08.2-9 NON-PERIODIC PATTERSON FUNCTION OF β -AgI. By <u>M. Tokonami</u> and N. Haga. Faculty of Science. University of Tokyo. Tokyo. Japan and K. Koto and A. Yoshiasa. ISIR. Osaka University. Osaka, Japan.

The experimental approach to find out the difference of the correlation of atomic fluctuations between the nearest neighbors from that between the sufficiently distant atoms has been done by means of X-ray single crystal diffractometry. Fourier transform of a square of continuous |F| distribution would give a kind of Patterson function map. which has pseudoperiodicity in the Patterson space. The peaks near the origin in that Patterson map should be very sharp caused by little influence of thermal motion or positional disorder. Whereas, those in the distant region from the origin should correspond to the ordinary Patterson peaks, which are averaged in the unit cell scale. They are considered to be the products of two atoms vibrating independently. β -AgI is well known as the crystal which has the characteristic diffuse streaks caused by the extreme thermal motion of atoms. The diffuse intensity mostly appears in the plane parallel to (1010)* and in the rods paral el to a^* and c^* axes. This wurtzite-type compound should be the convenient specimen for executing the experiment for the above purpose.

Integrated intensities around the points corresponding to one fifth of the basic reciprocal cell vectors were measured on a four-circle diffractometer using monochromatized Mo Ka radiation. The effect of thermal motion on atomic correlations according to distance will be discussed based on the Patterson synthesis of the observed intensity distribution.

08.2-10 CRYSTAL STRUCTURE DETERMINATION OF CoBo₂ (8₃O₆)₂ BY POWDER DIFFRACTION METHOD. BY <u>Huang Dingzhen</u>, Zhou Zicong, Liang Jingkui, Fujian Institute of Research on the Structure of Matter, Academia Sinica, China.

The pseudo-binary system 8a820,-Ca820, has been studied by means of thermal analysis and X-ray diffraction. A compound CoBo₂ (B₃O₆)₂ has been formed in the system, which melts conqruently at 1117±3°C. There exist eutectic horizontals from BaB_2O_4 to $CaBa_2(B_3O_5)_2$ at 1076 ± 3 °C and from $CaBa_2(B_3O_5)_2$ to meter respectively. The density measured is $R=3,\,64\,g/\,cm^3$. The IR absorption spectra show that there may be the $(B_3\,O_6)^{3-}$ rings of boroxol type in CaBo₂ (B₃D₆)₂. A powder SHG test has been corried out, but no SHG effect has been observed. The indexing of the X-ray powder diffraction pattern indicates that Ca8a₂(B_3O_8)₂belongs to the trigonal system with the unit cell dimensions: a=b=7.157Å, c=35.298Å in hexagonal lattice.There are 6 formula units in one hexagonal unit cell. &=3.64q/cm³. Only the reflections of type $-h^{+}h^{+}J^{-}$ ore observed while the reflections hal with odd / are absent. This limits the possible space groups to R3c and R3c. From the fact that no SHG effect can be observed, the space group is assumed to be $R\overline{\Im}c.$ A further structure refinement has been conducted by the comparison between the calculated intensities l_c and the observed intensities I_o . $I_c=M\cdot k\cdot PL\cdot f^{-2} \exp{(-2Bsin^2\theta/\Lambda^2)}$, where M is the multiplicity factor, PL the Lorentz polarization factor, the multiplicity factor, P_{L} the Lorentz polarization factor, the atomic scattering factor in F is calculated by relativistic Hartree-Fack wave function (D.T. Cromer, J. T. Wober, "International Table for X-ray Crystallagraphy." Vol. IV, PP. 71. 184. Birmingham, Kynoch, 1974.) refined by anomalous scattering. B, the average temperature factor, and k, the proportional constant. By using 2B = 3. \mathbb{R}^{2} , we obtain the refinement result $R = \sum |I_{c} - I_{d}| / \sum I_{a} = 117$ with 75 possible reflections of $I_{c} > 0.25$ (the strongest one is 100.). The final atomic parameters.

interatomic distances and band angles of $(B_3 D_6)^{3-}$ rings are listed in Tables 1 and 2. The $(B_3 D_6)^{3-}$ ring is shown in Fig.1.

The result of structure determination indicates that both $CaBa_2(B_3D_6)_2$ and high temperature phase BaB_2D_4 (A. D. Mighel], A. Perloff, S. Block. Acta Cryst. 2D, 1966, 819.) are isostructural compounds. The high temperature structure of BaB_2D_4 is stabilized by the substitution of Ca^{2*} for Ba^{2*} , as is the case in $Ba_{1\times}Sr_{\times}B_2D_4$ (x<0.35) solid solution in the BaB_2D_4 -SrD system (Wang Guofu, Huang Dingzhen, Liang Jingküi. Acta Chimica Sinica, 42, 6, 1984, 503.). The other isostructural compound $CdBa_2(B_3D_6)_2$ (a=b=7.144 Å, c=34.562 Å, Z=6.) had been found by using Cd^{2*} . It can be seen that the high temperature phase structure of BaB_2D_4 is stabilized by the substitution of $Divalent cations with smaller ionic radius for <math>Ba^{2*}$. But no isostructural compounds can be formed by the substitution of $^2A^{2*}$ and Mg^{2*} with much smaller ionic radius.

Tab.	le I. The fin	0(1)			
	rusitions	x/u	y/ 0	2/0	φų
0())	36(f)	D.159	0.213	D. 0449	× 12,
0(2)	36(f)	D. 420	0.104	0.0403	4 122 B 0 (12)
B	36(f)	0.944	0.162	0.0425	
Ba	12(c)	0	0	0.3515	Fig. 1. The (8305) ³⁻
Ca	6 (a)	0	۵	0.25	ring.
Tab.	le 2. The in	nterato	mic dista	ances and (8-0-13-)	band angles.

(8 ₃ 0 ₆) - 1	Uistances	(8306) * Angles
B-D(1)	1.40 Å	B-D(1')-B 122°
B-D(1')	1.39 Å	D(1')-B-D(1) 118°
8-0(2)	1.31 Å	0(1')-B-D(2) 120°
		0(2)-B-0(1) 122°
8-8	2.43 Å .	8a-0 and Ca-0 distances
		Bo-D(2) 2.97 Å
0(1)-0(1') 2.37 Å	Ba−D(1) 2.99 Å
0(1)-0(2)	2.36 Å	Ba−D(2′) 2.68 Å
0(1')-0(2) 2.35 Å	Co-D(2) 2.28 Å

08.2-11 INTERACTION PRODUCTS OF CONDENSED PHOSPHORIC ACIDS WITH VARIOUS AMINES AND AMINO ACIDS. By M.T. Averbuch-Pouchot and <u>A. Durif</u>, Laboratoire de Cristallogaphie, associé à l'U.S.T.M.G., C.N.R.S., 166 X, 38042 Grenoble Cedex (France).

Interactions between amines, amino acids and various condensed phosphoric acids $(H_4P_2O_7, H_3P_3O_9, H_5P_3O_{10}, H_4P_4O_{12}...)$ have been investigated. Being given the instability of the condensed phosphoric acids a metathesis reaction between the amine chlorohydrates and the phosphoric silver salts has been used for their preparation. A typical reaction of this kind is :

 $3[(CH_3)_2CH.NH_3]CI+Ag_3P_3O_9 \rightarrow [(CH_3)_2CHNH_3]_3P_3O_9+3AgCI$

We describe here the first results obtained in the case of the cyclic phosphoric acid : $H_3P_3O_9$.

Formules	a	ь	с	ß	S.G.	z
(CH3.NH3)3P3O9	12.144	15.362	7.203	97.32	P2 ₁ /n	4
(C ₂ H ₅ .NH ₃) ₃ P ₃ O ₉	16.284	19.17	12.425	118.62	C2/c	4
[(CH ₃) ₂ CH.NH ₃] ₃ P ₃ O ₉	25.22	12.225	15,451	123.90	C2	8
(CH2.COOH.NH3)3P3Og	12.31	14.62	10.146	100.40	P21/c	4

To the difference with previous investigations made with monophosphoric acid : H_3PO_4 the phosphoric group is here a non acidic one : P_3O_9 . The three dimensional cohesion is assured by hydrogen bonds between the external oxygen atoms of the P_3O_9 ring anion and the hydrogen of the NH₃ radicals.