A Detailed Refinement of the Crystal and Molecular Structure of Thiourea.

By N. R. Kunchur and Mary R. Truter.

A three-dimensional X-ray crystal-structure analysis of thiourea has The anisotropic thermal motion has been analysed and the bond lengths corrected for systematic errors arising from rotational oscillations of the molecule. The sulphur, carbon, and nitrogen atoms in each molecule are coplanar, with bond lengths C-N = 1.33 \pm 0.01 Å and C-S = 1.71 \pm 0.01 Å, and angles S-C-N = $122\cdot2^{\circ}\pm0\cdot6^{\circ}$ and N-C-N = $115\cdot6^{\circ}\pm1\cdot1^{\circ}$.

THIOUREA is conventionally written S=C(NH₂)₂ but its reactions show that the zwitterion -S-C(=NH₂+)NH₂ must also be considered. The Raman spectrum ¹ of the solid has been interpreted as indicating that the ionic form is predominant; on the other hand the approximate bond lengths, 2 C-S = 1.64 and C-N = 1.35 ± 0.1 Å, are confusing because both seem to show a good deal of double-bond character. A refinement of the X-ray analysis of this simple molecule has therefore been undertaken in order to provide more accurate bond lengths for comparison with theory and also with the bond lengths in co-ordination compounds containing thiourea.³

Wyckoff and Corey 2 found that there were four molecules in the orthorhombic unit cell (a = 7.68, b = 8.57, c = 5.50 Å) with space group $Pnma-D_{2b}^{16}$. (For convenience Wyckoff and Corey's b and c axes have been interchanged to conform with the space-group orientation in the International Tables.4) In each molecule the carbon and the sulphur atom lie on a mirror plane with the two equivalent nitrogen atoms on either side of it. The co-ordinates determined by two-dimensional analysis were C, 0.10, $\frac{1}{4}$, -0.14; S, -0.007, $\frac{1}{4}$, 0.12; N, 0.13, 0.125, -0.27₈. The molecule was planar within the limits of experimental error.

Isotropic Refinement by Three-dimensional Methods.—The intensities of the 260 reflections observable with copper radiation were measured; few spots of high values of sin θ were observed so that very little additional information could have been obtained by the use of radiation of shorter wavelength at room temperature. Preliminary calculation showed good agreement between the observed structure amplitudes, |Fobs|, and those calculated from Wyckoff and Corey's co-ordinates which were therefore taken as a starting point for three-dimensional analysis. The atomic scattering factors used were those given by McWeeny ⁵ for carbon and nitrogen and by James and Brindley for sulphur, ⁶ multiplied by a temperature factor $\exp(-B\sin^2\theta/\lambda^2)$. This theoretical scattering factor for sulphur was chosen in preference to empirical values 7 determined more recently because the latter are not independent of the anisotropic thermal motion and bonding in the structures from which they were derived.

The co-ordinates were refined in three successive stages by calculation of differential syntheses 8 with a back-shift correction for finite-series effects, and at each stage the

Kohlrausch and Wagner, Z. phys. Chem., 1940, B, 45, 229.
 Wyckoff and Corey, Z. Krist., 1932, 81, 386.
 Truter, Acta Cryst., 1957, 10, 785.

^{4 &}quot;International Tables for X-ray Crystallography," 1952, Kynoch Press, London.

⁵ McWeeny, Acta Cryst., 1951, 4, 51.

^{6 &}quot;International Tables for the Determination of Crystal Structures," Borntraeger, Berlin, 1935.
7 Cox, Gillot, and Jeffrey, Acta Cryst., 1949, 2, 356; Abrahams, ibid., 1955, 8, 661.
8 Booth, Trans. Faraday Soc., 1946, 42, 444.

2552

scale and temperature factors were altered to make the observed and calculated mean curvatures and peak values of the electron-density distributions more nearly equal. (Because of the uncertainty in their structure amplitudes, planes which gave reflections too weak to be observed were omitted from the refinement.) These refinement operations and their results are summarised below $\{R \text{ is the agreement index}, \}$

		$B(\mathbf{A^2})$	Max. co-ordin		
Cycle	sulphur	nitrogen	carbon	$oldsymbol{R}^+$	shift (Å)
1	4.0	4.0	4.0	0.20	0.18
2	3.5	4.5	4.5	0.16^{3}	0.04
3	3.5	4.5	4.24	0.15_{8}	0.01

 $\sum |(|F_{\text{obs}}| - |F_{\text{calc}}|)|/\sum |F_{\text{obs}}|\}$. The smallness of the co-ordinate shifts in the last cycle showed that no further improvement was possible by isotropic refinement, even though (as the figures below show) the mean B factor for carbon had not yet been adjusted to its best value. However, the need for anisotropic adjustment of the parameters was shown by the variation with direction of the observed curvature of the electron density in each atom; in the Table below r_x , r_y , and r_z are respectively the ratios of the observed curvatures, $\partial^2 \rho / \partial x^2$, $\partial^2 \rho / \partial y^2$, and $\partial^2 \rho / \partial z^2$, to the corresponding calculated curvatures; ρ is the peak electron density:

	Pobs/Peale	' x	' y	' z
C	1.01	1.06	1.10	1.14
S	0.98	0.92	0.99	1.09
N	0.94	0.85	1.03	1.01

It can be seen that for all atoms r_x is smaller than the other two ratios, which indicates that the thermal motion is greatest in the direction of the a axis, i.e., the axis which is most nearly normal to the planes of the molecules.

Determination of the Anisotropic Thermal Parameters.—A method of successive refinement for determining anisotropic thermal-motion parameters has been described by Cruickshank, who also gives the refinement equations.

After three cycles of anisotropic refinement the agreement index was reduced to 0·112, the following parameters, which resulted from the third cycle, being used to calculate the structure factors:

				v_{33}	023	v_{22}	013	012	011
	x/a	y/b	z/c	$(\times 10^5)$					
C ·	0.09151	0.25000	0.16278	1161	0	1331	2115	0	1140
S	0.00726	0.25000	0.11400	1770	0	1069	266	0	1789
N	0.13160	0.12011	0.27742	3885	819	1011	3026	173	2941

The figures given are those actually used, although, as shown below, they are not all significant. The parameters b_{ij} for each atom were used in the expression $\exp(b_{11}h^2+b_{12}hk+b_{13}hl+b_{22}k^2+b_{23}kl+b_{33}l^2)$ to give the temperature factor for each hkl. A list of observed and calculated structure factors is given in Table 1; the scale factor, obtained by the criterion given above, also made $\sum |F_{\rm obs}|$ equal to $\sum |F_{\rm calc}|$ where the summations are over all planes allowing for multiplicity. The poor agreement for the planes (020), (200), and (101) may be due to extinction; if these planes are omitted the scale factor is altered by 0.3% and the agreement index R is reduced to 0.109.

A fourth differential synthesis was carried out and the maximum change required in the atomic co-ordinates was 0.005 Å (in x_N). The final co-ordinates and their standard deviations (calculated from Cruickshank's formula ¹⁰) are:

	x/a	y/b	z/c	(Å)	$egin{array}{c} \sigma(x) \ (10^3 \ { m \AA}) \end{array}$	<i>y</i> (Å)	$egin{array}{c} \sigma(y) \ (10^3 \ { m \AA}) \end{array}$	$(\mathring{\mathbf{A}})$	$egin{array}{c} \sigma(z) \ (10^3 \ { m \AA}) \end{array}$
C	0.0916	0.2500	0.1632	0.701	8	$2 \cdot 134$	0	$\overline{0}.\overline{9}\overline{0}\overline{1}$	13
S	ō∙ōō 7 3	0.2500	0.1138	0.056	2	2.134	0	0.628	2
N	0.1322	0.1201	$\mathbf{\overline{0} \cdot \overline{2767}}$	1.012	11	1.025	10	1.527	11

The standard deviation of the electron density is 0.2e/Å3.

[°] Cruickshank, Acta Cryst., 1956, 9, 747.

¹⁰ Idem, ibid., 1949, 2, 65.

2553

TABLE 1.

			IADL			
F	F	F F	F F	F F	F F	F F
	(obs.) hkl		hkl (calc.) (obs.)	hkl (calc.) (obs.)	hkl (calc.) (obs.)	hkl (calc.) (obs.)
000 (160)	— 111		113 — —	125 13 14	102 2 3	273 10 14
020 -66	44 121		123 24 24	135 — —	$\frac{102}{112} - 34 41$	283 — —
040 17	17 131		133 -7 7	145 -9 10	122 -9 7	$\frac{203}{293}$ -7 7
060 -31	27 141		143 - 22 24	140 -3 10	132 19 17	200
080 28	20 151		153 — —	305 -10 12	142 9 10	403 6 7
0, 10, 0 -9	7 161		163 11 14	315 — —	152 - 17 17	413 -13 14
0, 10, 0 0	171		173 — —	325 12 12	162 — —	423 — —
200 51	43 181		183 - 6 7	335	172 16 15	433 21 20
210 -38	37 191			345 -11 14	182 — —	443
220 - 46	42 1, 10,		303 -16 17		192 -8 10	453 -16 17
230 - 14	90		313	505 -8 7		
240 35	97 301		323 11 10	515	302 6 7	603 6 10
250	- 311		333 —9 10	525 10 7	312 - 29 30	613 - 11 14
260 - 22	20 321		343 - 8 10	535 — —	322 -12 10	623
270 10	10 331		353 5 7	545 -10 10	332 28 27	633 7 7
280 14	10 341		363 8 10		342 12 14	
290 -4	3 351 3 361			011 14 14	352 -20 20	104 — —
2, 10, 0 -9	7 371		503 -8 10	031 39 38	362 — —	$\frac{114}{2}$ -8 7
	381			051 -23 20	372 12 14	124 — —
400 —	_		004 -19 17	201 12 15	382 — —	134 13 10
410 —	501	15 14	024 26 27	201 50 47	392 -9 8	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$
420 -21	20 511		044 - 28 27	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	502 6 7	154 -9 9
430 —	521	-11 10	064 15 14		$\begin{array}{cccccccccccccccccccccccccccccccccccc$	304
440 29	27 531		084 -7 7	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	512 — 21 20 522 — —	304
450 — 460 —11	- 541 10 551		204 24 20	251 -13 11	532 23 20	324 — —
470 —			214 — —	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	542 — —	334 — —
480 4	3 561	−7 8	224 21 20	271 10 9	552 -17 15	344 -7 8
490	701	7 7	234 — —	281 10 10	562	
4,10,0-6	3 711	4 5	244 - 15 15	291 -6 7	572 10 10	015 7 7
1, 10, 0	721		254 — —		•••	
600 13	10 731		264 13 7	401 6 8	702 — —	106 — —
610 10	10 741			411 -33 30	712 -12 10	116 10 11
620 -10	10		404 - 18 17	421 -9 10	722	126 — —
630	002		414 — —	431 9 10	732 10 10	136 - 12 10
640 7	7 022		424 12 14	441 8 10		146 — —
650 	042	18 17	434 — —	451 -9 10	013 - 37 41	156 9 7
660 - 7	7 202	5 7	444 7 7	461 -4 3	033 12 14	
670 — 5	5 919		454 — —	471 14 14	053 - 11 14	306 — —
680 5	3 222		464 8 3		073 16 17	316 13 13
000 5-	929			601 -4 3	093 -9 8	700
800 10	(949		604 -6 7	611 -13 14	000 11 14	506 — —
810 6	5 959		614 — —	621 — — 631 14 15	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	516 9 8
820 -7	7 262		624			
830 -6	7 272		634 — — 644 —8 8	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	
840 — 850 5	3 402		644 —8 8	661 — -	243 12 14	
990 9	3 402	17 17	105 -20 17	671 — —	253 -17 17	
101 41	34 103	-18 17	115 — —	681 -2 3	263 — —	

The peak value of the electron density for each atom is given below together with the ratios of the observed and calculated values of the electron densities and curvatures.

	$ ho_{ m obs}~(e/{ m A^3})$	$ ho_{ m obs}/ ho_{ m calc}$	r_x	r_y	r_z
C	7.0	0.95	1.00	1.00	0.97
S	28.0	1.00	1.00	0.99	1.02
N	$7 \cdot 2$	1.01	1.00	0.99	0.98

The ratios still differ slightly from unity, and the final thermal parameters (all $\times 10^3$) quoted below have been obtained by calculating the changes corresponding to the differences between the observed and calculated curvatures.

		b_{33}	b_{23}	$b_{f 22}$	b_{13}	$b_{f 12}$	b_{11}	$\sigma(b_{f 33})$	$\sigma(b_{f 22})$	$\sigma(b_{11})$
С		14	0	13	28	0	11	8	2	3
S	• • • • • •	18	0	11	3	0	18	2	0.4	0.6
N		40	8	10	32	Ī	29	8	2	3

The standard deviations, $\sigma(b_{ii})$, are derived from the expression:

$$\sigma^2(b_{ii}) = \{\frac{6}{5}\sum h_i^4 \sigma^2(F)\}/(\sum h_i^4 f)^2$$

given by Cruickshank,9 in which f is the scattering factor and $\sigma^2(F)$ is taken as $(|\mathbf{F}_{\text{obs}}| - |\mathbf{F}_{\text{calc}}|)^2$.

Analysis of the Anisotropic Thermal Motion.—Cox, Cruickshank, and Smith 11 have 11 Cox, Cruickshank, and Smith, Nature, 1955, 175, 766.

2554 Kunchur and Truter: A Detailed Refinement of the

shown that rotational oscillations of molecules can give rise to systematic errors in bond lengths determined by X-ray methods, so that an analysis of the thermal motion of the molecule is necessary before the final corrected bond lengths can be given.

The mean square amplitudes of vibration of an atom in any direction can be represented by the radius of an ellipsoid; the magnitudes and orientation of the axes of this ellipsoid can be calculated from the six numbers b_{ij} . In the direction of the unit cell edge of length a_i the mean square amplitude of vibration, μ_{ii}^2 , is given by $\mu_{ii}^2 = b_{ii}a_i^2/2\pi^2$. The cross terms, μ_{ij}^2 ($\mu_{ij}^2 = b_{ij}a_ia_j/4\pi^2$, where $i \neq j$), are required with the three values of μ_{ii}^2 to determine the dimensions and orientation of the ellipsoid; only if the values of b_{ij} are zero will the axes of the ellipsoid be parallel to the crystallographic axes. From the six values of μ^2 for all atoms we can derive (a) the r.m.s. amplitudes and directions of maximum and minimum thermal vibration of each atom with respect to the crystallographic axes, (b) the same quantities with respect to a set of molecular axes, and (c) the r.m.s. amplitudes of angular oscillation and translational vibration of the molecule as a whole assuming it to be a rigid body.

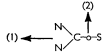
(a) The maximum and minimum r.m.s. amplitudes of vibration, μ_{max} and $\mu_{\text{min.}}$, derived from the lengths of the major and minor axes of the ellipsoids, and their direction cosines are given in Table 2 with, for comparison, the direction cosines of the normal to the molecular plane and of the bonds.

		TABLE 2.		
C	$\mu_{ ext{max}}$ (Å) 0.232 0.231 0.333 Normal	Direction (0.785, 0, 0.620) (0.992, 0, 0.129) (0.812, ±0.057, 0.581) (0.896, 0, 0.444)	μ_{\min} (Å) 0.040 0.163 0.169 C-S C-N	Direction $(0.620, 0, \overline{0}.78\overline{5}, (0.129, 0, \overline{0}.78\overline{5}, (0.129, 0, \overline{0}.59\overline{2}), (0.383, \pm 0.699, \overline{0}.60\overline{4}), (0.444, 0, \overline{0}.89\overline{6}, (0.237, 0.846, \overline{0}.47\overline{5}), (0.237, 0.846, \overline{0}.47\overline{5}, \overline{3}.78\overline{5})$
			C-N′	$(0.237, \ \overline{0}.\overline{846}, \ \overline{0}.\overline{478})$

The angles between the various directions are

between $\mu_{\max}(C)$ and normal, 12° between $\mu_{\min}(S)$ and C-S bond, 19° between $\mu_{\max}(S)$ and normal, 19° between $\mu_{\min}(N)$ and C-N bond, 14° between $\mu_{\max}(N)$ and normal, 10°

(b) Molecular axes were chosen thus:



with axis (3) as the normal to the molecular plane through the mass centre, o. Tensor transformations (Cruickshank 12) gave the following mean square amplitudes U_{ij} (all in Å2) referred to these axes:

	U_{11}	${U}_{f 12}$	U_{13}	${U}_{22}$	U_{23}	$U_{f 33}$
C	0.004	0	ō∙ ō 13	0.049	0	0.051
S	0.030	0	0.008	0.041	0	0.051
N	0.039	0.010	0.014	0.038	0.003	0.109
N'	0.039	0.010	0.014	0.038	0.003	0.109

From these we obtain the following r.m.s. amplitudes (in Å) of vibration:

along the normal, C, 0·227; S, 0·225; N, 0·330 along the C-S bond, C, 0·061; S, 0·172 along the C-N bond, C, 0·190; N, 0·173 perpendicular to the C-S bond in the molecular plane, C, 0·221; S, 0·202 perpendicular to the C-N bond in the molecular plane, C, 0·129; N, 0·211 Cruickshank, Acta Cryst., 1956, 9, 754.

2555

(c) For the molecule as a whole the tensors T (in $Å^2$) and ω (in radians²) for the mean square amplitudes of the translational and rotational motions can be obtained by suitable transformations.12 The values are:

T_{11}	T_{12}	T_{13}	T_{22}	T_{23}	T_{33}
0.0234	0	ŏ∙ōō 98	0.0332	. 0	0.0515
ω_{11}	ω_{12}	ω_{13}	ω_{22}	ω_{23}	ω_{33}
0.0470	0	0.0020	0.000	0	0.0077

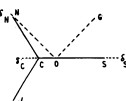
giving a r.m.s. angular oscillation of 5.0° about axis (3), 0° about (2), and 12.5° about (1), i.e., the main oscillation is about an axis through the C-S bond. The r.m.s. amplitudes of translation are 0.153 Å along axis (1), 0.182 Å along (2), and 0.227 Å along (3).

The effect of rotational oscillation is to make an atom, P, appear closer to the centre of oscillation, O (taken as the mass centre), by a small distance & which can be determined from the relation: 13

$$\delta = \frac{1}{2r} \left\{ \frac{s^2}{1 + s^2/q^2} + \frac{t^2}{1 + t^2/q^2} \right\}$$

where r is the uncorrected distance from O to P, $q^2 = \rho/(\delta^2 \rho/\delta x_{ti}^2)$, and s^2 and t^2 are the mean square amplitudes of oscillation about two axes through O orthogonal to OP.

For carbon and sulphur $t^2 = \omega_{22}r^2$ and $s^2 = \omega_{33}r^2$, whence $\delta_C = 0.0016$ Å and $\delta_S =$ 0.0045 Å and the correction to the C-S bond length is +0.006 Å. For nitrogen $s^2 = \omega_{33} r^2$



and $t^2 = r^2 \omega_{OG}$, where OG is perpendicular to ON in the plane of the molecule. ω_{OG} , calculated from ω_{11} , ω_{12} , ω_{22} , and the direction cosines of OG, has the value 0.0231 radians², whence δ_N = 0.0189 Å. The correction to the C-N bond is the component of δ_N along the bond minus the component of δ_C along the bond and is 0.018 Å.

Hydrogen Atoms.—When the anisotropic thermal motion for the other atoms had been determined, attempts were made to

locate the hydrogen atoms. First the intermolecular N-S distances were calculated; the shortest N-S distance (see Figure) is 3.42 Å which, being longer than the sum of the van der Waals radii (3·35 Å), indicates that there is no hydrogen bonding, so that no deductions could be made in this way about the positions of the hydrogen atoms.

By analogy with urea 14 the hydrogen atoms would be expected to lie in the plane of the heavy atoms forming, with the carbon atom, a trigonal arrangement about the nitrogen atom; these positions have been designated H₁ and H₂. Another position consistent with trigonally hybridised nitrogen would be with the H-N-H plane perpendicular to the S-C-N-N' plane; these positions are designated H₃ and H₄. However, three-dimensional values for $\rho_{obs} - \rho_{calc}$ at all four postulated hydrogen positions did not differ significantly from the standard deviation in the electron density $(0.2e/\text{Å}^3)$ and calculation of structure factors with hydrogen atoms at either H₁ and H₂ or H₃ and H₄ with an isotropic $B = 5.0 \text{ Å}^2$ gave no change in the agreement index.

Hydrogen atoms in any position should be shown by a $\rho_{\text{obs}} - \rho_{\text{calc}}$ projection, the most favourable one being along the [010] axis on to the mirror plane. This projection was calculated and is shown in the Figure where the positions of the heavy atoms and the previously postulated hydrogen atoms are marked. In fact the only appreciable electron density, $1.7e/\text{Å}^2$, is at x/a = 0.183, z/c = 0.400 and a line difference synthesis along y through this point showed peaks of 0.4 and $0.5e/Å^3$ at v/b = 3/4 and 1/4 respectively, and some additional peaks of $0.2e/Å^3$. The individual peaks are not of significant height and are 2 Å from the nearest atom (S).

These results suggest that the thermal motion of the hydrogen atoms is so large that

¹³ Cruickshank, Acta Cryst., 1956, 9, 757.

¹⁴ Worsham, Levy, and Peterson, ibid., 1957, 10, 319; Vaughan and Donohue, ibid., 1952, 5, 530.

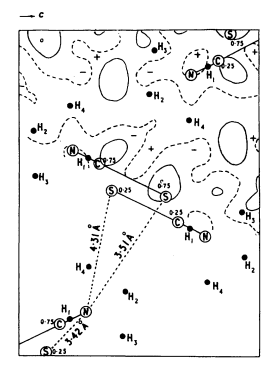
2556 Kunchur and Truter: A Detailed Refinement of the

the peak electron density has been reduced to an insignificant level; completely free rotation, however, is not consistent with some preliminary nuclear magnetic resonance observations.¹⁵

Dimensions of the Molecule.—For comparison the bond lengths and angles and their standard deviations are given below after (a) isotropic refinement, (b) anisotropic refinement, and (c) correction for rotational oscillation:

	C-S (Å)	σ (Å)	C-N (Å)	σ (Å)	S-Ĉ-N	σ	N−Ĉ−N	σ
(a)	1.710	0.016	1.319	0.015	123·0°	0.8°	114·0°	1·6°
(b)	1.707	0.012	1.311	0.012	$122 \cdot 2$	0.6	115.6	1.1
(c)	1.713	0.012	1.329	0.012	$122 \cdot 2$	0.6	115.6	1.1

The molecule is accurately planar, the deviation from the plane being less than 0.0003 Å for all atoms other than hydrogen. The values of σ are derived from the estimated



Difference projection along [010] with contour intervals of 1 e/Å². The zero contour is dotted and contours are drawn over half the area only. For the carbon and sulphur atoms y/b is given. The shortest N-S separations are indicated.

standard deviations in the atomic co-ordinates and do not allow for possible uncertainties in the corrections for rotational oscillation. Final values for the bond lengths are S⁻C = 1.71 ± 0.01 Å and C⁻N = 1.33 ± 0.01 Å, and for the angles S⁻C⁻N = $122.2^{\circ} \pm 0.6^{\circ}$ and N⁻C⁻N = $116^{\circ} \pm 1^{\circ}$.

Discussion

The carbon-sulphur (1.71 Å) and carbon-nitrogen (1.33 Å) bond lengths are significantly shorter than the generally accepted single-bond values, 1.82 and 1.475 Å, respectively. The double-bond lengths are not known accurately. For the carbon-nitrogen bond in s-triazine the order of which is at least 1.50, Lancaster and Stoicheff 16 obtained a very accurate value of 1.338 ± 0.001 Å, which is the same, within the limits of error, as in thiourea. This suggests that structures (I) and (II) must make a large contribution to the bonding, but the short carbon-sulphur bond with a bond order greater than 1.0

¹⁵ Emsley and Smith, Proc. Chem. Soc., 1958, 53.

¹⁶ Lancaster and Stoicheff, Canad. J. Physics, 1956, 34, 1016.

shows that there is also a contribution from the conventional form (III). Thus the apparent bonding capacity of the carbon atom is more than 4.0, a result which has been obtained for other π -bonded molecules and which indicates the need for a molecular-orbital treatment.

Davies and Hopkins ¹⁷ have recently carried out molecular-orbital calculations on urea, all the atoms of which, hydrogen included, have been shown experimentally ¹⁴ to be coplanar. They obtain the following results: (i) complete delocalisation of the two lone pairs of electrons on the nitrogen atoms being assumed, the bond order of both C⁻O and C⁻N is 1·50, and the agreement between the calculated delocalisation of resonance energy and Pauling's ¹⁸ empirical value is good; (ii) delocalisation of the lone pairs being assumed to be sufficient only to make the C⁻O bond order 1·60, which is the value found from force constants, the bond order calculated for C⁻N is 1·40 and the calculated resonance energy is smaller than Pauling's empirical value.

Since the carbon-nitrogen bond length in thiourea does not differ significantly from that in urea (1.35 Å) we may infer that the lone-pair electrons are delocalised to about the same extent in the two substances, but a detailed theoretical treatment of sulphur compounds is much more difficult than that of the corresponding oxygen compounds. Accordingly, and because few suitable data for comparison are available, further discussion will be deferred to a later paper in which accurate bond lengths for thioacetamide will be presented.

EXPERIMENTAL

Crystals of thiourea were grown from alcohol. The unit cell dimensions, determined by Straumanis's method from the reflections (501), (404), and (0, 10, 0), were found to be a=7.655, b=8.537, c=5.520 Å all ± 0.007 Å (v=360.7 ų). The density calculated for four molecules per unit cell is 1.399 g./c.c.; the observed value is 1.405 g./c.c. The absorption coefficient is 59.6 cm. and no absorption correction was made because very small crystals (diam. 0.1—0.15 mm.) were used.

Equi-inclination Weissenberg photographs were taken with Cu-K radiation about the three principal axes for all layer lines up to (hk4), (4kl), and (h3l). A multiple-film technique was used and the intensities were estimated visually by comparison with a calibration slip. After corrections for Lorentz and polarisation factors had been applied, the intensities from different layer lines were correlated and their square roots extracted to give structure factors on an arbitrary scale; the scale factor required to make these values absolute was one of the parameters refined. For structure factors determined by each of us independently R was 0.076.

We thank Professor E. G. Cox, F.R.S., and Dr. D. W. J. Cruickshank for helpful discussions, and Miss D. E. Pilling for carrying out the computations on the Manchester University electronic computer. Some of the equipment used was purchased with grants from the Royal Society and Imperial Chemical Industries Limited, and part of the cost of the computing work was borne by a grant from the D.S.I.R.

DEPARTMENT OF INORGANIC AND STRUCTURAL CHEMISTRY,
THE UNIVERSITY OF LEEDS.

[Received, Febru

[Received, February 11th, 1958.]

¹⁷ Davies and Hopkins, Trans. Faraday Soc., 1957, 53, 1563.

¹⁸ Pauling, "The Nature of the Chemical Bond," Cornell Univ. Press, 1939, p. 197.