Materials Formed between Tin(II) Fluoride and Alkaline Earth Metal Fluorides and Chlorides

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This report deals with compounds of tin(II) fluoride, also commonly called stannous fluoride, SnF2, the structure of which is related to the fluoride (CaF2) type. The fluorides are exceptionally high performance fluoride-ion conductors, particularly when SnF2 is combined with PbF2. This has been attributed to Pb2+ having a 6s2 lone pair, making the ion softer than alkaline earth metal ions. The phases can be either ordered or disordered, or with gradual change from fully disordered to fully ordered. Some of the disordered phases are nanocrystalline. In MSnF4 (M = Pb and Ba), a change from crystalline to nanocrystalline, simultaneous with the formation of two types of disorder, a positional disorder and an orientational disorder, is observed to take place upon an unusually short ball-milling time. One of the SnF2/PbF2 phases, PbSn4F10, deserves a special attention: with full Pb/Sn disorder and the undistorted fluoride-type β-PbF2 structure, just its existence seems to be a miracle, and as a consequence it is metastable with a fairly short life time. Tin(II) alkaline earth metal chloride fluorides were obtained only with barium, and one has a disordered form of the BaClF structure, that is itself derived from the fluoride-type structure of BaF2 by Cl/F ordering in the form of alternating planar sheets of fluoride ions and corrugated sheets of chloride ions perpendicularly to the c axis of the tetragonal unit-cell. The Ba/Sn(II)/Cl/F phase, doubly disordered Ba1−xSnxCl1+xF1−y solid solution was obtained, that has the BaClF structure, with two types of unique disorders: (i) a limited disorder between Cl and F, the limits of the solid solution being a function of the method of preparation (precipitation or solid state reactions), (ii) a very unique disorder between Ba2+ ions, Sn2+ ions and Sn(II) covalently bonded on the same site. Disorder between the two types of bonding of divalent tin on the same site had never been observed before and is in direct violation with the normal criteria for substitution. In addition, the presence of Sn2+ on the much larger Ba2+ site results in Sn2+ being very much loose in the much oversized site, resulting in considerable rattling, that is frozen at cryogenic temperatures. Furthermore, the Ba1−xSnxCl1+xF1−y solid solution obtained by precipitation undergoes a disorder-disorder phase transition on heating, i.e. from one form of disorder to another. The study of these phases and their properties was made possible by a combined use of X-ray powder diffraction and 119Sn Mössbauer spectroscopy.

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1. Introduction

The crystal structure of alkali metals fluorides is dependent on the r+2/r− ratio of the radius of the cation over that of the anion [1]. For r+2/r− ≤ 0.414 the metal ion coordination is tetrahedral (BeF2, SiO2 type), for 0.414 < r+2/r− ≤ 0.732, it is distorted octahedral (MgF2, rutile type), and for r+2/r− > 0.732, it is cubic, fluoride-type structure (CaF2, SrF2 and BaF2) [1]. For β-PbF2 (cubic) the structure is also the fluoride-type, while α-PbF2 has the orthorhombic structure of PbCl2 [1, 2]. The crystal structure of BaCl2 is also fluoride-type [1]. While the fluoride-type structure and BaCl2 are typically ionic, the structures of the three phases of SnF2 are covalent. Monoclinic α-SnF2 has a molecular structure; it is made of SnF4 tetramers with strong intermolecular interactions and it contains two kinds of tin atoms structurally different: Sn(1) is in a highly distorted SnF3E (E = lone pair) trigonal pyramidal coordination with a tetrahedral electron pair geometry, while Sn(2) is in a highly distorted SnF3E (E = lone pair) square pyramidal coordination with an octahedral electron pair geometry [3, 4]. The tetragonal high temperature phase γ-SnF2 has the structure of TeO2 with a seesaw SnF4E coordination, with a trigonal bipyramidal electron pair geometry, which is, in contrast with other similar coordinations, a distortion of a square antiprism [5, 6]. On cooling, a second order displacive γ ⇆ β transition takes place to give ferroelastic β-SnF2. In β-SnF2, the coordination of tin is intermediate between a SnF4E seesaw and a SnF4E square pyramidal [5, 6]. The tin valence orbitals are hybridized in all phases of SnF2 and bonding is covalent. Tin(II) has not been found to form ionic bonding with fluorine, however with chlorine, it can form either covalent bonding, for example in SnCl2 · 2H2O [7] or ionic bonding, like in anhydrous Ba2SnCl6 [8]. In the fluoride chlorides, such as SnClF [9] and Sn2ClF3 [10], tin(II) makes covalent bonds either with fluorine only to form cations that make ionie bonds with chloride ions, or with both fluorine and chlorine to form a polymeric covalent structure.

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The purpose of this work was to prepare new phases that would be hybrids of ionically bonded barium fluoride or chloride, or of lead(II) fluoride, and covalently bonded tin(II) fluoride, and see how the two modes of bonding would accommodate each other to create new materials. Only the phases, the structures of which are related to the fluorite-type, are described in details here. The structure and bonding of the compounds formed were studied by elemental analysis, X-ray powder diffraction and $^{119}$Sn Mössbauer spectroscopy.

2. Experimental

2.1. Materials preparation

2.1.1. Starting materials

The following reactants were used for the syntheses: SnF$_2$ 99% from Ozark Mahoning, α-PbF$_2$ 99.9% from Alfa, BaCl$_2$·2H$_2$O analytical grade from American Chemicals, BaF$_2$ 99% from Allied Chemicals and Dye Corporations, Ca(NO$_3$)$_2$·4H$_2$O ≥ 99.0% from Sigma-Aldrich, HF 40% aqueous solution from Mallinckrodt, and doubly distilled or deionized water. Anhydrous BaCl$_2$ was prepared by dehydrating the dihydrated salt. Crystalline reactants were checked by X-ray powder diffraction and were found to have only the expected peaks. In addition, SnF$_2$ was checked by DTA and identified by the α → γ transition at 150°C–160°C and melting point at 215°C [11]. The degree of hydration of BaCl$_2$·2H$_2$O was checked by TGA and found to be 2.04.

2.1.2. Precipitation reactions

For all precipitation reactions, the molar fraction $X_A$ (A = any reactant) was varied from 0.10 to 0.90 by increments of 0.10 for a coarse screening in order to identify reactions mixtures that result in new compounds being formed, and then a fine screening was performed in the regions of interest, as needed, by use of smaller increments. In all cases, the solution of one of the reactants was added dropwise to a solution of the other upon stirring. Aqueous solutions at ambient temperature were used in all cases. The precipitates were filtered, washed with a minimum amount of cold water and allowed to dry at ambient temperature.

2.1.3. Reactions at high temperature in dry conditions

The dehydration temperature of BaCl$_2$·2H$_2$O was reported to be 113°C [12]. This was checked by thermogravimetric analysis (TGA), and it was found to be fully dehydrated at 140°C, and the mass loss was measured to be 14.71% (theoretical for 2H$_2$O = 14.75%). To ensure that dry anhydrous BaCl$_2$ was obtained, BaCl$_2$·2H$_2$O was heated at 200°C for two hours in air, and then placed under vacuum while still hot to prevent rehydration.

Stoichiometric amounts of the dry reactants were intimately mixed by grinding in a mortar with a pestle. Then, the mixture was placed in a copper tube reactor that was pumped to remove air and filled with nitrogen in a glove box. Pumping and filling with nitrogen was done twice to ensure that residual air is reduced to a minimum. Temporary sealing of the copper tube reactor was carried out in the glove box, while final sealing was performed outside. The full details of the copper tube reactor method have already been described [13]. Then, the tubes were heated in an oven to the appropriate temperatures, and after the adequate heating time, they were taken out and quenched in air or in cold water, and then the tubes were opened according to the procedure described in Ref. [13].

Heating temperatures to obtain the phases of interest here were the following: PbSn$_4$F$_{10}$: 250°C, α-PbSn$_4$F$_{14}$: 250°C, β-PbSn$_4$F$_{14}$: 500°C, the Pb$_{1-x}$Sn$_x$F$_2$ solid solution: 500°C, BaSn$_4$: 500°C, the Ba$_{1-x}$Sn$_x$Cl$_{1+y}$F$_{1-y}$: 350°C, the latter according to the following reaction scheme:

$$\frac{1-2x}{2} \text{BaF}_2 + x\text{SnF}_2 + \frac{1+y}{2} \text{BaCl}_2 \rightarrow \text{Ba}_{1-x}\text{Sn}_x\text{Cl}_{1+y}\text{F}_{1-y}.$$  

2.1.4. Phase transitions by ball-milling and further treatments

Some of the phases prepared by the above methods were subjected to ball-milling using a Wig L Bug® amal- gamator from Crescent Dental MFG Co, with two steel balls. In some cases, the ball-milled phases were annealed under dry nitrogen in a copper tube reactor for two hours [13]. Others were stirred in water for 12 h, then they were filtered, washed and allowed to dry according to the same procedures as for precipitates.

2.2. Materials characterization

2.2.1. X-ray powder diffraction

X-ray powder diffraction was carried out by use of a Philips PW1050 diffractometer that had been automated with the Scipow Sietronics® system from DiffracTech. This allowed a phase identification of phases already known, by comparison with the diffraction patterns of starting materials and other possible side products already collected in our laboratory and by use of the µPDSM Micro Powder Diffraction Search Match® from Fein-Marquet. Only the phases of interest in this study were subjected to further analysis.

2.2.2. Mössbauer spectroscopy

The Mössbauer spectra were recorded using the following setup. The source was a nominally 25 mCi Ca$^{119m}$SnO$_3$-γ-ray source from Ritverc GmbH. Isomer shifts were referenced relative to a standard CaSnO$_3$ absorber at ambient temperature. The counting system was a scintillation counter from Harshaw, equipped with a 1 mm thick (Tl)NaI crystal. A palladium foil was used to absorb the 25.04 keV and the 25.72 keV X-ray lines generated by the source decay from the metastable 11/2 spin level of the $^{119m}$Sn precursor to the 3/2 spin of the first excited state. The Doppler velocity ($±10$ mm/s) was generated by the use of an Elscint driving system, including a Mössbauer MVT-4 velocity transducer, a Mössbauer MDF-N-5 waveform generator, and a MFG-N-5 driver. The amplifier, the single channel analyzer, and
the conductivity of terstitials, although that interpretation cannot explain type structure, by a mechanism of the Frenkel defects at the metal ion vacancies has been held responsible for (Fig. 1b). The presence of a large number of empty sites occupied alternate parallel to the axes of the unit-cell of fluoride ions occupied by a metal ion and those non-cubic, while that of the anion is tetrahedral. The cubes is a metal ion vacancy. The coordination of the cation is significant amount of covalency, according to the Fajans ing cations to be polarized sufficiently to give bonding a oride and chloride ions require much more highly polariz- large to have a negligible polarizing power, while the flu- structures can be considered to be purely ionic since the cubes 

3. Theoretical considerations

3.1. Crystal structures and bonding type

3.1.1. Ionic bonding in the fluorite-type structure

Figure 1 shows the relationship between the crystal structure of BaF₂ and those of BaClF and BaSnF₄. In unsubstituted BaF₂ (Fig. 1b), the crystal system is cubic and the Ba²⁺ ions are in cubic [MF₈] coordination, however half of the cubes of fluorides ions do not contain any metal ions and are therefore vacant [□F₈], where □ is a metal ion vacancy. The coordination of the cation is cubic, while that of the anion is tetrahedral. The cubes of fluoride ions occupied by a metal ion and those non-occupied alternate parallel to the axes of the unit-cell (Fig. 1b). The presence of a large number of empty sites at the metal ion vacancies has been held responsible for the significant fluoride ion conductivity of the fluorite-type structure, by a mechanism of the Frenkel defects using the metal ion vacancies for lodging fluoride ion interstitials, although that interpretation cannot explain why the conductivity of β-PbF₂ is higher than that of BaF₂ [15]. The structure of BaClF (Fig. 1a) is obtained from the structure of BaF₂ (Fig. 1b) by ordering fluoride and chloride ions according to a F–Cl sequence, parallel to the c axis of the unit-cell. This results in a tetragonal distortion with a large increase of the unit-cell parameter and a lattice change from face centered to primitive, however, there is no superstructure. These structures can be considered to be purely ionic since the Ba²⁺ ion has a noble gas configuration and is sufficiently large to have a negligible polarizing power, while the fluoride and chloride ions require much more highly polarizing cations to be polarized sufficiently to give bonding a significant amount of covalency, according to the Fajans rules [16–18].

3.1.2. The covalent structure of SnF₂

In contrast with the typically high melting point (Mp) of the fluorite-type MF₂ (Mp = 1355°C for BaF₂ and 855°C for PbF₂), the low melting point of SnF₂ (215°C) suggests that its structure is molecular [11, 12]. The crystal structure of the phase of SnF₂ stable at ambient conditions, α-SnF₂, confirms this: it is made of Sn₄F₈ tetramers (Fig. 2) with covalent intramolecular Sn–F bonds, while the intermolecular Sn...F interactions are much weaker, although quite stronger than normal [3, 4]. The structure of the β- and the γ-phases of SnF₂ is a covalent network type, with clearly strongly covalent bonding [5, 6]. The three phases of SnF₂ have been shown to have a fluoride ion conductivity similar to that of β- PbF₂ [19]. While the coordination of the metal ion in the fluorite-type is as high as allowed by the r⁺/r⁻ ratio of the ionic radii, and it is not distorted, that of tin in all phases of SnF₂ is reduced by one, two, or three, and its highly distorted (see Sect. 3.2.1 below).

Fig. 1. Projection of the structures of (a) BaClF (F-Cl order), (b) non-substituted BaF₂ and (c) BaSnF₄ (Ba- Ba-Sn-Sn order), all three in the system of axes of BaF₂.

Fig. 2. The crystal structure of α-SnF₂ consists of Sn₄F₈ tetramers. The Sn–F bonds are covalent. The numbers 1/4 and 3/4 are the z coordinate of the average height of each tetramer.

3.2. Tin(II) stereochemistry and Mössbauer spectroscopy

3.2.1. Tin(II) electronic structure and consequences regarding bonding and coordination

Tin belongs to group 14 and period 5 of the periodic table and its electronic structure is [Kr] 4d¹⁰5s²5p². It has therefore four valences electrons and it can exist in the +4 full oxidation state (tetravalent) and in the +2 suboxidation state (divalent), both of which are stable at ambient conditions. In the latter case, only the two 5p electrons are lost to form the Sn²⁺ ion or used to form covalent bonding, while the two 5s electrons are unused and form a non-bonding electron pair, most commonly called a lone pair since, being unshared, it belongs to tin only. This can generate the following two bonding situations.

(a) The Sn²⁺ ion is formed. This situation is found mostly with large and weakly electronegative metalloids or non-metals, such as selenium, tellurium, and iodine, in rare cases chlorine, and in some cases bromine. In this
situation, the Sn\(^{2+}\) electronic structure is [Kr] 4d\(^{10}\)5s\(^2\), i.e. the tin lone pair remains on the native 5s orbital. Bonding is ionic and the tin orbitals are not hybridized. Since s orbitals are spherical, the lone pair being purely 5s\(^2\) makes the stannous ion spherical, therefore its coordination can be highly symmetrical, like tetrahedral, octahedral, or cubic. Small distortions can occur, as allowed by local crystal symmetry. When the stannous ion is present, the lone pair is said to be "non-stereoactive", since being on an unhybridized spherical orbital, it does not modify the stereochemistry, i.e. it does not modify the tin coordination [20].

(b) Tin(II) is covalently bonded. This is found for small and highly electronegative non-metals, such as fluorine, oxygen, sulfur, and most of the time chlorine and bromine. In this bonding mode, the tin valence orbitals are hybridized. The following types of hybridization schemes have been observed, where the lone pair \(E\) occupies one of the hybrid orbitals:

- \(sp^3\): tetrahedral electron pair geometry, trigonal pyramidal molecular geometry, \(SnX_3E\), similar to the \(NH_3\) molecule;
- \(sp^3d\): trigonal bipyramidal electron pair geometry, seesaw molecular geometry, \(SnX_4E\), similar to \(SF_4\);
- \(sp^3d^2\): octahedral electron pair geometry, square pyramidal molecular geometry, \(SnX_5E\), similar to \(ClF_5\).

In all of the above cases, the geometry is in agreement with the expectations of the valence shell electron pair repulsion (VSEPR) model of Gillespie and Nyholm [21]. The VSEPR model accounts for the molecular geometry in nearly all cases for main group elements in free molecules. Deviations are observed sometimes in solids because of the contribution from the lattice energy. For example, in SnO and red PbO, the coordination \(MO_2E\) is square pyramidal, instead of seesaw expected by the VSEPR model [22, 23]. Being located on a hybrid orbital, a stereoactive lone pair occupies a position in the valence shell of the tin atom that would otherwise be occupied by a bonding pair, connected to an atom in the tin sphere of coordination. Therefore, a stereoactive lone pair decreases the coordination number by at least one. In addition, in agreement with the VSEPR model, a lone pair occupies more room in the valence shell of an atom than a bonding pair, therefore lone pair–bonding pair repulsions are larger than the repulsion between two bonding pairs, and it results in an additional distortion of the polyhedron of coordination of covalently bonded tin, making the bond angles smaller. Two models have been used to analyze the decrease of coordination and distortions that result from the presence of a stereoactive lone pair. Both are based on electron pair repulsions such as described by the VSEPR model [21].

(a) The Galy–Andersson model is based on the fact that the stereoactive lone pair of the period 5 elements In(I), Sn(II), Sb(III), Te(IV), I(V), and Xe(VI), occupies a volume similar to that of an oxygen or a fluorine atom [20]. The coordination model thus obtained is tetrahedral for \(SnX_3E\) (Fig. 3a), trigonal bipyramidal for \(SnX_4E\) (Fig. 3b) and octahedral for \(SnX_5E\) (Fig. 3c), where \(X\) is F or O and E is the lone pair, considered like if it were F or O. These geometries are the same as what is commonly called the “electron pair geometry”. In this model, the polyhedra are taken as being regular and the central atom is shifted away from the center of the polyhedron, toward the lone pair. This makes sense since the lone pair is unshared and is therefore closer to the central atom than the bonding pairs.

![Fig. 3. Tin(II) polyhedra according to the Galy–Andersson model: (a) SnF\(_3\)E, (b) SnF\(_4\)E, and (c) SnF\(_5\)E (E = lone pair) [20].](image)

(b) The Brown model is based on the fact that the \(r_+/r_-\) cation/anion ratio of ionic radii for the period 5 elements In(I), Sn(II), Sb(III), Te(IV), I(V), and Xe(VI) bonded to fluorine or oxygen is in the range to expect an octahedral coordination [24]. Brown’s model examines how the stereoactive lone pair disturbs the coordination that would otherwise be octahedral, by (i) reducing the coordination number, since no bonding can take place in the direction of the lone pair, and (ii) by distorting the...
reduced coordination because the lone pair occupies more room in the valence shell of an atom than a bonding pair (2nd VSEPR rule) (Fig. 4) [24]. The following models are observed:

- **C model** (Fig. 4c): three X (F or O) atoms are pushed away by the lone pair, to give a SnX$_3$X$'_3$ coordination, which is actually a SnX$_3$ trigonal pyramid (tetrahedral electron pair geometry taking the lone pair into consideration), the three X$'$ being pushed away so far that they form only very weak interactions with the central atom. The three Sn–X interactions are strong covalent bonds, very significantly shorter than the sum of the ionic radii, indicating a substantial orbital overlap. In addition, the X–Sn–X angles are smaller than the 109.5$^\circ$ expected for a tetrahedral electron pair geometry, due to intense lone pair–bonding pair repulsions. This is the same as the SnX$_3$E tetrahedral model in Galy–Andersson’s model [20];

- **A model** (Fig. 4a): two X atoms are pushed away by the lone pair, to give a SnX$_2$X$'_2$ coordination. With the two X$'$ pushed away by the lone pair, the real polyhedron is SnX$_4$ with a seesaw shape (triangular bipyramidal electron pair geometry, including the lone pair). Distortions by the lone pair–bonding pair repulsions are of two types: (i) two kinds of bond lengths: the two equatorial Sn–X$_{eq}$ bonds are shorter than the axial Sn–X$_{ax}$ bonds, and (ii) angular distortion: the equatorial X$_{eq}$–Sn–X$_{eq}$ angle is smaller than 120$^\circ$ and the X$_{aq}$–Sn–X$_{aq}$ angle is smaller than 180$^\circ$, values expected in the regular trigonal bipyramid. This is the same as the SnX$_4$E trigonal bipyramidal coordination in Galy–Andersson’s model [20];

- **E model** (Fig. 4e): only one X atom is pushed away by the lone pair and it results in a SnX$_3$X$'_3$ coordination, that is a SnX$_5$ square pyramid (octahedral electron pair geometry, including the lone pair), since the X$'$ atom is pushed far away from being a bonding distance. High repulsions by the lone pairs result in distortions on bond lengths and bond angles: (i) the Sn–X$_{eq}$ bond is very significantly shorter than the four equatorial Sn–X$_{eq}$ bonds, and (ii) the X$_{ax}$–Sn–X$_{eq}$ and non-diagonal X$_{eq}$–Sn–X$_{eq}$ bond angles are lower than the 90$^\circ$ expected for the regular octahedral electron pair geometry. This corresponds to the SnX$_2$E octahedral geometry in Galy–Andersson’s model [20];

- **B and D models**: the large disturbance created by the lone pair creates strictly the regular C, A, or E model only if the site symmetry of the central atom requires it. The regular C model requires a 3-fold axis along the Sn–E axis, and it has three equal bond lengths and three equal bond angles. The regular A model requires a 2-fold axis along the Sn–E axis, along the intersecting line of two normal mirrors, and it has two equal equatorial bond lengths, two equal axial bond lengths, and four equal axial to equatorial X$_{ax}$–Sn–X$_{eq}$ bond angles. The regular E model requires a 4-fold axis along the Sn–E axis, and it has four equal equatorial bond lengths, four equal axial to equatorial X$_{ax}$–Sn–X$_{eq}$ bond angles and two equal non-diagonal X$_{eq}$–Sn–X$_{eq}$ equatorial angles. Lower local symmetry results in further distortions. Some of these can be small and the coordination can be considered to be close enough to one of the regular models. However, distortions can be much larger and even the number of X$'$ atoms really pushed away by the lone pair can be difficult to define. Often, one or more atoms are sufficiently far to be counted as not being a covalent bond but still not far enough to be neglected and they form “weaker bonds”. In order to account for many of these unclear cases, Brown’s description includes two additional models, B and D, that are intermediates between the C, A, or E models. The B model (Fig. 4b) accounts for cases intermediate between models C and A, i.e. between the SnX$_3$ trigonal pyramid and the SnX$_4$ seesaw, therefore the coordination, SnX$_3$, is intermediate between these, where the last X is neither close enough to tin to be a covalent bond nor far enough to be neglected. Similarly, the D model (Fig. 4d) for SnX$_4$X is intermediate between the SnX$_4$ seesaw and the SnX$_2$ square pyramid. The uncertainty on the number of non-metal atoms that form a covalent bond to the central atom also makes it difficult to assign a hybridization scheme. Should a non-negligible “weak-bond” count to be using a hybrid orbital, although it is clearly much weaker than a normal covalent bond, yet much too close to be neglected? In addition, the presence of one or more extra atoms forming weak bonds changes dramatically the value of the bond angles. Some coordinations do not obey the VSEPR model, such as in SnO [23], no electron pair geometry can be defined, and neither can be the symbols in the Galy–Andersson model or in the Brown model. In such cases, the lattice energy compensates for the higher local energy at tin due to the VSEPR model not being obeyed.

### 3.2.2. Tin(II) electronic structure, bonding type and Mössbauer spectroscopy

Tin-119 has the same nuclear spin as iron-57, 1/2 in the ground state and 3/2 in the first excited state therefore, in the absence of magnetic interactions, only one line per site is observed if there are no quadrupolar interactions. This is the case of Fig. 5a and c. On the contrary, in the case of Fig. 5b, a large quadrupole splitting is observed, due to the interactions of the quadrupole moment of the $^{119}$Sn nucleus in the 3/2 spin state with the electric field gradient (e.g.) generated by the electric environment of tin. Calcium stannate CaSnO$_3$ has the perovskite structure, hence tin is present in the form of Sn$^{4+}$ stannic ions in octahedral coordination, i.e. there is no e.g. generated by the surrounding atoms/ions (|(\tilde{V}_{zz})_{latt}| = 0). The Sn$^{4+}$ ion has the [Kr] 4d$^{10}$ electronic structure. All orbitals that contain electrons are full and are in the core, and all the valence electrons have been lost, therefore there is no e.g. due to valence electrons (|(\tilde{V}_{zz})_{val}| = 0), hence the total \(\tilde{V}_{zz} = 0\). In addition, CaSnO$_3$ is taken as the reference of isomers shifts therefore, with in addition no quadrupolar interactions, it gives a single line at 0 mm/s (Fig. 5a). CsSnBr$_3$ has also the perovskite structure, thus tin is in the form of Sn$^{2+}$ stannous ions located in octahedral coordination, and it results that...
\((V_{zz})_{\text{latt}} = 0\). However, in contrast with \(\text{CaSnO}_3\), in \(\text{CsSnBr}_3\) tin is in the divalent state and the stannous ion has the \([\text{Kr}] 4d^{10}5s^2\) electronic structure, the \(5p\) electron being lost by ionization. In addition to the core electrons, there are two valence electrons, however they are on the \(5s\) orbitals, and since \(s\) orbitals are spherical, they generate no e.f.g. \((V_{zz})_{\text{val}} = 0\); hence the total \(V_{zz} = 0\) and there results a single line (Fig. 5c). The spherical lone pair is non-stereoactive, bonding at tin is ionic, and this generates no quadrupole splitting. The \(5s\) orbitals are not shared since there is no orbital hybridization and it does not participate in bonding. This makes it that the valence \(s\) electron density acting at the tin nucleus is high and this results in a large isomer shift, hence a single line (since \(V_{zz} = 0\)) at \(\approx 4.1\) mm/s (Fig. 5c). When bonding in \(\text{tin(II)}\) is covalent, the lone pair is stereoactive, therefore the coordination is incomplete and highly distorted and this results in a large \((V_{zz})_{\text{latt}}\). In addition, since the tin orbitals are hybridized, the non-spherical orbitals \(5p\) and \(5d\) when the octet is exceeded, contribute to form a large \((V_{zz})_{\text{val}}\). Since the contribution of the valence electrons to the quadrupolar interactions is much larger than that of the lattice, i.e. \(|(V_{zz})_{\text{val}}| \gg |(V_{zz})_{\text{latt}}|\), the former predominates, regardless of the value and sign of the Sternheimer shielding and antishielding factors. Therefore, a large quadrupole doublet centered at 3.0–3.6 mm/s is observed (Fig. 5b).

4. Results and discussion

4.1. The ordered structure of \(\alpha\)-\(\text{PbSnF}_4\) and \(\text{BaSnF}_4\)

The X-ray powder diffraction pattern of \(\alpha-\text{PbSnF}_4\) (Fig. 6b) is easily indexed to be a tetragonal distortion of the cubic structure of \(\beta\)-\(\text{PbF}_2\) (Fig. 6a). It should be noted that further lattice distortion to orthorhombic can also be obtained (Fig. 6c), however this will not be studied further here. The relationship between the Miller indices of the Bragg peaks of \(\beta\)-\(\text{PbF}_2\) and of \(\alpha\)-\(\text{PbSnF}_4\) leads to the following conclusions:

- there is a tetragonal distortion;
- The F (face centered) lattice translations of the fluorite-type structure have disappeared to give a P (primitive) lattice;
- the periodicity along the c axis of the unit-cell is doubled;
- there is a 45° rotation of the a and b axes around the c axis.

The relationship between the unit-cell axes of the two structures is shown in Fig. 7. They are related as follows: \(\beta\)-\(\text{PbF}_2\) or \(\text{BaF}_2(a,b,c) \Rightarrow \alpha\)-\(\text{PbSnF}_4\) or \(\text{BaSnF}_4(a/\sqrt{2},a/\sqrt{2},2c)\).

It results that \(^{119}\text{Sn}\) Mössbauer spectroscopy is a very useful tool for probing the divalent tin bonding type, lone pair stereoactivity, and orbital hybridization. It is particularly helpful in cases when the information obtained from the crystallographic methods have limitations, like the cases studied in this work, in order to attempt to understand how divalent tin, expected to give covalent bonding to fluorine, can combine with fluorite-type ionic structures to give stable structures. In such cases, much more complete conclusions can be obtained by combining the results of both techniques.
ferred orientation to use the Rietveld method. The X-ray or a polycrystalline sample with sufficiently reduced pre-
suitablesinglecrystalforsinglecrystalX-raydiffraction
ferred orientation [25]. It was therefore impossible to get
nanophase instead of the same phase with less or no pre-
this results in a phase transition giving a disordered
ble to reduce the preferred orientation by milling since
dimensional shape in the form of large sheets that crum-
ferred orientation, as expected by the crystallite two-
The samples suffer from a very acute case of pre-
or a polycrystalline sample with BaSnF$_4$ (primitive (a, b) face with F at the corners and Ba on two faces). The direction of the c axis, perpendicular to the figure, is unchanged however the periodicity is doubled.

Fig. 7. Relationship between the unit-cell axes of BaF$_2$ (face centered (a, b) face with Ba at the corners) and of BaSnF$_4$ (primitive (a, b) face with F at the corners and Ba on two faces). The direction of the c axis, perpendicular to the figure, is unchanged however the periodicity is doubled.

Fig. 8. Ambient temperature Mössbauer spectra of highly oriented $\alpha$-PbSnF$_4$: change of the asymmetry of the quadrupole doublet with the orientation of the sample relative to the $\gamma$-ray beam direction: (a) $\theta = 0^\circ$ ($\gamma$-ray beam perpendicular to the sample), (b) $\theta = 45^\circ$. $\theta$ is the angle between the $\gamma$-ray beam and the normal to the plane of the sample.

The samples suffer from a very acute case of preferred orientation, as expected by the crystallite two-dimensional shape in the form of large sheets that crumble as soon as one tries to isolate some. It is not possible to reduce the preferred orientation by milling since this results in a phase transition giving a disordered nanophase instead of the same phase with less or no preferred orientation [25]. It was therefore impossible to get a suitable single crystal for single crystal X-ray diffraction or a polycrystalline sample with sufficiently reduced preferred orientation to use the Rietveld method. The X-ray powder diffraction pattern (Fig. 6b) shows that all (00l) peaks are strongly enhanced while the (h00) and (hk0) peaks are much depressed, suggesting that the cleavage planes are perpendicular to the c axis of the unit-cell, hence parallel to the (a, b) plane.

The preferred orientation of a sample of $\alpha$-PbSnF$_4$ was enhanced by filtering the precipitate by suction and allowing it to dry in the Buchner funnel, and then it was removed from there without any disturbance and placed in the Mössbauer holder. The spectrum was recorded versus the orientation of the sheet in the $\gamma$-ray beam. The spectrum is a large quadrupole doublet with an isomer shift $\delta = 3.24(1)$ mm/s and a quadrupole splitting $\Delta = 1.57(2)$ mm/s. These spectra are very much like that of Fig. 5b and confirm that the tin lone pair is stereocative and that bonding is covalent with the lone pair being located on a hybrid orbital. It can be seen that when the $\gamma$-ray beam is perpendicular to the sheet, the Mössbauer doublet is highly asymmetric (Fig. 8a). When the sample is rotated at 45° from the $\gamma$-ray beam, the asymmetry is reduced dramatically. Such a behavior is observed in single crystals and in highly oriented polycrystalline samples [26]. The ratio of the intensity of the two lines in a single crystal is the following:

$$\frac{I_{3/2}(\theta)}{I_{1/2}(\theta)} = \frac{1 + \cos^2 \theta}{2/3 + \sin^2 \theta} \neq 1,$$

whereas $\theta$ is the angle between the $\gamma$-ray beam and $V_{zz}$, the main axis of the e.f.g. acting at tin. The spectrum will be symmetrical when the following condition is met:

$$I_{3/2}(\theta) = I_{1/2}(\theta) \rightarrow 1 + \cos^2 \theta = 2/3 + \sin^2 \theta \rightarrow \theta = \arcsin(1/\sqrt{3}) = 54.74^\circ.$$

For $\theta = 54.74^\circ$, called the magic angle, the two lines will have the same intensity. It was not possible to measure a Mössbauer spectrum at $\theta = 54.74^\circ$ because most of the $\gamma$-ray beam was absorbed by the sample and the sample holder at such a high grazing angle, however the spectrum measured at 45° (Fig. 8b) is clearly much more symmetrical than at $\theta = 0^\circ$. Obviously, at 54.74°, it would be close to symmetrical. This shows that at $\theta = 0^\circ$, the $\gamma$-ray beam is parallel to $V_{zz}$, and since this direction is normal to the sheet, $V_{zz}$ is parallel to the c axis. The main contributor to the e.f.g. is $(V_{zz})_{val}$, the imbalance in non-spherical valence electrons (p and d electrons), d due to the contribution of one or two 5d orbitals to hybridization, and the lone pair, on a hybrid orbital, is the main contributor (Fig. 9). This proves that the tin lone pair axis is parallel to c, thereby creating sheets of lone pairs that are at the origin of the extreme case of two-dimensionality and the resulting extreme case of preferred orientation (Fig. 6b).

Finally, combining together the crystallographic and the Mössbauer data, a realistic model of the $\alpha$-PbSnF$_4$ structure can be made. The relationship between the unit-cell axes of the fluorite-type structure and of $\alpha$–PbSnF$_4$, and the doubling of the c unit-cell parameter, require to have M (Pb or Ba) and Sn layers parallel to...
Fig. 9. Sample orientation of highly oriented α-PbSnF₄ in the γ-ray beam: (a) the c axis is normal to sheet-like crystallites, (b) the c axis is parallel to the lone pair axis and to \( V_{zz} \), (c) \( \theta = 0 \) in highly oriented α-PbSnF₄ when the γ-ray beam is normal to the sheets.

Fig. 10. M/Sn ordering in MSnF₄: (a) MF₂, (b) MSnF₄ first hypothesis (M Sn M ordering), (c) MSnF₄ second hypothesis (M M Sn Sn M M ordering).

Fig. 11. Ordering of the type ... Pb Pb Sn Sn Pb Pb ... required to double the c parameter, with lone pairs squeezed between the two tin layers in order to give highly efficient cleavage, and still agree with the tetragonal symmetry of the unit-cell. Unit-cell axes: \( \beta \)-PbF₂ cubic unit-cell \( (a,b,c) \); \( \alpha \)-PbSnF₄ tetragonal unit-cell \( (a',b',c') \).

Fig. 12. Barium and tin coordination in BaF₂ and in BaSnF₄: (a) BaF₂ cube in BaF₂, (b) SnFF'₂E pseudo-octahedron in BaSnF₄, (E = lone pair), (c) BaF₂F'₂F₄ unit in BaSnF₄, and (d) top view of the Barium coordination in BaSnF₄ (F' and F'' are superimposed).

4.2. Disordered structures of SnF₂ with CaF₂ and with PbF₂

The X-ray powder diffraction pattern of the following new phases discovered in our laboratory is cubic fluorite-type, despite containing more than one metal. Some of these phases are stoichiometric, such as \( \gamma \)-PbSnF₄, a non-quenchable high temperature phase, PbSnF₁₀, metastable at ambient conditions, and nanocrystalline \( \mu \gamma \)-PbSnF₄. Others are solid solutions and are therefore non-stoichiometric, \( M_{1-x}Sn_xF_2 \) (M = Ca and Pb). Figures 13 and 14 show two examples of X-ray powder diffraction patterns. Figure 13 shows the diffraction pattern of the precipitate obtained when a solution of calcium nitrate is added to a solution of stannous fluoride,
versus the molar ratio \( n \) of calcium nitrate in the reaction mixture. At low \( n \) values, i.e. for reactions mixtures rich in SnF\(_2\), a stoichiometric phase that was determined by chemical analysis to be CaSn\(_2\)F\(_6\) was obtained. Its X-ray powder diffraction pattern (Fig. 13a) is not related to that of the fluoride-type structure, therefore it will not be discussed further here. At high \( n \) values, for reactions mixtures rich in calcium ions, the X-ray diffraction pattern (Fig. 13b) is the same as that of fluoride-type CaF\(_2\). (Fig. 14a), with significantly broadened lines, while for medium values of \( n \), a mixture of the two phases is obtained (Fig. 13c).

Chemical analysis of several precipitates that have the diffraction pattern of Fig. 13b showed that, for all of them, they contain no nitrate and that 20 to 30% of the calcium has been replaced with tin. Therefore, the precipitate of Fig. 13b is a Ca\(_{1-x}\)Sn\(_x\)F\(_2\) solid solution. Applying the Scherrer formula to the linewidth corrected from instrumental broadening using the Warren formula, it was determined that the average diameter of the crystallites is 9 nm, and therefore the material is nanocrystalline [28]. A wide Pb\(_{1-x}\)Sn\(_x\)F\(_2\) solid solution \( 0 \leq x \leq 0.50 \) was prepared by heating mixture of the two fluorides at 500°C. It was found to have the cubic fluoride-type structure for \( 0 \leq x \leq 0.30 \) and the tetragonal \( \beta\)-PbSnF\(_4\) type for \( 0.30 < x \leq 0.50 \). The Ca\(_{1-x}\)Sn\(_x\)F\(_2\) solid solution could not be prepared by the same method. Instead, it decomposes to a mixture of CaF\(_2\) and SnF\(_2\) when heated. When \( \alpha\)-PbSnF\(_4\) is ball-milled, the superstructure peaks ((001) and (002)) and the peak splitting due to the tetragonal distortion (e.g. (110) and (004)) disappear rapidly to give the same list of peaks of the F lattice as for fluoride-type \( \beta\)-PbF\(_2\), while all peaks become broad (Fig. 15). The average crystallite diameter calculated from the line width at half height was 9 nm. Therefore, the following order-disorder phase transition takes place rapidly (50–60 s) on ball-milling (b.m.):

\[
\text{crystalline ordered } \xrightarrow{(b.m.)} \text{nanocrystalline disordered }
\]

\[
\alpha\text{-PbSnF}_4 \quad \mu\gamma\text{-PbSnF}_4
\]

The absence of superstructure peaks and of peak splitting that would show there is a lattice distortion in both the solid solutions and the stoichiometric phases that have the cubic F lattice of the fluoride-type show that tin and the other metal ion (Ca\(^{2+}\) or Pb\(^{2+}\)) are randomly disordered on the same Wyckoff site of the \( Fm\overline{3}m \) space group. If the two metals are disordered on the same site, they should have a similar size and the same bonding type, i.e. ionic bonding in the fluorite structure, in order to abide by the Hume–Rothery rules [29]. The size rule is violated in both cases, by 17% with Ca\(^{2+}\) and by 35% for Pb\(^{2+}\). If Sn substitutes Ca\(^{2+}\) or Pb\(^{2+}\) ions, Sn must be present in the form of Sn\(^{2+}\) ions in order to have the same bonding mode. However, divalent tin is known to form covalent bonding with fluorine. As shown in Fig. 5, Mössbauer spectroscopy can provide an unambiguous answer about the nature of Sn–F bonding. The spectrum of the Ca\(_{1-x}\)Sn\(_x\)F\(_2\) solid solution (Fig. 14c-2) and that of Pb\(_{1-x}\)Sn\(_x\)F\(_2\) (Fig. 16b) show clearly a quadrupole characteristic of covalent bonding. The same is true for nanocrystalline \( \mu\gamma\)-PbSnF\(_4\) obtained by ball-milling (Fig. 17c). It should be noted that the broad peak at ca. 0 mm/s is due to amorphous SnO\(_2\).
Fig. 15. X-ray powder diffraction pattern of $\alpha$-PbSnF$_4$(aq) subjected to ball-milling, versus ball-milling time: (a) not milled, (b) 10 s, (c) 20 s, (d) 50 s, (e) 60 s.

Figures 18 and 19 show attempts to fit covalent bonded tin in the MF$_2$ fluorite structure, on the M$^{2+}$ site, while abiding with the $m3m/O_h$ point symmetry of the site, i.e. in cubic or octahedral symmetry. In both cases, the tin lone pair is repeated by the site symmetry to generate eight lone pairs (Fig. 18) or six lone pairs (Fig. 19). Since tin has enough electrons for only one lone pair, none of these models is realistic, and in addition, none of these models provides opportunities for tin to make bonds with fluorine. It is very likely that each tin atom is located inside a [F$_8$] cube of the fluoride ion sublattice present in the fluorite structure, and forms bonds to one face only, to form four Sn–F bonds, with a square pyramidal coordination and a square pyramidal electron pair geometry (Fig. 20a). This is similar to the tin coordination in the tetragonal MSnF$_4$ (M = Ba and Pb), except that the axial fluorine has moved to the corner of the [F$_8$] cube that are vacant in MSnF$_4$ (Fig. 20b). The square pyramidal electron pair geometry is not favored by the VSEPR model for five electron pairs, however it can be stabilized by the lattice energy in order to satisfy the requirements of the crystal symmetry, like in SnO, and for lead(II), in red PbO, litharge [22, 23]. However, in order to avoid creating six lone pairs (Fig. 19), the tin atoms must be shifted away from the center of the [F$_8$] cube, along the tin–lone pair axis, thereby changing its point symmetry from $m3m$ to $4 mm$. In addition, the shift of the tin atom must be toward the face of the cube it form bonds to, in order to give room for the lone pair on the other side. This results in the model of Fig. 21. In order to agree with the cubic symmetry of the lattice and the absence of superstructure, two types of disorders must be present:

- a *positional disorder*: the two metals, Sn and Ca or Pb must be randomly disordered on the metal ion site;
- an *orientational disorder*: the tin–lone pair axis must point randomly towards any of the six faces of the [F$_8$] cube in order to avoid creating a privileged direction and to maintain an averaged cubic symmetry.

due to surface oxidation of the particles [30]. The contribution of this peak to the total spectrum is much higher than the proportion of SnO$_2$ in the sample, because its recoil-free fraction is about 20 times as high as that of tin(II) fluorides. The X-ray diffraction and the Mössbauer results appear to give contradictory results: on the one hand, the structure is ionic fluorite-type (diffraction), and on the other hand, Sn–F bonds are covalent (Mössbauer), and in addition, the Hume–Rothery rules state that both metals disordered in the structure, Sn and Ca (or Pb) must use the same mode of bonding.
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4.3. The doubly disordered $Ba_{1-x}Sn_{x}Cl_{1+y}F_{1-y}$ solid solution

When an aqueous solution of BaCl$_2$·2H$_2$O is added to a solution of SnF$_2$ (Ba$ightarrow$Sn), or vice-versa (Sn$ightarrow$Ba) upon stirring, versus the molar ratio X of barium chloride in the reaction mixture, four different types of precipitates were obtained:
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X ≤ 0.10: pure BaSnF₄, already known [31];
0.34 ≤ X ≤ 0.40: pure BaSn₂Cl₂F₄, a new phase;
0.52 ≤ X ≤ 0.80: pure BaSnClF₃ · 0.8H₂O, a new phase;
X > 0.85: a pure “BaClF” like phase, a new phase.

Mixtures of phases were obtained for X values in between these. The stoichiometry of each pure phase was established by elemental analysis and the degree of hydration of BaSnClF₃·0.8H₂O by thermogravimetry. Only the “BaClF” like phase is of interest to us here. Its X-ray powder diffraction pattern contains the same set of the Bragg peaks as that of BaClF with some lineshifts due to the “BaClF” like phase. Its X-ray diffraction pattern contains the same set of the Bragg peaks as that of BaClF with some lineshifts due to the “BaClF” like phase. Its X-ray diffraction pattern contains the same set of the Bragg peaks as that of BaClF with some lineshifts due to the “BaClF” like phase.

The chemical composition was found to be variable for both the metals and the halogens, therefore it is a non-stoichiometric solid that has the BaClF structure, hence its chemical formula can be written in the form of a doubly substituted solid solution, Ba₁₋ₓSnₓCl₁₋ₓF₁₋ₓ. Element analysis of several samples showed the following variation of the chemical composition: 0 < x < 0.15 and 0 ≤ y ≤ 0.15. The ¹¹⁹Sn Mössbauer spectra (large quadrupole doublet at an isomer shift around 3 mm/s) show that tin is covalently bonded in BaSn₂Cl₂F₄ and in BaSnClF₃ · 0.8H₂O, while the broad line near 4 mm/s for the Ba₁₋ₓSnₓCl₁₋ₓF₁₋ₓ solid solution shows that it contains tin(II) with ionic bonding in a distorted environment (Fig. 23). The Ba₁₋ₓSnₓCl₁₊yF₁₋ₓ solid solution was also prepared by the dry method at high temperature, by reaction of appropriate amounts of BaF₂, BaCl₂ and SnF₂. This method makes it possible to control the stoichiometry of the solid solution and no elemental analysis is necessary. Figure 24 shows the domain of a pure solid solution in the (x, y) plane. A pure Ba₁₋ₓSnₓCl₁₊yF₁₋ₓ solid solution was obtained within the following limits: 0 ≤ x ≤ 0.25 and −0.15 ≤ y ≤ 0.15. All give the same X-ray diffraction pattern as that of BaClF, although hardly with any significant line shift, hence no change of unit-cell parameters. The ambient temperature Mössbauer spectrum of a large number of samples was recorded, versus x and y. Figure 25 shows the influence of y at a fixed x value. It is observed that for negative y values (F/Cl > 1), a highly asymmetrical spectrum is obtained (Fig. 25a). When y increases to less negative values, i.e. there is a smaller excess of F, the asymmetry of the spectrum increases, and at positive y values (Cl/F > 1), only one line at ca. 4 mm/s is observed (Fig. 25b). In addition, the spectrum becomes weaker and weaker, such as nearly to escape detection at the most positive y values. Cooling the sample of Fig. 25b to 17 K results in a very dramatic increase of the intensity of the spectrum (Fig. 25c). These observations lead to the following model of the Ba₁₋ₓSnₓCl₁₊yF₁₋ₓ solid solution (Fig. 26):

(a) X-ray diffraction shows that the structure is the same as that of non-substituted BaClF (since it is the same diffraction pattern), with no further lattice distortions.

Fig. 21. Structural model of the disordered M/Sn/F phases: the tin lone pair points randomly toward any of the six faces of the [F₈] cube.

Fig. 22. X-ray powder diffraction pattern of: (a) unsubstituted BaClF, (b) precipitated Ba₁₋ₓSnₓCl₁₋ₓF₁₋ₓ solid solution. The arrows show the line shift relative to non-substituted BaClF.

Fig. 23. Ambient temperature ¹¹⁹Sn Mössbauer spectrum of precipitated (a) BaSn₂Cl₂F₄, (b) BaSnClF₃ · 0.8H₂O, (c) Ba₁₋ₓSnₓCl₁₋ₓF₁₋ₓ solid solution.
Fig. 24. Diagram of the Ba$_{1-x}$Sn$_x$Cl$_{1+y}$F$_{1-y}$ solid solution in the $(x,y)$ plane, obtained by reaction at 350°C for 43 h: (a) pure Ba$_{1-x}$Sn$_x$Cl$_{1+y}$F$_{1-y}$ phase, a mixture of Ba$_{1-x}$Sn$_x$Cl$_{1+y}$F$_{1-y}$ with other phases.

Fig. 25. $^{119}$Sn Mössbauer spectrum of Ba$_{1-x}$Sn$_x$Cl$_{1+y}$F$_{1-y}$ solid solution prepared by solid state reaction at 350°C for 43 h, $x = 0.225$ with (b) $y = 0.25$ at 298 K, (c) $y = 0.25$ at 17 K, and (a) $y = -0.15$ at 298 K. Spectra (b) and (c) were measured on the same sample, hence containing the same amount of tin, using the same γ-ray source and the same setup. Only the temperature was changed. It took 21 days to get spectrum (b) and only 2 days to get spectrum (c).

(c) At negative $y$ values ($F/Cl > 1$), there is more chance of forming covalent bonding like in SnF$_2$ (Fig. 2), in MSnF$_4$ (Fig. 1c) and in $M_{1-x}$Sn$_x$F$_2$ (Fig. 21), and in addition, at large $x$ values, there is a significant chance of a few tin atoms being neighbors and forming –F–Sn–F–Sn–F– oligomers since polymerization is quite common when tin binds to fluorine (Fig. 2). This gives rise to the model of Fig. 26b. The strength of the Sn–F bonds and the room taken by the stereoactive lone pair provides enough stability to prevent excess thermal vibrations and keep the recoil-free fraction at a value similar to that of SnF$_2$, hence a spectrum strong enough to be easily detected at ambient temperature. The doublet of Fig. 25a shows the presence of covalent bonding like the one proposed in the model of Fig. 26b. However, the asymmetry of the spectrum increases rapidly when $y$ increases, to give a single line characteristic of ionic tin. This can be explained only by the overlap of a quadrupole doublet and a single line, with the high velocity line of the doublet and the single line overlapping nearly perfectly. This shows that the Ba$_{1-x}$Sn$_x$Cl$_{1+y}$F$_{1-y}$ solid solution...
contains either only Sn$^{2+}$ (low $x$ and high $y$) or a mixture of covalently bonded tin(II) and Sn$^{2+}$, (high $x$ and low $y$) however all the tin is never present with covalent bonding. The presence of covalently bonded tin(II) violates the Hume–Rothery rule about the same bonding type, and the similar size rule is also grossly violated. The stereoactive lone pair compensates, at least in part, for the size difference.

(a) It has shown that an appropriate combination of X-ray powder diffraction and the Mössbauer spectroscopy results can be very helpful in understanding how tin(II) bonding is the most complex by far. It is doubly non-stoichiometric, for metals (Sn/Ba) and for the halogens (Cl/F). It contains tin(II) being either all ionic or a mixture of covalently bonded tin(II) and Sn$^{2+}$, depending on the amount of tin ($x$) and on the relative amounts of Cl and F ($y$). This solid solution is the first example of the presence of covalently bonded tin(II) and ionic Sn$^{2+}$ in the same compound, and moreover disordered with one another on the same site, and furthermore both are disordered with the much larger Ba$^{2+}$ ions.

(b) In the case of PbSn$_4$F$_{10}$ is highly unusual: despite the random replacement of 80% of the Pb$^{2+}$ in cubic coordination by covalently bonded tin forming softer bonding only on one side of each tin atom, the stereoactive lone pair being on the other side exerting repulsions on bonding pairs, the compound is stable for a time sufficiently long to be studied (from a few hours to a few days). It is the same as removing 80% of the bricks from all over a wall, at random, and the wall still stands up.

(c) In the case of the Ba$_{1-x}$Sn$_x$Cl$_{1+y}$F$_{1-y}$ solid solution the VSEPR model, hence stabilized by the lattice energy.

(d) The disordered fluorides contain covalently bonded tin(II) in the square pyramidal coordination and with the square pyramidal electron pair geometry, unfavored by the VSEPR model, hence stabilized by the lattice energy.

(e) The case of PbSn$_4$F$_{10}$ is highly unusual: despite the random replacement of 80% of the Pb$^{2+}$ in cubic coordination by covalently bonded tin forming softer bonding only on one side of each tin atom, the stereoactive lone pair being on the other side exerting repulsions on bonding pairs, the compound is stable for a time sufficiently long to be studied (from a few hours to a few days). It is the same as removing 80% of the bricks from all over a wall, at random, and the wall still stands up.

(f) The case of PbSn$_4$F$_{10}$ is highly unusual: despite the random replacement of 80% of the Pb$^{2+}$ in cubic coordination by covalently bonded tin forming softer bonding only on one side of each tin atom, the stereoactive lone pair being on the other side exerting repulsions on bonding pairs, the compound is stable for a time sufficiently long to be studied (from a few hours to a few days). It is the same as removing 80% of the bricks from all over a wall, at random, and the wall still stands up.

Fig. 26. Tin(II) bonding in Ba$_{1-x}$Sn$_x$Cl$_{1+y}$F$_{1-y}$: (a) ionic bonding in a solid solution dilute in tin ($x$ low) and rich in Cl ($y > 0$), (b) clusters of covalently bonded tin in systems richer in tin ($x$ high) and rich in F ($y < 0$). The dashed lines show weak interactions.

5. Conclusion

The present study has provided some major results concerning the materials containing divalent tin and that have a structure closely related to the fluorite-type structure.

(a) It has shown that an appropriate combination of X-ray powder diffraction and the Mössbauer spectroscopy results can be very helpful in understanding how tin(II) fits in the typically ionic fluorite structure.

(b) In the case of PbSnF$_4$, the very extreme case of preferred orientation made it possible to determine the orientation of the stereoactive lone pair from the variation of the asymmetry of the Mössbauer quadrupole doublet with the orientation of the sample in the γ-ray beam, in relation with the orientation of the unit-cell axes.

(c) In the case of disordered structures, fluorides or chloride fluorides, stoichiometric or non-stoichiometric, stable structures and metastable PbSn$_4$F$_{10}$ were formed despite blatant violations of the Hume–Rothery rules.

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