The Structure of Braggite and Palladium Sulphide.

By T. F. Gaskell in Cambridge (England).

The investigation of the structure of Palladium Sulphide PdS is of double interest. In the first place it is important in order to determine the coordination of palladium, and in the second, because the identification of Braggite (Pt, Pd, Ni) S, which is isomorphous with palladium sulphide, by F. A. Bannister¹), was the first example of a mineral determination carried out solely by X-ray methods.

The structure of braggite was complicated by the large replacement of palladium by platinum and nickel (approx. 4 atoms Pt, 2Pd, 2Ni, 8S). The best method of attack was clearly an investigation of the isomorphous synthetic compound PdS, of which, through the kindness of F. A. Bannister, two single crystals were available. These were small black, metallic, tetragonal crystals, with elongation along the c-axis, and a flattening perpendicular to one a-axis, producing a thick plate parallel to the (400) face. Oscillation photographs about the c and b axes give $a = 6.43 \pm 0.02 \text{ Å}$, $c = 6.63 \pm 0.02 \text{ Å}$. Assuming a density of 7, there are eight molecules of PdS in the unit cell, giving the X-ray density 6.69. Oscillation and Weissenberg photographs (CuK radiation) show a pseudo-cubic character, and the only regular absent reflections to be those for which 00l is odd. There are no planes of symmetry parallel to the principal axes (hk0 + hk0) and hk0 + kh0, so that the space group is most probably $P4_2/m$ or $P4_2$.

The determination of the positions of the palladium atoms was made by the direct F^2 Patterson method²). The F^2 's were obtained by estimating visually intensities on Weissenberg and oscillations photographs, and connecting these intensities by their respective polarisation factors $-\Theta = \frac{1+\cos^2 2\theta}{\sin 2\theta}$. The measurement of intensities was made by fixing the strongest spot as 100 and estimating the intensities of others on this scale. Cross checks of one spot against another ensure that the right order of intensities is obtained, and an overlap of 2° on the oscillation photographs allows of comparison of different photographs. The results from oscillation and rotation photographs agreed well, as also the F^2hk0 and $F^2k\bar{h}0$, which should be equal. No account was taken of absorption, but this should not affect the hk0 reflections much, because the thickness

¹⁾ See preceding note.

²⁾ Patterson, Z. Kristallogr. (A) 90 (1935) 517-542.

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(1) and (2) peaks are larger than the others because of the effect of two Pd atoms at $(0, 0, \frac{1}{4})$ and $(0, 0, \frac{3}{4})$ lying on the four-fold screw axis. The peak (4) is not accounted for with these positions for palladiums, and is therefore assumed to be due to sulphur atoms in eight-fold positions, the peak indicating x = .19 y = .32. Although the sulphur has an atomic scattering factor only one third that of palladium, a peak on the F^2 projection may arise because the vector represents the interatomic distance between two S's (one vertically above the other) and two Pd's (on fourfold screw axis).

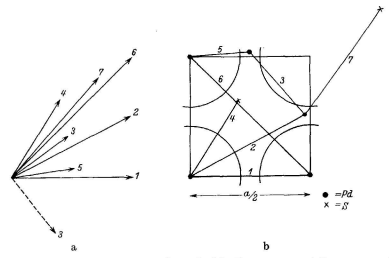


Fig. 2. (a), (b) Showing the vectors from the hk0 Patterson and the corresponding interatomic distances.

The $hk0 F^2$ projection shows that two two-fold positions are occupied by the palladium atoms. For these the z-co-ordinate is fixed. For the palladium atoms in the four-fold position the z-co-ordinate is fixed for the space group $P 4_2/m$, but not for the space group without the plane of symmetry, $P 4_2$. An F^2 projection on the h0l plane was made, and to the limits of accuracy obtained the z-co-ordinates for the four-fold position were 0 and $\frac{1}{2}$, showing that there was no need to assume the lower $P 4_2$ symmetry. All coordinates, will, therefore, be expressed as of the $P 4_2/m$ space group. (The unit cells are $\frac{1}{4}c$ apart in the two space groups.) The F^2 projection on h0l confirmed the results of the hk0 projection (its similarity showed the pseudo-cubic character) and indicated that the z-parameter for the sulphur was approximately 0.23, that is, just a little off the $z = \frac{1}{4}$ position.

F values were calculated for the hk0 and h0l reflections using the value:

```
2 Pd at (0, 0, \frac{1}{4})
2 Pd at (0, \frac{1}{2}, 0)
4 Pd at (x, y, 0) where x = .48, y = .25
8 S at (x, y, z) where x = .19, y = .32, z = .23
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These gave in most cases unambiguous values of the sign of F, so that the Fourier projection could be made on the hk0 and h0l planes.

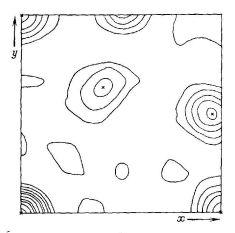


Fig. 3. Fourier projection on hk0. Contours every 25 units.

Fig. 4. Fourier projection on hol. Contours every 20 units.

The hk0 gave very regular palladium peaks, and quite a good sulphur peak (due to 2S atoms), giving the parameters:

for
$$Pd$$
 (4-fold) $x = .475$, $y = .250$
for S $x = .20$, $y = .32$

The h0l projection is not so regular, the palladium peaks being elongated in the c-direction, due probably to absorption. It does, however, confirm the z=0 parameter for the palladiums in four-fold positions. The two sulphur peaks are not so well defined as the single S peak in the hk0 projection, but are consistent with the parameters from the hk0 projection. The z parameter is indicated approximately as z=.22. It will be noticed that in both hk0 and h0l projections slight diffraction maxima round the large peaks are observable, although most diffraction effects were removed by decreasing the effect of the higher order reflections by means of an artificial temperature factor $(F=F_0e^{-B}(\sin\theta/\lambda)^2, B=3)$.

An attempt was made to obtain more accurate values of the parameters for the sulphur atoms—the exact position of the sulphur being required to give the distortion of the Pd-S co-

ordinations. Very few reflections could be found for which the total F of the palladium atoms was zero. Those for which $F_{Pd} = 0$ were, further, found to be very insensitive to change of the S-position. However, the table I of intensities, (and more clearly the graphical representation of these Fig. 5) for h00 and hk0reflections does show that the position of the sulphur atom indicated by the hk0 Fourier (x = .20, y = .32) is more probable than that for which the four Pd-S distances are equal. (x = .20, y = .30).

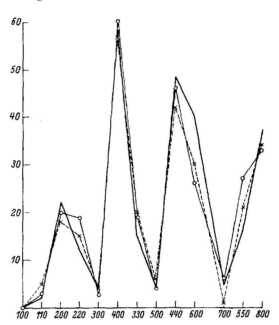


Fig. 5. Plot of h00 and hh0 observed and calculated intensities.

Table I. F's observed and calculated for S positions 1. x = .20, y = .30; 2. x = .20, y = .32

hk0	F_1	F_2	$F_{ m obs.}$	hk0	F_1	F_2	$F_{ m obs.}$
100	0	0	0	700	6	1	7
200	18	20	28	800	33	34	62
300	3	4	5	110	3	4	2
400	60	57	73	220	19	15	15
500	4	6	9	330	49	20	19
600	26	30	50	440	47	42	60
				550	27	21	20 -

There is slightly better agreement here with the (.20, .32) position, indicating a certain amount of distortion. The (.20, .32) has also the weight of the hk0 Fourier result.

It will be seen, however, that the change in F values for 0.02 change in parameter is small, and in no case really significant. This must be expected, since the eighth order reflections are the highest obtainable, so that the resolving power of the crystal is small. The comparatively large palladium atoms also give a masking effect to the sulphur F contributions, especially as only half of the Pd atoms are in special positions, giving, therefore, no regularities in F contributions for palladiums. It will be assumed, then that the statistical result of the hk0 Fourier (which gives an almost regular sulphur peak) is correct, and that the sulphur parameters are x = .20, y = .32, z = .22. The z-parameter from the h0lFourier has a possible accuracy of ± 0.045 and the x and $y \pm 0.04$. The parameters for the palladium atoms in (x, y, 0) positions are estimated to be correct to ± 0.005, the Fourier peak (and the Patterson peak also) being very regular and sharply defined. A possible approach to a more accurate value of the sulphur parameters is the three dimensional method of Patterson¹), which will bring in the extra weight of a large number of hkl reflections. A study of the selenide of palladium (which may be isomorphous with the sulphide) would be even more promising, because the selenium would give an F contribution comparable with that of palladium.

Table II. F's observed and calculated for hol.

h0l	$F_{ m calc.}$	$F_{ m obs.}$	hot	$F_{ m calc.}$	$F_{ m obs.}$	h0l	$F_{ m ealc.}$	$F_{ m obs.}$
101	2	0	103	4	0	305	40	9
204	39	20	203	33	24	405	0	2
304	1	0	303	7	3	505	11	18
401	1	4	403	3	3	506	13	24
501	6	16	503	2	4	106	24	26
604	19	32	603	22	30	206	4	8
701	4	9	703	8	19	306	19	24
804	4	10	104	1	0	406	19	24
102	38	20	204	19	20	506	17	21
202	12	14	304	2	0	107	5	4
302	32	20	404	47	40	207	21	. 22
402	25	28	504	4	0	307	12	12
502	20	36	604	23	28	407	5	8
602	9	25	704	3	23	108	0	3
702	12	20	105	4	3	208	17	20
802	14	26	205	23	24			

¹⁾ Patterson, J. Phys. Chem. Soc. June 1936.

hk0hk0 $F_{\mathrm{calc.}}$ hk0Fobs. Fcalc. Fobs. $F_{\mathrm{obs.}}$ $F_{\mathrm{calc.}}$

Table III. F's observed and calculated for hk0.

The basis of the structure may be considered to be a cubic β -tungsten type lattice of palladium atoms. If the four palladiums in the general

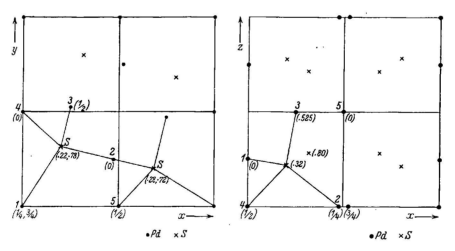


Fig. 6. Plan of structure on hk0. (The Pd-atoms are numbered as in the text.)

Fig. 7. Plan of structure on h0l.

position (x, y, 0) are moved to $(\frac{1}{2}, \frac{1}{4}, 0)$ and the unit cell considered from the atom $(0, \frac{1}{4}, 0)$, as origin, a body centred cell, plus two atoms on each face, is at once apparent. This cubic arrangement, with the sulphur in

the symmetrical position $(\frac{1}{4}, \frac{1}{4}, \frac{1}{4})$ (referred to the tetragonal axes to avoid confusion of diagrams) gives a five-fold coordination about the sulphur atom, there being three neighbours at $\sqrt{2}a/4$ and two at $\sqrt{3}a/4$. The arrangement of sulphurs is square about six of the palladiums, cubical about the other two. To avoid the five-fold coordination the sulphur in PdS has moved away from one palladium atom ((5) in the Fig. 6) and the four palladiums about the centre move slightly also. The cell clongates

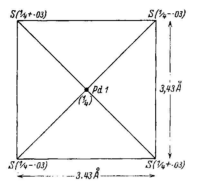


Fig. 8.

Arrangement of sulphur atoms about Pd(4), $(0, 0, \frac{1}{2})$.

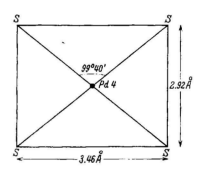


Fig. 9. Arrangement of sulphur atoms about Pd (4), $(0, \frac{1}{2}, 0)$. The S's and Pd are in the same vertical plane due to the symmetry.

slightly and the sulphur moves up, still further away from (5) at $(\frac{1}{2}, 0, \frac{1}{2})$ Fig. 7. We have for the Pd-S distances:

$$\begin{array}{lll} d_1 = 2.43\,\text{\AA} & d_3 = 2.29\,\text{Å} & d_5 = 2.94\,\text{Å}. \\ d_2 = 2.34\,\text{Å} & d_4 = 2.26\,\text{Å} & \end{array}$$

The sides of the palladium tetrahedron, with the angles subtended at the centre are:

The tetrahedron of palladium atoms about each sulphur is therefore much deformed. There are eight tetrahedra of palladiums in the unit cell, of similar shape because of the symmetry of the crystal. The arrangement of the tetrahedra is by sharing of one side (the short (4) (2) side in the notation above), and connecting with four tetrahedra at both of the other corners. The short edge common to both tetrahedra is that between the sulphur atoms of closest approach, and may be considered to be a consequence of the repulsion of sulphur from sulphur, and the consequent

drawing in of the palladiums until the Pd-S attraction and the Pd-Pd repulsion give equilibrium. The presence of palladium atom (5), which was pushed away when the cubic structure was discarded, causes a flattening of the tetrahedron in its direction, so that the face (1) (2) (3) is quite the largest tetrahedron face. This is a forcing of the palladium atoms out of position by the attraction between Pd (5) and the sulphur atom.

The arrangement of sulphur atoms around palladium atoms is of three types. Type (4) $(0,0,\frac{1}{4})$ has a "buckled" square of sulphurs about the palladium atom Fig. 8. The side of the square is $3.43\,\text{Å}$ in projection on hk0, or $\sqrt{3.43^2+(0.06c)^2}=3.45\,\text{Å}$ in actual length. If the square

is to be made flat the sulphurs have to be put on the same level. This would deform the tetrahedra even more than they are, since the distance between palladiums (4) and (2) would become shorter.

Type (4) (0, ½, 0) has a plane rectangle of sulphur atoms round it, this plane being vertical (c-axis vertical) Fig. 9. The sides are 2.92Å (the nearest approach of sulphur atoms in the structure) and 3.46Å. This is farremoved from a square, and to make it a square large deformation of the palladium tetrahedron would be necessary.

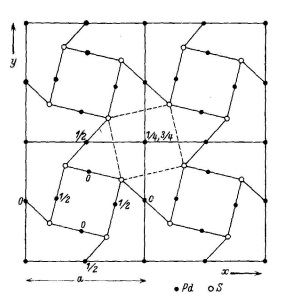


Fig. 40. Arrangement of sulphur atoms about Pd (2), (.475, .250, 0). The S's and Pd are in the same vertical plane within the limits of measurement.

The palladiums in the general positions (atoms (3) and (2)) have within the limits of error a plane arrangement of sulphur around them (the S-S distance in the hk0 projection = 3.48Å, Fig. 10 the sum Pd - S + S - Pd = 3.49Å), the sulphurs being 2.92Å and 3.71Å apart at the two ends as shown in the diagram. The planes are vertical, and form a square prism around the central four-fold half screw axis. These prisms are joined by the vertical rectangle about the palladium atom on the two-fold axis,

and by the horizontal square about the palladium on the corner four-fold axis (Fig. 44).

The structure is of considerable interest in showing that although

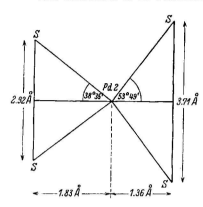


Fig. 41. Plan on hk0 plane, showing the arrangement of the PdS_4 groups. Four unit cells are shown.

square planar arrangements of sulphur about palladium are approached—these arrangements are readily deformed. This deformation is especially easy in the plane of the sulphur atoms, the angle between the Pd-S links varying from 77° 40′ to 99° 40′. This is slightly greater deformation than in cooperite for which the smaller angle subtended at a platinum atom by two sulphur atoms in a plane rectangular arrangement is 82° 36′. The deformation of the plane square out of the plane seems more difficult, the angle between the Pd-S

link and the horizontal being only 4° 40'. The deformation out of the plane occurs by two sulphurs moving up, two moving down, as might

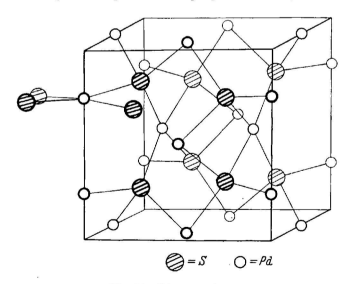


Fig. 12. Diagram of unit cell.

be expected if a structure rigid in the plane of the palladium atom were buckled by a large force. The palladium sulphur co-ordination is, then, the square type generally assumed, but this square can be easily deformed while kept plane, and deformed less easily out of the plane.

I wish to thank Mr. Bernal for giving me this problem and for his constant help in the work — also Dr. Fankuchen for taking the photographs and for much advice.

Summary.

Palladium sulphide is tetragonal, space group $P4_2/m$ $a=6.43\,\text{Å}$, $c=6.63\,\text{Å}$, eight molecules in the unit cell. The structure was determined by direct Patterson and Fourier analysis on visually estimated intensities of reflections. The arrangement of the palladium atoms closely resembles the β -tungsten structure. The sulphur atoms lie almost at the corners of a cube of half the linear dimensions of the unit cell, and twisted a little about the central half-screw axis. The structure shows deformed tetrahedra of palladium about sulphur atoms, and three types of deformed square in the arrangement of sulphur about palladium atoms. This latter deformation in two cases retains the plane arrangement, but is not square, in the third, retains the square arrangement but is not planar.

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