Crystal structure of tetramethylformamidinium disulfide μ -Oxo-bis-(trichloroferrate(III))

メタデータ	言語: eng
	出版者:
	公開日: 2017-10-03
	キーワード (Ja):
	キーワード (En):
	作成者:
	メールアドレス:
	所属:
URL	http://hdl.handle.net/2297/3911

Crystal Structure of Tetramethylformamidinium disulfide μ-oxo-bis-
trichloroferrate(III)]
Hitoshi SENDA, Soh-ichi KITOH, Takumi SUGANAMI and Ko-Ki KUNIMOTO †
Department of Chemistry and Chemical Engineering, Faculty of Technology,
Kanazawa University, Kakuma-machi, Kanazawa 920-1192, Japan
† To whom correspondence should be addressed.

Oxidation of thioureas has been known to give a variety of reaction products depending on the substituents of the thioureas, the type of the oxidizing agent, the polarity of the reaction medium and other reaction conditions.¹ We have shown that oxidation of 1,1-diphenylthiourea and N-methylthiourea with iron (III) leads to 1,2,4-thiadiazoline derivatives.^{2,3} Kinoshita et al. studied the mechanism of oxidation reaction of substituted thioureas with BPO and suggested that 1,2,4-thiazolidine heterocyles are formed through abstraction of thiourea hydrogen and sulfur atoms.⁴ In this respect, it is interesting to examine oxidation products of thioureas having no urea hydrogen atoms. In this work we report on the crystal structure of the oxidation product of tetramethylthiourea with iron (III) chloride.

Tetramethylthiourea (1.32 g: 0.01 mol) and anhydrous iron (III) chloride (1.62 g: 0.01 mol) dissolved in absolute ethanol (30 ml) were reacted at room temperature for 1 h.

Dark-red gel-like precipitate was formed, which was subsequently separated from the mother liquor by decantation. The residue was re-dissolved in anhydrous acetone dried with molecular sieve (60 ml). The solution was kept at 4 °C in refrigerator for a few days. Precipitated crystals were filtered off and dried in a desiccator. Two different types of crystals were obtained from the same acetone solution; diamond shaped petal-like crystals (Form I) and rhomboid platelet-like crystals (Form II). X-ray analyses were carried out on these two crystal modifications.

The detailed measurement conditions and crystal data are listed in Table 1. The atomic scattering factors and anomalous dispersion terms were taken from the International Tables for X-ray Crystallography, Vol. IV. All calculations were performed using the program TEXSAN crystallographic software package. The final atomic parameters are listed in Table 2. The molecular structure is shown in Fig. 1, together with the atomic labeling scheme. Selected bond distances and bond angles are listed in Tables 3 and 4.

Forms I and II turn out to be polymorphic forms of formamidinium salts of μ -oxobis-[trichloroferrate(III)] ions where two tetramethylthiourea molecules are linked by a disulfide bond. Similar formamidinium disulfide salts were also derived from thiourea. There are two independent μ -oxo-bis-[trichloroferrate(III)] ions in the asymmetric unit of the Form I crystal, whereas only one ion exists in that of Form II. This ion occupies a

special position in the unit cell of the Form I crystal; the O(1) atom is situated on the mirror plane and the O(2) atom on the inversion center. Significant differences are observed in the Fe-O-Fe bond angles (Fe(1)-O(1)-Fe(2) and Fe(3)-O(2)-Fe(3) are 146.4(2)° and 180° for Form I and Fe(1)-O(1)-Fe(2) is 167.9(1)° for Form II). In consistent with variation in the Fe-O-Fe bond angles, the Fe-O-Fe antisymmetric vibrations are observed in IR spectra at 845 cm⁻¹ and 890 cm⁻¹ for Form I crystals and 875 cm⁻¹ for Form II crystals. Molecular geometries of the formamidinium disulfide part are generally similar for Form I and II crystals. The only significant differences are in the torsional angle around the C-S bond as shown in Table 4.

References

- 1. C. Christophersen, T. Ottersen, K. Seff and S. Treppendahl, J. Am. Chem. Soc., 97, 5237 (1975).
- 2. H. Senda and J. Maruha, Acta Cryst., C41, 1329 (1985).
- 3. H. Senda and J. Maruha, Acta Cryst., C41, 1626 (1985).
- 4. T. Kinoshita, S. Sato and C. Tamura, Bull. Chem. Soc. Jpn., 49, 2236 (1976).
- 5. "International Tables for X-ray Crystallography", vol. IV, The Kynoch Press, Birmingham, England, 1974.
- 6. "TEXAN", TEXRAY Structure Analysis Package, Molecular Structure Corporation,
 The Woodlands, TX, USA 1985.
- 7. O. Foss, J. Johnsen and O. Tvedten, Acta. Chim. Scand., 12, 1782 (1958).

Table 1 Crystal and experimental data

Table 1 Crystal and experimental data					
	Form I	Form II			
Formula: Formula weight: Crystal system: Space group:	C ₁₀ H ₂₄ Cl ₆ Fe ₂ N ₄ OS ₂ 604.86 orthorhombic Pnma Z=8	monoclinic P2 ₁ /n Z= 4			
Radiation: T $2\theta_{max}$	Mo Kα (0.71069 Å) 296 K 55.0°				
Scan type:	ω/2θ 2448	1224			
F(000)	a=15.209(3) Å b=31.741(4) Å c=10.360(1) Å	a=11.096(1) Å b=12.367(3) Å c=17.984(1) Å $\beta=92.761(8)^{\circ}$			
V= Dcalc= μ	5001(1) Å ³ 1.61 g/cm ³ R=0.038, R _w =0.043 19.77 cm ⁻¹	2465.0(6) Å ³ 1.63 g/cm ³ R=0.027, R _w =0.036 20.06 cm ⁻¹			
No. of reflections used No. of parameters Goodness-of-fit $(\Delta/\sigma)_{max}$	3509 (I>3.00σ(I)) 331 1.58 0.39	4222 (I>3.00o(I)) 322 1.44 0.24			
(Δ/ρ) _{max} Measurement: Program system: Structure determination: Refinement:	0.58 e/ Å ³ Rigaku AFC-5R TEXSAN direct method full-matrix least-square	0.36 e/ Å ³			

ydrogen atoms	u-uou 10	cooldinates	Ltactional	Table 2

			$_{l}v_{i}v_{i}$ g_{l} Ξ_{i} $\Xi(8)$	Beq=(4/
4.8(1) 3.81(3) 3.97(3) 3.27(8) 3.27(1) 4.6(1) 4.6(1) 4.6(1) 5.2(2) 5.2(2) 3.1(1) 4.8(1) 4.8(1)	(1)275(1) (4)3711(4) (6)31176,0 (1)21310,0 (1)2183(1) (1)2182,0 (1)2182,0 (2)2184(2) (2)4568(2) (2)456(2) (2)45(2) (2)419(2) (2)1310,0 (2)1310,0 (2)1310,0 (2)1310,0	0.1953(2) 0.17084(6) 0.17084(2) 0.16160(5) 0.16160(2) 0.3503(2) 0.3624(2) 0.3624(2) 0.3624(3) 0.4093(3) 0.4093(3) 0.4093(3) 0.4093(3)	(2)882.0 (2)882.0 (6)19234(6) (1)0490(6) (2)1490(6) (2)2459(2) (2)287(2) (3)287(2) (4)287(3) (4)287(3) (5)210.0 (4)287(3) (5)210.0 (5)210.0 (6)211.0 (6)3821.0 (7211.0 (8)3821.0	Bed=(4) C(10) C(10) C(3) C(4) C(6) C(6) C(6) C(1) C(1) C(1) C(1) C(1) C(1) C(1) C(1
5.0(3) 5.0(3) 7.8(4) 9.4(6) 5.73(4) 5.73(4) 5.46(4) 5.46(4) 5.55(4) 4.92(3)	(2)6262.0 (2)626.0 (3)450.0 (1)824.0 (2)2750.0 (2)4760.0 (2)4760.0 (2)4250.0 (3)4531.0 (4)5276.0 (4)16531.0 (4)16531.0	0.1561.0 0.1074(2) 0.1074(2) 0.1898(3) 0.1264(5) 0.1269(3) 0.2889(3) 0.1689(6) 0.1689(7) 0.1689(7) 0.1689(7)	(4)7278.0 (5)9429.0 (6)4117.0 (5)65139.0 (6)7852.0 (7)46747.0 (8)28210.0 (8)2724.0 (8)2724.0 (6)0822.0	C(8) C(8) C(8) C(10) C(10) C(10) C(10) C(10) C(10) C(10) C(10) C(10)
7.3(3) 3.78(5) 3.95(5) 3.9(2) 3.4(2) 4.9(2) 4.9(2) 4.8(3) 4.8(3) 5.1(3) 4.8(3)	2/I (1)1941.0 (1)4941.0 (2)8800.0- (2)800.0- (4)804.0 (2)800.0- (7)0700.0 (7)8120.0 (4)8876.0	0.08614(4) 0.08614(3) 0.06971(3) 0.0857(1) 0.1986(1) 0.1081(1) 0.0327(2) 0.127(2) 0.1463(2)	(7)06490(7) 0.66490(7) 0.74706(8) 0.7223(2) 0.7223(2) 0.7015(2) 0.700(2) 0.8909(3) 0.8041(4) 0.8659(4) 0.767(3)	(1) (1) (1) (1) (1) (1) (1) (1) (1) (1)
2.74(3) 3.44(4) 3.44(1) 5.15(9) 9.4(1) 4.88(6) 4.88(6) 3.9(2)	(7)48541.0 (8)18721.0- (8)18721.0- (8)818720.0 (2)0828.0 (1)1041.0 (2)8282.0- (1)8820.0 (1)8820.0 (1)8018.0 (4)0210.0	7/I (2)99520.0 (2)99520.0 (4)06990.0 (4)06940.0 (4)06940.0 (4)06940.0 (4)06940.0 (4)06940.0 (5)52670.0	x (2)19096.0 (3)70440.1 (4)(6)70440.1 (4)(6)70440.1 (1)(6)7778.0 (1)106.1 (1)2889.0 (1)2889.0 (1)620.1 (1)620.1 (1)620.1 (2)620.1 (2)620.1	Atom Form I Fe(1) Fe(3) Cl(1) Cl(2) Cl(3) Cl(4) Cl(5) Cl(5) Cl(5)
٠, ١, ٩		An mon to commi		7 210p I

Table 3 Selected bond lenghts (Å)

		Distance			
Atom	Atom	Form I	Form II		
Fe(1)	O(1)	1.757(4)	1.760(2)		
Fe(2)	O(1)	1.767(4)	1.757(2)		
Fe(3)	O(2)	1.7402(6)			
S(1)	S(2)	2.048(2)	2.059(1)		
S(1)	C(1)	1.784(4)	1.792(2)		
S(2)	C(6)	1.780(4)	1.795(2)		
N(1)	C(1)	1.323(5)	1.313(3)		
N(1)	C(2)	1.457(5)	1.468(4)		
N(1)	C(3)	1.465(6)	1.466(3)		
N(2)	C(1)	1.312(5)	1.325(3)		
N(2)	C(4)	1.476(6)	1.471(4)		
N(2)	C(5)	1.467(6)	1.471(4)		
N(3)	C(6)	1.306(5)	1.310(3)		
N(3)	C(7)	1.471(6)	1.470(4)		
N(3)	C(8)	1.474(6)	1.472(4)		
N(4)	C(6)	1.319(5)	1.326(3)		
N(4)	C(9)	1.464(9)	1.468(4)		
N(4)	C(10)	1.48(1)	1.467(4)		

Estimated standard deviations in the least significant figure are given in parentheses.

Table 4 Selected bond angles (°) and torsion angles (°).

given in natentheses.					
Estimated standard deviations in the least significant figure are					
26.2(2)	(5)4,551-	N(3)	C(1)	(1)S	(2)8
-154.4(2)	50.2(3)	N(1)	C(1)	(1)S	S(S)
24.3(2)	(4)6.64	N(4)	(9)D	S(2)	(1)S
-128.1(2)	(5)0.251-	N(3)	(9)D	S(2)	S(1)
			S	ou sugle	
153.2(2)	123.1(4)		N(t)	C(6)	N(3)
(2)7.911	120.1(3)		N(4)	C(6)	S(2)
117.1(2)	(5)8.911		N(3)	(9)D	S(2)
154.4(2)	153.1(4)		N(5)	C(1)	N(1)
118.4(2)	(E)E.TII		N(5)	C(1)	S(1)
117.2(2)	(5)2.911		N(1)	C(1)	(I)S
113.5(3)	(7)8.711		C(10)	N(4)	(6)2
123.6(3)	(7)1.121		C(10)	N(4)	C(9)
155.8(2)	151.1(5)		(6)ጋ	(4)N	(9)D
113.5(2)	113.2(4)		(8)D	N(3)	(Y)O
124.1(2)	153.5(4)		(8)D	N(3)	C(9)
155.1(2)	122.9(4)		$C(\lambda)$	N(3)	C(9)
115.4(3)	(4)9.6[1		C(2)	N(5)	C(4)
122.1(3)	(4)6.121		C(2)	N(5)	C(I)
122.5(3)	124.2(4)		C(4)	N(S)	C(1)
(E)7.EII	114.2(4)		C(3)	N(I)	C(5)
155.4(2)	122.6(4)		(£)O	N(I)	C(I)
153.5(2)	123.2(3)		C(5)	(Ì)N	(ĭ)́O
(8)20.101	102.6(1)		(9)D	(7)S	(I)S
(8)74.001	102.5(1)		C(i)	(I)S	S(2)\
	180.00		Fe(3)	(2)0	(£) ₉ H
(1)6.731	146.4(2)		Ee(2)	(I)Ŏ	Fe(1)
			_		Bond a
II m10		moiA	moiA	motA	толА
ə	gnA				

given in parentheses.

Figure Captions

Fig. 1 Molecular structure with the numbering of the atoms, (a) Form I and (b) Form II. Primed atoms are symmetrically related either via the mirror or the inversion center. Thermal ellipsoids of the non-hydrogen atoms scaled to enclose 50 % probability.



