

Visualizing lone pairs in compounds containing heavier congeners of the carbon and nitrogen group elements

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Abstract. In this mini-review, I discuss some recent work on the stereochemistry and bonding of lone pairs of electrons in divalent compounds of the heavier carbon group elements (Sn^{II}, Pb^{II}) and in trivalent compounds of the heavier nitrogen group elements (Bi^{III}). Recently developed methods that permit the real-space visualization of bonding patterns on the basis of density functional calculations of electronic structure, reveal details of the nature of *s* electron lone pairs in compounds of the heavier main group elements – their stereochemistry and their inertness (or lack thereof). An examination of tetragonal *P4/nmm* SnO, α -PbO and BiOF, and cubic *Fm $\bar{3}m$* PbS provides a segue into perovskite phases of technological significance, including ferroelectric PbTiO₃ and antiferroelectric/piezoelectric PbZrO₃, in both of which the lone pairs on Pb atoms play a pivotal rôle.

Keywords. Main group elements; lone pairs; stereochemistry; ferroelectricity.

1. Introduction

The valence shell electron pair repulsion (VSEPR) theory, first suggested by Sidgwick and Powell¹, to be refined later by Gillespie and Nyholm², is the stuff of high school chemistry. The sequence of decreasing repulsion, lone pair–lone pair > lone pair–bond pair > bond pair–bond pair, is a familiar thumb rule that permits thousands of school and college students to hazard the geometry of molecules such as XeO₃. While it is usual to assume that these thumb rules have a basis in hard quantum chemical theory, in reality, even the experimental structures of these simple molecules are sometimes a matter of controversy.

Regarding the disposition of lone pairs in extended solids, Sten Andersson and coworkers have in a series of papers³, come up with modifications of VSEPR theory that permit a better prediction of coordination polyhedra. The new rules are described in the *magnum opus* of Hyde and Andersson⁴ and include, crucially, the definition of a typical atom–lone pair distance, and the notion that in the space of the unit cell, the lone pair occupies as much volume as an oxide or fluoride anion.

At the same time, the question of what constitutes a chemical bond has received renewed attention, with key contributions from Richard Bader in his Atoms In Molecules approach⁵. The electron localization function (ELF), of Becke and Edgecombe⁶ measures the extent of the Pauli repulsion, i.e. the conditional probability of finding an electron in the vicinity of another electron of like spin, and has seen much use in the demarcation of the space of a molecule or solid into regions which are bonding, non-bonding or antibonding. Such analysis has been further extended by the work of Silvi and

Savin⁷ who in effect, have coupled the topological approach of Bader with the evaluation of ELF.

In this contribution, I present some recent attempts to understand structure in compounds that contain stereochemically active lone pairs of *s* electrons. The basis for this is density functional calculations of the electronic structure followed by ELF visualizations. It is found that the anion and the degree of covalency are important for the manner in which the lone pairs dispose themselves. These studies also throw light on so-called ‘inert pair’ effects⁸ and support the view that these effects have rather complex origins that necessitate an analysis beyond the usual atomistic interpretations.

The outline of the article is as follows. First, I perform density functional electronic structure calculations as implemented in the scalar relativistic linear muffin–tin orbital method⁹, on some typical lone pair compounds including SnO, *a*PbO and BiOF. This permits plots of the ELF to be made, which when plotted in the real space of the crystal structure, permits the lone pair to be visualized, even in structures where it does not form a lobe – an example of such a structure being cubic PbS. I then consider the structures of more complex materials, namely the perovskites PbTiO₃ and PbZrO₃, and inquire into the rôle of the lone pair in driving (respectively) the ferroelectric and antiferroelectric phase transitions.

2. SnO, *a*PbO, BiOF, PbS

SnO and *a*PbO adopt a tetragonal structure⁴ with double slabs of the cation which sandwich O, each four-coordinate with the other and stacked along the *c* axis. In BiOF, the BiO network forms similar double slabs except that there are F[−] ions arranged in the interslab space above and below the cations. Interaction across the slabs is important for the semi-metallicity of SnO and for the transformation of *a*PbO to a black γ phase under pressure, before it transforms to the red massicot phase¹⁰.

The structures of these three compounds are respectively displayed in figures 1a through 1c. The large gray spheres are the cations and the orange spheres are O. In BiOF, fluorines have been represented by cyan spheres in the interslab region. In the panels of figure 1, the beige lobes in the vicinity of the cations in the inter-slab space are the lone pairs, visualized as isosurfaces with an ELF value of 0.85, corresponding to rather strong localization. In a plane midway through the structure, the ELF is presented as a map, following the colour coding indicated in the bar at the bottom. Regions of poor localization (nearly free electrons) are indicated by deep blue and regions of strong localization (corresponding to an ELF value of one) are indicated by white.

First, this figure illustrates that it is now routinely possible to visualize in the real space of the crystal structure, the disposition of lone pairs from *reliable* first-principles calculations. This obviates the need to position lone pairs by guess work. Second, rather subtle features emerge, that cannot *possibly* be guessed. For example, one observes that there is some interaction between the lone pairs and O within the slabs revealed (particularly for SnO) as a lightening in the colour in the appropriate region of the map. Additionally, the lone pairs in PbO are more compact when compared with SnO, in keeping with the empirical findings of Andersson and coworkers⁴, that the lone pair–atom distance is larger for Sn^{II} than it is for Pb^{II}. In conjunction with the fact that the Sn–O system is much more covalent¹⁰, the more compact lone pair on Pb reveals the reluctance of Pb^{II} toward oxidation to Pb^{IV}. This in fact is a manifestation of the inert pair effect – the propensity of the post-lanthanoid main-group elements to retain a valence

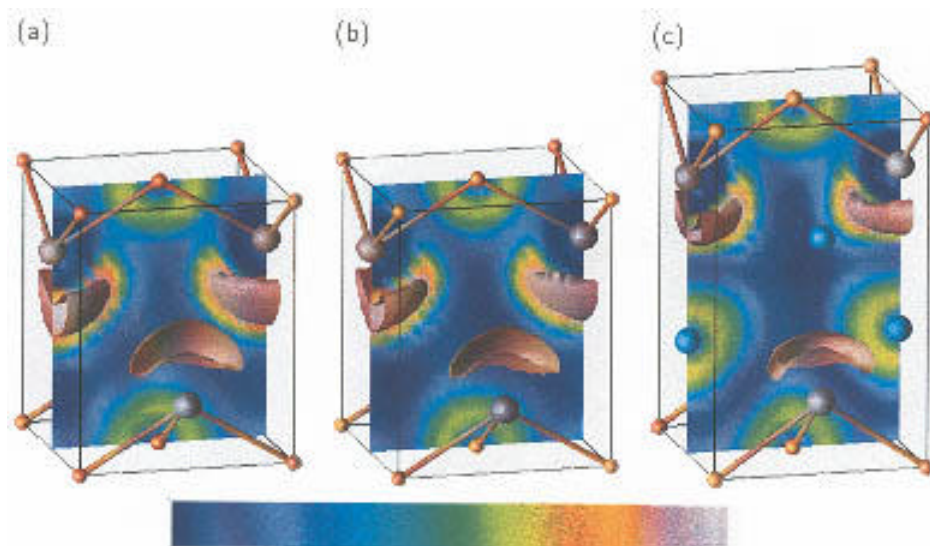


Figure 1. Crystal structures of (a) SnO, (b) α -PbO, and (c) BiOF. Gray spheres are cations, orange O and cyan F. In the space of the structures, isosurfaces of the electron localization function (ELF) with a value of 0.85 reveal the lone pairs as lobes near the cations. The map halfway through the structure is the ELF, represented by colour coding following the bar at the base of the figure. The deep blue end corresponds to poor localization and the white end to strong localization.

that is less than the group valence by two electrons. It is known that such a reluctance to attain the group valence cannot be explained merely from trends of ionization energies⁸. Electron localization in BiOF visualized in figure 1c suggests a much more ionic system with little or no interaction between F and the lone pairs, and this is confirmed by an analysis of the densities of state.

The lone pairs are expected to have primarily 5s (Sn) or 6s (Pb, Bi) character. If they are to form lobes, there must be some *p* admixture. The question is whether the *p* character arises from mixing *s* and *p* on the cation or whether it is the effect of covalency, arising from cation *s*, anion *p* mixing. Watson and Parker¹¹ have recently suggested that it is due to *p* character from the anions that lone pairs are allowed to form lobe-shaped structures. The anion is clearly implicated since in a homologous series of compounds, changing the anion often greatly affects the stereochemistry of the lone pair. In the case of PbO, the density of Pb *s* and Pb *p* electronic states are well-separated (figure 2a) in energy, while there seem to be *p* states of O in the region of energy where *s* states of Pb are found (figure 2b). Crystal Orbital Hamiltonian Populations (COHPs) permit the decomposition of densities of state into specific pairwise interactions between atoms¹². Positive COHPs correspond to bonding interactions and negative to antibonding. The Pb–O COHP for α -PbO is plotted in figure 2c. It is seen that the coincidence of O *p* and Pb *s* densities of state at an energy of –8 eV (referenced to the top of the valence band) does indeed correspond to strongly covalent interactions.

If lone pair *s* must overlap with anion *p* levels in order to attain the typical lobe shape, then conversely, in a situation where there is no such overlap, or when the overlap is

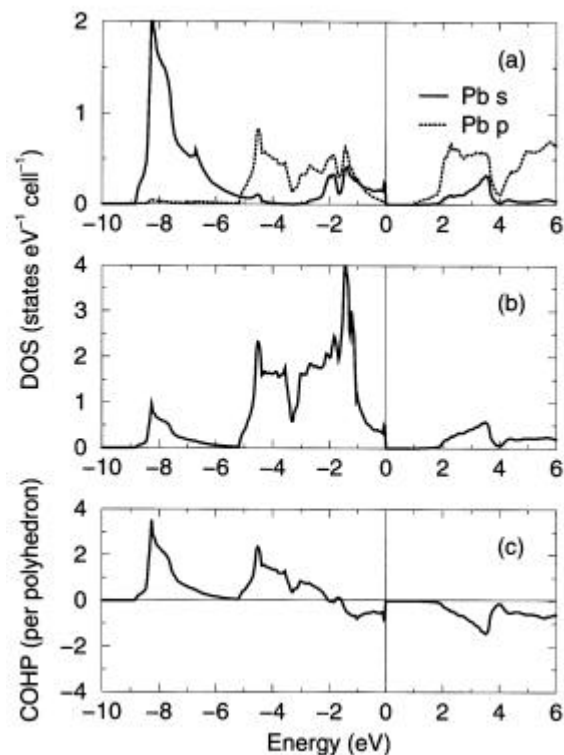


Figure 2. (a) Orbital projected density of Pb *s* and *p* states in *a*-PbO. (b) Density of O *p* states in *a*-PbO. (c) Crystal orbital Hamiltonian population (COHP) of Pb–O bonding in *a*-PbO. Positive values correspond to bonding interactions and negative values to antibonding interactions. The origin on the energy axis is the top of the valence band.

weak, the lone pair could retain its *s* character and remain spherical. PbS is a strongly covalent system in terms of Pb-*p* S-*p* overlap, but (i) the *p* levels of S are not as deep in energy as O *p* to bond strongly with Pb *s* and (ii) the intrinsically longer Pb–S distance (due to larger atomic radii) may lessen overlap. The densities of state and the COHP of cubic (NaCl) PbS are displayed in figure 3 to support this view. It is seen that while there is some overlap of Pb *s* with anion *p* states, it is much weaker than what is seen in *a*-PbO. One octant of the rock-salt structure of PbS is displayed in figure 4. The yellow spheres are S and the gray spheres Pb. A map of the ELF is displayed at one end of the structure. In order to visualize the lone pairs around Pb, it was necessary to select an isosurface with an ELF value of 0.6 signifying rather poor localization of the lone pair. At this value, spheres of localization around S atoms are also seen. As anticipated from the discussion, the lone pairs around Pb are nearly spherical in shape.

3. PbTiO₃

On cooling below 766 K, PbTiO₃ undergoes a displacive phase transition (with a textbook soft mode) from an ideal cubic perovskite (*Pm* $\bar{3}$ *m*) to a ferroelectric tetragonal

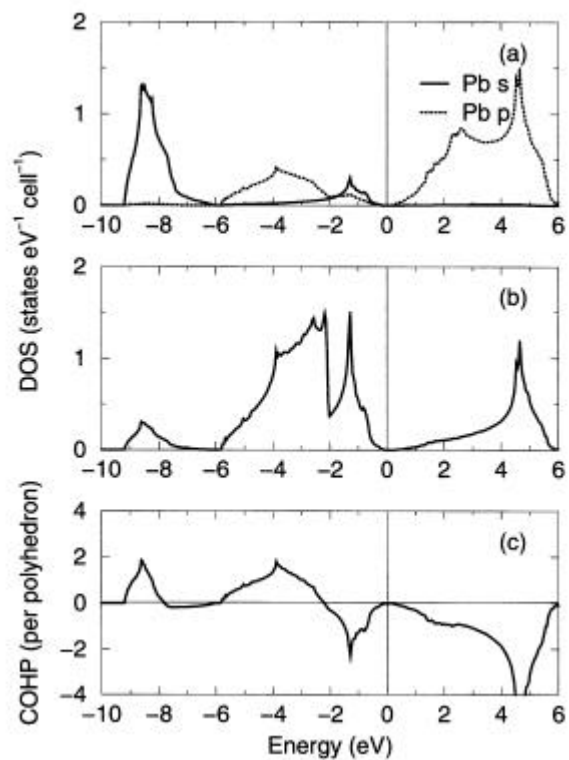


Figure 3. (a) Orbital projected density of Pb *s* and *p* states in PbS. (b) Density of S *p* states in PbS. (c) COHP of Pb–S bonding in PbS.

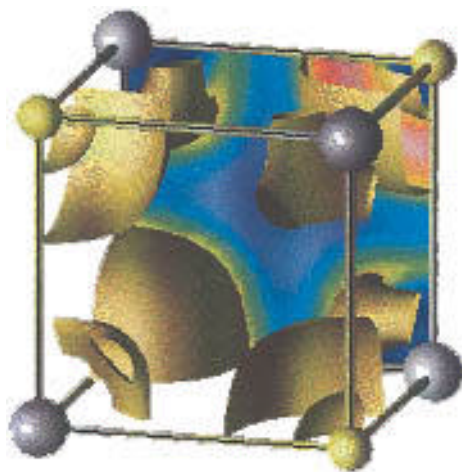


Figure 4. One octant of the rock salt structure of PbS. Gray spheres are Pb and yellow spheres are S. A map of the ELF is displayed at one end of the octant. An ELF isosurface corresponding to a value of 0.6 is displayed, and indicates localization around both the cation and the anion.

perovskite with the space group $P4mm$ ¹³. There are two aspects of the $PbTiO_3$ structure that need to be considered. The first is the Goldschmidt tolerance factor which is calculated (in the cubic phase) to be about 1.03. This means that the distance in the cubic phase between Pb and O is a little too small, and one expects a tendency for the structure to undergo a distortion that would in effect permit a larger volume for Pb. Secondly, d^0 transition metal ions (particularly Ti^{IV} , V^V and Cr^{VI}) display a strong tendency to so-called second order Jahn–Teller distortions¹⁴ that result in off-centric coordination polyhedra around these ions. These two factors cooperate in obtaining the rather high Curie temperature of $PbTiO_3$. By the same token, the tolerance factor of $BaTiO_3$ is 1.07, and one would perhaps expect it to possess an even higher Curie temperature. In fact, the Curie temperature (again going from a cubic to a tetragonal phase) of $BaTiO_3$ is only 393 K¹³. The distinct differences between $PbTiO_3$ and $BaTiO_3$ have been suggested as arising from the involvement of Pb states in bonding near the Fermi energy¹⁵.

Figure 5 displays the structure of cubic $PbTiO_3$ ¹⁶ with gray Pb atoms in the corners of the cube and octahedral Ti (black) in the centre. Midway through the structure, a map of the ELF (following the usual colour code) is displayed. As in the case of PbS, the lone pair (visualized at an ELF value of 0.65) forms a nearly spherical object around Pb. In the tetragonal phase (two unit cells shown in figure 6) it is seen that the distortion of the TiO_6 octahedra act in concert with the localization of the lone pairs on Pb into lobes (visualized at an ELF value of 0.7). The manner in which the lone pairs dispose themselves suggests a greater tendency to the ferroelectric distortion and a reason for the higher Curie temperature.

In figure 7(a and b) I compare the densities of state in cubic (displayed as solid lines) and tetragonal (displayed as dashed lines) $PbTiO_3$, focusing on Pb s and p states in panel (a) and O p states in panel (b). The separation of Pb s and Pb p in both the cubic and tetragonal phases is similar to what is seen in α -PbO, except perhaps that the Pb s states are less disperse. Reflecting the smaller dispersion of the Pb s states are correspondingly fewer O p states in this region of energy (figure 7b), -8 eV with reference to the Fermi energy. Once again, the COHPs (figure 7c) reveal details that are not evident in the DOS.

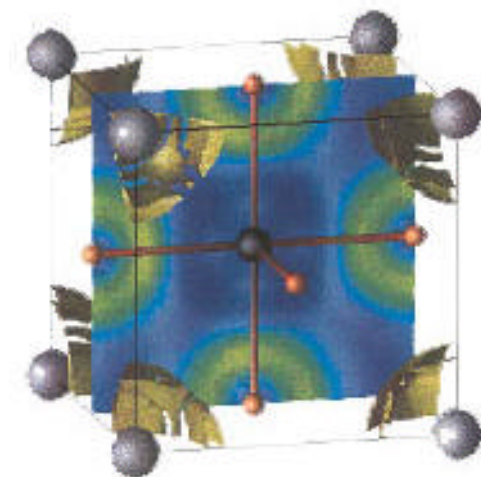


Figure 5. The structure of cubic $PbTiO_3$. Gray spheres are Pb, black Ti and orange O. The isosurface is at an ELF value of 0.65 and permits the nearly spherical (octahedral, in a stricter sense) lone pair to be visualized.

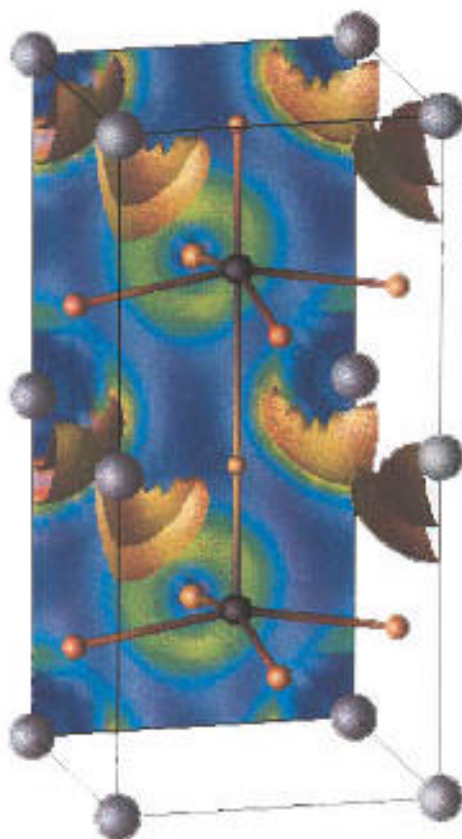


Figure 6. The structure of tetragonal, ferroelectric PbTiO_3 . Two unit cells are displayed. The lone pairs are visualized at an ELF isosurface of 0.7. If the frame of reference is the O in the plane perpendicular to c , both Pb and Ti are observed to be displaced in the same direction, due respectively to the lone pair and the second-order Jahn–Teller distortion.

It is seen that while Pb s –O p bonding is not very strong, it is at least as important as the Pb p –O p interaction. An interesting change across the phase transition is that the slightly stronger bonding between Pb and O in the tetragonal phase in the energy range corresponding to the lone pair, is compensated for by the development of antibonding states just below the Fermi energy. These antibonding states are missing in the cubic compound. Once again, the formation of the lone pair lobe is favoured by overlap between Pb s and O p .

4. PbZrO_3

Zr^{IV} is significantly larger than Ti^{IV} and as a result, the tolerance factor of PbZrO_3 is about 0.97. A tolerance factor that is less than 1 suggests that (unlike the case of PbTiO_3) Pb is in a coordination polyhedron that is a little too large. If the octahedra tilt in a concerted manner, it is possible to slightly squeeze the polyhedron around Pb in order to achieve a more comfortable Pb–O bond distance. Being a d^0 transition metal ion, Zr^{IV}

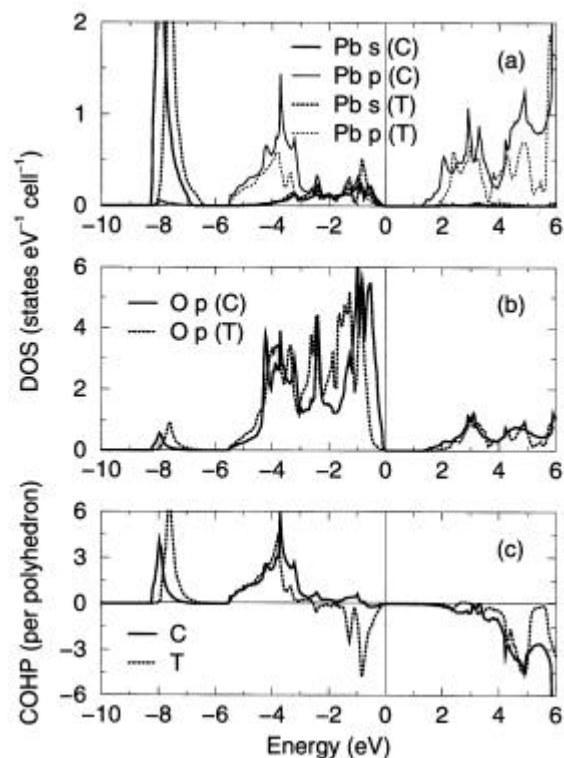


Figure 7. (a) Pb *s* and *p* densities of state in the cubic (C) and tetragonal (T) structures of PbTiO₃. (b) O *p* states in the cubic (C) and tetragonal (T) structures. Pb–O COHP in the cubic (C) and tetragonal (T) structures.

would like to undergo an off-centric distortion within its oxygen octahedron though the tendency is reduced when compared with Ti^{IV}¹⁴. Finally, the Pb^{II} ions on the A site of the structure possess a tendency to move away from the centre of their coordination polyhedron in order to permit the lone pair to take on the shape of a lobe. The two overwhelming driving forces for the phase transition in PbZrO₃ at 503 K turn out to be the tendency of the ZrO₆ octahedra to tilt in a cooperative manner (driven by a tolerance factor less than 1) and the tendency of the Pb atom lone pairs to localize into lobes. As a result of the Pb atoms displacement and the tilting not being cooperative, the resulting structure is antiferroelectric rather than ferroelectric¹³. Waghmare and Rabe¹⁷ have been able to reproduce the rather complex experimental structure in a completely *ab-initio* manner on the basis of density functional theory.

Electronic structure calculations using the crystal structure of orthorhombic PbZrO₃ determined by Glazer and coworkers¹⁸ once again permit the disposition of Pb lone pairs within the space of the crystal structure to be visualized. Figure 8 displays the crystal structure with gray spheres representing Pb atoms, black Zr and orange O. Isosurfaces of the ELF with a value of 0.7 permit reveal the lobe-like lone pairs possessed by the Pb atoms. These lone pairs are seen to arrange themselves in an antiferrodistortive manner resulting in the antiferroelectric properties of this orthorhombic phase.

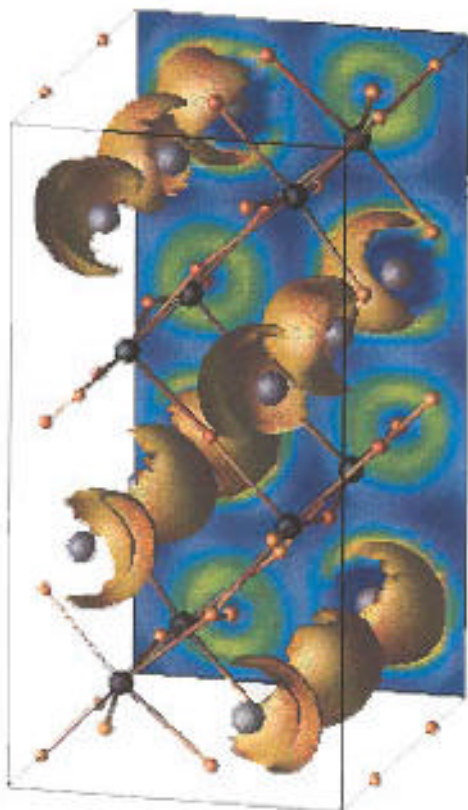


Figure 8. The structure of orthorhombic, antiferroelectric PbZrO_3 . The lone pairs are visualized at an ELF isosurface of 0.7. The displacements of the Pb atoms are such that their effects cancel out suggesting an antiferrodistortive structure.

5. Prospects

In brief, I have attempted to show that the stereochemistry of lone pairs in structures containing the heavier congeners of the carbon and nitrogen groups can be visualized on the basis of reliable density functional calculations of the electronic structure. Such visualization obviates the necessity to guess the role and disposition of lone pairs in determining structure. What is interesting is that even complex pyroelectric systems such as PbTiO_3 and PbZrO_3 turn out to be amenable to such analysis. The visualization of the lone pairs suggests reasons for the high Curie temperature of PbTiO_3 and also suggests why PbZrO_3 is antiferroelectric. Elsewhere, it has been suggested on the basis of such visualizations that lone pairs on Pb atoms are important in driving the cubic–orthorhombic transition in Pb_2MgWO_6 ¹⁹. In the magnetic, possibly multiferroic BiMnO_3 ²⁰, the lone pairs on Bi are crucial for stabilizing ferromagnetism and in driving the off-centric distortion²¹.

An important aspect of the present work is the identification of conditions corresponding to when lone pairs localize and to when they remain spherical or nearly so. An interesting system in which these factors are manifest are the nine binary

chalcogenides of the divalent carbon group elements, including GeS, GeSe ... SnSe, ... PbTe. If these are represented as a matrix of compositions, the upper left diagonal is unstable in the cubic rock-salt structure. We are currently examining how lone pairs help sort this structural field.

Acknowledgements

I thank O K Andersen and O Jepsen for generously making their LMTO codes available. I have greatly profited from discussions and continuing collaboration on the stereochemistry of lone pairs with G Baldinozzi, C Felser, N A Hill and U Waghmare. I am particularly grateful to J Gopalakrishnan for sharing his insights into crystal structures of compounds with d^0 elements, and to N A Hill for critically reading this manuscript. The Laboratoire SPMS and the École Centrale Paris are acknowledged for their kindness in inviting me for two months in the summer of 2001, when this article was written. This work has been supported by the Council of Scientific and Industrial Research, New Delhi.

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