hydrogen bonding system also involves all six NH₃ hydrogen atoms as donors as well as the six O atoms belonging to the two saccharinate anions as acceptors [10].

Infrared spectra

The assignment of some of the most characteristic IR bands of the investigated compound is shown in Table 3. For comparative purposes we have also measured the spectrum of the free amine, and known data for the thiosaccharin molecule are also included [11, 12]. Standard references [13, 14] were used in the analysis and assignment of the spectra. Some aspects of the proposed assignments are commented as follows:

• In the case of the free amine a relative broad band is observed in the high-frequency range and only one of the two expected N–H stretching bands of the NH_2 groups could be observed. The same behavior was observed for the $v(NH_3^+)$ vibrations in the case of the salt.

• The characteristic deformational bands of the primary amine groups are absent in the salt, instead of which those of the ammonium groups ($R-NH_3^+$) are observed.

• The stretching vibrations of the water molecules generate a very broad and ill-defined band, centered at about 3490 cm^{-1} , pointing to a participation in hydrogen bridges of medium length [15]. The respective bending motion causes only a very weak shoulder at

d (D–H)	<i>d</i> (H···A)	Z DHA	<i>d</i> (D…A)	Α	Symm. operation
0.890	2.089	158.98	2.937	O22	[2-x, 2-y, 2-z]
0.890	2.344	161.40	3.201	S21	
0.890	2.183	142.97	2.942	O21	[2-x, 1-y, 2-z]
0.890	2.376	132.99	3.053	O22	[x-1/2, 3/2-y, z]
0.890	1.956	161.17	2.813	O1w	[3/2 - x, 1/2 + y, 1 - z]
0.890	2.070	150.53	2.878	O2w	[1/2 + x, 3/2 - y, z]
0.890	2.501	118.34	3.027	O12	
0.890	2.254	158.95	3.101	011	[2-x, 1-y, 1-z]
0.860	1.987	168.39	2.835	N1	[3/2 - x, 1/2 + y, 1 - z]
0.859	2.417	158.13	3.230	S11	[x - 1/2, 1/2 - y, z]
n H_2	$_{2}N-(CH_{2})_{6}-N$	~ .		NH ₃]	Assignment
		3	490 s, br		v(O-H) H ₂ O
33	38 vs				v (N–H)
			· · · · · · · · · · · · · · · · · · ·	VS	$v_{\rm as}({\rm NH_3^+})$
					$v_{\rm as}({\rm CH_2})$
		2	866 s		$v_{s}(CH_{2})$
16	37 s				$\delta(\mathrm{NH}_2)$ sciss.
		1	630 sh		$\delta({ m H_2O})$
		1	587 vs		$\delta_{\rm as}({\rm NH_3^+})$
		1	506 vs		$\delta_{\rm s}({\rm NH_3^+})$
14	52 s	1	456 vs		$v(CC), \delta(CH_2)$ sciss.
		1	348 vs, 1321 :	3	$v_{\rm as}({\rm SO}_2)$
		1	234 vs		v(NC), v(CC)
		1	115 vs, 1068	m	$v_{\rm s}({\rm SO}_2), \delta({\rm CCC})$
11	44 m	1	142 vs		V(CC)
10	61 m	1	047 w		v(CN)
		1	011 vs		v(CS), v(CC)
82	3 s, 796 s				$\delta(NH_2)$ out of plane
W		7	85 m, 700 w		$v(NS), \delta(CCC)$
	3 m, 712 s		56 s, 742 m		$\rho(CH_2)$
	0.890 0.800 0.	0.890 2.089 0.890 2.344 0.890 2.183 0.890 2.376 0.890 2.376 0.890 2.070 0.890 2.501 0.890 2.501 0.890 2.254 0.860 1.987 0.859 2.417 n H ₂ N-(CH ₂) ₆ -N 3338 vs 2943 s 2860 s 1637 s 1452 s 1144 m 1061 m 823 s, 796 s	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$

Table 2. Hydrogen bonds of $[H_3N-(CH_2)_6-NH_3](tsac)_2 \cdot 2H_2O$ (in A, angles in degrees).

Table 3. Assignment of some of the most characteristic IR bands of $[H_3N-(CH_2)_6-NH_3](tsac)_2 \cdot 2 H_2O, H_2N-(CH_2)_6-NH_2$, and thiosaccharin^a.

^a vs: very strong; s: strong; m: medium; w: weak; sh: shoulder; br: broad.

1630 cm⁻¹, at the higher energy side of the very strong $\delta_{as}(NH_3^+)$ band located at 1587 cm⁻¹.

• Some of the characteristic v(CC), $\delta(CCC)$ and CH_2 bands could also be unambiguously identified. Interestingly, the CH_2 -rocking mode shows the typical splitting [19] in both, the free amine and the adduct.

• Three of the most typical thiosaccharinate bands, *i. e.* the two SO₂ stretchings, and the v(CS) mode are clearly displaced to lower energies in $[H_3N-(CH_2)_6-NH_3](tsac)_2 \cdot 2 H_2O$ in comparison to their position in free thiosaccharin, surely as a consequence of its ionization [4] and involvement in hydrogen bonding, as commented above. Besides, both the $v_{as}(SO_2)$ and the $v_s(SO_2)$ vibrations appear as a singly split absorption band. On the other hand, the v(NC) and v(NS) bands of thiosaccharin are displaced to lower frequencies in the salt.

• The v(CN) vibration of the amine is also affected by the salt formation as are also some of the characteristic vibrations of the CH₂ groups. To conclude, this brief spectroscopic analysis is clearly compatible and consistent with the structural characteristics of the compound and reflects nicely its peculiarities.

Experimental Section

Synthesis of the adduct

Thiosaccharin (1.2-benzisothiazol-3(2H)-thione-1.1-dioxide) was obtained by heating of a mixture of 0.1 mol of saccharin (Aldrich, 99%) and 0.11 mol of P₂S₅ (Merck) under an N₂ atmosphere, according to the procedure described by Meadow and Cavagnol [16]. The reaction product was extracted with benzene and the crude orange-yellow product was recrystallized from benzene. Its purity was confirmed by the melting point (181–182 ⁻C) and by IR and Raman spectra [11].

The adduct was obtained by addition of 1.0 mmol of hexamethylenediamine to an aqueous solution (15 mL) containing 2.0 mmol of thiosaccharin. The mixture was heated with stirring for 10 min at 80 °C. The resulting solution was filtered, and the filtrate was slowly evaporated

Table 4. Crystal data and details of the structure refinement of $[H_3N-(CH_2)_6-NH_3](tsac)_2 \cdot 2 H_2O$.

Formula	$C_{20}H_{30}N_4O_6S_4$
Formula weight	550.72
Crystal system	monoclinic
Space group	$P2_1/a$ (Nr.14)
a, A	16.385(2)
b, Ă	7.927(1)
<i>c</i> , A	20.280(1)
β , deg	93.95(1)
$V(A^3)$	2627.8(5)
$D_{\rm c} ({\rm g} \cdot {\rm cm}^{-3})$	1.392
Z	4
F(000)	1160
θ Range for data collection	$2.18-67.99^{\circ}$
Index ranges hkl	$-19 \le h \le 0;$
	$-1 \le k \le 9;$
	$-24 \le l \le 24$
Reflections collected	5473
Independent reflections	4515
R(int)	0.050
Observed reflections $[I \ge 2\sigma(I)]$	3371
Data/restraints/parameters	4515/3/323
Goodness-of-fit on F^2	1.036
$R1/wR2[I \ge 2\sigma(I)]$	0.0671/0.1807
R1/wR2 (all data)	0.086/0.211
Largest diff. peak and hole, e A^{-3}	0.80/-0.39

at r.t. The obtained yellow microcrystalline product was recrystallized twice from methanol, and after the slow evaporation of the solvent a few, very small crystals adequate for crystallographic studies could be selected under the microscope.

Crystal structure determination

Measurements were performed at 296(2) K, with an Enraf-Nonius CAD-4 diffractometer employing graphite monochromated Cu K_{α} radiation ($\lambda = 1.54184$ Å). A colorless prismatic crystal of dimensions $0.31 \times 0.17 \times 0.08$ mm was used. Crystal data, collection procedures and refinement results are summarized in Table 4. The thiosacchari-

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nate and hexamethylenediammonium hydrogen atoms were located in calculated positions and refined with the riding model. The H positions of the $H_3N-(CH_2)_2-NH_3$ terminal ammonium groups were optimized by treating them as rigid groups which were allowed to rotate during the refinement around the corresponding N–C bonds. As a result, both NH₃ groups converged to staggered conformations. The H atoms of one water molecule were located in a difference Fourier map and refined isotropically with O–H and H···H distances restrained to target values of 0.86(1) and 1.36(1) Å, respectively. The other water molecule showed positional disorder and therefore its H atoms could not be determined reliably to be included in the final molecular model.

Intensity data were corrected for Lorentz and polarization effects as well as for absorption (PLATON [9]). The unit cell dimensions were obtained by least-squares refinement of the angular settings for 23 reflections in the range 16.54 $< \theta < 41.21^{\circ}$. Programs used were CAD4 [17], XCAD-4 [18], and SHELXS-97 [19] for data reduction and correction and structure solution and SHELXL-97 [20] for structure refinement.

CCDC 653328 contains the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre *via* www.ccdc.cam.ac.uk/data_request/cif.

Spectroscopic measurements

The infrared spectra of the compound as well as that of free hexamethylenediamine were recorded in the spectral range between 4000 and 400 cm⁻¹ on a FTIR Bruker IFS 66 spectrophotometer, using the KBr pellet technique.

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