Correlation between crystal structures and polar (ferroelectric) properties of hybrids of haloantimonates(III) and halobismuthates(III)

R. Jakubas, M. Rok, K. Mencel, G. Bator and A. Piecha-Bisiorek

Halogenoantimonates(III) and halogenobismuthates(III) are a highly versatile class of organic–inorganic hybrid materials, applicable in optoelectronics and switchable dielectric devices. In this review, we discuss the rich chemistry of molecular–ionic halide complexes of Bi(III) and Sb(III) focusing on the correlations between their crystal structures and ferroelectric properties as well as on an explanation of the molecular mechanism of the paraelectric–ferroelectric phase transition. This review summarizes the current state of the art in the field of ferroelectricity among organic–inorganic hybrids based on Bi(III) and Sb(III) halides, which has become one of the key exploration areas of modern materials chemistry.

1. Introduction

Materials whose physical properties can be controlled by external stimuli have attracted considerable interest due to their potential applications in substrate-film interfaces, high-strain states, electrocaloric circuits, nanotubes and nanowires, ferroelectric random access memories (FeRAMs), dynamic random access memory (DRAM) capacitors, electron emitters, solar cell components, and weak-magnetic field sensors. Ferroelectrics are of importance in the design of thermo-sensitive multifunctional switching materials because they are often accompanied by multiple switchable physical properties, e.g. piezoelectricity, pyroelectricity, dielectric constant and second harmonic generation (SHG). Ferroelectrics are polar substances in the solid state (crystalline or polymeric) or liquid-crystalline state (liquid crystals), in which a spontaneous polarization ($P_s$) is generated and it is reversible in an external electric field ($E$). The relationship between an electric shift ($D$) or polarization ($P$) and the electric field intensity leads to a hysteresis loop ($D–E$ loop) between the opposite polarities. Such an electrical bistability can be used in non-volatile memory parts. It refers in particular to the ferroelectric computer memory type, FeRAM, which is crucial for...
Ferroelectricity is no exception. From this point of view, there are 230 space groups and 32 point groups, which describe both the macro- and microscopic symmetries of crystal structures. Among the 32 point groups, 11 belong to the centrosymmetric classes, while the remaining 21 are noncentrosymmetric. One should note, that the lack of spatial-inversion symmetry observed in the noncentrosymmetric group of crystals is most sought for in applications. It is important for the ferroelectric phase to adopt one of the 10 polar point groups, from which 68 polar space groups are formed (see Table 1).

The most important functionalities that make ferroelectrics so useful are the high values of the real part of complex dielectric permittivity, \( \varepsilon' \), spontaneous polarization, \( P_s \), good mechanical and thermal resistance and small dielectric losses.\(^{18,19}\) This set of traits determines the transition from the

### Table 1 68 ferroelectric space groups belonging to the 10 polar point groups\(^{17}\)

<table>
<thead>
<tr>
<th>Crystal system</th>
<th>Polar point group</th>
<th>Space group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Triclinic</td>
<td>1</td>
<td>( P1 )</td>
</tr>
<tr>
<td>Monoclinic</td>
<td>2 ( m )</td>
<td>( P2, P2_1, C2 )</td>
</tr>
<tr>
<td>Orthorhombic</td>
<td>( mm2 )</td>
<td>( Pmmm, Pnc2_1, Pcc2, Pma2, Pca2_1, Pn2_1, Pna2_1, Pnn2, Cmcm, Cmc2_1, )</td>
</tr>
<tr>
<td>Tetragonal</td>
<td>4 ( 4mm )</td>
<td>( P4, P4_1, P4_2, P4_3, I4, I4_1 )</td>
</tr>
<tr>
<td>Trigonal</td>
<td>3 ( 3m )</td>
<td>( P3, P3_1, P3_2, P3_3, R3, P6, P6_1, P6_2, P6_3, P6_4 )</td>
</tr>
<tr>
<td>Hexagonal</td>
<td>6 ( 6mm )</td>
<td>( P6mm, P6cc, P6_1, P6_2, P6_3, P6_4 )</td>
</tr>
</tbody>
</table>

Klaudia Mencel was born in 1992 in Ostrów Wielkopolski, Poland. She received her M.Sc. degree at the Faculty of Chemistry of the University of Wrocław in 2016. She is a Ph.D. student in the Ferroelectrics and Liquid Crystals Team, Faculty of Chemistry (University of Wrocław).

Grażyna Bator was born in 1958 in Pleszew, Poland. She completed her Ph.D. in 1986 at the Faculty of Chemistry of the University of Wrocław, Poland. After presenting her habilitation dissertation in 2000, she obtained the title of Professor in 2007. She took scientific internships at KU Leuven in Belgium and ZIBJ in Dubna, FR, and carried out experiments on neutron scattering in cooperation with FRM II, Garching, Germany. She is the co-author of about 160 articles in the field of materials chemistry. She is the head of the specialization ‘Materials Chemistry for Modern Technologies’ at the Faculty of Chemistry of the University of Wrocław.
paraelectric phase (which is usually high-temperature, high-symmetry state) to the ferroelectric one (low-temperature, low-symmetry state) in the close vicinity of the Curie temperature ($T_C$). The paraelectric–ferroelectric PT is related to the symmetry change (e.g. some symmetry elements of the paraelectric phase are lost, when the ferroelectric phase is reached), and leads to the structurally modified (ordered) ferroelectric phase. This structural transformation (described often as symmetry-breaking), is reflected in the macroscopic properties of the material, e.g. the appearance of $P_e$ in the ferroelectric phases.

One of the classes of molecular–ionic materials are organic–inorganic hybrid ferroelectrics which are currently of particular interest, due to the designable and modifiable characteristics of their organic and inorganic components, for exploring ferroelectric-based multifunctional materials such as multiferroics, semiconductors and photovoltaic materials.\(^{20-23}\)

Of note, the symmetry, size, and value of the dipole moment and finally, the reorientations of the organic cations play an important role in triggering the structural PT. Therefore, control of the ‘order–disorder’ motion of the organic counterions in a molecular crystal is a key issue for the design of hybrid molecular–ionic ferroelectrics.

Haloantimonates(III) and halobismuthates(III), which are the subject of the present review, defined by the general formula $R_nM_X X_{3-n}$ (where $R$ – organic cations, $M$ = Sb(III) or Bi(III) and $X = Cl, Br, I$) create one of the well-recognizable groups of ferroelectric compounds, essential mainly from an application point of view as they combine many desirable features, e.g. facile synthesis and processing, and cost-effectiveness with promising electrical and optical properties.\(^{24-29}\) A characteristic feature of the compounds under consideration is that they possess the lone electron pair (5s\(^2\) (Sb) and 6s\(^2\) (Bi)). However, its presence, which has a moderate impact on the structural properties of these salts, seems to have a tremendous effect on the physical properties of crystals. The lone electron pair deforms very easily a spherical symmetry of the orbital, on which they are located. This leads to an increase in the length of the $M-X$ bonds, and thus to a considerable deformation of the $MX_6$ octahedra.

The polyanionic structure of the lattice is determined by the size of the organic cation, its symmetry and ability to create hydrogen bonds of the $N-H \cdots X$ type; the size of the halogen atom, which jointly creates the lattice is also of crucial importance. The alkylammonium cations belong to a wide variety of organic cations, which crystallize with various anionic sub-layers. They are characterized by the different chain lengths, from methyl- to butyl-ammonium, and they may form five- or six-membered rings in the aromatic cations (e.g. imidazolium, pyridinium and its substituted derivatives) as well as saturated heterocyclic amines (e.g. pyrrolidine) (Fig. 1).

An analysis of these compounds has revealed a richness of the cationic unit movements. The motions depend on three basic factors such as: (i) the symmetry of the cation; (ii) the value of the dipole moments of the cation; (iii) the structure of the anionic network surrounding the cation. The reorientation ability of the dipolar units occupying vacancies inside the anionic sub-layers is determined mainly by the symmetry and size of these spatial vacancies (a reorientational disorder), while the dynamic properties of the anionic networks are limited primarily by the formation of polymeric structures. In the organic part, the dynamic properties change significantly when the temperature is being lowered since the cation mobility decreases. This process is accompanied by a significant distortion of the anionic lattice. It must be mentioned that one of the most significant features of this lattice is its flexibility and, at the same time, its facility of adaptability of symmetric cations, which is related to the effect of the electron lone pair on the Sb(III) and Bi(III) atoms. The presence of hydrogen bonds of the $N-H \cdots X$-type in the crystal structure is significant for the lattice formation. The strength of these bonds increases proportionally to the size of atom radius of the corresponding halogen. A rearrangement of these bonds takes place during the PT, which significantly affects the dynamic properties as well as the ordering of the cations.

In the present paper, we would like to focus on the ferroelectric structures among halogenoantimonates(III) and halogenobismuthates(III)), which can be found in the literature and on the correlation between their crystal structures and polar properties. For this reason, the structural analysis of the compounds with the organic cations, a description of the cationic dynamics and the deformation of the inorganic layers have been undertaken. Moreover, macroscopic dielectric properties are discussed in detail.

### 2. Structural diversity of the anionic networks of haloantimonates(III) and halobismuthates(III)

Haloantimonates(III) and halobismuthates(III) are characterized by a significant diversity of the anionic networks (from zero-, through one-, two-, and three-dimensional architectures).\(^{30-44}\) Taking into account the molar ratio of the amine ($R$) and metal ($M$) ($Sb(III)/Bi(III)$) the following four basic stoichiometries are observed.
2.1. RMX₄ (R:M = 1:1)

The anionic sublattices observed in this group may be divided into two types: (i) polyanionic chains; (ii) isolated units.

As for type (i) three different types of polyanionic chain arrangements are observed:

- chains of the edge-sharing octahedra (with two terminal and four bridging halogen atoms) (Fig. 2(a))
- the [MX₆]³⁻ octahedra linked as follows: three non-equivalent M atoms possess three bridging and three contact contacts, whereas the remaining one possesses only one terminal and five bridging contacts (Fig. 2(b));
- chains of the corner-sharing square pyramids (Fig. 2(c));

With regard to the separated units (ii) four different assemblies may be recognized.

Four octahedra linked with each other in two different ways:

- one type of central atom with three bridging and three terminal halogens (Fig. 2(d))
- two central atoms surrounded by three terminal and three bridging halogens and two others coordinated by two terminal and four bridging halogens (Fig. 2(e));
- face-sharing octahedra (Fig. 2(f));
- edge-sharing two square pyramids (Fig. 2(g));

2.2. R₃MX₉ (R:M = 1.5:1)

- infinite, one-dimensional (1D) zig-zag double chains (pleated ribbon structures) (Fig. 3(a));
- two-dimensional (2D) layers (honeycomb like corrugated sheets) (Fig. 3(b));
- discrete bicontinuous (0D) (Fig. 3(c));
- four octahedral units (0D) ([M₄X₁₂]⁻) (Fig. 3(d)).

2.3. R₂MX₅ (R:M = 2:1)

- one-dimensional (1D) chains (cis- or trans-type) (Fig. 4(a) and (b))
- isolated [M₂X₁₀]⁴⁻ units (two edge-sharing octahedra) (Fig. 4(c)).
tetrameric polyanion with corner-sharing octahedra (Fig. 4(d)).

2.4. R₅M₂X₁₁ (R : M = 2.5 : 1)
- Corner-sharing octahedra (0D) ([M₂X₁₁]⁻⁻) (Fig. 5(a)).

2.5. R₃MX₆ (R : M = 3 : 1)
- Isolated [MX₆]³⁻ octahedra (Fig. 5(b)).

2.6. Different modifications of the anionic substructures
The richness of the anionic structures among halobismuthates (III) and haloantimonates(III) is related also to the more exotic type of stoichiometry (see Fig. 6), e.g. RM₇X₇, RM₉X₁₀, or RM₈X₁₂.

3. Ferroelectricity among haloantimonates(III) and halobismuthates(III)
The ferroelectricity among halobismuthates(III) and haloantimonates(III) is limited only to four different types of chemical...
compositions: RMX₄, R₃M₂X₉, R₂MX₅ and R₃M₂X₁₁. Below, the analysis of the physical properties is presented and a correlation between the crystal structure and the origin of their polarity (ferroelectricity) is discussed. In our review we focus on pure organic–inorganic hybrids based on Bi(III) and Sb(III) (mixed anionic networks were not analyzed); however, due to the discovery of the first ferroelectric among halobismuthates (m) characterized by the mixed organic networks (R'R₂'MX₆) a single exception has been made.

3.1. RMX₄ type compounds

Ferroelectricity in the RMX₄ type compounds is a unique feature. Based on the literature, only two examples of ferroelectrics, characterized by the 1D polymeric anionic structure, are known: (4-aminopyridinium)[SbCl₄]₅¹–₅₃ ((4-NH₂C₅H₄NH)[SbCl₄], 4-APCA) and (trimethylamino-N-methyl stilbazolium) [Bi₂Cl₆]₅₄ ([C₁₇H₂₄N₂][Bi₂Cl₆], [TAMS][Bi₂Cl₆]). The detailed X-ray structure analysis indicates that in both compounds, the ferroelectricity is related mainly to a strong deformation of the [MX₄]₁⁻ type of substructure (see Fig. 2(a) and Table 2).

The crystal structure of 4-APCA is built up of the polyanionic chains of [SbCl₄]⁻ forming a tunnel-like structure and disordered 4-aminopyridinium cations connected via weak hydrogen bonds. Phase situation is complex and characterized by rich polymorphism in the solid state (see Scheme 1). The ac calorimetry studies have shown that the high temperature PT at 304 K is of the ‘order – disorder’ type and leads to a incom-
mensurate modulated phase (II). At 270.5 K (II → III) a ‘lock-in’ transition to a commensurate modulated phase is observed, which, in turn, becomes a ferroelectric one. The anomaly at 248 K leads to the next polar, but modulated incommensurate phase, while below 240 K the crystal becomes nonpolar again. The ferroelectric transformation at 270.5 K is characterized by a ‘displacive’ type of mechanism related to the displacement of the organic cations in relation to the anionic chains. Dielectric dispersion measurements carried out in the radio-frequency region around $T_c$ (270.5 K) revealed a critical slowing-down assigned mainly to the domain-like wall motion of the incommensurate phase (Fig. 7(a)). The relaxation process was found to be nearly monodispersive, which is rare among incommensurate ferroelectrics. Dielectric measurements confirm the lack of PT in a wide temperature range and indicate the existence of the dielectric relation process (Fig. 7(b)). The expected effect of the slowing down of the macroscopic relaxation time, related to the dynamics of the ferroelectric domains, is not observed when approaching the para-ferroelectric transition, which might be connected to the huge contribution of the electric conductivity to the dielectric response. It should be noted that the estimated relaxation time for the relaxation process in [(TAMS)][Bi$_2$Cl$_8$] becomes shorter on heating, which is a quite opposite effect in comparison with that observed in the other ferroelectrics. This is characteristic of the systems with dipolar group reorientations without any long-range dipole–dipole interactions. The activation energy of the relaxation process calculated from the Arrhenius relationship, $E_a$, equals to 280 kJ mol$^{-1}$ and may be a result of some dipole cooperative motions or steric hindrance originating from the huge organic molecule as for the single dipole motion.

3.2. $\text{R}_3\text{M}_2\text{X}_9$ type compounds

Within the chemical stoichiometry $\text{R}_3\text{M}_2\text{X}_9$ the ferroelectricity is limited to two types of anionic sublayers among four possible ones (Fig. 3): 2D (two-dimensional layers) (Fig. 3(b)) and 0D-discrete bioctahedral units (Fig. 3(c)). The 2D structure is usually characteristic of the chloride and bromide ferroelectrics containing only small alkylammonium moieties (methyl-, dimethyl and trimethyl-ammonium) or non-substituted rings (e.g. pyrrolidinium cation) (Table 3). The bulky organic cations due to the steric effect cannot occupy small vacancies within the layers. On the other hand all known iodide ferroelectrics possess the 0D anionic network (Table 4) thus the restriction with respect to the size and symmetry of the organic moieties is not so obvious.
The characteristic feature of the paraelectric phase in 2D R₃M₂X₉ ferroelectrics is the presence of two crystallographically independent alkylammonium cations in the crystal structure (N(1) and N(2)). The cation N(1) is located in voids inside the polyanionic layers, whereas the N(2) one between the layers (Fig. 8). All cations are connected to the anionic sublattice by N–H⋯X hydrogen bonds. Both types of cations are highly disordered in the paraelectric phase.

The disorder of N(1) is usually described by a two-site model with an occupancy factor of the N(1) atom, equal to 0.5. The other cation N(2) is also split between two positions, N(21) and N(22), however, with different occupancy factors, which change with temperature. Since the N(2) cation is in the general position of the crystal structure, this change does not lead to any changes in the lattice symmetry in contrast to the N(1) cation dynamics. The paraelectric–ferroelectric transitions lead to the distortion of the anionic network, which results in the loss of the symmetry center. Thus the N(1) cations now occupy a single position in a ferroelectric domain and their long-range order is responsible for an appearance of $P_s$. In the

### Table 3 Phase diagram of R₃M₂X₉ with a 2D anionic architecture

<table>
<thead>
<tr>
<th>Compound</th>
<th>Phase</th>
<th>Temperature</th>
<th>$P_s$</th>
<th>$\varepsilon_{max}$</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>(CH₂NH₃)₂[B₁₂Br₉]</td>
<td>II: C₂/k</td>
<td>164 K, 1st/2nd</td>
<td>0.15 $\mu$C/cm²</td>
<td>80</td>
<td></td>
</tr>
<tr>
<td>(CH₂NH₃)₂[Sb₂Br₉]</td>
<td>III: P2₁</td>
<td>164 K, 1st/2nd</td>
<td>0.15 $\mu$C/cm²</td>
<td>81</td>
<td></td>
</tr>
<tr>
<td>[(CH₂NH₃)₂Br]₂Br</td>
<td>III: P2₁</td>
<td>164 K, 1st/2nd</td>
<td>0.15 $\mu$C/cm²</td>
<td>86 and 87</td>
<td></td>
</tr>
<tr>
<td>[(CH₂NH₃)₂Br]₂Cl</td>
<td>IV: P4</td>
<td>242 K, 2nd</td>
<td>0.004 $\mu$C/cm²</td>
<td>80</td>
<td></td>
</tr>
</tbody>
</table>

1st-first order PT, 2nd-second order PT

The characteristic feature of the paraelectric phase in 3D R₃M₂X₉ ferroelectrics is the presence of two crystallographically independent alkylammonium cations in the crystal structure (N(1) and N(2)). The cation N(1) is located in voids inside the polyanionic layers, whereas the N(2) one between the layers (Fig. 8). All cations are connected to the anionic sublattice by N–H⋯X hydrogen bonds. Both types of cations are highly disordered in the paraelectric phase.

The disorder of N(1) is usually described by a two-site model with an occupancy factor of the N(1) atom, equal to 0.5. The other cation N(2) is also split between two positions, N(21) and N(22), however, with different occupancy factors, which change with temperature. Since the N(2) cation is in the general position of the crystal structure, this change does not lead to any changes in the lattice symmetry in contrast to the N(1) cation dynamics. The paraelectric–ferroelectric transitions lead to the distortion of the anionic network, which results in the loss of the symmetry center. Thus the N(1) cations now occupy a single position in a ferroelectric domain and their long-range order is responsible for an appearance of $P_s$. In the

### Table 4 Phase diagram of R₃M₂X₉ with a 0D anionic architecture

<table>
<thead>
<tr>
<th>Compound</th>
<th>Phase</th>
<th>Temperature</th>
<th>$P_s$</th>
<th>$\varepsilon_{max}$</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>(CH₂NH₃)₂[B₁₂Br₉]</td>
<td>II: C₂/k</td>
<td>145 K, 1st</td>
<td>7.96 $\mu$C/cm²</td>
<td>147 and 148</td>
<td></td>
</tr>
<tr>
<td>(C₄H₉NH₃)₂[Sb₂Br₉]</td>
<td>II: R₃c</td>
<td>322 K, 1st</td>
<td>3.0 $\mu$C/cm²</td>
<td>149 and 150</td>
<td></td>
</tr>
<tr>
<td>(C₄H₉NH₃)₂[Sb₂Cl₉]</td>
<td>II: R₃c</td>
<td>323 K, 1st</td>
<td>5.2 $\mu$C/cm²</td>
<td>151</td>
<td></td>
</tr>
<tr>
<td>(NH₄CH₂NH₃)₂[B₁₂Br₉]</td>
<td>III: Cmc₂</td>
<td>134.4 K, 2nd</td>
<td>176.9 K, 2nd</td>
<td>152</td>
<td></td>
</tr>
<tr>
<td>(NH₄CH₂NH₃)₂[Sb₁₂Br₉]</td>
<td>IV: Cmc₂</td>
<td>328 K, 2nd</td>
<td>0.005 $\mu$C/cm²</td>
<td>153</td>
<td></td>
</tr>
</tbody>
</table>

1st-first order PT, 2nd-second order PT

The characteristic feature of the paraelectric phase in 2D R₃M₂X₉ ferroelectrics is the presence of two crystallographically independent alkylammonium cations in the crystal structure (N(1) and N(2)). The cation N(1) is located in voids inside the polyanionic layers, whereas the N(2) one between the layers (Fig. 8). All cations are connected to the anionic sublattice by N–H⋯X hydrogen bonds. Both types of cations are highly disordered in the paraelectric phase.

The disorder of N(1) is usually described by a two-site model with an occupancy factor of the N(1) atom, equal to 0.5. The other cation N(2) is also split between two positions, N(21) and N(22), however, with different occupancy factors, which change with temperature. Since the N(2) cation is in the general position of the crystal structure, this change does not lead to any changes in the lattice symmetry in contrast to the N(1) cation dynamics. The paraelectric–ferroelectric transitions lead to the distortion of the anionic network, which results in the loss of the symmetry center. Thus the N(1) cations now occupy a single position in a ferroelectric domain and their long-range order is responsible for an appearance of $P_s$. In the

The characteristic feature of the paraelectric phase in 2D R₃M₂X₉ ferroelectrics is the presence of two crystallographically independent alkylammonium cations in the crystal structure (N(1) and N(2)). The cation N(1) is located in voids inside the polyanionic layers, whereas the N(2) one between the layers (Fig. 8). All cations are connected to the anionic sublattice by N–H⋯X hydrogen bonds. Both types of cations are highly disordered in the paraelectric phase.

The disorder of N(1) is usually described by a two-site model with an occupancy factor of the N(1) atom, equal to 0.5. The other cation N(2) is also split between two positions, N(21) and N(22), however, with different occupancy factors, which change with temperature. Since the N(2) cation is in the general position of the crystal structure, this change does not lead to any changes in the lattice symmetry in contrast to the N(1) cation dynamics. The paraelectric–ferroelectric transitions lead to the distortion of the anionic network, which results in the loss of the symmetry center. Thus the N(1) cations now occupy a single position in a ferroelectric domain and their long-range order is responsible for an appearance of $P_s$. In the

The characteristic feature of the paraelectric phase in 2D R₃M₂X₉ ferroelectrics is the presence of two crystallographically independent alkylammonium cations in the crystal structure (N(1) and N(2)). The cation N(1) is located in voids inside the polyanionic layers, whereas the N(2) one between the layers (Fig. 8). All cations are connected to the anionic sublattice by N–H⋯X hydrogen bonds. Both types of cations are highly disordered in the paraelectric phase.

The disorder of N(1) is usually described by a two-site model with an occupancy factor of the N(1) atom, equal to 0.5. The other cation N(2) is also split between two positions, N(21) and N(22), however, with different occupancy factors, which change with temperature. Since the N(2) cation is in the general position of the crystal structure, this change does not lead to any changes in the lattice symmetry in contrast to the N(1) cation dynamics. The paraelectric–ferroelectric transitions lead to the distortion of the anionic network, which results in the loss of the symmetry center. Thus the N(1) cations now occupy a single position in a ferroelectric domain and their long-range order is responsible for an appearance of $P_s$. In the
case of the 2D ferroelectrics the polar direction is maintained within the corrugated layer structure. This means that the layers enhanced the polarizability of the system, which is coupled with the dynamics of the N(1) type cations.

An enhanced value of the electric permittivity close to \( T_c \) is observed for the 2D ferroelectrics, because easily polarizable anionic layers favor the long-range order of the dipole–dipole interactions (Fig. 9).

In spite of the structural similarities of the 2D type ferroelectrics their dielectric responses, character of PTs and phase situations are quite diverse. The dielectric response of the majority of the 2D analogs shows a critical slowing down of the macroscopic relaxation time, which indicates an ‘order–disorder’ mechanism of ferroelectric transition (Fig. 9(a)). In turn in the case of MABB the dielectric response (\( \varepsilon(T) \)) is characteristic of ‘improper’ ferroelectrics (1st order PT).

![Fig. 8](image_url) Comparison of the crystal structures of DMACA in the paraelectric and ferroelectric phases (ref.88).

![Fig. 9](image_url) Temperature dependence of the real part of the dielectric permittivity (\( \varepsilon' \)) for (a) \((\text{CH}_3\text{NH}_2)_3[\text{Sb}_2\text{Cl}_9]\) (DMACA)\textsuperscript{131} (b) \((\text{CH}_3\text{NH}_3)_3[\text{Bi}_2\text{Br}_9]\) (MABB)\textsuperscript{84,85} (c) \((\text{CH}_3\text{NH}_3)_3[\text{Sb}_2\text{Cl}_9]\) (TMACA)\textsuperscript{131} around \( T_c \).
(Fig. 9(b)), while for the TMACA the dielectric results indicate the existence of a narrow-range intermediate metastable phase (incommensurate) between the ferroelectric and paraelectric phases (Fig. 9(c)).

The only parameter that correlates well with the temperature scope of the ferroelectric phases is the cation size (Tables 3 and 4). It is clearly seen that an increase of the cations size leads to the shift of the Curie point towards high temperatures. The large cations usually have less freedom of movement in the anion layer (N(1)-type) gaps, which is why they are easier to freeze at higher temperatures in cooling cycles in comparison with the ferroelectrics with the small cations.

Haloantimonates(III) and halobismuthates(III), characterized by the 2D anionic structure, usually possess a smaller value of $P_s$ in contrast to that for the 0D analogs, in spite of the fact that the organic cations of the presented crystals have a larger value of dipole moments, $\mu$. Nevertheless it should be noted that in the formamidinium analogs the cations possess a quite small value of $\mu = 0.23$ D and the dielectric anomalies are rather intermediate. It seems that the relationship between structural parameters of the crystals and the macroscopic properties ($P_s$ and $\varepsilon_{\text{max}}$) is rather complex and its recognition requires further studies on $R_2MX_5$-type compounds. It is interesting that most 0D ferroelectrics possess polar properties above room temperature, except for $(\text{NH}_2\text{CHNH}_2)_3[\text{Bi}_2\text{I}_9]$\textsuperscript{151} and $(\text{NH}_2\text{CHNH}_2)_3[\text{Sb}_2\text{I}_9]$\textsuperscript{152}. The paraelectric phase exhibits usually high symmetry (trigonal or hexagonal), which proves a high dynamic disorder of the cationic networks. Generally, the mechanism of ferroelectric phase transitions for the 0D compounds is quite complex because the ‘order–disorder’ and ‘displacive’ contributions are possible.

3.3. $R_2MX_5$ type compounds

A Cambridge Structural Database survey (version 5.40 March 2019) permits us to predict the probability of the appearance of acentric/ferroelectric compounds within $R_2MX_5$-type stoichiometry. 192 entries, adopting either polymeric (1D) or isolated (0D) anionic substructures, have been found.

The results presented in Tables 5 and 6 confirm that the ferroelectricity in the $R_2MX_5$-type compounds is adopted in the 1D anionic network. The ferroelectricity does not depend on the symmetry and size of the organic cations (heteroaro-
matic, heterocyclic aliphatic) but is strongly affected by the dipole moment values.

A characteristic feature of this subclass is the construction of the anionic lattice, which is created by 1D infinite chains of different configurations (cis- or trans-mode), however, the cis-conformation (and its modifications) is very common in the analyzed subgroup. Generally, one can state that the distortion of the anionic chains through the PT is a driving force, which contributes mainly to the \( P_s \) value. This effect is coupled with the dynamics of the polar organic cations, which are usually disordered in the paraelectric phase to varying degrees. Distortion of the anionic substructure is treated as a 'displacive' contribution to the molecular mechanism of the ferroelectric transition, whereas a change in the dynamics of the organic moieties is an 'order–disorder' contribution (Fig. 10).

The molecular mechanism of the paraelectric-ferroelectric transformation varies from 'displacive' through mixed up to purely 'order–disorder', however, which of these mechanisms is dominant depends strongly on the subtle differences in the development of the polyanionic chains. In the case of the 'displacive' \( R_2MX_5 \) subclass we should take into account the ns\(^2\) electron lone pair effect (5s\(^2\)-Sb and 6s\(^2\)-Bi), which seems to play a key role in the distortion of the MX\(_6\) octahedron. The derivatives, characterized by trans-connected octahedra containing bulky organic methyliogen MV amine: (MV) [BiBr\(_3\)],\(^{157}\) (MV)[Bi\(_3\)Cl\(_4\)], are an example of such a distorted structure.\(^{107}\) The MV\(^{2+}\) cations have stabilized unprecedented regular [BiBr\(_5\)]\(_2\) chains of trans-connected octahedra.

Through PT, the trans Bi–Br bonds differentiate markedly and the result is the chain polarity. Moreover, the electronic lone pair is stereochemically activated below 243 K leading to an acentric polar phase. The mixed-halide analog, (MV) [Bi\(_3\)Cl\(_4\)], reaches one of the highest room-temperature \( P_s \) values (>15 \( \mu \)C cm\(^{-2}\) at 298 K) in the field of organic–inorganic hybrid ferroelectrics. It should be added that trans-connected isomers are much rarer than cis-connected ones. It is difficult to indicate a different correlation between the crystal structure and dynamics and the scope of polar phases in this group of compounds.

Compared to \( R_3M_2X_9 \) connections, the organic cations in \( R_2MX_5 \) show less freedom of movement in the paraelectric phases (usually a two-site model) thus the compounds adopt

---

**Table 6** The results of CSD analysis

<table>
<thead>
<tr>
<th>Anionic substructure</th>
<th>All compounds</th>
<th>Piezoelectric</th>
<th>Ferroelectric</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isolated pyramids [MX(_5)](^{2-})</td>
<td>35</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>Isolated bioctahedra [M(<em>4)X(</em>{10})](^{4-})</td>
<td>81</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Isolated tetrameric forms [M(<em>4)X(</em>{30})](^{8-})</td>
<td>4</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Infinite chains [MX(<em>5)](</em>\infty)</td>
<td>72</td>
<td>30</td>
<td>9</td>
</tr>
<tr>
<td>Summary</td>
<td>192</td>
<td>41</td>
<td>9</td>
</tr>
</tbody>
</table>

---

**Fig. 10** Comparison of the crystal structures of (NH\(_2\))(CH\(_2\))\(_3\)NHCH\(_3\)][BiCl\(_5\)]\(^{155}\) in the paraelectric and ferroelectric phases.
lower symmetry (mainly orthorhombic). Thus most of the para-electric-ferroelectric transitions in the R₂MX₅-subclass is described by the mmmFmm2 Aizu relationship (non ferroelastic transitions), in turn methylviologen analogs characterized by trans-connected-modes of the anionic chains exhibit also non-ferroelastic transition between monoclinic phases described by 2/mFm or 2/mP2 species (Fig. 11).

As it is presented (Table 5) for the hybrids based on very similar organic cations (N-methyl-1,3-diaminopropane) and (N, N-dimethyl-1,3-diaminopropane) the differences in the Tc values (existence of polar phases) reach about 230 K (376 and 143 K, respectively), however, the values of Pₑ are comparable (2.38 and 1.36 μC cm⁻², respectively). In turn, the organic–inorganic compounds based on the ethylammonium cation indicate similar values of Tc (190 and 160 K), while Pₑ differs almost by three orders 1.4 and 5 x 10⁻³ μC cm⁻². In conclusion, within R₂MX₅-type ferroelectrics the amount of the deformation of the anionic chains is crucial, when it comes to the value of Pₑ and a character of the dielectric response close to Tc (Fig. 12).

3.4. R₃M₂X₁₁ type compounds

Complexes of R₃M₂X₁₁ stoichiometry are unique in crystalchemistry of haloantimonates(III) and halobismuthates(III). Currently, scientific literature gives only seven examples of the compounds marked by this type of chemical composition and all of them exhibit ferroelectric properties (see Table 7). A typical trait of the compounds in question is the presence of

![Fig. 11](image1.png)

**Fig. 11** Comparison of the crystal structures of (C₁₂H₁₄N₂)₃[BiBr₅]¹⁵⁷ in the paraelectric and ferroelectric phases.

![Fig. 12](image2.png)

**Fig. 12** Temperature dependence of the real part of the complex dielectric permittivity for: (a) (C₃H₅N₂)₂[SbCl₅]¹⁵⁸ (b) (C₁₂H₁₄N₂)₃[BiBr₅]¹⁵⁷
discrete biocahedral units, \([\text{M}_2\text{X}_{11}]^{5-}\) in their crystal structure, in which two octahedra are connected with each other by one bridging halogen atom (see Fig. 13).

A cationic subnetwork consists in turn of four kinds of nonequivalent types of organic species marked by varied dynamic disorder. Phase situation for all \(\text{R}_5\text{M}_2\text{X}_{11}\)-ferroelectrics is presented in Table 7.

A detailed analysis of the phase diagrams and structural data shows that the size of the organic cation and its shape and symmetry, have a significant influence on PTs’ sequence, on the symmetry of each phase, on the polar properties of compounds as well as on the temperature range of the existence of the polar (ferroelectric) phases. Thus, this group may be divided into two subgroups: the first one created by the hybrids based on the aliphatic cations, e.g.: \((\text{CH}_3\text{NH}_3)^5[\text{Bi}_2\text{Br}_{11}],^{116-118} (\text{CH}_3\text{NH}_3)^5[\text{Bi}_2\text{Cl}_{11}]^{119-122}\) and \((\text{NH}_2\text{CHNH}_2)^5[\text{Sb}_2\text{Br}_{11}]^{163}\) (abbreviated as aliphatic analogs) and the second one formed by aromatic counterions:

![Phase diagram of \(\text{R}_5\text{M}_2\text{X}_{11}\)]

Table 7  Phase diagram of \(\text{R}_5\text{M}_2\text{X}_{11}\)

<table>
<thead>
<tr>
<th>Compound</th>
<th>PT Type</th>
<th>(T_c) (K)</th>
<th>(P) (μC/cm²)</th>
<th>(\varepsilon_{\max})</th>
</tr>
</thead>
<tbody>
<tr>
<td>((\text{CH}_3\text{NH}_3)^5[\text{Bi}<em>2\text{Br}</em>{11}])</td>
<td>1st-first order PT, 2nd-second order PT</td>
<td>77 K, 2nd</td>
<td>II: Pca2,</td>
<td>312 K, 2nd</td>
</tr>
<tr>
<td>((\text{CH}_3\text{NH}_3)^5[\text{Bi}<em>2\text{Cl}</em>{11}])</td>
<td></td>
<td>170 K, 2nd</td>
<td>III: P2₁</td>
<td>250 K, 2nd</td>
</tr>
<tr>
<td>((\text{NH}_2\text{CHNH}_2)^5[\text{Sb}<em>2\text{Br}</em>{11}])</td>
<td></td>
<td>163 K, 2nd</td>
<td>II: P2₁/n</td>
<td>200 K, 2nd</td>
</tr>
<tr>
<td>((\text{C}_5\text{H}_5\text{N})^5[\text{Bi}<em>2\text{Cl}</em>{11}])</td>
<td></td>
<td>166 K, 2nd</td>
<td>II: P2₁/n</td>
<td>360 K, 1st</td>
</tr>
<tr>
<td>((\text{C}_5\text{H}_5\text{N})^5[\text{Bi}<em>2\text{Br}</em>{11}])</td>
<td></td>
<td>155 K, 2nd</td>
<td>II: P2₁/n</td>
<td>355 K, 1st</td>
</tr>
<tr>
<td>((\text{C}_5\text{H}_5\text{N})^5[\text{Sb}<em>2\text{Br}</em>{11}])</td>
<td></td>
<td>135 K, 2nd</td>
<td>II: P2₁/n</td>
<td>354 K, 1st</td>
</tr>
<tr>
<td>((\text{C}_5\text{H}_5\text{NH})^5[\text{Bi}<em>2\text{Br}</em>{11}])</td>
<td></td>
<td>118 K, 2nd</td>
<td>II: P2₁/n</td>
<td>405 K, 1st</td>
</tr>
</tbody>
</table>

A detailed analysis of the phase diagrams and structural data shows that the size of the organic cation and its shape and symmetry, have a significant influence on PTs’ sequence, on the symmetry of each phase, on the polar properties of compounds as well as on the temperature range of the existence of the polar (ferroelectric) phases. Thus, this group may be divided into two subgroups: the first one created by the hybrids based on the aliphatic cations, e.g.: \((\text{CH}_3\text{NH}_3)^5[\text{Bi}_2\text{Br}_{11}],^{116-118} (\text{CH}_3\text{NH}_3)^5[\text{Bi}_2\text{Cl}_{11}]^{119-122}\) and \((\text{NH}_2\text{CHNH}_2)^5[\text{Sb}_2\text{Br}_{11}]^{163}\) (abbreviated as aliphatic analogs) and the second one formed by aromatic counterions:

**Fig. 13** Independent parts of the unit cell of \([\text{C}_5\text{H}_5\text{N})^5[\text{Bi}_2\text{Br}_{11}]\) in the paraelectric and ferroelectric phases \((\text{M} = \text{Bi}(\text{III})/\text{Sb}(\text{III}), \text{X} = \text{Cl}, \text{Br}, \text{I})^{113,114}\)
ferroelectrics are much stronger than those in the aromatic R5M2X11 type: the following elements are common for the compounds of the di
towards the low temperatures. Despite obvious structural
differences between these two subgroups (aliphatic/aromatic),
the following elements are common for the compounds of the R5M2X11 type:
- PTs are continuous in nature;
- PTs are classified as an ‘order–disorder’ type;
- Close to Tc, a critical slowing down of the macroscopic relaxation time is observed;
- Molecular mechanism of the ferroelectric PT is dominated by the dynamics of the organic cations.

In the paraelectric phase, two out of five cations, being in a general position, are ordered, while the remaining three, (located at special positions) reveal an orientational disorder (two-site model, 180° reorientation). In the ferroelectric phase, all cations are ordered, however, the key input to Ps is given only by three out of five organic cations.

Another element that distinguishes the analyzed materials is the long-range dipole–dipole interaction. It determines both the values of Ps and the electric permittivity close to Tc. Methylammonium ferroelectrics mark themselves out by the higher values of Ps in the range of 2–3 μC cm⁻² and εmax 1–2 ×10⁴, than those for the aromatic analogs. The value of Ps for the latter compounds is by almost one order, and for ε’ almost by 1.5 order, lower than that for the former ones. This means that the long-range dipole–dipole interactions in the aliphatic ferroelectrics are much stronger than those in the aromatic ones. This in turn has an impact on the values of the PT temperatures. The stronger the dipole–dipole interaction the higher the values of Tc (Fig. 14).

The other property, which deserves to be compared, is the dynamic property of ferroelectrics, analyzed using the dielectric relaxation data. All R5M2X11 ferroelectrics indicate the critical slowing down of the macroscopic relaxation time around Tc, a phenomenon typical only of the ferroelectrics with an ‘order–disorder’ mechanism of PT. What matters for these crystals is a difference in the values of the macroscopic dielectric relaxation time. The dielectric dispersion in a case of the aromatic analogs is limited to the kilohertz frequencies (10⁻²–10⁴ Hz), while in the aliphatic ones it is shifted to the microwave range (10⁻⁸–10⁻¹⁰ Hz). One of the reasons for such a difference may be the size of the inertia moments of the organic dipoles as well as the way, in which the cations are bound through the hydrogen bound networks. The imidazolium cations have much higher inertia moments than methylammonium and formamidinium.

The most mysterious phenomenon in the R5M2X11 ferroelectric group is the molecular mechanism of PT. In this regard, two possible contributions must be taken into account:
(i) ‘order–disorder’
(ii) ‘displacive’

The first contribution (i) is related to the dynamics of the organic cations, i.e. orientational disorder. The typical feature of the ferroelectrics with a continuous PT is the critical slowing down of the macroscopic relaxation time and a significant entropy effect (ΔS > R ln 2) indicating the mechanism of type (i).

The other contribution, of the ‘displacive’ type (ii), is not proved by experimental methods; however, based on the structural data, there are reasons to believe that the distortion of the discrete biotahedral units is likely to appear and, therefore, should be taken into account. This effect is well visible among the aliphatic ferroelectrics but it is marginal in aromatic ones. The distortion of the biotahedra units causes a

Fig. 14 Temperature dependence of the real part of the dielectric permittivity (ε’) for (a) (CH₃NH₃)₅[Bi₂Br₁₁]¹¹⁶–¹¹⁸ (MAPBB); (b) (C₃H₇N₂)₅[Bi₂Br₁₁]¹¹⁴; (c) (C₂H₅NH₃)₅[Bi₂Br₁₁]¹¹⁵ around Tc.
change in the negative charge distribution in the crystal structures, related to the halogen atoms; this subsequently leads to a change of the 

\[ P_e \] values. There is one factor more necessary to analyze, namely, the electron lone pair effect. This effect should be attributed to the presence of the 6s^2 electrons of Bi (\(\text{Bi}\)) atom, which is less polarizable than 5s^2 electrons of Sb (\(\text{Sb}\)) atoms. All that leads to the conclusion that the anionic sublattice plays an important role in the generation of ferroelectricity in the R_5M_2X_{11}-type hybrids. Flexibility of the biotahedral units (corner-sharing octahedra) may be a decisive parameter in the origin of spontaneous polarization.

### 3.5. R_3MX_6 type compounds and their modification (e.g. R’-R’’-MX_6)

Recently a first halobismuthate(\(\text{Bi}\)) compound characterized by mixed organic networks has been reported by Wang et al.\(^{161}\)

The cationic substructure consists of two different molecules: one methylammonium (R’) and two benzylammonium (R’’) ones as well as by an isolated octahedral unit [BiBr_6]^{-}.\(^{162}\)

\[ \text{[CH}_3\text{CNH}_2]_2\text{H}_3\text{CH}_2\text{CH}_2\text{NH}_2\text{BiBr}_6 \] appears to be a RT multiaxial ferroelectric with \( P_e = 1.0 \mu \text{C cm}^{-2}. \) The symmetry-breaking paraelectric–ferroelectric transformation is regarded in the Aizu notation as \( 3\text{m} \parallel \text{m} \) at 386 K. The molecular mechanism of the ferroelectric PT is assigned to the change in dynamics of the organic cations thus the transition is classified as an ‘order–disorder’ type. The discovery of a new type of connection of halobismuthates(\(\text{Bi}\)) with a mixed anionic substructure creates a new path to design molecular multiaxial ferroelectrics.

Although R_3MX_6-type compounds seem to be the ‘poorest’ with regard to their ferroelectricity, an interesting phenomenon, related to the imidazolium based hybrid, has been detected. The most spectacular result was the preparation of a new crystalline ferroelectric material (\(\text{[(CH}_3\text{H}_2\text{N}_3)_2\text{SbBr}_6]\)) using an \textit{in situ} solid-state chemical reaction, where the starting substrate was \(\text{(C}_3\text{H}_5\text{N}_2)_5\text{[Sb}_2\text{Br}_11\text{]}\)}.\(^{162}\)

### 4. Conclusions

In this review we present the attempts to find a correlation between crystal structures and polar/ferroelectric properties of haloantimonates(\(\text{Bi}\)) and halobismuthates(\(\text{Bi}\)). The materials under consideration are characterized by a rich diversity of the anionic sublattice (more than 40 types), nevertheless, the ferroelectric properties are limited only to four specific compositions: RMX_4, R_3MX_9, R_3MX_5, R_5M_2X_{11}. Of importance is that within the mentioned stoichiometry, the anionic network dimensions are also changed. One should note that so far already 31 ferroelectrics have been synthesized and fully characterized from the above mentioned family, which put them on one line with a well-known family of perovskite-type ferroelectric crystals (\(\text{ABX}_3, \text{A}_2\text{BX}_4\)).\(^{164–167}\)

The RMX_4-type ferroelectrics (2 examples) indicate a 1D anionic structure, while the PTs are classified as a ‘displacive’ one. In the case of R_3MX_9 hybrids the ferroelectric ordering is observed only in the 0D (face-sharing biotahedral species) and 2D inorganic lattice (corrugated layer structure and small organic cations). The molecular mechanism of the paraelectric–ferroelectric PTs for the 2D ferroelectrics is of the ‘order–disorder’ type. In turn for the 0D compounds the mechanism is quite complex because the ‘order–disorder’ and ‘displacive’ contributions are possible. Moreover, in the case of the R_5M_2X_{11} complexes of the 2D-type the ferroelastic structure in both paraelectric and ferroelectric phases is observed (possible biferroics), however, the coupling between the corresponding order parameters, polarization and deformation, is very weak.

With regard to R_3MX_6 complexes, the ferroelectricity is conditioned by the presence of the 1D anionic structures with different modifications and the majority of PTs is of the ‘displacive’-type with a minor ‘order–disorder’ contribution.

The R_5M_2X_{11} hybrids are the most attractive from the viewpoint of polar properties. In this subclass, the inorganic part is created by the discrete anionic structures of the corner-sharing octahedra (0D-type). All of the R_5M_2X_{11} crystals reveal ferroelectric properties. The molecular mechanism of the paraelectric–ferroelectric PT is not well recognized yet, however, the ‘order–disorder’ contribution, related to the dynamics of the organic cations, seems to play a key role. On the other hand the strong distortion of the anionic units as a result of the electron lone-pair effect (\(5s^2\) (\(\text{Sb}\)) and \(6s^2\) (\(\text{Bi}\))) influences additionally the total polarization of the material. Of great importance is that aliphatic analogs, \(\text{e.g.} \ [\text{CH}_3\text{NH}_3]_5\text{[Bi}_2\text{Cl}_{11}\text{]}, \ [\text{CH}_3\text{NH}_3]_5\text{[Bi}_2\text{Br}_{11}\text{]}\) and \(\text{[NH}_2\text{CHNH}_2]_5\text{[Sb}_2\text{Br}_{11}\text{]}\) indicate dielectric properties similar to the well-known TGS (triglycine sulphate) family and thus make them the most promising materials for applications.

It appears that not all stoichiometries have been analysed in terms of possible ferroelectric properties and R’R’’BiBr_4 is the best example. It should be also added that the complexes of iodobismuthates(\(\text{Bi}\)), both with the organic cations (\text{e.g.} methylammonium) and with the monovalent metals (mixed cationic sublattice), are the most advantageous in the case of absorbing materials in solar cells. We anticipate that the broadening group of molecular–ionic ferroelectrics will help to develop rational structure–property relationships, leading to materials with precisely tunable characteristics.

### Conflicts of interest

There are no conflicts to declare.

### Acknowledgements

The paper is a result of the realization of the project no. UMO-2016/21/B/ST3/004640201/2078/17 financed by the National Science Centre, Poland (Prof. G. Bator).

### References

1. G. Rijnders, S. Curra’s, M. Huijben, D. H. A. Blank and H. Rogalla, Influence of Substrate-Film Interface


30. S. A. Adonin, I. D. Gorokh, D. G. Samsonenko, I. V. Koroïkov, M. N. Sokolov and V. P. Fedin, Crystal Structures of Binuclear Bi(m) Chloride and Bromide
55 A. Lipka, Preparation and Crystal structures of 2,2′ Bipyridinium Pentachloroantimonate (C₁₀H₈N₂H₂)SbCl₅
61 U. Geiser, E. Wade, H. H. Wang and J. M. Williams, Structure of a new iodobismuthate: tetra(n-butyl)ammonium 1,2;1,2;1,2;2,3;2,3;2,3-hexaμ-iodo-1,1,1,3,3,3-hexadiotribismuthate(III) (3:1), Acta Crystallogr., 1990, C46, 1547.
65 S. Pohl, W. Saak, R. Lotz and D. Haase, On the Existence of Weak Interactions between Sb(n) and Phenyl Groups: SbBr3(SPMMe2Ph)4, Sb4i4(SePPh3)2,2CH2Cl2, SbBr3(iSPMe2Ph)4 and (Ph4P)2Sb2Br6·CH3CN, Z. Naturforsch., 1990, 45b, 1355.
Inorganic Chemistry Frontiers

Review

81. J. Mróz and R. Jakubas, Ferroelectric and Ferroelastic Phase Transitions in \((\text{CH}_3\text{NH}_3)_2\text{Sb}_2\text{Br}_9\) Crystals, \textit{Ferroelectr. Lett. Sect.}, 1994, 17, 73.


88. J. Zaleski and A. Pietraszko, Structure at 200 and 298 K and X-ray Investigations of the Phase Transition at 242 K of \([\text{NH}_4(\text{CH}_3)_3]_2\text{Sb}_2\text{Cl}_6\) (DMACA), \textit{Acta Crystallogr.}, 1996, B52, 287.


90. F. Lazarini, Tetraphenylphosphonium Enneabromodibismuthate(III), \textit{Acta Crystallogr.}, 1977, 33, 2686.


92. M. Hall, M. Nun, M. J. Begley and D. B. Sowerby, Nonahalogenodiantimonate(III)ates; their preparation and the crystal structures of [Hpy]_3[SbCl\textsubscript{6}], [nMe\textsubscript{4}][SbBr\textsubscript{6}], and [nMe\textsubscript{4}][SbBr\textsubscript{5}Cl\textsubscript{6}], \textit{J. Chem. Soc., Dalton Trans.}, 1986, 1231.

93. M. Węclawik, A. Gągor, R. Jakubas, A. Piecha-Bisiorek, W. Medycki, J. Baran, P. Zieliński and M. Gałązka, Structure-Property Relationships in Hybrid \((\text{C}_3\text{H}_7\text{NH}_3)_2\text{Sb}_2\text{I}_6\) and \((\text{C}_3\text{H}_5\text{NS})_2\text{Sb}_2\text{I}_6\) Isomorphs, \textit{Inorg. Chem. Front.}, 2016, 3, 1306.

94. P. Szklarz, R. Jakubas, A. Piecha-Bisiorek, G. Bator, M. Chański, W. Medycki and J. Wuttke, Organic-Inorganic Hybrid Crystals, \((2,4,6-\text{CH}_3\text{PyH})_3\text{Sb}_2\text{Cl}_6\) and \((2,4,6-\text{CH}_3\text{PyH})_3\text{Bi}_2\text{Cl}_6\). Crystal Structure Characterization and Tunneling of \(\text{CH}_3\) Groups Studied by \(^1\text{H}\) NMR and Neutron Spectroscopy, \textit{Polyhedron}, 2018, 139, 249.


96. A. Piecha, R. Jakubas, V. Kinzhybalo and W. Medycki, Crystal Structure, Dielectric Properties and Molecular Motions of Molecules in Thiazolium Halometalates(III): \((\text{C}_3\text{H}_7\text{NH})_6\text{M}_4\text{Br}_{18}·2\text{H}_2\text{O} (\text{M} = \text{Sb}, \text{Bi})\), \textit{J. Mol. Struct.}, 2012, 1013, 55.

97. A. Piecha, R. Jakubas, V. Kinzhybalo and T. Lis, Structural and Dielectric Properties of Thiazolium Chlorobismuthate (III) and Chloroantimonate(III), \textit{J. Mol. Struct.}, 2008, 887, 194.


108. M. Owczarek, P. Szklarz, R. Jakubas and A. Miniewicz, MX\textsubscript{2}: A New Family of Morpholinium Nonlinear Optical Materials among Halogenoantimonate(III) and


118 M. Polomska and R. Jakubas, Dynamics of Ferroelectric Domains in (CH3)NH2Bi2Br11 Studied by Liquid Crystal Decoration Technique, *Ferroelectrics*, 1990, 106, 57.


