

## XPS AND ELECTRONIC STRUCTURE OF FERROELECTRIC $\text{Sn}_2\text{P}_2\text{S}_6$ CRYSTALS

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Electronic properties of uniaxial ferroelectric  $\text{Sn}_2\text{P}_2\text{S}_6$  single crystal are studied by X-ray photoelectron spectroscopy. X-ray photoelectron spectra (XPS) of the valence band (VB) and of the principal core levels (CL) are obtained from different crystallographic planes in both paraelectric and ferroelectric phases. The XPS were measured with monochromatized Al K $\alpha$  radiation in the energy range 0–1400 eV. The VB consists of five bands with the maxima between 3.3 and 14.5 eV below the Fermi level. Experimental energies of the VB and core levels are compared with results of theoretical *ab initio* calculations of molecular model of the  $\text{Sn}_2\text{P}_2\text{S}_6$  crystal. Electronic structure of the VB is revealed. Ferroelectric phase transition changes atom's charge and bonds strength, VB electronic structure, CL lines width, and chemical shifts for the Sn, P, and S states which are crystallographic plane-dependent.

**Keywords:**  $\text{Sn}_2\text{P}_2\text{S}_6$ , ferroelectric, crystals, XPS, electronic structure

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### 1. Introduction

X-ray photoelectron spectroscopy is widely used for investigations of the electronic structure of solids. Due to reduced co-ordination number, the surface atoms experience a different potential than the bulk atoms. Therefore, the core-electron binding energies are different for surface and bulk atoms. XPS enables one to study the surface valence changes, which affect the core-level shift with respect to the bulk atoms. Form and electronic structure of the VB and CL were studied both theoretically and experimentally in the quasi-one-dimensional  $\text{Sb}_2\text{S}_3$  [1] and  $\text{Bi}_2\text{S}_3$  [2],  $\text{SbSI}$  [3],  $\text{BiSI}$  [4], and  $\text{SbSeI}$  [5] single crystals. XPS revealed huge crystallographic plane-dependent splitting of the core-level binding energies in ferroelectric  $\text{SbSI}$ , and smaller core-level shifts in non-ferroelectric  $\text{BiSI}$  and  $\text{SbSeI}$ . Theoretical *ab initio* calculations confirmed these findings. The valence band and core levels of these quasi-one-dimensional crystals are extremely sensitive to changes of the chemical environment of atoms.

$\text{Sn}_2\text{P}_2\text{S}_6$  is a well-known uniaxial ferroelectric semiconductor, which exhibits a number of prominent

strongly coupled semiconductive and ferroelectric properties that are discussed in detail in the monograph [6]. The room-temperature ferroelectric (FE) phase is a monoclinic one, with the space group  $Pc$ . Above  $T_c = 337$  K, the crystal undergoes a second-order ferroelectric phase transition into paraelectric (PE) monoclinic space group  $P2_1/c$ . The shapes of the coordination polyhedrons of chalcogen atoms do not differ significantly. Below  $T_c$ , all four Sn atoms are shifted significantly in the  $[100]$  direction, with respect to the locations in the centrosymmetric state. Two of them are shifted along the  $[010]$  direction by  $0.04 \text{ \AA}$ , while the remaining two are shifted by the same distance along  $[0\bar{1}0]$ . The two non-equivalent Sn atoms are shifted by  $0.325 \text{ \AA}$  along  $a$ ,  $0.044 \text{ \AA}$  along  $b$ , and  $0.094 \text{ \AA}$  along  $c$  and by  $0.225 \text{ \AA}$  along  $a$ ,  $0.044 \text{ \AA}$  along  $b$ , and  $0.033 \text{ \AA}$  along  $c$  direction, respectively. Two pairs of Sn atoms have different shifts in the FE phase. This is related to nonlinear interaction of soft  $B_u$  and full symmetrical mode  $A_g$  modes in the PE phase. This interaction is very important for the phase transition nature. Changes in the position of P and S atoms at  $T_c$  are small. Thus,

Sn atoms shift mainly along the [100] direction and this allows considering the Sn sublattice as ferroactive.

Ferroelectricity in ferroelectric semiconductors is closely related to their electronic structure [7]. However, electronic properties of this crystal were less studied than structural and phonon properties [6]. Electronic structure of  $\text{Sn}_2\text{P}_2\text{S}_6$  was studied by X-ray photoelectron and X-ray fluorescence spectroscopies in combination with the full-potential linearized augmented plane wave (FLAPW) band structure calculations only in ferroelectric phase [8]. Due to a relative complexity and low symmetry of the crystal, one can expect different XPS from the different crystal planes (as in SbSI [3]) as well as for the PE and FE phases. We failed to find any studies of the ferroelectric phase transition influence on the electronic structure of this crystal.

The purpose of this paper is to study XPS and electronic structure of the ferroelectric  $\text{Sn}_2\text{P}_2\text{S}_6$  crystals in both FE and PE phases, and to reveal the influence of the ferroelectric phase transition on electronic structure of valence band and on shifts in core-level binding energies.

Organisation of the paper is as follows. A brief description of experimental details is given in Section 2. Section 3 presents a molecular model of the  $\text{Sn}_2\text{P}_2\text{S}_6$  crystal and the Hartree–Fock–Roothan (HFR) method of *ab initio* calculations of photoionization energies. The calculated bonds strength and atom's charge in the PE and FE phases as well as the calculated binding energies are presented and compared with the experimental findings. In Section 4, the results of calculated VB electronic structure and XPS of VB and CL for  $\text{Sn}_2\text{P}_2\text{S}_6$  crystal from different crystallographic planes in the both – FE and PE – phases are presented. It is shown that the ferroelectric phase transition changes atom's charge and bonds strength, VB electronic structure, and CL lines width, as well as chemical shifts for the Sn, P, and S states, which are crystallographic plane-dependent. Finally, conclusions are given in Section 5.

## 2. Experimental details

The  $\text{Sn}_2\text{P}_2\text{S}_6$  crystal was grown by vapour transport. The crystal of good optical quality was 2 cm long and 1 cm<sup>2</sup> in area. The XPS of valence band and principal core levels were measured using a PHI 5700/660 Physical Electronics spectrometer with monochromatic Al  $K_\alpha$  radiation (1486.6 eV) of 0.3 eV full width at half maximum. The photoelectron spectra as functions of

Table 1. Atomic concentration at different planes in FE (RT) and PE (360 K) phases.

Peak	Atomic concentration					
	<i>yz</i> plane		<i>xy</i> plane		<i>xz</i> plane	
	RT	360 K	RT	360 K	RT	360 K
P 2p	22.4	21.8	22.3	22.3	22.5	23.6
S 2p	56.1	56.4	56.3	57.3	55.8	54.7
Sn 3d <sup>5</sup>	21.5	21.8	21.4	20.4	21.7	21.7

Table 2. Chemical composition at different planes in FE (RT) and PE (360 K) phases.

Element	Chemical composition					
	<i>yz</i> plane		<i>xy</i> plane		<i>xz</i> plane	
	RT	360 K	RT	360 K	RT	360 K
P	2.2	2.2	2.2	2.2	2.3	2.4
S	5.6	5.6	5.6	5.7	5.6	5.5
Sn	2.2	2.2	2.1	2.0	2.2	2.2

kinetic energy were analysed in the 0–1400 eV energy range by a hemispherical mirror analyzer. All spectra are obtained using 400  $\mu\text{m}$ -diameter analysis area. The measurements were performed from the crystallographic planes perpendicular to ferroelectric *x* axis, *z* axis, and non-ferroelectric *y* axis, cleaved *in situ* in the low  $10^{-10}$  torr range vacuum to obtain the clean surfaces. The survey spectra taken directly after breaking the crystal showed small contamination by oxygen and a rather low one by carbon that should have no influence on the VB and CL spectra. The angle was 45° between the sample and X-ray incident beam. A charging effect was observed. All spectra were corrected for this charging effect using the carbon 1s line of adsorbed carbon ( $E_b = 285$  eV).

The most intensive core-level lines are selected to specify the stoichiometry of the crystal. The atomic concentration specified by the peak areas is shown in Table 1. The experimentally determined atomic concentration is in good agreement with the ideal stoichiometry of the crystal. The chemical composition is given in Table 2. This indicates that the crystal has a high quality with regard to the chemical purity. However, the presented quantities slightly depend on the crystallographic plane and the phase.

## 3. Molecular model of $\text{Sn}_2\text{P}_2\text{S}_6$ crystal and *ab initio* calculation of the energy levels

The electronic structure of VB and CL was calculated by the method based on HFR equations solutions, in the Linear Combinations of Atomic Orbitals

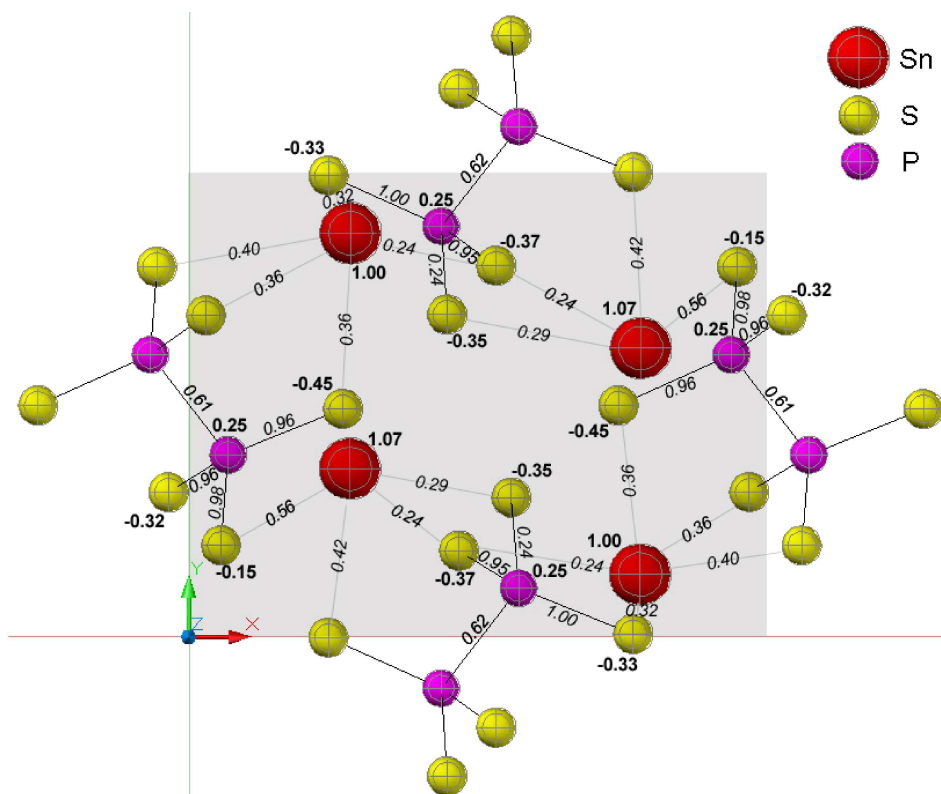


Fig. 1. Sn<sub>8</sub>(P<sub>2</sub>S<sub>6</sub>)<sub>8</sub> cluster as a molecular model of Sn<sub>2</sub>P<sub>2</sub>S<sub>6</sub> crystal on the *xy* plane. The calculated bond strengths and atom's charges in the PE phase are shown in the picture.

(LCAO) approach for the molecular orbitals (MO). According to the Koopmans' theorem, the one-particle energies obtained from the canonical HFR equations correspond to the approximate ionization potentials. This would be an exact solution if the ionisation process were instantaneous and the correlation energy contribution could be completely neglected.

For calculating the energy levels we need a molecular model of the crystal. The model must be a cluster composed from an even number of molecules. However, the interaction between the clusters is not as weak as it should be. Figure 1 shows a fragment of the crystal structure on the *xy* plane. The unit cell is shadowed. This selected Sn<sub>4</sub>(P<sub>2</sub>S<sub>6</sub>)<sub>4</sub> cluster gives the stable HFR solution. Such a cluster was used for theoretical calculations.

The molecular orbital (MO) is a linear combination of atomic orbitals (AO). The MO ( $\varphi_i$ ) can be expanded in the AO ( $\chi_\mu(r)$ ) basis:

$$\varphi_i(\mathbf{r}) = \sum_{\mu=1}^M C_{i\mu} \chi_\mu(\mathbf{r}), \quad (1)$$

where  $\mu$  is the number of the AO, or the set of quantum numbers  $nlm$ . For simplicity, a Gaussian 3G orbital

basis set [9] is used. The **C** matrix is obtained by solving Hartree–Fock matrix equation

$$\mathbf{F} \mathbf{C} = \mathbf{S} \mathbf{C} \boldsymbol{\varepsilon}. \quad (2)$$

Equation (2) is solved by diagonalization as described in [1–3]. As the matrix **F** nonlinearly depends on unknown coefficients  $C_{i\mu}$ , the Eq. (2) is solved by iterations method. The diagonal matrix  $\boldsymbol{\varepsilon}$  gives the energies of MO levels. For the core levels they nearly correspond to the energy of the AO levels. However, due to interaction they are slightly diffused. Method of finding the overlap integrals  $S_{\mu\nu}$  and the Fock's matrix (**F**) elements is also described in [1–3].

The coefficients  $C_{i\mu}$  allow us to calculate the electronic structure of molecules and to interpret the experimental results. They also allow us to find the matrix of the electron distribution density, according to Mulliken, which is given by

$$P_{\mu\nu} = 2 \sum_{i=1}^{N/2} C_{i\mu} C_{i\nu}. \quad (3)$$

Here  $N$  is the number of electrons. For the closed shells  $N$  is even.

Table 3. Sn–S bonds strength in FE and PE phases. Numbers of atoms are given in Fig. 1.

SnI–S	FE	PE	Sn2–S	FE	PE	Sn3–S	FE	PE	Sn4–S	FE	PE
1–9	0.41	0.56	2–10	0.55	0.40	3–11	0.19	0.24	4–13	0.23	0.36
1–15	0.10	0.24	2–11	0.48	0.36	3–17	0.19	0.56	4–15	0.35	0.36
1–30	0.42	0.42	2–19	0.45	0.36	3–26	0.05	0.29	4–18	0.32	0.40
1–33	0.38	0.24	2–31	0.36	0.32	3–27	0.07	0.24	4–27	0.46	0.24
1–35	0.43	0.29	2–33	0.10	0.24	3–32	0.28	0.42	4–29	0.33	0.32

The bond strengths  $P_{AB}$  between the atoms A and B

$$P_{AB} = \sum_{\mu \in A} \sum_{\nu \in B} P_{\mu\nu} \quad (4)$$

and the charge of the atoms is given by

$$q_A = Z_A - \sum_{\mu \in A} P_{\mu\mu}. \quad (5)$$

This method of calculations is implemented in GAMESS program [10], which can deal with the molecular clusters up to  $M \leq 2000$  orbitals or 250 atoms in MINI base. MIDI base is twice bigger for valence electrons and better describes the VB. We used both MINI and MIDI 3G orbital basis sets. The calculations are done using the experimental lattice parameters [6]. The calculated bond strengths and the Mulliken charge of the atoms in the PE phase are shown in Fig. 1. In this model, Sn atom's charge is close to +1 and the crystal is ionic  $\text{Sn}_2^+(\text{P}_2\text{S}_6)^-$ . At the FE phase transition, the bonds strength and charge of atoms change. The greatest changes occur in Sn3 surroundings. These changes are given in Table 3. Those results show that a rearrangement of most of the bonds takes place at the

ferroelectric phase transition. The valence of ferroactive Sn ions also changes: Sn1 is 3.03 and 2.99, Sn2 is 2.81 and 2.72, Sn3 is 1.92 and 2.99, and Sn4 is 2.81 and 2.72 in the PE and FE phase, respectively. The valence and charge of Sn3 ions change at phase transition most of all, i. e. they decrease in the FE phase (charge decreases from 1.07 to 0.95).

However, the cluster  $\text{Sn}_4(\text{P}_2\text{S}_6)_4$  obviously does not reflect the symmetry of  $\text{Sn}_2\text{P}_2\text{S}_6$  crystal unit cell, which contains two formula units –  $\text{Sn}_4(\text{P}_2\text{S}_6)_2$ . This determines different charges of Sn1 and Sn3 ions in the paraelectric phase.

Table 4 presents theoretical values of binding energies of the  $\text{Sn}_4(\text{P}_2\text{S}_6)_4$  cluster without taking into account the spin–orbit interaction. The quantum mechanical method and the chosen cluster give higher negative CL as well as VB energies than their experimental values are (see Table 5). Nevertheless, the cluster reflects the electronic structure and binding energies of the crystal.

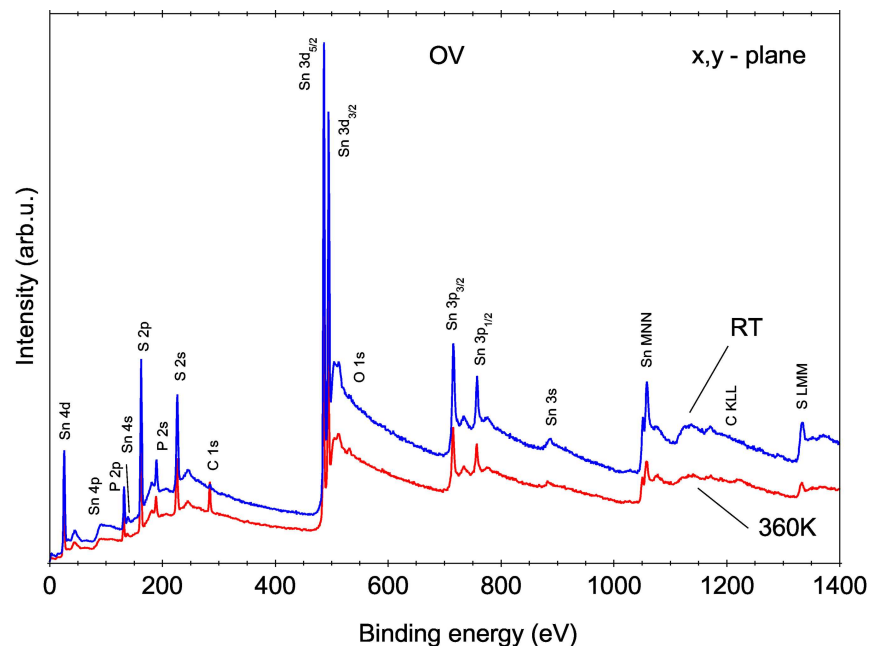


Fig. 2. Overview spectra of  $\text{Sn}_2\text{P}_2\text{S}_6$  crystal from the  $xy$  plane in the PE (360 K) and FE (RT) phases.

Table 4. Values of the calculated binding (negative) energies (eV).

State	FE phase	PE phase
Sn 3s	860.7–859.7	860.3–860.2
Sn 3p	741.2–740.1	740.8–740.7
Sn 3d	520.2–519.1	519.8–519.7
S 2s	246.3–240.1	244.9–241.4
P 2s	210.1–207.9	209.5–208.8
S 2p	182.3–175.9	180.8–177.2
P 2p	153.1	152.7
Sn 4s	149.7	150.6
Sn 4p	111.2–110.1	110.7
Sn 4d	40.5–39.4	40.1–39.9

## 4. Results of XPS and electronic structure

### 4.1. Survey spectra

Figure 2 shows the XPS of the Sn<sub>2</sub>P<sub>2</sub>S<sub>6</sub> crystal in the energy range from 0 to 1400 eV below the Fermi level in the both, PE (360 K) and FE (RT), phases from the crystallographic *xy* plane. Only small amounts of oxygen (O 1s) and carbon (C 1s) have been detected. The spectra from other crystallographic planes are similar. However, CL peaks in the ferroelectric phase are higher. Inelastically scattered electrons give the background. Auger spectra of Sn MNN, S LMM, and C KLL are also seen in the high energy range. The strongest peaks of Sn 3d, Sn 4d, Sn 3p, P 2p, S 2p were

Table 5. Binding energies and chemical shifts of atoms at different planes in FE (RT) and PE (360 K) phases.

Peak	Binding energy (eV)					
	<i>yz</i> plane		<i>xy</i> plane		<i>xz</i> plane	
	RT	360 K	RT	360 K	RT	360 K
P 2p <sub>3/2</sub> in compound	132.0	131.8	132.6	132.1	132.3	131.8
P 2p <sub>3/2</sub> in literature			130.0			
chemical shift	2.0	1.8	2.6	2.1	2.3	1.8
P 2p <sub>1/2</sub> in compound	132.9	132.6	133.5	133.0	133.2	132.6
P 2p <sub>3/2</sub> in literature			131.0			
chemical shift	1.9	1.6	2.5	2.0	2.2	1.6
S 2p <sub>3/2</sub> in compound	162.3	161.9	162.7	162.3	162.4	161.7
S 2p <sub>3/2</sub> in literature			164.0			
chemical shift	−1.7	−2.1	−1.3	−1.7	−1.6	−2.3
S 2p <sub>1/2</sub> in compound	163.5	163.0	163.9	163.5	163.6	162.9
S 2p <sub>3/2</sub> in literature			165.0			
chemical shift	−1.5	−2.0	−1.1	−1.5	−1.4	−2.1
Sn 3d <sub>5/2</sub> in compound	486.7	486.5	486.9	486.3	486.8	486.3
Sn 3d <sub>5/2</sub> in literature			485.0			
chemical shift	1.7	1.5	1.9	1.3	1.8	1.3
Sn 3d <sub>3/2</sub> in compound	495.1	494.9	495.3	494.7	495.5	494.6
Sn 3d <sub>3/2</sub> in literature			493.0			
chemical shift	2.1	1.9	2.3	1.7	2.5	1.6

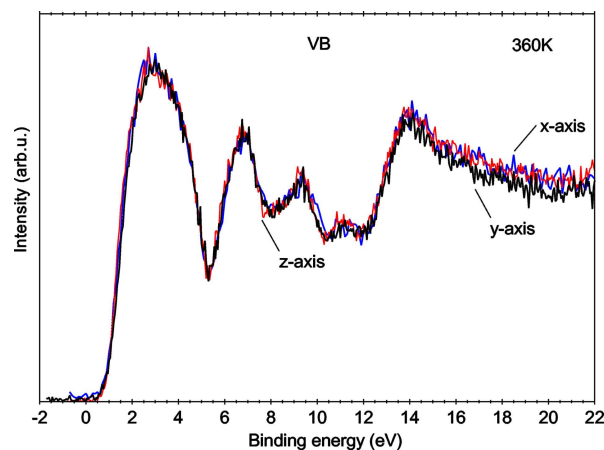


Fig. 3. XPS of VB in the PE phase from the crystallographic planes perpendicular to all the three axes.

chosen for investigation of peculiarities of the crystallographic plane-dependent core-level XPS in this ferroelectric semiconductor.

### 4.2. Valence band

X-ray photoelectron spectroscopy of the valence band provides data on the occupied total density of states. Figure 3 shows the VB spectra in the PE phase from the crystallographic planes perpendicular to all three axes. The VB consists of five distinguishable bands with the maximum intensity at 2.8, 6.7, 9.2, 11.0,

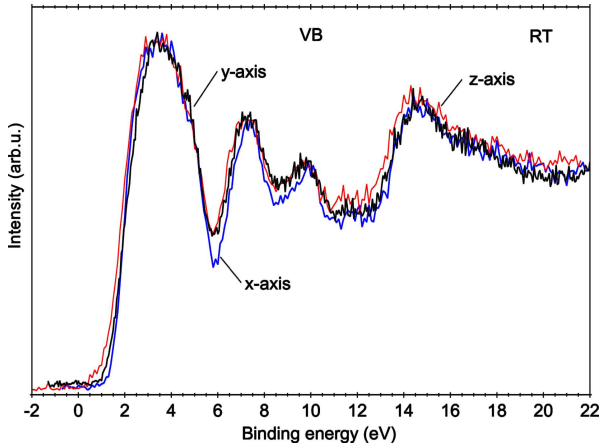


Fig. 4. XPS of VB in the FE phase from the three crystallographic planes.

and 14.0 eV. The similar but more outspread bands are in the FE phase (Fig. 4). Their energies are shifted by approximately 0.5 eV to higher values and coincide with those obtained in [6]. The 11.0 eV band in this phase is nearly erased. In the PE phase, the VB is separated by a gap of about 1.1 eV from the Fermi level. In the ferroelectric RT phase, the VB is located at about 1.5 to 22 eV below  $E_F$ . In these crystals, the optical band gap at room temperature is 2.3 eV. It means that the Fermi level is pinned nearly in the middle of the band gap.

Figure 5 shows the calculated PE phase VB form and electronic structure of  $\text{Sn}_2\text{P}_2\text{S}_6$  crystal. The spectrum is referred to the Fermi level  $E_F$ . The  $E_F$  was determined experimentally within the accuracy of 0.3 eV. The intensities of the XPS were described:

- (a) by the energy states band  $\varepsilon_i$  from the characteristic Eq. (2);
- (b) by the peaks of the density of states:

$$D(\varepsilon) = \frac{1}{N_M} \frac{1}{\Delta\varepsilon}, \quad (6)$$

where  $N_M$  is the number of molecules in the cluster;

- (c) by the Gaussian broadening [11]:

$$D(\varepsilon) = \frac{1}{\sqrt{2\pi}\sigma} \sum_i \exp\left[-\frac{(\varepsilon - \varepsilon_i)^2}{2\sigma^2}\right], \quad (7)$$

where the summation is performed over the states  $i$ ,  $\varepsilon_i$  are the corresponding energy levels, and  $\sigma$  is the half-width of the Gaussian function.

The Gaussian broadening method is the simplest approach to the Brillouin-zone spectral integrations. The  $k$ -points are distributed as evenly as possible throughout the Brillouin zone, and the DOS  $D(\varepsilon)$  is formed according to Eq. (6). This method is designed for efficient

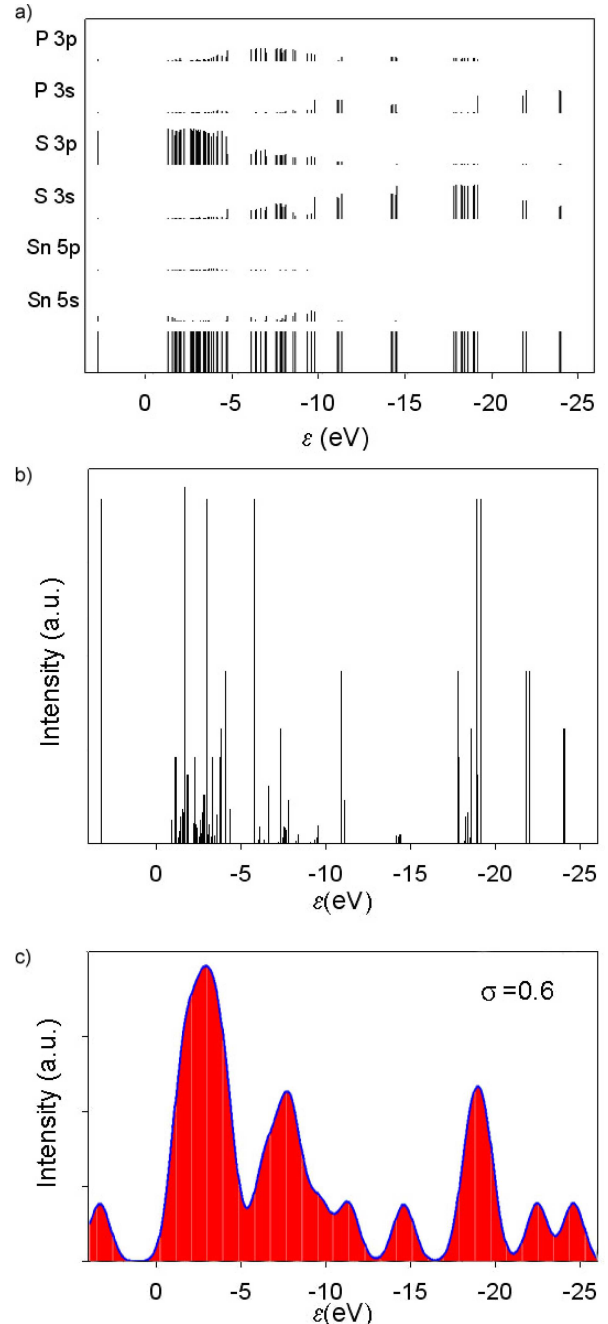


Fig. 5. Electronic structure of VB in PE phase: (a) DOS band and contribution of states (%); (b) intensity of DOS ( $\Delta\varepsilon = 0.27$  eV taken in Eq. (6)); (c) calculated VB for  $\text{Sn}_4(\text{P}_2\text{S}_6)_4$  cluster and approximated by the Gaussian broadening method (Eq. (7)).

calculation of experimentally observed crystal spectral properties where experimental and lifetime broadenings place a limit on the resolution required.

Further, all experiments have a finite energy resolution and frequently concentrate on the shifts, distortions, and changes in weight in the spectral features. In the approach used (Eq. (7)), the aim was to calculate the DOS only at the high density of  $k$ -points and



then to smear the resulting spectrum to the experimental resolution. However, the application of the Gaussian smearing to the DOS results in good convergence at low  $k$ -point sampling densities.

The form of the VB can be explained by the analysis of MO population. Knowing the MO coefficients  $C_{i\mu}$  (Eq. (1)), we can evaluate the contribution of the A atom electrons to  $\varepsilon_i$  state:

$$p_{iA} = \sum_{\mu \in A} C_{i\mu}^2. \quad (8)$$

From the partial density of states (PDOS) (in %) calculated by Eq. (8) and the total density of states (a), the representation  $D(\varepsilon)$  for the peaks (b) of the DOS (Eq. (6)), and approximation of the bands spectra (c) by the Gaussian broadening method (Eq. (7)) for  $\text{Sn}_4(\text{P}_2\text{S}_6)_4$  molecular cluster (Fig. 5) it follows that from 4 to 2 eV there is a conduction band (CB), the right edge of which is formed by S 3p electrons. Unlike  $\text{Sb}_2\text{S}_3$ -type [1, 2] and SbSI-type [3–5] crystals, the VB consists of five bands. It is dominated by Sn 5s, Sn 5p, S 3s, S 3p, P 3s, and P 3p states. Five DOS peaks are degenerate. The main contribution to the most intensive band between  $-1$  and  $-5$  eV is given by S 3p states. The left edge of this band is formed by S 3p states with 13% Sn 5s and few percent P 3p states. The main contribution to the next intensive band between  $-5$  and  $-12$  eV comes from S 3s, S 3p, P 3p, and Sn 5s states. Sn 5s and 5p states give only small contribution to the first two bands. The low-energy part of VB is formed mainly by S 3s and P 3s states. S 3s electrons form band at  $-14$  eV, while P 3s electrons form band at  $-18.5$  eV. Above  $-25$  eV dominate P 3s with mixture of S 3s states. The five experimental bands at  $-3.3$ ,  $-7.2$ ,  $-9.7$ ,  $-11.5$ , and  $-14.5$  eV are in good agreement with [8].

The calculated bands are shifted to higher energies, however. It means that electron–electron interactions are overestimated. CL electrons extend the VB to higher energies. This is also confirmed experimentally. Higher-energy edges of all the bands are more outspread (Figs. 3, 4). The last three high-energy bands experimentally are seen as one diffused band. Tin atoms are strongly ionized. The electron density from Sn goes primarily to the neighbouring S atoms, yielding nearly full occupation of the S 3p band. The calculated PDOS in this approach differs slightly from the FLAPW calculations [8]; however, the main VB features are the same. Intensity of DOS is higher and minima are deeper than in FE phase. Due to the quasi-

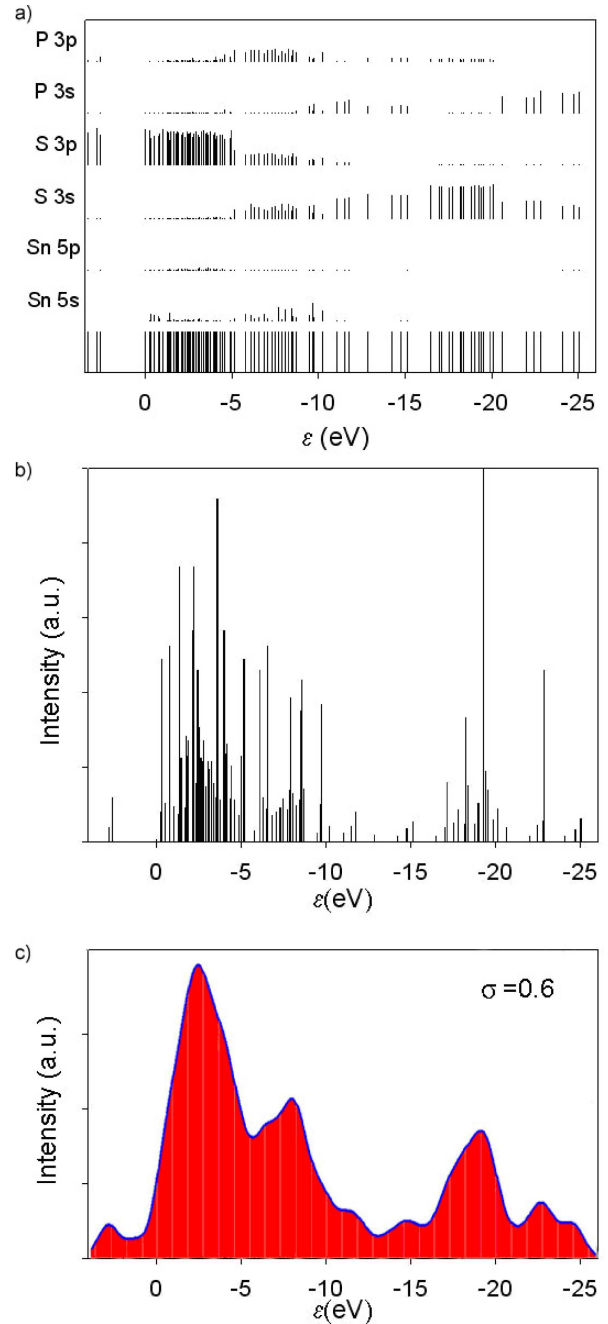


Fig. 6. Electronic structure of VB in FE phase. Designations are the same as in Fig. 5.  $\Delta\varepsilon = 0.021$  eV taken in Eq. (6), therefore the intensities are lower and minima are shallower than in PE phase.

degenerate levels, the PE phase DOS band is rarefied in comparison with FE phase DOS.

Figure 6 shows the calculated VB form and electronic structure of FE phase. The gap between the VB and CB decreases from 4.1 eV (in PE phase) to 2.6 eV. The calculated width of the VB is 24.9 eV, while in PE phase it is 23.1 eV. In FE phase, there is only one quasi-degenerate state instead of five in PE phase. Disappearance of centro-symmetry lifts degeneracy of the

electronic states. The left edge of the most intensive band between  $-1$  and  $-5$  eV is formed by S 3p states with few percent of P 3p states. Sn 5s states disappear. The phase transition does not change essentially the electronic structure of the crystal.

Mechanism of the tin cations electron lone pair formation, which is related to the appearance of the spontaneous polarization [12], could be used for explaining the VB spectra evolution at transition from the PE phase to FE one. The hybridization of  $sp^2$  type determines peculiarities of the density of states in VB. Antibonding mixing of Sn 5s – S 3p orbitals gives states at the top of VB. Bonding interaction of Sn 5p orbitals with previous antibonding Sn 5s – S 3p states generates lower-energy filled states Sn 5p + (Sn 5s – S 3p),  $sp^2$  for short. By this the gain of electronic subsystem energy is determined. Lowering of these  $sp^2$  states energy is proportional to acentricity of the Sn ions surrounding (similarly to the known Jahn–Teller effect). Such formation of the Sn lone pair electron “cloud” together with deformation of nearest sulphur atoms polyhedron determine the origin of spontaneous polarization. So, at transition to the FE phase and increase of spontaneous polarization the discussed  $sp^2$  hybridization becomes stronger and by this the density of electronic states at top of the VB is lowered. The observed lowering of energy of the band with maximum near 3 eV in the VB XPS on cooling from 360 K to RT could be interpreted in such manner.

#### 4.3. Core levels

Figures 7–9 show the spectra of Sn 3d, Sn 4d, and S 2p spin–orbit doublets from the  $xy$  plane in the FE and PE phases. In the FE phase, all the CL bands are broader than in the PE phase. This indicates stronger interactions and higher bond strengths. However, even in the PE phase the peak width, i. e. the interaction, is also crystallographic plane-dependent (Fig. 10). The strongest interactions are observed for the polar  $yz$  plane.

Sn 3d and 4d spectra could be described by two lines in PE phase. Indeed, in PE phase all Sn ions are equivalent. In FE phase, two non-equivalent pairs of Sn ions already exist.

Sn 4d spectral lines in PE phase could be described by two Voigt contours with 1.04 eV splitting and 1.02–0.85 eV width. Such spectra obviously are related to four equivalent Sn atoms. In the FE phase, the spectra are best fitted by three doublets with near to 1.04 eV splitting. One doublet is placed at energy position similar to observed for doublet in the PE phase.

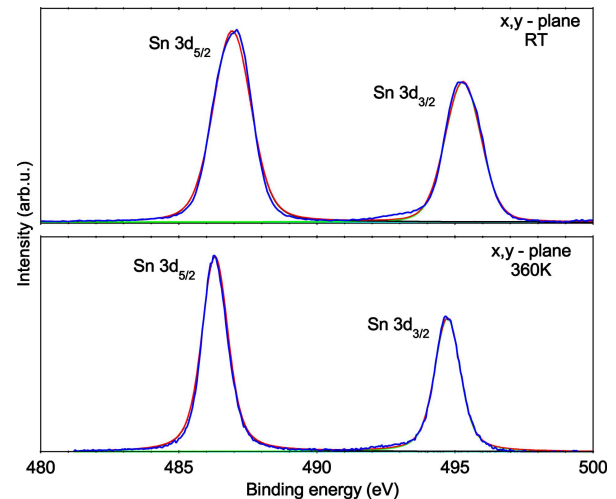


Fig. 7. XPS of Sn 3d spin–orbit doublet in FE (RT) and PE (360 K) phases.

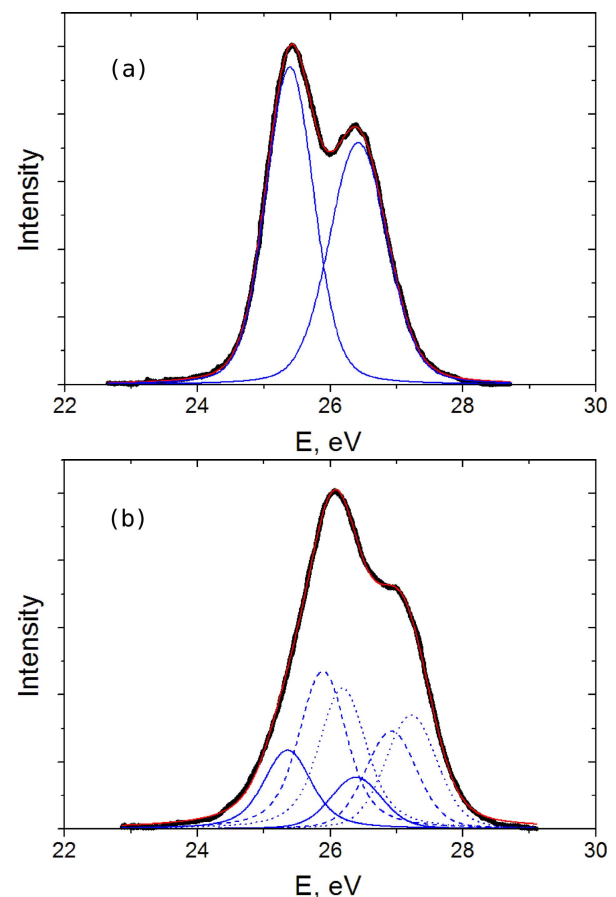


Fig. 8. XPS (in arb. units) of Sn 4d spin–orbit doublet in (a) PE and (b) FE phases.

Another two doublets are shifted to higher energies. Line width is smaller (0.85–0.91 eV) than that found in the PE phase. Such spectra could be explained taking into account the triple-well potential energy surface for  $\text{Sn}_2\text{P}_2\text{S}_6$  ferroelectrics [12]. This lowest energy



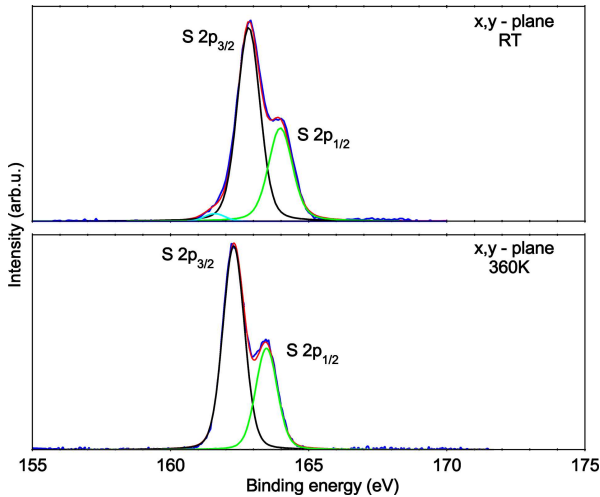


Fig. 9. XPS of S 2p spin-orbit doublet in FE (RT) and PE (360 K) phases.

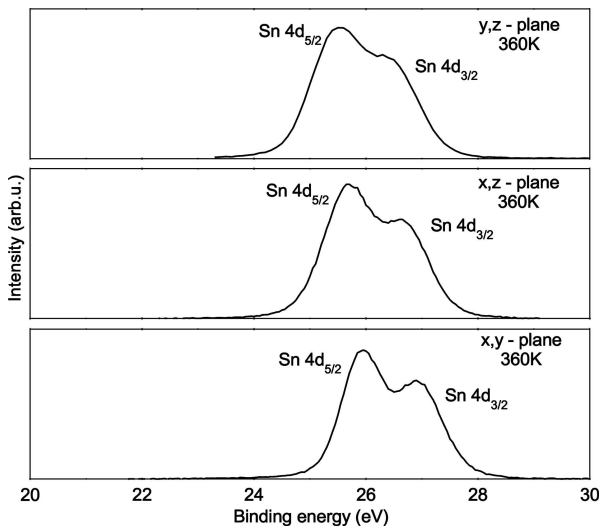


Fig. 10. XPS anisotropy of Sn 4d spin-orbit doublet in PE phase.

doublet could be related to Sn atoms in non-polar elementary cells. Such cells are placed in domain walls of the FE phase. Another two higher energy doublets are related to two non-equivalent pairs of Sn atoms in the FE phase.

Anisotropy of Sn 4d spectra, which were registered from surfaces perpendicular to  $z$ ,  $y$ , and  $x$  directions, in the PE phase is not so high. This anisotropy could be determined by different properties of near-surface layers. In the FE phase, the anisotropy of XPS is clearly seen. For example, Sn 4d  $y$ -cut RT spectra could be fitted by two doublets, which are similar to the two higher-energy doublets in the Sn 4d  $z$ -cut RT spectra. For  $y$ -cut the low-energy doublet, which is traced from the PE phase, is not seen. But for  $x$ -cut in the FE phase this low-energy doublet has the highest intensity. The observed difference for the Sn 4d XPS registered from

Table 6. Anisotropy of the line width in FE (RT) and PE (360 K) phases.

Peak	FWHM (eV)					
	$yz$ plane		$xy$ plane		$xz$ plane	
	RT	360 K	RT	360 K	RT	360 K
P 2p <sub>3/2</sub>	1.0	1.1	1.0	1.0	1.1	1.1
P 2p <sub>1/2</sub>	1.0	0.9	1.0	0.9	1.1	0.9
S 2p <sub>3/2</sub>	1.1	1.1	1.0	0.9	1.2	1.3
S 2p <sub>1/2</sub>	1.0	1.1	1.1	0.9	1.3	1.4
Sn 3d <sub>5/2</sub>	1.3	1.3	1.8	1.3	1.4	1.3
Sn 3d <sub>3/2</sub>	1.3	1.3	1.7	1.3	1.3	1.4

$x$ -,  $y$ -, and  $z$ -cuts in the FE phase could be related to the shape of the domain walls and to their different concentration on crystal surfaces. Also, for the perpendicular to  $y$  direction planes (010), with variation of their position, different non-equivalent pairs of Sn atoms are placed near the surface. For the  $z$ -cut both types of the Sn non-equivalent atoms simultaneously are located at the surface. Probably this structure peculiarity determines greater difference in the spectral area for the doublets in Sn 4d  $y$ -cut RT spectra.

Anisotropy could be a result of different type Sn ions placed at surface. In FE phase, the macroscopic field of spontaneous polarization also modifies surface properties [3] in different manner for the polar  $yz$  and non-polar planes. On the Sn/Ge (111) surface [13] three components in the line shape of the Sn 4d core level also reflect the three inequivalent positions of the Sn atoms that form the unit cell.

The experimental binding energies and chemical shifts of atoms at different planes in both the FE (RT) and PE (360 K) phases are given in Table 5. The electronic structure measurements revealed the chemical shifts of Sn and P states to a higher binding energy and shifts of S states to a lower binding energy. This shift suggests a charge transfer from Sn and P to S atoms. However, the binding energies and chemical shifts are crystallographic plane-dependent. Also, they change at the ferroelectric phase transition. In the FE phase, the chemical shifts of Sn and P atoms are higher while those of S atoms are smaller. Due to the non-equivalent positions of the Sn atoms in FE phase the line width in FE (RT) phase significantly increases (Table 6) mainly in  $xy$  plane.

## 5. Conclusions

X-ray photoelectron spectra of the valence band and of the principal core levels of the semiconductor ferroelectric Sn<sub>2</sub>P<sub>2</sub>S<sub>6</sub> crystal are presented in the energy

range from 0 to 1400 eV. A molecular model of the crystal is used for *ab initio* theoretical calculations of binding energies. Theoretical values of the binding energies are close to the experimental ones. The structure of valence band is calculated and confirmed experimentally. The XPS studies revealed the crystallographic plane-dependent binding energies and chemical shifts which also change at the ferroelectric phase transition. Appearance of non-equivalent positions of Sn atoms in FE phase split the spin-orbit doublets of Sn 3d and 4d spectra.

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## FEROELEKTRINIŲ $\text{Sn}_2\text{P}_2\text{S}_6$ KRISTALŲ RENTGENO FOTOELEKTRONINIAI SPEKTRAI IR ELEKTRONINĖ SANDARA

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### Santrauka

Rentgeno fotoelektronų spektroskopija ištirtos elektroninės vienąsnių feroelektrinių  $\text{Sn}_2\text{P}_2\text{S}_6$  monokristalų savybės. Gauti valentinės juostos (VJ) ir svarbiausių gilių lygmenų paraelektrinės ir feroelektrinės fazių spektrai nuo įvairių kristalografinių plokštumų. Fotoelektronų sužadavimo šaltinis buvo  $\text{Al K}_\alpha$  1486,6 eV monochromatinė spinduliuotė. Sužadintų fotoelektronų spektrai matuoti energijos ruože nuo 0 iki 1400 eV. Eksperimentiškai gautos fotoelektronų energijos yra palygintos su teorinių *ab initio* skaičiavimų

rezultatais molekuliniam  $\text{Sn}_2\text{P}_2\text{S}_6$  kristalo modeliui. Apskaičiuota ir eksperimentiškai patvirtinta kristalo VJ sandara abiejose fazėse. VJ sudaro penkios juostos, o jų smailės yra nuo 3,3 eV iki 14,5 eV žemiau Fermio lygmens. Įvertinti Sn, S ir I atomų cheminiai poslinkiai. Ištirta feroelektrinio fazinio virsmo įtaka VJ sandarai ir gilių lygmenų spektrams. Nustatyta, kaip fazinis virsmas keičia atomų krūvius, ryšių stiprius, VJ elektroninę sandarą ir gilių lygmenų juostų pločius bei Sn, S ir I atomų cheminius poslinkius, kurie priklauso nuo kristalografinės plokštumos.