Electronic Structure and Phase Transition in Ferroelectric Sn$_2$P$_2$S$_6$ Crystal

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(Dated: June 7, 2018)

An analysis of the P$_2$S$_6$ cluster electronic structure and its comparison with the crystal valence band in the paraelectric and ferroelectric phases has been done by first-principles calculations for Sn$_2$P$_2$S$_6$ ferroelectrics. The origin of ferroelectricity has been outlined. It was established that the spontaneous polarization follows from the stereochemical activity of the electron lone pair of tin cations what is determined by hybridization with P$_2$S$_6$ molecular orbitals. The chemical bonds covalence increase and rearrangement are related to the valence band changes at transition from the paraelectric phase to the ferroelectric one.

PACS numbers: 31.15.A-, 77.80.B-, 71.20.-b, 71.70.Ej

INTRODUCTION

For such perovskite ferroelectrics as BaTiO$_3$, the main origin of spontaneous polarization is commonly related to the hybridization interaction between the transition-metal and oxygen ions. Another mechanism involves cations with "lone pair" electrons which have a formal ns$^2$ valence electron configuration. In the same manner as for the d$^0$ transition-metal ions, these p$^0$ ions (as example Pb$^{2+}$ for PbTiO$_3$, or Bi$^{3+}$ for BiMnO$_3$) contain some p-charge density which contribute to the displacive distortions. If the lowering of energy associated with the hybridization interaction is larger than the interionic repulsion opposing the ion shift, then a ferroelectric distortion appears. This "stereochemical activity of the lone pair" is the driving force for off-center distortion in ferroelectrics. Both named origins of ferroelectricity (the first – d$^0$-"ness", and the second – "lone pair" activity) are familiar to the second-order Jahn-Teller (SOJT) effect. This effect is determined by a balance of positive and negative contributions to the total energy. The first one describes short range repulsive forces and is related to the rigid ions (with frozen electronic configuration) shifts from original high symmetry positions. Such term is small for the cases of "closed-shell" d$^0$ or p$^0$ cations. The second, negative, contribution describes the relaxation of electronic configuration in response to the ions displacements through covalent bonds formation. This term favors the ferroelectric distortion. For full picture, the geometrical (or hybrid improper) mechanism, which is related to the rotational modes that trigger instability of polar mode, could also be considered at study of ferroelectricity nature.

The cubic crystal lattices of ABO$_3$ compounds are built by covalent bonds A – O and B – O with considerable contribution of ionicity for former ones. Naturally more complex bounding evolution could be supposed at the ferroelectric phase transition in ion-covalent crystal Sn$_2$P$_2$S$_6$ with monoclinic lattice. For this compound the Sn$^{2+}$ cations and the (P$_2$S$_6$)$^{4-}$ anion clusters are joined by mostly ionic Sn – S bonds at covalent P – S and P – P bonding. Sn$_2$P$_2$S$_6$ uniaxial ferroelectric undergoes the second order phase transition at $T_0 \approx 337$ K ($P2_1/c \rightarrow Pc$, two formula units in the elementary cell for both phases (Fig. 1)) in a crossover displacive-order/disorder region. Ferroelectric instabil-

![FIG. 1. The crystal structure of Sn$_2$P$_2$S$_6$ ferroelectric phase. The tin atoms positions in the paraelectric phase are shown by red. The shape of the Brillouin zone with denoted symmetrical points is shown for primitive monoclinic lattice.](image-url)
the only one lattice mode could determine dynamical instability related to the ferroelectric phase transition\cite{15}. In general, all the 13 \( B_{\alpha} \) and 15 \( A_{\rho} \) optic modes were accounted in the frozen phonons approximation for construction of effective Hamiltonian for \( \text{Sn}_2\text{P}_2\text{S}_6 \) crystal what was applied for the MC simulation of the ferroelectric phase transition and their behavior under hydrostatic pressure\cite{14}. A system with a three-well potential was early considered by Lines\cite{12,13} with application for the \( \text{LiNbO}_3 \) and \( \text{LiTaO}_3 \) crystals. Such system can be generally described by two order parameters (related to dipole and quadruple moments), and as a result, a variety of stable, metastable and unstable states can be realized on a phase diagrams\cite{14,15}.

The strong anharmonicity of \( \text{Sn}_2\text{P}_2\text{S}_6 \) crystal lattice is obviously joined with effective electron-phonon interaction, that appears as a stereochemical activity of the tin cations electron lone pair \( 5s^2 \), and in fact, it is a reflection of the SOJT effect. Possible leading role of the cations’ stereovacuity for \( \text{Sn}_2\text{P}_2\text{S}_6 \) crystals was early noted at X-ray structure investigations of their paraelectric phase in comparison with structure data for the ferroelectric phase\cite{2}. Structural evidences of the tin cation stereoactivity was also analyzed in details by structure refinement of the paraelectric and the ferroelectric phases for \( \text{Sn}_2\text{P}_2\text{S}_6 \) selenide analogs\cite{21}. By Mössbauer effect investigations for \( \text{Sn}^{119}_2 \) nucleus\cite{18} and by NMR spectroscopy for isotopes \( 31\text{P} \) and \( 119\text{Sn} \)\cite{19,15}, the important changes of chemical bonding at the ferroelectric phase transition in \( \text{Sn}_2\text{P}_2\text{S}_6 \) were found. The X-ray photoelectron spectroscopy confirms growth of the chemical bonds covalence in the ferroelectric phase\cite{20}.

\( \text{Sn}_2\text{P}_2\text{S}_6 \) crystals are ferroelectric-semiconductors with promising photorefractive\cite{22}, photovoltaic\cite{23}, electrooptic\cite{24} and piezoelectric\cite{25} characteristics. Their ferroelectric properties are effectively influenced by state of electronic subsystem\cite{26}. Influence of the sulfur and tin vacancies on semiconductive and optic properties of \( \text{Sn}_2\text{P}_2\text{S}_6 \) crystals was studied recently\cite{20}. These data motivate the electronic structure investigation for \( \text{Sn}_2\text{P}_2\text{S}_6 \) crystals in the paraelectric and the ferroelectric phases.

The first-principles calculations in LDA approach of Density Functional Theory (DFT) for \( \text{Sn}_2\text{P}_2\text{S}_6 \) ferroelectric phase were carried out by several groups\cite{22,23}. By Grigas et al\cite{20,21}, the electronic structure of both paraelectric and ferroelectric phases of \( \text{Sn}_2\text{P}_2\text{S}_6 \) was calculated in the cluster approach. For the \( \text{Sn}_2\text{P}_2\text{S}_6 \) selenide analog, the electronic structure have been investigated\cite{21} by first principles calculations only for the paraelectric phase. The electronic structure and phonon spectra pressure dependence for acenctic layered rhombohedral crystal \( \text{Sn}_2\text{P}_2\text{S}_6 \) were investigated theoretically in LDA approach\cite{22}. For this compound, the tin cations are almost fully ionized (Sn\textsuperscript{4+} charge state) what exclude possibility of stereochemical activity of their \( 5s^2 \) electron lone pair. The electronic structure of high charged (\( \text{P}_2\text{S}_6 \))\textsuperscript{2–} and (\( \text{P}_2\text{S}_6 \))\textsuperscript{4–} anion clusters was discussed in papers\cite{23,21,22}.

At analysis of \( \text{Sn}_2\text{P}_2\text{S}_6 \), \( \text{Sn}_2\text{P}_2\text{Se}_6 \) and \( \text{SnP}_2\text{Se}_6 \) electron energy spectra. Analysis of \( \text{P}_2\text{S}_6 \) cluster chemical binding was also done in Hartree-Fock approach\cite{33}. These anions arrangement have been investigated experimentally and theoretically in different approximation for the layered crystals like \( \text{M}_2\text{P}_2\text{S}_6 \) (\( \text{M} = \text{Fe}, \text{Ni}, \text{Mn}, \ldots \)\cite{34–38}). For the \( \text{CuInP}_2\text{Se}_6 \) layered compound with two differently charged cations, the SOJT effect was established as an origin of cooper ferrielectric ordering\cite{39}.

In this paper, the first-principles calculations in LDA approach of DFT for electronic structure of \( \text{Sn}_2\text{P}_2\text{S}_6 \) crystal in the paraelectric and the ferroelectric phases were used for analysis of chemical bonds transformation at the spontaneous polarization appearance and for establishing of the ferroelectric state origin in the phosphorus containing chalcogenides. As a background of the investigations, the free \( \text{P}_2\text{S}_6 \) structure group (atomic cluster) electron spectra and peculiarities of their molecular orbitals were considered. The electron structure and chemical bonding nature in the paraelectric phase will be also analyzed. The stereochemical activity of the electron lone pair of tin cations is examined in detail. The growth of covalence and ions recharging are related to the structure spontaneous polarization. Finally, the influence of tin by lead and sulfur by selenium substitution on crystals properties is discussed.

## I. METHOD OF CALCULATIONS

The calculations of the band structure of both phases of \( \text{Sn}_2\text{P}_2\text{S}_6 \) crystal as well as energy levels of \( \text{P}_2\text{S}_6 \) molecule has been performed by means of the package program ABINIT\cite{40} (total and projected densities of states where calculated using SIESTA\cite{41} software) in the framework of the DFT, using the local density approximation for representing the exchange-correlation interaction. A basis set of \( 28000 \) plane waves, restricted by the kinetic energy \( E_{\text{cut}} = 25 \) Hartree has been used. The tin, sulphur and phosphorus atoms had the following electron configurations: Sn: [Kr] 5s\textsuperscript{2}5p\textsuperscript{6}; S: [Ne] 3s\textsuperscript{2}3p\textsuperscript{4}; and P: [Ne] 3s\textsuperscript{2}3p\textsuperscript{3}, respectively. The 'frozen' core electron configurations for each atom is shown in brackets. The first-principles pseudopotentials in the Hartwigsen-Goedecker-Hutter scheme\cite{42} have been applied and the integration over irreducible part of the Brillouin zone has been done by means of the tetrahedron method using the \( 4 \times 4 \times 4 \) Monkhorst-Pack mesh\cite{43} of \( k \)-points. The chosen parameters were sufficient for a good convergence in the calculations. Prior to commence the calculation of physical properties of \( \text{Sn}_2\text{P}_2\text{S}_6 \) crystal, we carried out the structural optimization, which minimized total energy of the system simultaneously with the forces\cite{44} acting on atoms. The spin-orbit interaction was not taken into account in our calculation. The parameters of \( \text{Sn}_2\text{P}_2\text{S}_6 \) crystal obtained after structural relaxation can be compared with experimental data, presented in\cite{8,9}. The comparison of experimental and calculated relaxed
values of lattice constants demonstrate the difference of about 3% in order of magnitude.

II. ELECTRONIC STRUCTURE OF P₂S₆ CLUSTER

The molecular orbitals of P₂S₆ cluster create covalent P — S and P — P bonds. Their hybridization with tin atomic orbitals determines electronic structure of Sn₃P₂S₆ crystal. The electronic energy spectra of this material could be analyzed by calculation of free P₂S₆ cluster electronic structure with following accounting of their molecular orbitals hybridization with atomic orbitals of tin. The calculated energy spectrum and partial densities of electron states for orbitals of tin illustrate the formation of different molecular orbitals at creation of covalent P — S and P — P bonds. The spatial electron density distribution for related energy levels reflects peculiarities of these bonds. It is seen that for cluster the energy level near −15 eV mostly is determined by hybridization of phosphorus s orbitals. Here it have been also found some contributions of sulfur s and p orbitals. The hybridization of these atomic orbitals (scheme (1) at (Fig. 2)) creates bonding P — P and P — S molecular orbitals. The level near −14 eV is determined by antibonding combination of two phosphorus s orbitals and by bonding hybridization of s orbitals (2) of phosphorus and sulfur atoms.

The group of the levels near −12 eV mostly is formed by sulfur s orbitals. These levels are related to the P — S bonding and P — P antibonding molecular orbitals (3,4,5,6). The levels near −6.5 eV and −8.5 eV appear as a replica of doublet of the levels near −15 eV and −14 eV in results of antibonding hybridization of phosphorus and sulfur atoms s orbitals. The molecular orbitals for levels near −8.5 eV are P — P bonding and P — S antibonding (7), the orbitals with energy near −6.5 eV are antibonding for all P — S and P — P bonds (8). In interval between −3.5 and −4.7 eV, the energy levels are created by bonding hybridization of phosphorus and sulfur p orbitals. These orbitals (9, 10, 11, 12, 13) are bonding for P — P and P — S bonds. In region from 0 till −1.5 eV, the energy levels of cluster are also formed by p orbitals of phosphorus and sulfur atoms. They are hybridized in P — P bonding and P — S antibonding molecular orbitals (14 – 23). Here the contribution from phosphorus d orbitals is also presented.

It is worth to be noted that upmost 23rd energy level of P₂S₆ molecule is double degenerated due to high symmetry but only half occupied. Such peculiarity can cause instability of this complex by means of Jahn-Teller like mechanism.

From detailed analysis of the electron energy spectrum, the important information about creation and character of the chemical bonds in P₂S₆ cluster could be found (Fig. 2). The bond P — P is determined by σ hybridization of phosphorus s orbitals (the level near −15 eV (1)) and by their replica near −8.5 eV (7). In P — P bond, the contribution from π hybridization of pₓ and pᵧ orbitals of phosphorus, that are oriented normally to the bond direction, is also presented. The levels of these orbitals (9) are placed near −4 eV. However, the essential contribution into energy of the P — P bond adds the σ hybridization of phosphorus p₂ orbitals which are oriented along the bond. The bonding combination σ(pₓ + pᵧ) (22) has filled by electrons with the energy level in the range 0 ÷ −1.5 eV. Also the nonbonding combination σ*(pₓ − pᵧ) which is related to the empty energy level has been found in this energy region. Some contribution to the bond between PS₃ structure pyramids of P₂S₆ cluster also came from hybridization of p orbitals of sulfur atoms that belong to different pyramids.

By σ hybridization of phosphorus and sulfur s orbitals, the P — S bonds are created. Also the hybridization of p orbitals of phosphorus and sulfur atoms is observed (Fig. 2). Such hybridization has obviously both σ and π character. Thus the next scheme for appearing of molecular orbitals that form P — S bonds in PS₃ structural pyramid could be proposed. The phosphorus atom realizes sp² hybridization from which three symmetrically oriented bonds with involving of sulfur p orbitals appear. Two electrons from surrounding cations in a crystal lattice and one s electron of phosphorus atom (excited on d orbital) supply filling of covalent P — S bonds.

In Sn₃P₂S₆ crystal, all energy levels for P₂S₆ clusters are occupied, and in the ionic bonding approach, the charge states S²⁻ for sulfur ions and P⁴⁺ for phosphorus ions are expected. However, as it follows from calculations for P₂S₆ molecular orbitals, enough high charge density is found at phosphorus atoms. By this matter, not high positive charge (drastically smaller than +4) is expected for the phosphorus ions. In addition, the cal-

![FIG. 2. The partial densities of states and space distribution of electron density for the molecular orbitals of P₂S₆ cluster in free state.](image-url)
culcations provide evidence about high electronic density at the middle of P — P bond.

III. ELECTRONIC STRUCTURE OF Sn₂P₂S₆ CRYSTAL

At building of Sn₂P₂S₆ crystal structure with two formula units in the elementary cell (Fig. 1), the quantity of energy levels of P₂S₆ clusters is doubled with their energies splitting in results of the inter-cluster interaction. Also, the energy levels of 5s² orbitals of four Sn²⁺ cations are added to the structure of the crystal valence band (VB). The cation’s 5p² orbitals participate in formation of the conduction band of crystal. The scheme of electron orbital hybridization in the crystal could be presented as additive combination of above described scheme of P₂S₆ cluster orbitals creation with scheme of these molecular orbitals hybridization with tin atomic orbitals. Generally, for the crystal, four tin atoms, four phosphorus atoms and 12 atoms of sulfur in the elementary cell have 108 valence electrons that are placed at 54 energy levels in the VB.

It’s known⁴⁵ that energy positions of the atomic orbitals of phosphorus, sulfur and tin are next: P 3p = −8.35 eV, P 3s = −17.13 eV; S 3p = −10.28 eV, S 3s = −20.8 eV; Sn 5p = −4 eV, Sn 5s = −11 eV. The scheme of hybridization (Fig. 3) could be proposed which is in agreement with calculated energy spectra for free P₂S₆ cluster (Fig. 2) and for Sn₂P₂S₆ crystal (Fig. 4). At the crystal structure formation, the energy of phosphorus valence orbitals almost doesn’t change while the bonding energy for sulfur valence orbitals strongly lowers (almost by 7 eV). This is in agreement with raised electronic density on the sulfur anions and with enough high electronic density surrounded the phosphorus atoms.

In accordance to the calculated energy spectra and densities of states of Sn₂P₂S₆ crystal (Fig. 4), its VB could be divided into eight subbands which are labeled at the hybridization scheme of atomic and molecular electronic orbitals. Remember, that for the energy spectrum of free P₂S₆ cluster only seven subbands were found (Fig. 2). For Sn₂P₂S₆ crystal, the additional levels of tin atomic s orbitals are placed near −8 eV and they are related to the sixth subband of the VB.

The subband I contains the two energy levels near −16.5 eV for which a contribution of phosphorus s orbitals is dominated (Fig. 4). Here, a small contribution of sulfur s orbitals is also presented, and a minor appearance of sulfur p orbitals is observed. The contribution of tin atoms valence orbitals in this subband is specific peculiarity of the crystal energy spectrum. Generally, the bonding orbitals of covalent P — P and P — S bonds are created in subband I (Fig. 4).

The subband II includes the two energy levels near −15.5 eV. It is formed by antibonding combination of two phosphorus atoms s orbitals and by bonding hybridiza-

FIG. 3. The hybridization scheme for electronic orbitals in Sn₂P₂S₆ crystal

FIG. 4. The electron energy spectrum of Sn₂P₂S₆ crystal in the paraelectric (top) and the ferroelectric (bottom) phases. The calculated total density of states is compared with experimental XPS data.⁴⁶
P – P covalent bonds in the crystal structure. For these levels also, a some contribution from phosphorus d orbitals is observed.

The subband III in region from $-13$ eV to $-13.8$ eV has eight energy levels for which electron charge density is mostly localized at sulfur atoms (Fig. 4). Here the contribution of phosphorus 3p and 4d orbitals is also presented. This subband is characterized by bonding hybridization for P – S and antibonding hybridization for P – P covalent bonds in the crystal structure.

The subbands IV and V near $-10$ eV and $-8.5$ eV (both of them contain two energy levels) are formed by s and p orbitals of phosphorus and sulfur atoms. They are the replica of the subbands II and I and originated from their hybridization with subband III. For the IV-th subband, the charge is mostly localized at phosphorus atoms and has bonding character for the P – P bonds and antibonding character for the P – S bonds. The subband V has antibonding character for both P – P and P – S bonds.

The subband VI with four energy levels in range from $-8$ up to $-6.5$ eV appears in the VB of Sn$_2$P$_2$S$_6$ crystal as a result of hybridization of tin atomic electron orbitals with P$_2$S$_6$ clusters molecular orbitals. This subband is mainly formed by tin s orbitals and by p orbitals of phosphorus and sulfur (Fig. 4). Charge of this subband’s hybridized orbitals is located between phosphorus atoms and around tin atoms and has Sn – S and P – P bonding character (Fig. 4). The peculiarities of orbitals hybridization for every of four levels from this subband will be analyzed in details later with the aim of interatomic interactions explanation that are related to the ferroelectric phase transition in Sn$_2$P$_2$S$_6$ crystal.

The subband VII located in energy region between $-6.5$ and $-4.5$ eV and contains ten energy levels. They are formed by phosphorus and sulfur p orbitals and also include small contribution of tin electronic orbitals. This subband has bonding character for P – S and P – P bonds and it is antibonding for the Sn – S bonds.

The VIII-th subband is situated near the top of crystal’s valence band and includes 24 energy levels in the energy range from $-4.5$ till $-0.5$ eV. The considered subband is mostly formed by lone pairs of sulfur p orbitals, with some participation of phosphorus p orbitals, and it has P – P bonding and P – S antibonding character. Here the hybridization of s and p orbitals of tin atoms what determines their stereoactivity (Fig. 5) is also reproduced. The nature of Sn$^{2+}$ cations stereochemical activity in the Sn$_2$P$_2$S$_6$ crystal structure will be analyzed in details later on.

IV. HYBRIDIZATION OF TIN ATOMIC ORBITALS WITH MOLECULAR ORBITALS OF P$_2$S$_6$ CLUSTERS

As it was mentioned above, the valence subband VI includes four levels in the region from $-8$ to $-6.5$ eV which are related to s orbitals of tin atoms (Fig. 4). These are levels from 17 to 20, and their spatial electronic charge distribution illustrates a bonding peculiarities of tin atomic orbitals with sulfur and phosphorus orbitals which create the clusters P$_2$S$_6$. The level 17 is characterized by enough strong bonding of tin atoms with two nearest sulfur atoms, the levels 18 and 19 demonstrate strong bonding of tin atom with one of the nearest sulfur atom. The spatial charge distribution of the 20th level is of special interest – here the electron density is elongated from tin atom to the middle of P – P bond inside P$_2$S$_6$ cluster (Fig. 4).

It can be easily seen that in the paraelectric phase in addition to Sn – S – P – P – S – Sn – sequence of chemical bonds the Sn – P – P – Sn – series also exists. Such sequence of direct bonds of tin atoms with phosphorus atoms appears due to anisotropy of the spatial charge distribution for the level 20 (Fig. 4). Named distribution has a form of layers that are oriented close to the plane (1011). It is important that directions of the tin atoms shifts at transition into ferroelectric phase and are also rather close to orientation of the mentioned plane.

In the ferroelectric phase, the two pairs of nonequivalent tin atoms are presented in the crystal structure what is clearly illustrated by the spatial distribution of the electron charge density for the 20th level (Fig. 3a). Near mentioned P$_2$S$_6$ cluster, one of the tin atoms, further Sn$_2$, is approached to the middle of P – P bond. At this, the electron charge distribution which surrounded the Sn$_2$ atom connects with charge distribution between two phosphorus atoms. Another tin atom (Sn$_1$) goes away from the middle of P – P bond, and surrounded Sn$_1$ atom electron charge distribution is oriented to one of the sulfur atoms. Such difference in hybridization of the electron orbitals for two types of tin atoms in the ferroelectric phase determines disappearance of the layer-
FIG. 6. The spatial charge density distribution (in plane which contains $S \cdots P \cdots P \cdots S$ bonds) for orbitals in the valence subbands I – VIII for the paraelectric phase of Sn$_2$P$_2$S$_6$ crystal.

V. TRANSFORMATION OF ELECTRON ENERGY SPECTRA AT TRANSITION FROM PARAELECTRIC PHASE INTO FERROELECTRIC PHASE

According to experimental data$^{17–20}$, the chemical bonds and electron energy spectra of Sn$_2$P$_2$S$_6$ crystal have noticeable changes at the ferroelectric phase transition. The calculated energy spectra demonstrates the changes in the energy gap and in positions of all energy levels of the VB – energies of electron states density peaks in the VB shifts approximately by 0.5 eV (Fig. 4,5). In acentric phase the degeneration of electron energy levels disappears in some regions of the Brillouin zone, what determine higher smearing of the energy distribution of electron density of states. It is important to mention that the lowering of the electron density of states near the top of the VB is also observed at transition into ferroelectric phase together with a rise of the energy gap.

The changes of electron energy spectra obviously reflect an important role of electron-phonon interaction in nature of the Sn$_2$P$_2$S$_6$ crystal spontaneous polarization. Such interaction is illustrated by a transformation of the spatial electron density distribution at change of the atoms coordinates in the crystal structure. The squares of wave functions for electron orbitals, which are summed for the energy levels of valence subband VI in the paraelectric and the ferroelectric phases of Sn$_2$P$_2$S$_6$ crystal, are shown at Fig. 9. In the elementary cell of centrosymmetric structure, a similar distribution of electron den-
sity around four tin atoms is observed. This distribution reflects the stereactivity of the lone electron pair $5s^2$ of cations $Sn^{2+}$. Also, the spatial charge distribution around phosphorus atoms is similar. This is in agreement with the presence of inversion center at the middle of $P — P$ bond. In the acentric structure, the two pairs of tin atoms, with different distribution of surrounding charge, appear. The nonequivalence of the electron density distribution near the phosphorus atoms is also seen. It should be mentioned that elevated electron density is located in the vicinity of neighboring tin atoms $Sn_2$ and phosphorus atoms $P_1$. Also, lowering of the surrounding charge is observed for the adjacent atoms $Sn_1$ and $P_2$. Acentricity of the $P_2S_6$ clusters is also reflected in deformation of the charge distribution along the $P — P$ bonds.

Let’s analyze peculiarities of the orbitals hybridization for study of the tin $5s^2$ electrons stereactivity and their contribution into lattice spontaneous polarization. In the crystal structure the tin atoms are placed in the polyhedrons that are created by eight sulfur atoms. At beginning we will consider the mixing of tin orbitals with valence orbitals of surrounding sulfur atoms. Further, it will be analyzed a role of hybridization between tin atomic orbitals and $P^2$ orbitals of cations. The electron pair in $5s^2$ configuration is definitely not spherical – almost rigid dipoles exist which are not aligned. At cooling the stereactivity of electron lone pair growths and the orientation of cation shifts in nearest elementary cells is correlated by dipole-dipole interaction. The spontaneous polarization appears with two contributions – "displaceive" and "order/disorder".

The experimental data of XPS spectroscopy about electron energy spectra near the top of VB in the paraelectric and the ferroelectric phases of $Sn_2P_2S_6$ crystal confirm the lowering of the electron energy states density at transition into acentric structure. The calculations of energy spectra show big enough contribution of tin $s$ orbitals into electron states density near the top of the VB and permit a possibility for tracing of their change at the phase transition (Fig 3). The calculated space distribution of total electron density illustrates an evolution of electron lone pair at transition from the paraelectric phase into the ferroelectric one (Fig. 10). The appearance of the tin cations nonequivalence is observed: two of them shifts in a direction of those sulfur atoms with which in the paraelectric phase they have the biggest overlap of electron orbitals; alternative two cations "flip" the biggest overlap of their electron orbitals in direction of other sulfur atoms (Fig 10).

The high coordination of tin cations could be obviously described by taking into account their $d$ orbitals. For symmetry requirements the hybridization of $sp^3d^5$ type satisfies and they could be related to the positioning of the tin cations inside of the eight caped polyhedron of sulfur anions. Indeed, the orbitals $s$, $p_x$, $p_z$, $d_{xz}$, $d_{xy}$, $d_{yz}$, $d_{zx}$, $d_{x^2-y^2}$ transform on the irreducible presentations $A_g$, $B_u$, $A_u$, $B_g$, $B_g$, $A_g$, $A_g$ (orientation of the monoclinic symmetry plane coincides with cartesian plane $XZ$) that have fully symmetrical combination.

In the elementary cell of the paraelectric phase, the four such dodecahedrons are equivalent – they pair-by-pair are related by second order screw axis or by glade mirror plane (Fig. 11). In the ferroelectric phase as a result of earlier discussed charge density redistribution, the inversion center and symmetry axis disappear. The two pairs of nonequivalent dodecahedrons appear. In one type of the dodecahedrons at the phase transition, the electron density switches between two nearest sulphur
atoms which corresponds to the strongest Sn — S bond. In the other type of dodecahedrons cooling from the centrosymmetric phase to acentric one, the "flipping" of the electron density between almost oppositely oriented Sn — S bonds is observed (Fig. 11).

The built by sulfur atoms pyramids could be divided in the coordination polyhedrons (Fig. 11). Localized at hybridized $sp^2$-like orbitals of tin atoms, the electron density is oriented in direction opposite to the base of named pyramids. This is a direction to the three sulfur atoms with weaker Sn — S bonds.

A characterization of the chemical bonds changes at phase transition could be found by compare of calculated data ( Mulliken charges and overlap parameters for the electron orbitals of neighbor atoms) with experimental structure data, with Mössbauer, XPS and NMR spectroscopies data. For the paraelectric phase of Sn$_2$P$_2$S$_6$ crystal, the following electron configurations were calculated: for four equivalence tin atoms Sn $- 5s^{1.865}p^{1.153}d^{0.022}$; for four equivalence phosphorus atoms P $- 3s^{1.187}3p^{2.508}3d^{1.172}$; for one of the three types of sulfur atoms S $- 3s^{1.833}3p^{4.231}3d^{0.179}$. It could be seen that $d$ orbitals are populated, what agrees with the explanation of high dodecahedral coordination of tin cations on background of $sp^2d^5$ type hybridization. The highest population of $d$ orbitals is appeared for phosphorus atoms. As was mentioned earlier, for the P$_2$S$_6$ clusters the bonds in PS$_3$ structure pyramids are determined by $sp^2$ hybridization of phosphorus atomic orbitals with their further $\sigma$ hybridization with sulfur $p$ orbitals. Thereafter one of two $3s^2$ electrons of phosphorus is involved into $sp^2$ hybridization of atomic orbitals, other electron is excited on the atomic $d$ orbital. The named $sp^2$ hybridization involves two $p$ electrons of phosphorus, third $p$ electron by $\sigma(p-p)$ hybridization creates covalence P — P bond.

In the ferroelectric phase, the electron configurations for two types of tin atoms are the following: Sn$_1$ $- 5s^{1.851}p^{1.163}d^{0.224}$; Sn$_2$ $- 5s^{1.850}p^{1.177}d^{0.226}$. It is seen that at transition into the ferroelectric phase, the quantity of $s$ electrons lowers by $\Delta n_s = 0.014$. Such calculations data are in agreement with observed lowering of isomer shift for spectral line of $^{119}$Sn Mössbauer effect at cooling from the paraelectric phase into the ferroelectric one.

The diminishing of $s$ electron quantity in the ferroelectric phase for two types of tin atoms is compensated in a different step (level) by growing of $p$ orbitals occupancy. Hence the total charge of tin atoms changes from 3.240e in the paraelectric phase to 3.239e (Sn$_1$) and 3.251e (Sn$_2$) in the ferroelectric phase. The calculated rise of electron density in the vicinity of Sn$_2$ atoms is in agreement with decreasing of resonance frequency in NMR spectrum for $^{119}$Sn from $-781.3$ ppm in the paraelectric phase to $-782.5$ ppm in the ferroelectric phase.

The increase of the resonance frequency till $-754.5$ ppm for Sn$_1$ is obviously determined by a occupancy redistribution from $s$ orbitals to $p$ orbitals at some lowering of total charge.

Determined by positions of XPS spectral lines, the energy of chemical binding for the $4d$ core orbitals in the ferroelectric phase differently increases for the two types of tin cations. This is obviously defined by growing of $5s^2$ electrons lone pair stereoactivity and by these electrons redistribution on more distanced from tin cores $p$ orbitals.

The changes in Sn$_2$P$_2$S$_6$ crystal structure at the spontaneous polarization appearance are characterized by the calculated values for the overlap parameters of electron orbitals of tin atoms and the nearest sulfur atoms. In the paraelectric phase such parameter with the biggest value about 0.054 has been found for the shortest bonds Sn — S in the dodecahedrons of sulfur atoms (Fig. 11). It must be mentioned that in the nearest dodecahedrons, which are related by a second order screw symmetry axis, such Sn — S bonds have opposite orientations of their projections onto plane (010), which contains the

![FIG. 9. The electron density space distribution for the VI valence subband in the paraelectric (a) and the ferroelectric (b) phases of Sn$_2$P$_2$S$_6$ crystal.](image-url)
spontaneous polarization. In the ferroelectric phase the nonequivalence of two pairs of cations Sn$_1$ and Sn$_2$ and surrounded them dodecahedrons of sulfur ions appear. Here the biggest overlap parameter for atomic orbitals (0.096) was found for the Sn$_2$ — S bond. This bond was strong already in the paraelectric phase and it is in addition enforced at spontaneous polarization appearance – the atom Sn$_2$ in the ferroelectric phase is shifted in the direction of sulfur atoms with the biggest content of chemical bonds covalency. For other coordination dodecahedron the strongest chemical bond Sn$_1$ — S has the overlap parameter 0.087. Here at transition from the paraelectric phase into the ferroelectric one, the "flipping" of maxima in the space distribution of electron density occurs between bonds of central metal atom and ligand atoms in the coordination polyhedron. As a result of such "flipping" in both types of the dodecahedron, the strongest bonds Sn$_1$ — S and Sn$_2$ — S have nearly oriented projections onto plane (010) (Fig. 11).

The changes in the electron density space distribution correlate with variations of the interatomic distances. For example, in the paraelectric phase the Sn — S bonds with the biggest overlap parameter (0.054) are strongest and they have the smallest length in the ferroelectric phase. For the dodecahedrons with tin atoms of Sn$_2$ type, the overlap parameter for strongest bonds increases to 0.096 and their length decreases by 0.21 Å. Hence the occupancy of XZ plane oriented $p$ orbitals raises – from the calculations follows: $\Delta p_x = 0.048$, $\Delta p_y = -0.002$, $\Delta p_z = 0.022$. For the dodecahedrons with tin atoms of Sn$_1$ type at the electron charge density "flipping" on almost oppositely oriented Sn — S bond, the overlap parameter, for the strongest bond in the paraelectric phase, lowers from 0.054 till 0.022 and length of this bond grows by 0.26 Å. The strongest Sn$_1$ — S bond in the ferroelectric phase is characterized by overlap parameter 0.087 and its length decreases by 0.35 Å.
Here the changes for occupancies of tin \( p \) orbitals are: \( \Delta p_x = 0.025, \Delta p_y = 0.036, \Delta p_z = 0.010 \). We could see that cations of Sn\(_2\) type donate the biggest electronic contribution into the spontaneous polarization and this contribution has "displacive" character. For the cations of Sn\(_1\) type, the electronic contribution is a little smaller and this one has "ordering" character.

The calculated changes of Mulliken charges and atomic orbitals overlap parameters coincide with the temperature dependence of the resonance frequencies in NMR spectra for \( ^{31}\text{P} \) phosphorus\(^{18,19}\). In the paraelectric phase all phosphorus atoms are equivalent (calculated charge is 4.869e) and here only one NMR line with frequency 92.12 ppm is observed. At the spontaneous polarization appearance, the inversion center vanishes what is associated with growth of phosphorus atoms nonequivalence in \( \text{P}_2\text{S}_6 \) clusters. For two types of phosphorus atoms the following electron configurations were calculated: \( \text{P}_1 = 3s^13p^23d^1.193; \text{P}_2 = 3s^13p^23d^1.183 \). For atoms of \( \text{P}_1 \) type with grown calculated charge (3.251e) in result of shielding effect, the NMR specter resonance frequency decreases to 89.2 ppm. This is in agreement with closeness of \( \text{P}_1 \) type atoms and Sn\(_2\) cations for which also a lowering of the NMR resonance frequency is observed\(^{18,19}\) in the result of electron density growing in their vicinity at cooling into the ferroelectric phase. For atoms of \( \text{P}_2 \) type in the ferroelectric phase, the resonance frequency rises till 93.7 ppm. In this case the weakening of the charge shielding effect is obviously determined by diminishing of the \( s \) orbitals occupancy. By the way, for neighbor Sn\(_1\) cation the quantity of \( s \) electrons also decreases, what induces growth of related resonance frequency in \(^{119}\text{Sn} \) NMR spectrum\(^{19}\).

### VII. DISCUSSION OF RESULTS

The calculated electron energy spectra, densities of electron states and their variation at transition from the paraelectric phase into the ferroelectric one coincide with the available structure data and results of experimental investigation of the chemical bonds nature. On this ground the next generalized description of sources of the spontaneous polarization appearance in Sn\(_2\)\( \text{P}_2\text{S}_6 \) crystal could be proposed.

The ferroelectric distortion is proportional to the difference between the Sn\(_1\) and Sn\(_2\) positions. For this distortion the short range (mostly Sn — S) repulsions must be sufficiently small in order to allow to shift the equilibrium Sn positions from the center of the chalcogen dodecalhedron. The effective charge of phosphorus has to be sufficiently small in order to allow the shift of the Sn cations in the direction of P — P bond. Such requirement could be satisfied by the following way. At fully symmetrical \( A_g \) lattice vibration, the important changes of charge gradient in the elementary cell occur — the charge partially is waged from anions \( \text{P}_2\text{S}_6 \) onto cations Sn. Hence the \( p \) orbitals of tin cations have to be occupied — the stereoreactivity of valence electrons of these cations is realized by their partial hybridization with \( p \) orbitals of neighbor sulfur atoms. Such hybridization lowers the short range repulsion between tin cations and the nearest sulfur atoms what govern their approaching. The charge transferring between tin and sulfur atoms manages some lowering of electrostatic interactions energy.

Generally, in the ground state (at 0 K) the metastable center-symmetric Sn\(_2\)\( \text{P}_2\text{S}_6 \) structure is possible, for which both opposite tin atoms are a little approached to the middle of the P — P bond. However, at low temperatures the acentric structure is energetically more favorable. At approaching of two tin cations, that are related by inversion center at middle of P — P bond, their Coulomb repulsion increases (such repulsion between cations of tin and phosphorus obviously don’t play important role because positive effective charge of phosphorus is not big). Hence energetically more advantageous could be approach of one tin cation to the center of \( \text{P}_2\text{S}_6 \) cluster at the repulsion of opposite tin cation. Indeed, in the center-symmetric structure both tin atoms are placed on distance 3.633 Å far from middle of P — P bond, in acentric structure such distances equal to 3.463 Å for Sn\(_2\) atom and 3.857 Å for Sn\(_1\) atom. The space between considered tin atoms increases from 7.266 Å in the paraelectric phase till 7.310 Å in the ferroelectric phase\(^{18,19}\).

Removing of Sn\(_1\) atom away from \( \text{P}_2\text{S}_6 \) cluster decreases hybridization of his valence electron orbitals with molecular orbitals of cluster. By this matter the electron charge in cluster moves onto \( \text{PS}_3 \) structural pyramid with \( \text{P}_1 \) atom at their top that is the nearest to Sn\(_2\) atom. At repulsion of Sn\(_1\) atom from the cluster, an important change of the hybridization character occurs, which is accompanied by localization of valence electrons near the tin ion core and by growth of their kinetic energy. Such processes have obviously activation character and they determine presence of the energy barrier between central and side minima in the three-well potential.

By such way, the induced by the fully symmetrical \( A_g \) vibration, the important changes of charge gradient in the elementary cell determine variation of the electron configuration for the ions of crystal lattice. The reconstruction of electron configuration modifies the balance of interatomic interactions what induce instability of \( B_u \) polar lattice vibration. Exactly by such manner, the mechanism of lattice modes \( A_gB_u^2 \) nonlinear interaction could be presented, and this one governs the tree-well potential presence for fluctuations of the order parameter of the ferroelectric phase transition in Sn\(_2\)\( \text{P}_2\text{S}_6 \) crystal.

The value of energy barrier in the three-well potential\(^{18,19}\) equals near 0.015 eV, and the energy difference between central and side minima is near 0.01 eV. Such energetic characteristics are in agreement with our calculation of the electron energy spectra of Sn\(_2\)\( \text{P}_2\text{S}_6 \) crystal in the paraelectric and the ferroelectric phases. Thus, for the ferroelectric phase the full energy was found by 0.0078 eV smaller in compare with calculated full energy.
for the paraelectric phase. It must be mentioned that at the crystal symmetry lowering the negative contribution of coulomb interactions into full energy decreases.

The appearance of spontaneous polarization in ferroelectric crystal is determined by variation of the chemical bonds covalency and by delicate balance between short range repulsion forces, which determine the relief of local potential for the phase transition order parameter, and long range displace forces that define energy of intercell interaction.

The hybridization of tin and sulfur atomic orbitals which defines the appearance of "partially rigid" electric dipoles (pseudo-spins) in result of tin cations valence electrons stereoactivity was early analyzed. Such hybridization could be described as $sp^2$ or $sp^2d^5$ combination of tin and sulfur atomic orbitals. This fact is clearly demonstrated by presence of enough high density of Sn $5s$ states near the top of VB. In addition to that, it have been found neediness of accounting for hybridization between tin atomic orbitals and $P_2S_6$ molecular orbitals what has argument by presence of $5s$ and $5p$ states of tin even at the bottom of VB -- in the energy range near $-17.5$ eV where the $s$ states of phosphorus atoms are dominated.

The obtained pictures of the electron density space distribution show (Fig. 7) the presence of $-Sn-S-P-P-S-Sn-$ short-range bonds chains in the paraelectric phase. The occurrence of the short-range interactions in chains of $-Sn-P-P-Sn-$ type also have been demonstrated. Such sequences of the short-range interactions together with long-distant coulomb interactions determine the mean-field, which induce correlation of the pseudo-spins at lowering of disordering influence of heat energy. At temperature down till 0 K, the chains of central symmetric structure groups $-Sn-P_2S_6-Sn-$, which are related to the pseudo-spins position in the central well of local potential, could obviously exist also. However, the correlated ordering of the structure motives ("dimers") like $-Sn-P_2S_6-$, which responses to the pseudo-spins standing in one of the side well of local potential, is energetically more favorable.

It is important to remark about rise of anisotropy of tin and phosphorus atoms surround in the ferroelectric phase (Fig. 8) what support increasing of their dynamic or Born effective charges. The dynamical transfers of charge are expected to be larger when such a hybridization involves $d$ states, for which the interactions parameters with other orbitals are particularly sensitive to the interatomic distances. Also, the amplitude of Born effective charges is not monitored by a particular interatomic distance but is dependent on the anisotropy of the Sn environment along the $-Sn-S-P-P-S-Sn-$ bonds. In the paraelectric phase, the $3p$ electrons are obviously widely delocalized and dynamical transfer of charge can propagate along the $-Sn-S-P-P-S-Sn-$ chains. In the ferroelectric phase, these chains behave as a sequence of "dimers" $-Sn-S-P-P-S-$ for which the electrons are less polarizable.

The anomalously large dynamical charges produce big LO-TO splitting for the ferroelectric soft phonon mode. Moreover, this feature is associated with the existence of an anomalously large destabilizing dipole-dipole interaction, sufficient to compensate the stabilizing short range forces and induce the ferroelectric instability. In materials where polar soft modes play a major role, the dynamical charge relate the electronic and structural properties. However, for the $Sn_2P_2S_6$ crystal the big LO-TO splitting for polar modes wasn’t observed in phonon spectra. Here, at $4.2$ K such splitting is in the range of $2 \pm 7$ cm$^{-1}$. At heating to the temperature of phase transition in the ferroelectric phase, the LO-TO splitting for the lowest energy optic mode of $B_u$ symmetry (soft mode) reaches only the value of $10$ cm$^{-1}$.

The low frequency dielectric susceptibility temperature anomaly in $Sn_2P_2S_6$ crystal don’t described only by dielectric contribution of the polar lattice vibrations. On the data of dielectric spectroscopy in the paraelectric phase, the dielectric contribution from polar lattice vibrations into static dielectric susceptibility reaches only near ten percent. Obviously in the range of phonon frequencies, a significant destabilizing dipole-dipole interaction doesn’t appear. The essential contribution into dielectric anomaly appears at frequency lowering into submillimeter diapason – here the relaxational dispersion have been observed, which is obviously determined by nonlinear dynamic excitations.

The above attention was accentuated on the mixing between states of tin valence electrons and orbitals of phosphorus and sulfur atoms across all energy range of $Sn_2P_2S_6$ crystal VB. In addition, the enough large density of phosphorus $s$ and $p$ states is presented near the top of VB also. Such phosphorus orbitals also create the conductivity band. The defined facts give evidence about enough strong mixing of diffusive $s$ orbitals in structure of $Sn_2P_2S_6$ crystal. Obviously, the effective occupation of phosphorus $d$ orbitals, which commonly are enough localized, also give evidence of their important role in the mechanism of electron-phonon interaction for this ferroelectrics.

For the $P_2S_6$ structure cluster, with symmetry $D_{3h}$ in free state, the $3s$ electron orbitals of phosphorus atoms, which are placed at bottom of the VB (near $-15$ eV) and their bonding $\sigma (p_2 - p_2)$ orbitals, which create $P-P$ bonds and have energy level near the top of VB (in range $-1.5 \pm 0$ eV), satisfy the transformation according the $A_g$ irreducible presentation. In addition to these orbitals of free anion cluster, the orbitals with energies near $-8.5$ and $-4$ eV are also involved at formation of $P-P$ bond (Fig. 9).

In the $Sn_2P_2S_6$ structure the symmetry of $P_2S_6$ anion clusters is lowered, however possibility of effective mixing for the wave functions of $s$ and $p$ orbitals of phosphorus atoms, which have identical symmetry, is obvious. To mentioned series of four sets of $P-P$ bonding orbitals, which create high electron density at middle of this bond (Fig. 10), the combination of tin atomic orbital related to the energy level 20 in VI subband (in range from $-8$ till...
—6.5 eV) is added. Obviously, in result of such hybridization the similar changes at the top and bottom of the VB are observed at appearance of spontaneous polarization in the crystal (Fig. 5).

Generally, the high effective charge and large polarizability of $P_2S_6$ anionic clusters together with stereoelectricity of tin electron lone pair determine large electronic contribution from all atoms into spontaneous polarization of $Sn_2P_2S_6$ crystal. Such situation is in agreement with earlier founded involving of all 13 $B_n$ modes and 15 $A_g$ modes to dynamic instability of investigated ferroelectrics. However, obviously important role also belongs to the nonlinear interaction with participation of $A_n$ and $B_n$ nonsymmetric modes. For the point group $2/m$ in addition to the $A^g_1 B^g_1$ fully symmetric combination, the invariants of $A_n B_n B_n$ type are also present. A significant role of such nonlinear mixing of lattice vibrations is reflected in strong internal deformations of $P_2S_6$ clusters at the phase transition. In the crystal electron structure, such invariants obviously replicate the hybridization of molecular orbitals of $P_2S_6$ clusters with participation of atomic $d$ orbitals. Such hybridization is clearly illustrated by the spatial distribution of electron density for the energy level number 20, which is aligned from tin atoms to middle of $P — P$ bond (Fig. 7).

The energy decreasing for the $s$ orbitals of phosphorus and sulfur at the VB bottom (their contribution is dominated in the lowest subbands – from I to V), and also lowering of energy for tin $s$ orbitals, with their contribution across whole range of the VB (Fig. 5), are obviously essential for energetic motivation of the transition into the ferroelectric phase. Also some lowering of the $p$ and $d$ orbitals energy occurs.

The electronic structure XPS measurements for $Sn_2P_2S_6$ crystal[20,21] revealed the chemical shifts of Sn and P electronic core states to a higher binding energy and of S states to a lower binding energy at the crystal lattice formation. This shift suggests a charge transfer from Sn and P to S atoms. The binding energies and chemical shifts strongly change at the phase transition. In the ferroelectric phase, the chemical shifts of Sn and P atoms are higher while for S atoms they are smaller. So, for all atoms of crystal structure at transition into the ferroelectric phase the binding energy for core orbitals increases. These data give evidence about localization of electron charge in space between atoms, or about enhancement of chemical bonds covalency. Such variation of the core orbitals energy agrees with experimentally observed and founded at calculations (Fig. 4) transformation of the VB structure and they support an energetic stability of the ferroelectric phase.

Since covalency increases, there might be a possibility for the drastic collapse of the sulfur ionic size (which is related to charge transfer). For the $S^{2-}$ ion the ionic radius is 1.84 Å, the covalent radius $\approx 1.02$ Å. If the size of sulfurs were small compared to the allowed space then the sulfur atoms would be weakly bounded in the lattice. In this case an imbalance between the decreased (due to small S radius) repulsive forces and the polarization forces, what tend to displace the ion from its position, also support structure rearrangement.

In whole, for $Sn_2P_2S_6$ crystal at transition from the paraelectric phase to the ferroelectric one complex evolution of electron and phonon spectra occurs, which could be presented as sequence of five steps. Evidently as first factor assists change of the electron density charge distribution in the elementary cell by fully symmetric breathing modes $A_g$. Such redistribution of the electron density prompts the stereochemical activity of tin cations electron lone pair and produces the covalence bonds of tin atoms with sulfur atoms (hybridization of $sp^2d^5$ type), and also with phosphorus atoms, what could be considered as second part.

As third stage could be considered the weakening of the short-range repulsion between cations of tin and phosphorus, in result of their charges lowering, and at significant coulomb repulsion of the nearest (related by the inversion center) tin cations. Mentioned the second and the third factors represent the nonlinear interaction of $A_g B^g_2$ type, they govern an anisotropy of polar shifting of atoms in the elementary cell and define appearance of the dipole structure motives ($\text{Sn} — P_2S_6$), which are related to the polar normal coordinates of $B_n$ symmetry. The fourth important factor is the dipole-dipole interaction which correlates orientation of local dipoles (pseudospins) and defines appearance of the spontaneous polarization in the crystal structure. And, at finish, as fifth circumstance must be accounted that all low symmetry modes participate in result of permitted nonlinear $A_n B_n B_n$ relation in the structure transformation. Such combination of structure deformations mirrors participation of phosphorus and sulfur $d$ orbitals in the covalent bonds of $Sn_2P_2S_6$ crystal.

It is interesting to compare the peculiarities of chemical bonds in $Sn_2P_2S_6$ sulfide and $Sn_2P_2Se_6$ selenide compounds, and also in the lead contained $Pb_2P_2S_6$ crystal. At first, we will consider the binary compounds $MX$, where $M$ – metal Ge, Sn, Pb, X – chalcogen O, S, Se, Te. The stereoactivity of electron lone pair for metal atoms is determined by $sp^2$ hybridization of their $s$ and $p$ orbitals with $p$ orbitals of chalcogen atoms. Such hybridization is determined by the positions of the energy levels of electron states and by width of the related energy bands in the crystal structure[47,54]. The smallest energy difference is present between positions of energy levels of germanium $s$ orbitals and oxygen $p$ orbitals. Consequently for the compound GeO, the largest stereoactivity of $4s^2$ electron orbitals of Ge is observed[47]. At transition from Ge to Sn and than to Pb, the energy of chemical bonding for the $s$ orbitals increases. Thus with transition from O to S, and further to Se and Te, the energy of chemical binding for their $p$ orbitals decreases. It is expected that the hybridization of Ge $4s$ orbitals and O $2p$ orbitals is the strongest, and hybridization for Pb $6s$ orbitals and Te $5p$ orbitals is the most weak. However the hybridization is also influenced by width of related energy bands in the
crystal structure. The increase of width for the s and p electron states energy bands and their overlap could partially compensate increase of the energy distance between related energy levels, what produce some level of the stereoactivity and covalency of M — X bonds.

Thus, at transition from Sn$_2$P$_2$S$_6$ to Pb$_2$P$_2$S$_6$ the binding energy for Pb 6s level increases, what make weaker the stereoactivity of the 6s$^2$ electron lone pair in the decahedron of sulfur atoms. Obviously by this matter, the observed ionicity for the Pb — S bonds is higher in compare with Sn — S bonds ionicity. The melting temperature and the energy gap both rise at substitution Sn by Pb. The paraelectric phase in Pb$_2$P$_2$S$_6$ is stable at cooling till 4.2 K.

The largest stereoactivity of Ge 4s$^2$ electron lone pair in surround of sulfur atoms gives natural explanation of absence of Ge$_2$P$_2$S$_6$ crystal structure. The Ge atoms couldn’t be placed in positions with high coordination of sulfur atoms. Obviously introducing into Sn$_2$P$_2$S$_6$ crystal impurity of germanium in the charge state Ge$^{2+}$ strongly elevates the temperature of ferroelectric phase transition what was observed by dielectric investigations. Certainly the impurity in charge state Ge$^{4+}$ will not be stereoactive and will not support rise of temperature interval for the ferroelectric phase existence.

According to just described tendencies, at transition from Sn$_2$P$_2$S$_6$ to Sn$_2$P$_2$Se$_6$ the stereoactivity of Sn 5s$^2$ electron lone pair in the decahedron of selenium will be smaller than in the case of sulfide compound. However, for the ternary compounds the ion-covalence bonds Sn — S and Sn — Se are modified depending on peculiarities of P — S and P — Se bonds in P$_2$S(Se)$_6$ anion clusters. On the Mössbauer spectroscopy data at substitution of sulfur by selenium, the isomer shift for $^{119}$Sn nucleus decreases what directly show on higher covalency of Sn — Se bonds. The NMR spectroscopy for $^{119}$Sn$^{19,55}$ shows increase of the resonance frequency from $−78.1$ ppm in Sn$_2$P$_2$S$_6$ to $−608$ ppm in Sn$_2$P$_2$Se$_6$. The NMR spectral line for $^{31}$P decreases its frequency from $92.12$ ppm in Sn$_2$P$_2$S$_6$ to $28.7$ ppm in Sn$_2$P$_2$Se$_6$. These data provide evidence about lowering of the electron density in vicinity of tin nucleus and about their rise at phosphorus nucleus at transition from sulfide to selenide compound. The mentioned tendency could be explained by smaller electronegativity of selenium. Obviously the bonds P — Se are less polar what improve higher electron charge of phosphorus atoms in anionic clusters. Such situation probably also support more effective hybridization between cluster molecular orbitals and tin atomic orbitals (mostly the bonding hybridization of tin orbitals with molecular orbitals that are localized in middle of P — P bond) and increases the stereoactivity of tin lone pair of electrons in Sn$_2$P$_2$Se$_6$ crystal.

The growth of covalency and weakening of electrostatic interactions determine lowering of the melting temperature for Sn$_2$P$_2$Se$_6$ crystal in compare with sulfide analog, define decrease of the energy gap and govern smaller temperature of the ferroelectric phase transition$^2$.

VIII. CONCLUSIONS

The appearance of the spontaneous polarization in Sn$_2$P$_2$S$_6$ compound is accompanied by the significant changes of electronic structure that are observed in all subbands of this crystal VB. At the transition from the paraelectric phase to the ferroelectric one, the significant changes also occur for the phonon spectra in the whole frequency range – for both external and internal vibrations of the crystal lattice. The complicate evolution of the energy spectra could be represented by following contributions. The fully symmetrical (‘breathing’) $A_g$ modes change the space distribution of the electron density in the elementary cell. This one initiates the stereochemical activity of the tin cations electron lone pair and support creation of their covalence chemical bonds with sulfur atoms (the hybridization of sp$^2d^P$ type), and with phosphorus atoms also. Thus in result of the ionic charges lowering, the short-range repulsion between tin and phosphorus cations decreases, however the coulomb repulsion between tin cations still remains strong enough. These factors reflect the nonlinear interaction of $A_gB_u^2$ type, they determine the anisotropy of polar deformations in the elementary cell and induce appearance of the dipole structure motives (— Sn — (P$_2$S$_6$) —), which are related to the polar normal coordinates of $B_u$ symmetry. The dipole-dipole interaction correlates orientation of the local dipoles (pseudospins) and governs the spontaneous polarization of the crystal structure. In the structure rearrangement, all low symmetry modes take part – the possibility of the nonlinear linking of $A_uB_2B_u$ type is obvious. This interaction correlates with involving of tin and phosphorus d orbitals into creation of the covalent bonds in Sn$_2$P$_2$S$_6$ crystal lattice. Small difference of the paraelectric and ferroelectric phase’s energies and activation redistribution of the electron charge at the spontaneous polarization appearance determine the presence of the three-well local potential in the Sn$_2$P$_2$S$_6$ ferroelectrics.

ACKNOWLEDGMENTS

Part of presented in this paper results of calculations was conducted with support of computational cluster of Institute of Condensed Matter Physics, Lviv. Authors would like to thank staff of this facility (T. Bryk, T. Pat-sahan) for technical assistance. Also authors are grateful to Dr. R. Yevych for helpful discussions.
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