# The Crystal Structure of Thiourea Nitrate 

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#### Abstract

The structure of thiourea nitrate has been determined by three-dimensional X-ray methods. Both $\mathrm{Cu} K \alpha$ and Mo $K \alpha$ data were obtained with a single-crystal diffractometer and the final $R$ values are $4.9 \%$ and $5.5 \%$ respectively. The space group is $P 2_{1} / m$ and there are two molecules in the unit cell. All atoms, including hydrogen, lie on mirror planes. The whole structure is built up of layers of atoms, the atoms within each layer being linked by a network of hydrogen bonds.


## Introduction

There are two possibilities for the structure of the protonated urea or thiourea cation:

(I)

(II)

Various attempts have been made to locate the position of the extra proton by infrared-spectroscopic methods, but the interpretation of the spectra has been contradictory; both structure (I) and structure (II) have been used to explain the same infrared spectra.

From the spectrum of urea nitrate Davies \& Hopkins (1957) concluded that the proton is attached to one of the two $\mathrm{NH}_{2}$ groups. Spinner (1959) came to the same conclusion. On the other hand, Janssen (1961) and Kutzelnigg \& Mecke (1961a,b,c) came to the conclusion that the proton is attached to the oxygen or sulphur atom.

The X-ray structure analyses of the salts urea phosphate (Sundera-Rao, Turley \& Pepinsky, 1957) and $N$-methylurea nitrate (Bryden, 1957) indicated that the proton of the acid becomes attached to the urea oxygen, but the results were not conclusive. The present X-ray analysis was carried out in an attempt to locate the position of the proton in thiourea nitrate and urea nitrate, and then to compare the bonding in these two compounds. (Three-dimensional intensity data have been collected for urea nitrate and work on this structure is in progress.)

## Experimental

Crystals grown from an aqueous solution were mainly long monoclinic needles in the [010] direction. They were extremely soft, having well-developed faces showing easy cleavage along (010). The crystals were un-

[^0]stable in air, and the specimens investigated were sealed in thin-walled capillary tubes.

Unit-cell dimensions obtained from a single crystal mounted on the Nonius automatic diffractometer are: $a=6.384 \pm 0.005, b=6.398 \pm 0.004, c=6.980 \pm 0.005 \AA ;$ $\beta=98.65 \pm 0.05^{\circ}$. The density was determined by flotation to be $1.65 \mathrm{~g} . \mathrm{cm}^{-3}$; the calculated density is 1.65 g. $\mathrm{cm}^{-3}$ for 2 molecules in the unit cell.

Intensities of a single crystal were measured at room temperature on the Nonius diffractometer; 448 reflexions ( $h k l$ and $h k l$ ) were recorded with filtered $\mathrm{Cu} K \alpha$ radiation and 662 with filtered Mo $K \alpha$ radiation.
The linear absorption coefficient for $\mathrm{Cu} K \alpha$ radiation is $45 \mathrm{~cm}^{-1}$ and the optimum thickness 0.045 cm . The needle-shaped crystal measured had a cross-sectional thickness of the order of 0.01 cm ; no absorption corrections were applied.

## Patterson function and space group determination

A three-dimensional sharpened Patterson function was evaluated using $I /\left(\Sigma f_{j}\right)^{2}$ as coefficients, the sum being


Fig. 1. Section of sharpened Patterson at $y / b=0$, showing configuration of the proposed molecule.
taken over the atoms in the unit cell. $I$ is the intensity of a reflexion corrected for Lorentz and polarization factors, and $f_{j}$ is the scattering factor of the $j$ th atom.

Precession photographs showed that the $0 k 0$ reflexions were absent for odd values of $k$. The allowed space groups are $P 2_{1}$ and $P 2_{1} / m$, the latter being centrosymmetric. The statistical test of Howells, Phillips \& Rogers (1950) indicated that the structure was centrosymmetric.

Furthermore, the Patterson function showed maxima only on the sections $y / b=0$ and $y / b=\frac{1}{2}$. This indicates that the space group is $P 2_{1} / m$, with the two molecules lying on mirror planes separated by a distance $b / 2$. The molecules are further related to each other by screw axes running parallel to the $b$ axis.

## Structure determination

Investigation of the Patterson section $y / b=0$ showed the configuration of the molecule outlined in Fig. 1.


Fig. 2. Difference electron density projection showing the position of the second molecule.


Fig 3. Section of the sharpened Patterson at $y / b=\frac{1}{2}$.

If the location of this molecule could be assumed, the problem was then to locate the second molecule, or what amounted to the same thing, the position of the screw axis in the centrosymmetric structure had to be determined. Structure factors $F_{c}$ were calculated for the $h 0 l$ reflexions, assuming that the molecule shown in Fig. 1 was part of a non-centrosymmetric structure. The phases thus obtained were used to make the difference electron density projection shown in Fig. 2. The second molecule is now revealed and the section $y / b=\frac{1}{2}$ of the Patterson function can easily be interpreted (Fig. 3).

## Structure refinement

The three-dimensional least-squares refinement was carried out with a modification of the Busing, Martin $\&$ Levy (1962) program ORFLS. First the $\mathrm{Cu} K \alpha$ data were refined, and when the Mo $K \alpha$ data were obtained, they were refined in exactly the same way. The method used was the following.

Three cycles of refinement were carried out in which the overall scale factor and the $x$ and $z$ coordinates of the eight independent atoms and their isotropic temperature factors were varied. The function minimized was $\sum w\left(\left|F_{o}\right|-(1 / s)\left|F_{c}\right|\right)^{2}$, where the summation is over all the observed reflexions.
$s=$ scaling factor
$F_{o}=$ observed structure factor
$F_{c}=$ calculated structure factor

The weighting scheme was:

$$
w=1 \cdot 0 \text { if }\left|F_{o}\right|<10
$$

and

$$
1 / V w=\left|F_{o}\right| / 10 \quad \text { if } \quad\left|F_{o}\right| \geq 10
$$

The $R$ index, $\Sigma\left|\left|F_{o}\right|-(1 / s)\right| F_{c}| | / \Sigma\left|F_{o}\right|$, was $14 \cdot 3 \%$ for the $\mathrm{Cu} K \alpha$ data.

Next, three cycles of refinement were carried out with the temperature factors of five atoms in anisotropic form. (The number of parameters was limited by the size of the computer used.) The $R$ value for $\mathrm{Cu} K \alpha$ data dropped to $7 \cdot 3 \%$. A difference electron density section at this stage is shown in Fig.4. The positions of the 5 hydrogen atoms are revealed; they all lie on the same plane with the other atoms.

The final refinement was carried out in several stages since not more than 30 parameters could be varied at a time. Besides the overall scale factor, the $x$ and $z$ coordinates of all the atoms and the anisotropic temperature factors of the non-hydrogen atoms were varied. The hydrogen atoms were assigned fixed isotropic temperature factors of $3 \cdot 5 \AA^{2}$ except $\mathrm{H}(5)$ which had $B=4.0 \AA^{2}$. The terms $\beta_{12}$ and $\beta_{23}$ were kept equal to zero in accordance with the space group restrictions (Peterse \& Palm, 1966).

In order to allow for the anomalous dispersion of the sulphur atom a hypothetical scatterer was superimposed over the 'normal' sulphur atom. This scatterer was assigned a constant scattering power of 0.3 elec-
trons for the $\mathrm{Cu} K \alpha$ data and 0.1 electron for the Mo $K \alpha$ data. All other scattering factors used were those tabulated by Hanson, Herman, Lea \& Skillman (1964).

Refinement was stopped when the changes in the parameters were less than the standard deviations of the parameters themselves. The positional parameters of $\mathrm{H}(5) \mathrm{kept}$ oscillating slightly, however. The final $R$ values were $4.9 \%$ for the $\mathrm{Cu} K \alpha$ data and $5 \cdot 5 \%$ for the Mo $K \alpha$ data.

## Results

The final positional and vibrational parameters are shown in Tables 1 and 2, and the bond lengths and angles in Tables 3 and 4. An investigation of the Tables shows that the two sets of data yield essentially the same results. It is interesting to note, however, that the estimated standard deviations are smaller for the Mo $K \alpha$ data, where more than 200 extra reflexions were

Table 1. Fractional coordinates of the atoms
Standard deviations in the last digits are shown in parentheses. $y / b=\frac{1}{4}$ for all atoms.

|  | $\mathrm{Cu} K \alpha$ data |  | Mo $K \alpha$ data |  |
| :---: | :---: | :---: | :---: | :---: |
|  | $x / a$ | $z / c$ | $x / a$ | $z / c$ |
| S | 0.5169 (2) | 0.6782 (2) | 0.5171 (2) | 0.6782 (2) |
| O(1) | $0 \cdot 8578$ (5) | 0.0747 (5) | 0.8562 (4) | 0.0739 (4) |
| O(2) | $0 \cdot 8011$ (6) | $0 \cdot 3683$ (5) | $0 \cdot 8008$ (5) | $0 \cdot 3678$ (5) |
| $\mathrm{O}(3)$ | $0 \cdot 1163$ (5) | $0 \cdot 3060$ (5) | $0 \cdot 1161$ (4) | $0 \cdot 3065$ (4) |
| $\mathrm{N}(1)$ | $0 \cdot 1212$ (6) | 0.7592 (6) | $0 \cdot 1220$ (5) | 0.7594 (5) |
| N(2) | $0 \cdot 3900$ (7) | 0.0146 (6) | 0.3907 (5) | 0.0153 (5) |
| N(3) | 0.9234 (6) | $0 \cdot 2507$ (5) | 0.9238 (5) | $0 \cdot 2502$ (4) |
| C | $0 \cdot 3250$ (7) | 0.8299 (6) | $0 \cdot 3245$ (5) | 0.8299 (5) |
| H(1) | $0 \cdot 310$ (9) | $0 \cdot 113$ (9) | 0.314 (8) | $0 \cdot 106$ (8) |
| H(2) | $0 \cdot 522$ (10) | 0.062 (8) | 0.526 (9) | 0.055 (7) |
| H(3) | 0.089 (10) | $0 \cdot 630$ (10) | 0.083 (9) | 0.639 (8) |
| H(4) | 0.042 (10) | 0.842 (9) | 0.041 (9) | 0.846 (8) |
| H(5) | 0.441 | 0.549 | $0 \cdot 452$ | 0.572 |

Table 2. Vibration parameters ( $\AA^{2}$ ) with the standard deviations in the last digits shown in parentheses

The temperature factor used was:

$$
\exp \left[-2 \pi^{2}\left(h^{2} a^{* 2} U_{11}+2 h k a^{*} b^{*} U_{12}+\ldots\right)\right]
$$

$\mathrm{Cu} K \alpha$ data

|  | $U_{11}$ | $U_{22}$ | $U_{33}$ | $2 U_{13}$ |
| :---: | :---: | :---: | :---: | :---: |
| S | $0 \cdot 0426$ (8) | 0.0809 (13) | 0.0439 (10) | $0 \cdot 0132$ (7) |
| O(1) | 0.0408 (18) | 0.0892 (29) | 0.0391 (19) | 0.0007 (15) |
| O(2) | 0.0638 (26) | 0.0942 (33) | 0.0565 (24) | 0.0305 (20) |
| $\mathrm{O}(3)$ | 0.0353 (22) | $0 \cdot 1392$ (44) | 0.0509 (22) | 0.0035 (18) |
| N(1) | 0.0345 (22) | 0.0637 (29) | 0.0439 (24) | 0.0024 (18) |
| N(2) | 0.0353 (22) | 0.0678 (31) | 0.0417 (24) | 0.0057 (18) |
| $\mathrm{N}(3)$ | 0.0392 (24) | 0.0527 (27) | 0.0393 (24) | 0.0055 (20) |
| C | $0 \cdot 0361$ (26) | 0.0407 (27) | 0.0401 (27) | 0.0062 (20) |
| Mo K $\alpha$ data |  |  |  |  |
|  | $U_{11}$ | $U_{22}$ | $U_{33}$ | $2 U_{13}$ |
| S | $0 \cdot 0412$ (6) | $0 \cdot 0774$ (8) | 0.0442 (5) | 0.0154 (4) |
| O(1) | 0.0361 (12) | 0.0890 (22) | 0.0437 (15) | 0.0020 (11) |
| O(2) | 0.0603 (18) | 0.0934 (27) | 0.0579 (17) | 0.0296 (16) |
| $\mathrm{O}(3)$ | 0.0347 (14) | 0.1296 (33) | 0.0507 (15) | 0.0033 (11) |
| N(1) | 0.0345 (14) | 0.0577 (21) | 0.0434 (15) | 0.0033 (13) |
| N(2) | 0.0339 (14) | 0.0606 (21) | 0.0429 (15) | 0.0068 (11) |
| N(3) | 0.0375 (14) | 0.0487 (19) | 0.0396 (15) | 0.0066 (11) |
| C | $0 \cdot 0301$ (14) | 0.0359 (19) | 0.0422 (17) | 0.0079 (13) |

recorded. This is in spite of the fact that the errors in the measurement of the $\mathrm{Cu} K \alpha$ intensities were smaller; there was much less background associated with the $\mathrm{Cu} K \alpha$ radiation and the count rate was appreciably higher.

Cruickshank's (1961) correction for rotational oscillation of molecules yielded values of between 0.01 and $0.02 \AA$, which were applicable to the bond lengths of

Table 3. Bond distances $(\AA)$
Standard deviations of Mo $K \alpha$ data shown in parentheses.


Fig. 4. Difference electron density section at $y / b=\frac{1}{4}$, showing positions of hydrogen atoms.
thiourea (Kunchur \& Truter, 1958) and thioacetamide (Truter, 1960). A similar analysis was attempted for the thiourea nitrate molecules, but there were large discrepancies between the observed and calculated vibration ellipsoids. It is possible that the hydrogen bonds linking the thiourea and nitrate groups prevent
either group from performing strictly rigid-body vibrations.

## Discussion of the structure

The structure as a whole consists of layers of planar molecules, the planes lying perpendicular to the crys-

Table 5. Bond lengths and angles in thiourea derivatives

| Compound and reference | S-C | $\mathrm{C}-\mathrm{N}$ | S-C-N |
| :---: | :---: | :---: | :---: |
| Thiourea* (Truter, 1967) | 1.720 (9) $\AA$ | 1.340 (6) $\AA$ | 120.5 (5) ${ }^{\circ}$ |
| Thiourea nitrate $\dagger$ | 1.739 (4) | $1 \cdot 312$ (5) | 121.3 (3) |
| (Present work) |  | $1 \cdot 300$ (5) | 116.9 (3) |
| Trimethylene thiourea* (Dias \& Truter, 1964) | 1.722 (7) | $1 \cdot 334$ (6) | $119 \cdot 8$ (4) |
| Thioacetamide* (Truter, 1960) | 1.713 (8) | $1 \cdot 324$ (12) | 121.7 (6) |
| Ethylenethiourea $\dagger$ (Wheatley, 1953) | 1.708 (8) | 1.322 (8) | $124 \cdot 8(1 \cdot 4)$ |
| Allylthiourea $\dagger$ (Dragonette \& Karle, 1965) | 1.66 | $1 \cdot 36$ | 120 |
| Thiourea dioxide $\dagger$ (Sullivan \& Hargreaves, 1962) | $1 \cdot 851$ (16) | $1 \cdot 310$ (8) | 117.6 (4) |
| $S$-Methylisothiourea sulphate $\dagger$ (Stam, 1962) | 1.743 (7) | 1.308 (9) | 114.5 (5) |
|  |  | $1 \cdot 333$ (9) | 123.0 (5) |
| * Correction for rotational oscillation applied. <br> $\dagger$ Correction not applied. |  |  |  |

Table 6. Observed and calculated structure factors ( $\times 10$ ). Mo K $\alpha$ data

\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|c|}
\hline \& \({ }_{1} 1 \begin{aligned} \& \text { Fo } \\ \& 0\end{aligned}\) \& \& \& \& \& \& \& \& \& \[
\begin{array}{l|l|}
\mathrm{k} \& \mathrm{Fdq} \\
\mathrm{~s} \& 24
\end{array}
\] \& \[
\begin{array}{cc}
0 \& F c \\
4 \& -23
\end{array}
\] \& | \(\begin{gathered}\text { h k } \\ 3\end{gathered}\) \& \[
\begin{array}{lll}
\mathrm{k} \& \mathrm{IFd} \\
5 \& 25
\end{array}
\] \& \& \& \& \& \& \& \& \& \[
\begin{aligned}
\& \text { Fod } \\
\& 9
\end{aligned}
\] \& \& \& \& \\
\hline \& 436 \& 45 \& \& 41 \& \& \& 21 \& -24 \& -5 5 \& 566 \& \& 2 36 \& 655 \& 54 \& 27 \& 17 \& -19 \& 15 \& 11 \& \& 22 \& 75 \& \& \& 27 \& 21 \\
\hline \& 16 \& -16 \& -1 6 \& 33 \& 37 \& 92 \& \& -21 \& 56 \& 632 \& \(2-30\) \& - -36 \& 624 \& 28 \& -2 \& 21 \& -23 \& 16 \& 16 \& -8 \& 23 \& 23 \& \& -5 \& \& 27 \\
\hline 08 \& 856 \& 53 \& 17 \& \& \& \& \& \& -5 6 \& 642 \& \(2-45\) \& 40 \& 048 \& -46 \& -3 0 \& 54 \& 158 \& - \& 215 \& -221 \& -2 \& 10 \& \& -5 \& 69 \& -72 \\
\hline \& - 19 \& \& -17 \& 22 \& 22 \& \& \& -365 \& 60 \& 092 \& \(2-97\) \& -4 0 \& 0175 \& 177 \& 30 \& 32 \& -32 \& -2 1 \& 185 \& -192 \& 2 \& 33 \& 39 \& -5 \& \& 54 \\
\hline 11 \& 55 \& -43 \& 20 \& 105 \& 107 \& \& \& -62 \& -60 \& 084 \& \& \& 83 \& 82 \& 3 \& 7. \& -66 \& 2 \& 10 \& -34 \& -2 \& 14 \& -12 \& -5 \& 18 \& 19 \\
\hline \& 17 \& \& -2 0 \& 285 \& -291 \& 2 \& 266 \& 264 \& \& 183 \& \& \& 33 \& -14 \& -3 \& \& \& \& \& \& \& \& \& \& \& 30 \\
\hline \& 318 \& 19 \& 2 \& 421 \& 418 \& 3 \& 43 \& 4 \& -6 1 \& 1 \& 16 \& 42 \& 147 \& 14 \& -3 \& 119 \& -122 \& -2 2 \& 172 \& 182 \& -3 \& 71 \& \& -6 \& 35 \& 40 \\
\hline \& 425 \& -24 \& -2 \& 42 \& -42 \& \& 124 \& 127 \& \& 2 \& \& \& 351 \& \& \& \& \& -2 \& 21 \& \& -3 \& 31 \& \& \& 55 \& 56 \\
\hline \& 515 \& -14 \& 22 \& 5 \& -s2 \& 5 \& 21 \& -23 \& -6 2 \& 273 \& -70 \& -4 4 \& 494 \& 9 \& 33 \& 356 \& 55 \& -2 \& 10 \& -109 \& -3 \& 50 \& -49 \& -6 \& \& \\
\hline \& 620 \& 1 \& -2 2 \& 232 \& 222 \& \& 51 \& \& \& 356 \& \& \& \(5{ }^{21}\) \& \& -3 \& 59 \& \& \& \& \& -3 \& 18 \& -12 \& \& 39 \& 41 \\
\hline \& 204 \& 210 \& \& 235 \& 233 \& \& \& \& \& \begin{tabular}{l}
4 \\
5 \\
\hline
\end{tabular} \& \& \& \& -41 \& \&  \& -14 \& \& \& 52 \& \& \[
25
\] \& \& \& \& \\
\hline 21 \& 187 \& 166 \& -2 3 \& 22 \& 21 \& 0 \& 329 \& 220 \& -6 \& 5 \& \& 56 \& 6 44 \& -43 \& \& 415 \& -14 \& -27 \& 25 \& 177 \& -40 \& \& 120 \& \& \& 42 \\
\hline \& 161 \& -156 \& 24 \& 33 \& 40 \& 1 \& 302 \& 311 \& \& \& \& -5 0 \& 013 \& \& -36 \& 22 \& -19 \& \& 175 \& \& \& \& \& \& \& \\
\hline 23 \& 3109 \& -102 \& -2 4 \& 析 \& 116 \& 1 \& 152 \& 15 \& \& 052 \& \& - \& 13 \& -10 \& -3 6 \& 22 \& -19 \& -3 0 \& 21 \& 25 \& 4 \& 59 \& -56 \& \& \& \\
\hline \& 84 \& 81 \& 25 \& 87 \& 92 \& 2 \& 30 \& \& \& \& \& \& cs \& \& \& \& \& \& \& \& \& \& \& \& \& \\
\hline \& 548 \& 49 \& -2 6 \& 49 \& 48 \& -1 2 \& 162 \& 52 \& -7 3 \& 3 \& -22 \& \& \& 25 \& 40 \& 81 \& -84 \& 3 \& 31 \& -24 \& -4 2 \& 95 \& 100 \& 0 \& 23 \& 8 \\
\hline \& 632 \& -33 \& , \& 116 \& 118 \& 3 \& 67 \& -164 \& \& 429 \& \& \& \& \& \& \& \& -3 \& 12 \& \& \& 33 \& \& \& 27 \& 23 \\
\hline \& 721 \& -22 \& \& 15 \& -24 \& 3 \& 394 \& \& \& \& \& \&  \& \& \& 203 \& 207 \& \& \& \& \& 43 \& \& \& \& \\
\hline 28 \& 816 \& \& -3 \& 1 \& -50 \& 4 \& 69 \& -63 \& -3 \& \& \& \& \& \& - \& 7 \& \& \& \& \& \& 23 \& -18 \& 10 \& \& \\
\hline 30 \& 0354 \& 58 \& \& 101 \& -94 \& \& 58 \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \\
\hline \& 94 \& -74 \& -3 2 \& 127 \& 2 \& 15 \& 42 \& \& -s \& 40 \& \& -5 6 \& 6 \& \& 4 \& 65 \& - 74 \& 3 \& \&  \& \& \[
24
\] \& -22 \& -1 \& 20 \& 22 \\
\hline \& 72 \& 264 \& 3 \& 31 \& 37 \& 6 \& 5 \& \& \& \& \& 61 \& 69 \& -68 \& 4 \& \[
\begin{array}{r}
138 \\
48
\end{array}
\] \& - 341 \& -3 \& \[
21
\] \&  \& \& \[
\begin{aligned}
\& 29 \\
\& 16
\end{aligned}
\] \& \[
\begin{aligned}
\& -29 \\
\& -14
\end{aligned}
\] \& \& \& \\
\hline \& 32 \& 28 \& -3 3 \& 36 \& \& \& \&  \& \& 237 \& \& -6 2 \& 240 \& \(-43\) \& \& 52 \& -49 \& 3 \& 22 \& -17 \& -5 \& 62 \& -66 \& \& \& 34 \\
\hline \& \(4{ }_{6} 121\) \& -r18 \& -34 \& 59 \& \& -20 \& 196 \& -207 \& 1 \& 132 \& 2 \& -6 \& \& \& \& 38 \& -43 \& 30 \& 41 \& -39 \& -5 \& 29 \& \& \& \& \\
\hline \& \(1: 122\) \& -113 \& 35 \& 24 \& -30 \& 21 \& 16 \& \& \& - \& \& 63 \& 352 \& 52 \& 4 \& 70 \& \& -40 \& 63 \& 64 \& -5 \& 43 \& 44 \& \& \& -17 \\
\hline 41 \& \(1: 38\) \& -140 \& -3 5 \& 18 \& -16 \& -2 1 \& 540 \& -5:1 \& \& 0383 \& \& 1-64 \& 423 \& 20 \& -4 5 \& 23 \& 21 \& 40 \& 30 \& -27 \& -5 \& 24 \& -21 \& \& \& 38 \\
\hline 42 \& 283 \& 80 \& 36 \& 34 \& -33 \& 22 \& 79 \& \& 01 \& 185 \& -177 \& \& 530 \& \& 4 \& 27 \& \& 4 \& 62 \& -59 \& \& 79 \& \& \& 19 \& -12 \\
\hline \& 38 \& 85 \& -3 6 \& 20 \& -1 \& -2 2 \& 156 \& 149 \& 02 \& 2279 \& 9-294 \& \(4^{-7} 0\) \& 079 \& \& \& 29 \& -28 \& 4 \& 30 \& \& \& \& -4 \& -2 \& 60 \& 56 \\
\hline 44 \& 433 \& -34 \& 40 \& 63 \& -67 \& 23 \& 20 \& 21 \& - \& 3103 \& 102 \& \(2-7\) \& 13 \& 115 \& 5 \& \& \& -4 \& 52 \& \& \& 17 \& -21 \& \& 20 \& \\
\hline \& 38 \& -37 \& \(-40\) \& 244 \& 24 \& -2 3 \& 301 \& 236 \& 4 \& 4129 \& 1291 \& \& 76 \& 36 \& 50 \& 30 \& -30 \& -4 \& 44 \& 41 \& -6 \& 43 \& \& \& 33 \& \\
\hline \& 1166 \& -159 \& 4 \& 160 \& 164 \& 24 \& 57 \& -59 \& 05 \& 36 \& \(36-40\) \& 72 \& 64 \& 67 \& 5 \& 20 \& 16 \& 44 \& 25 \& -22 \& -7 \& 51 \& 51 \& -2 \& 43 \& \\
\hline \& 106 \& 109 \& \& 0 \& 115 \& 24 \& 60 \& -63 \& \& \& \& \& \& \& -5 \& \& -84 \& \& \& \& \& \& \& \& 16 \& \\
\hline \& 2135 \& 131 \& 42 \& 6 \& 48 \& 25 \& 29 \& -24 \& -1 0 \& 0216 \& 6-217 \& \(7^{-7} 4\) \& 43 \& -42 \& -5 \& 23 \& -2 \& \& 65 \& 62 \& 00 \& 13 \& -14 \& \& 53 \& \\
\hline 53 \& 378 \& -79 \& \(-4\) \& 6 \& 207 \& -2 5 \& 5109 \& \& \& \& \& \& 548 \& \& \& \& \& \& \& \& \& 66 \& \& -3 \& 48 \& 45 \\
\hline \& 477 \& -77 \& 43 \& 73 \& -69 \& 6 \& 630 \& \& -1 1 \& 143 \& \(3{ }^{3}-34\) \& 80 \& 044 \& \& -5 \& 60 \& 60 \& -5 \& 85 \& 85 \& \& 51 \& -51 \& -3 \& 45 \& 4 \\
\hline \& 546 \& 45 \& 44 \& 16 \& -16 \& -2 6 \& 19 \& 17 \& 1 \& 194 \& 4 \& \(5-8\) \& 1 i8 \& \& -5 \& 29 \& -31 \& 5 \& 25 \& 25 \& \& 27 \& \& \& 34 \& 34 \\
\hline 56 \& 36 \& 36 \& 44 \& 131 \& -127 \& 27 \& 21 \& \& \& 2 16 \& \& \& \& -41 \& \& \& \& \& \& \& \& \& \& \& \& \\
\hline \& 716 \& -21 \& 45 \& 38 \& 38 \& -27 \& 33 \& 3 \& -1 2 \& 2160 \& 169 \& \({ }^{8} 2\) \& 233 \& -35 \& 6 \& 24 \& 27 \& -5 \& 56 \& -5 \& \& 17 \& -17 \& \& \& 0 \\
\hline \& 063 \& 63 \& -4 5 \& 26 \& 29 \& 30 \& 0243 \& 246 \& -1 3 \& \& 9 \& \& \& -15 \& \& 137 \& \& -5 \& \& \& \& 94 \& \& \& 31 \& 0 \\
\hline 61 \& 1 \& -6 \& 46 \& 63 \& \& -30 \& \& -15 \& 13 \& 366 \& \(66-66\) \& \& \& \& -6 \& 104 \& 106 \& \& \& -21 \& \& 56 \& 52 \& -4 \& \& \\
\hline 6 \& 251 \& -53 \& 50 \& 4 \& -13 \& \& 95 \& \& \& 465 \& 55 \& \& \& \& \& 320

23 \& -18 \& \& 18 \& \& -1 \& 11 \& \& \& 2 \& <br>
\hline 64 \& 431 \& 32 \& -5 0 \& 14 \& 117 \& -31 \& 184 \& -182 \& -1 4 \& \& \& $2^{-9}$ \& \& \& -6 \& \& \& -5 4 \& 18 \& 26 \& -1 \& 13 \& \& \& \& <br>
\hline \& $6 \quad 15$ \& -14 \& -5 \& 59 \& -55 \& \& \& -187 \& -1 5 \& $\begin{array}{lll}5 & 18 \\ 5 & \end{array}$ \& 8-12 \& $2-9$ \& \& -16 \& \& \& -65 \& \& \& \& \& 66 \& \& \& 34 \& <br>
\hline 70 \& 036 \& -32 \& 52 \& 13 \& 14 \& 33 \& \& -58 \& \& 35 \& \& ${ }^{-9} 4$ \& \& -26 \& \& \& \& \& \& \& \& 36 \& \& \& \& <br>
\hline \& 171 \& 75 \& -5 2 \& 102 \& -99 \& -3 3 \& \& 13 \& \& 638 \& \& \& \& \& \& 20 \& -18 \& \& \& 45 \& \& 30 \& ${ }^{-3}$ \& \& \& <br>
\hline \& 27 \& 29 \& 53 \& 81 \& 35 \& 34 \& \& \& -1 6 \& 629 \& \& \& 30 \& 28 \& -7 \& 72 \& -70 \& -6 \& , \& -22 \& \& 13 \& -2 \& 5 \& 19 \& <br>
\hline 73 \& 49 \& -49 \& -5 3 \& 49 \& 49 \& -3 4 \& 4 \& \& 17 \& 719 \& 19 -19 \& 10 \& 38 \& -80 \& -7 2 \& 20 \& -39 \& -6 5 \& 27 \& -25 \& -2 \& 64 \& 62 \& \& 26 \& -30 <br>
\hline \& 18 \& -20 \& 54 \& 14 \& -12 \& 35 \& 28 \& 2 \& 20 \& 0194 \& 1946 \& 0 \& 27 \& 25 \& -7 \& \& \& \& \& 55 \& -2 \& 72 \& \& \& 34 \& -31 <br>
\hline \& 20 \& 21 \& -5 \& 64 \& 6 \& -3 5 \& 86 \& 80 \& -2 0 \& \& 13139 \& 19 \& \& \& -7 \& 28 \& 25 \& \& \& -40 \& \& 28 \& 29 \& \& 23 \& -30 <br>
\hline \& 26 \& -22 \& -5 5 \& 33 \& -33 \& 36 \& 31 \& $1 \begin{array}{ll}-30\end{array}$ \& -2 \& \& \& \& \& 20 \& \& \& 27 \& \& \& ¢ \& \& 20 \& 23 \& \& \& <br>
\hline 8 \& 239 \& 35 \& \& 0 \& \& -37 \& \& 8 -37 \& \& \& \& -10 \& \& 128 \& -8 \& 15 \& 23 \& \& \& \& -2 \& 51 \& -51 \& \& - \& <br>
\hline 9 \& 11 \& \& -5 6 \& 5 \& -28 \& \& \& -13 \& \& 2145 \& 25-150 \& d 1 \& \& \& -8 \& 227 \& \& \& \& -56 \& -2 \& 44 \& -47 \& \& 21 \& <br>
\hline \& 31 \& 30 \& 57 \& 14 \& 16 \& -4 0 \& 81 \& $1-79$ \& -2 2 \& 2122 \& 22-124 \& 4-1 \& \& \& \& \& \& \& 5 \& -52 \& \& 26 \& -25 \& \& 0 \& 19 <br>
\hline \& 1-1 \& \& -6 0 \& 70 \& -70 \& \& 50 \& -47 \& -2 \& 357 \& 57-49 \& \& \& -235 \& - \& \& -20 \& \& 52 \& 55 \& \& 29 \& 30 \& 10 \& \& -14 <br>
\hline \& $0 \quad 59$ \& 61 \& 61 \& 47 \& 48 \& -4 \& 1126 \& -122 \& 23 \& 313 \& $35-13$ \& $3-13$ \& 3168 \& \& \& $1=5$ \& \& 0 \& 35 \& 3 \& -2 \& 22 \& 19 \& -1 \& 43 \& 1 <br>

\hline 01 \& 46 \& 55 \& -6 1 \& 136 \& 136 \& 42 \& \& \& \& 475 \& \& 5-1 3 \& 369 \& 68 \& \& 94 \& -87 \& 04 \& 32 \& 33 \& 2 \& $$
7
$$ \& 16 \& \& 41 \& <br>

\hline 0 \& 252 \& -44 \& 62 \& 23 \& -22 \& 42 \& 63 \& 3 \& -2 4 \& 490 \& \& -14 \& 465 \& 65 \& 01 \& 116 \& \& 0 \& 17 \& -19 \& \& 32 \& -26 \& \& 32 \& 24 <br>
\hline 0 \& 350 \& -51 \& -6 2 \& 59 \& 55 \& 3 \& 32 \& 232 \& -2 5 \& 5 \& 23.20 \& $2{ }^{1}$ \& 66 \& -68 \& 0 \& 65 \& 66 \& 06 \& 16 \& -15 \& -3 \& \& 119 \& \& \& 34 <br>
\hline 04 \& 14 \& 21 \& 63 \& 40 \& -40 \& 4 \& 87 \& 783 \& \& 513 \& \& \& \& -32 \& \& 83 \& 85 \& 1 \& \& $-228$ \& -3 \& 38 \& 36 \& 20 \& 44 \& 42 <br>
\hline \& 44 \& 45 \& 63 \& 96 \& -95 \& \& \& 9 -39 \& \& 636 \& \& -16 \& 631 \& -3 \& 04 \& 29 \& -29 \& 10 \& 91 \& -92 \& 3 \& 29 \& -29 \& 21 \& 39 \& 88 <br>
\hline 07 \& 726 \& -24 \& -64 \& 25 \& -27 \& \& \& -16 \& -2 6 \& 649 \& \& $\mathrm{O}^{-2} 0$ \& O 12 \& -7 \& \& 47 \& 47 \& 11 \& 10 \& \& 3 \& 23 \& 22 \& 22 \& 38 \& -36 <br>
\hline \& 320 \& 237 \& \& 28 \& 26 \& \& \& 3 \& \& 0228 \& -241 \& 120 \& 0114 \& 113 \& 07 \& 19 \& 22 \& -1 \& 114 \& 108 \& -3 \& 100 \& -100 \& -3 \& 46 \& -44 <br>
\hline -10 \& 289 \& -267 \& -6 \& 47 \& 50 \& 50 \& - 125 \& 126 \& -3 0 \& 0220 \& -219 \& \& 112 \& 114 \& 10 \& 38 \& 35 \& -1 \& 180 \& 189 \& 33 \& \& 19 \& \& 42 \& <br>
\hline - \& 542 \& -540 \& -6 \& 21 \& -22 \& \& O 203 \& 201 \& -31 \& 1219 \& \& ${ }^{3}$ \& \& 105 \& -1 0 \& ${ }^{-14}$ \& \& \& 76 \& 87 \& ${ }^{-3}$ \& 63 \& 62 \& \& 23 \& 26 <br>
\hline -1 \& 384 \& -365 \& -7 0 \& 59 \& 55 \& 5 1 \& \& \& \& \& \& ${ }^{\text {O-2 }}$ \& \& 13 \& 1 \& 134 \& 43 \& -1 3 \& \& 74 \& ${ }^{-3}$ \& 12 \& -9 \& \& 36 \& -32 <br>
\hline 1 \& 2238 \& -212 \& \& 33 \& 33 \& 5 \& \& \& \& \& \& \& \& \& 1 \& \& 119 \& \& \& \& \& 32 \& -30 \& \& 33 \& -36 <br>

\hline -1 \& 2219 \& 188 \& 72 \& 23 \& 19 \& \& \& -106 \& -3 | - |
| :--- |
| -3 | \& 2165

3125 \& \& -2 4 \& \& \& - \& \& \& \& \& \& \& 48 \& \& \& \& <br>
\hline \& 303 \& 295 \& 7 \& \& -47 \& \& \& \& \& 31
3 \& \& - $0^{2} 24$ \& \& \& \& \& \& -1 \& 47 \& \& 4 \& 59 \& 58
51 \& \& \& <br>

\hline 1 \& 3202 \& 190 \& \& \& \& \& \& \& -3 \& 483 \& 83-78 \& 8 \& $$
\begin{aligned}
& 56 \\
& 47
\end{aligned}
$$ \& 46 \& \& \[

$$
\begin{array}{ll}
3 & 35 \\
3 & 25
\end{array}
$$

\] \& -24 \& \& 29 \& \& -4 \& \[

$$
\begin{aligned}
& 57 \\
& 38
\end{aligned}
$$

\] \& 36 \& $1{ }_{1} 1$ \& \[

$$
\begin{aligned}
& 37 \\
& 34
\end{aligned}
$$
\] \& <br>

\hline \& 4108 \& 107 \& \& $\stackrel{29}{7}$ \& \& \& \& \& 34 \& 4113 \& 13-116 \& $16-25$ \& 557 \& 62 \& 1 \& 425 \& 21 \& 2 \& 95 \& 96 \& 4 \& 44 \& -4! \& -2 \& \& 26 <br>
\hline \multirow[b]{3}{*}{} \& \& \& \& 35 \& 32 \& -5 4 \& \& \& -3 5 \& 546 \& \& 50-26 \& $6 \quad 16$ \& \& 4 \& 14 \& 14 \& 21 \& 34 \& -30 \& 43 \& 38 \& 35 \& \& \& <br>
\hline \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& -1 \& 31 \& 23 <br>
\hline \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& \& -30 \& 38 \& 2 <br>
\hline
\end{tabular}

tallographic $b$ axis; the inter-layer spacing is $3 \cdot 2 \AA$. Screw axes parallel to $\mathbf{b}$ relate molecules within adjacent layers.

Within the layers themselves, an extensive network of hydrogen bonds links thiourea and nitrate groups together (Fig.5). The atoms $\mathrm{N}(1)$ and $\mathrm{N}(2)$ are in the trigonally hybridized state and form $\mathrm{N}-\mathrm{H} \cdots \mathrm{O}$ bonds of lengths 2.97 and $2.88 \AA$ with $\mathrm{O}(1)$ and $\mathrm{O}(3)$ respectively. The other two hydrogen atoms attached to $\mathrm{N}(1)$ and $N(2)$ form bifurcated hydrogen bonds with two other nitrate groups. These bonds are not equal in length, the $\mathrm{N} \cdots \mathrm{O}$ distances being $3 \cdot 31,2 \cdot 94,3 \cdot 16$ and $3 \cdot 16 \AA$. The fifth hydrogen atom also lies in the plane formed by the rest of the atoms. It is attached to the sulphur atom and lies approximately equidistant from two oxygen atoms. The line $H(5) \cdots S$ makes an angle of $105^{\circ}$ with the C-S bond.

Theoretical considerations lead us to expect the S-C bond to be lengthened and the $\mathrm{C}-\mathrm{N}$ bonds to be shortened when the $S$ atom becomes protonated. Referring
to Table 5, we see that the S-C bond length in thiourea nitrate is indeed greater than that found in thiourea, but this difference is not large enough to be statistically significant.
The two $\mathrm{C}-\mathrm{N}$ bond lengths in thiourea nitrate are not significantly different. Assuming an average C-N length of $1.306 \AA$, we see that this value is significantly smaller than the $1.340 \AA$ determined for thiourea, with $\Delta / \sigma=6 \cdot 0$.
There is a large difference in the two $\mathrm{S}-\mathrm{C}-\mathrm{N}$ angles. A similar effect has been discovered in $s$-methylisothiourea sulphate by Stam (1962).
The values of the $\mathrm{S}-\mathrm{C}$ and $\mathrm{C}-\mathrm{N}$ bond lengths and the $\mathrm{S}-\mathrm{C}-\mathrm{N}$ bond angles determined for several thiourea derivatives are shown in Table 5.
The dimensions and symmetry of the nitrate anion have been discussed in several publications (Ferrari, Braibanti, Manotti-Lanfredi \& Tiripicchio, 1967 and references quoted therein). The results for thiourea nitrate show no significant differences in the $\mathrm{O}-\mathrm{N}-\mathrm{O}$


Fig. 5. Bond lengths and bond angles in thiourea nitrate
angles, but one of the $\mathrm{N}-\mathrm{O}$ distances is significantly shorter than the other two. The effect of the N-H . . O hydrogen bonds appears to be to 'stretch' the $\mathrm{N}(3)-$ $\mathrm{O}(1)$ and $\mathrm{N}(3)-\mathrm{O}(3)$ bonds.

We wish to express our gratitude to Mrs R. OlthofHazekamp for preparing the input tape for the diffractometer.

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# Die Kristallstruktur des $\alpha-\mathrm{KZnBr}_{\mathbf{3}}^{\mathbf{2}} \mathbf{2 \mathbf { H } _ { \mathbf { 2 } } \mathrm { O }}$ 

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(Eingegangen am 6. April 1967 und wiedereingereicht am 14. November 1967)
$\alpha-\mathrm{KZnBr}_{3} .2 \mathrm{H}_{2} \mathrm{O}$ crystallizes from aqueous solutions (mole proportion $\mathrm{KBr}: \mathrm{ZnBr}_{2}=1.75: 1$ ), with unit-cell dimensions $a_{0}=6 \cdot 71, b_{0}=12 \cdot 42, c_{0}=13 \cdot 02 \AA, \beta=130,3^{\circ}$. Space-group $P 2_{1} / c$ with $Z=4$. The structure was solved by Patterson and Fourier syntheses and by direct methods. By least-squares methods with anisotropic temperature factors the structure was refined to $R_{1}=5 \cdot 1 \% . \mathrm{Zn}$ is tetrahedrally coordinated by three bromine atoms and one water molecule.

## Einleitung

Im Rahmen einer systematischen Untersuchung der Kristallstrukturen von Alkalihalogenozinkaten wurde die Struktur des $\alpha-\mathrm{KZnBr}_{3} .2 \mathrm{H}_{2} \mathrm{O}$ bestimmt.

In den bisher bekannten Strukturen der Alkalihalogenozinkate wird Zink jeweils tetraedrisch umgeben, wobei bislang nur isolierte Tetraeder bekannt geworden sind. In den Verbindungen vom Typ $\mathrm{A}_{2} \mathrm{ZnCl}_{4} . x \mathrm{H}_{2} \mathrm{O}$ (e.g. Jacobi \& Brehler, 1968) wird Zn von vier Cl , im $\mathrm{KZnCl}_{3} .2 \mathrm{H}_{2} \mathrm{O}$ (Brehler \& Süsse, 1963) von drei Chlor und einem Wassermolekül umgeben. Zweck der vorliegenden Arbeit war es festzustellen, welche Koordination im $\alpha-\mathrm{KZnBr}_{3} .2 \mathrm{H}_{2} \mathrm{O}$ vorliegt.

## Experimentelles

Aus wässrigen Lösungen mit einem Molverhältnis beider Komponenten von 1:1 entsteht bei Zimmertem-
peratur das $\beta-\mathrm{KZnBr}_{3} .2 \mathrm{H}_{2} \mathrm{O}$, das zuerst von Ephraim (1908) angegeben und dessen Struktur von Brehler \& Holinski (1967) aufgeklärt worden ist. Bei einem Molverhältnis $\mathrm{KBr}: \mathrm{ZnBr}_{2}$ von 1,75:1 entsteht das $\alpha-$ $\mathrm{KZnBr}_{3} .2 \mathrm{H}_{2} \mathrm{O}$. Bei dieser Lösungszusammensetzung wurden ausserdem Kristalle der Zusammensetzung $\mathrm{K}_{2} \mathrm{ZnBr}_{4} \cdot \mathrm{H}_{2} \mathrm{O}$ und zwei Modifikationen des $\mathrm{K}_{2} \mathrm{ZnBr}_{4} \cdot 2 \mathrm{H}_{2} \mathrm{O}$, von denen eine ebenfalls bei Ephraim (1908) aufgeführt ist, erhalten.

Das $\alpha-\mathrm{KZnBr}_{3} .2 \mathrm{H}_{2} \mathrm{O}$ bildet farblose nach [001] gestreckte langprismatische monokline Kristalle, die stets nach (100) verzwillingt sind. Aus Weissenbergaufnahmen solcher Kristallnadeln wurde das in Fig. 1 dargestellte reziproke Gitter erhalten. Durch Aufnahmen von Teilindividuen eines längs der Zwillingsebene gespaltenen Kristalls konnte gezeigt werden, dass es sich dabei um eine Überlagerung von zwei um $180^{\circ}$ gedrehten bzw. gespiegelten reziproken Einzelgittern handelt.


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