THE ELECTRONIC STRUCTURE OF THE MIXED VALENCE COMPOUND Pb₃O₄

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Abstract—Ab initio self-consistent calculations of the electronic structure of Pb₃O₄ are presented. The calculations show that Pb₃O₄ is a semiconductor. The calculated bandgap of 1.1 eV is smaller than the observed gap of 2.1–2.2 eV. The calculations show strong hybridization between Pb(6s) and O(2p) states.

For one type of lead atom, this leads to a distribution of Pb(6s) states over two occupied energy bands, similar to the situation in PbO. For the other type of lead atom in Pb₃O₄, the Pb(6s) states are distributed over an occupied and an unoccupied band, similar to the situation in β-PbO₂. This clearly demonstrates that Pb₃O₄ is a mixed valence compound, i.e. Pb(II)₂Pb(IV)O₄. According to the calculations, the conductivity of p-type Pb₃O₄ is due to holes in a Pb(II)(6s) valence band and the conductivity in n-type Pb₃O₄ is due to electrons in a Pb(IV)(6s) conduction band. © 1997 Elsevier Science Ltd. All rights reserved

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1. INTRODUCTION

Trilead tetroxide Pb₃O₄ is a bright orange-red compound which occurs naturally as the mineral minium. Pb₂O₃ was historically used as a paint and the word miniature for illustrations in old manuscripts is derived from minium. More recently, Pb₂O₄ has been used in batteries and ceramics and was also widely used in corrosion resistant oil-based paints. With iron, the oxidizer Pb₃O₄ forms a layer of (Pb, Fe) oxides, which do not oxidize further. For environmental reasons Pb₃O₄ is now mostly replaced by other corrosion resistant compounds.

Although Pb₂O₄ is technologically a very important material, little is known about its physical properties. Pb₂O₄ is a semiconductor with a high resistivity, ρ ~ 10⁶ Ω·cm for a pressed powder [1] and ρ ~ 10⁴ Ω·cm for a Pb₂O₄ film [2]. The infrared spectra show the presence of Pb–O vibrations between 300 and 700 cm⁻¹ [3] and for the dielectric constant, values between 13 and 17 have been reported [4, 5]. The bandgap was measured by reflectance and photovoltage technique [6–11]; the spectra show an absorption edge, indicating a bandgap of 2.1–2.2 eV. Benschop [12, 13] measured nuclear magnetic resonance spectra of lead oxides; for Pb₂O₄ two signals were observed, one similar to the signal observed for PbO and one similar to the signal observed for the insulator SrPbO₃.

Several authors have reported photoelectron spectra of Pb₂O₄. Kim et al. [14] deconvoluted the Pb 4f peaks into two peaks with an intensity ratio of 1:2, with the most intense peak at the highest binding energy and suggested that the two peaks are due to the two types of Pb atoms in Pb₂O₄. In other studies [15, 16] no doublet structure for the Pb core levels was observed. According to Morgan and Van Wazer [16], the binding energies for the Pb 4f core levels of PbO, PbO₂ and Pb₂O₄ are all the same. Thomas and Tricker [15] reported differences, with the lowest binding energies found for β-PbO₂ (142.0 and 137.3 eV for Pb(4f₅/₂ and Pb(4f₇/₂), respectively) and for the tetragonal and orthorhombic modifications of PbO, values of 143.3 eV for Pb(4f₅/₂) and 138.6 eV for Pb(4f₇/₂) were found. For Pb₂O₄, intermediate values of 142.3 and 137.5 eV were found. We conclude from these data that there is no clear evidence for a splitting of Pb(4f) core levels in Pb₂O₄ as a result of the presence of two types of Pb atoms.

At room temperature Pb₂O₄ crystallizes in a tetragonal structure with space group P4₁/mmc-D₄h [17]. At lower temperatures Pb₂O₄ undergoes a sequence of ferroelastic and ferroelectric phase transitions [18–21]. The low temperature phase, stable below 170 K, has an orthorhombic structure with space group Pbam-D₂h, which is a subgroup of the space group of the high temperature phase. The orthorhombic distortion decreases with increasing temperature and has values b/a = 0.927 at 5 K and b/a = 0.968 at 140 K. At 170 K the low temperature structure transforms into a noncentrosymmetric pseudo-quadratic structure with symmetry 2mm; the orthorhombic distortion of this phase is very small, with a value b/a = 0.998 at 180 K. At
195 K there is a transition to a non-centrosymmetric tetragonal phase with symmetry 4mm. Finally, this phase transforms into the centrosymmetric tetragonal room temperature structure at 225 K. In all these phases, the atomic positions are close to the positions in the high temperature tetragonal phase; the phase transitions correspond to small shifts of the atoms.

Two types of lead atoms with different coordination can be distinguished in the room temperature tetragonal structure. One type is coordinated by six oxygen atoms in the form of a slightly distorted octahedron. The other type of atom is coordinated in a very asymmetric way by an irregular pyramid of four oxygen atoms, with the lead atom at the vertex. The coordinations are similar to those observed in the oxides PbO, with Pb in a pyramidal coordination of four oxygen atoms and PbO₂, with Pb in an octahedral coordination. This correspondence is traditionally interpreted as (indirect) evidence for the mixed valence character of Pb₃O₄. Therefore, the octahedrally coordinated Pb atoms in Pb₃O₄ are lead atoms with a (formal) valency of IV, like in PbO₂. The lead atoms coordinated by the pyramid have a formal valency of II, like in PbO. The formula for Pb₃O₄ can thus be written as Pb(II)₂Pb(IV)O₄.

In this paper we present ab-initio calculations of the electronic structure of Pb₃O₄ using the augmented spherical wave (ASW) method. The results are compared with electronic structure calculations for β-PbO and β-PbO₂. This comparison shows that the electronic structures of the two types of lead atoms in Pb₃O₄ correspond very closely to the electronic structures of lead atoms in β-PbO and β-PbO₂. This is direct evidence that Pb₃O₄ is indeed a mixed valence compound.

2. CRYSTAL STRUCTURE

The room temperature phase of Pb₃O₄ has a tetragonal crystal structure with space group P4₁/mmb-D₁₆, with Z = 4 formula units in the crystallographic unit cell [17]. The unit cell axes at 293 K are a = 0.8811 nm and c = 0.6563 nm. One type of lead atom, with formal valency Pb(IV), occupies the 4d positions with coordinates \((x, y, z; z, z, \frac{1}{2})\). The other type with formal valency Pb(II), occupies the 8h positions with coordinates \((x, y, z; z, z, \frac{1}{2})\), with \(x = 0.140\) and \(y = 0.163\). The oxygen atoms occupy the 8g position with coordinates \((x, x + \frac{1}{2}, \frac{1}{2}; x, x + \frac{1}{2}, \frac{1}{2})\), with \(x = 0.671\) and the 8h position with \(x = 0.096\) and \(y = 0.637\).

The structure consists of chains of PbO₆ octahedra which share common edges and which extend in the direction of the c-axis (see Fig. 1). Additional Pb(II) atoms form bridges between the chains of octahedra. Each Pb(II) atom is close to one oxygen in an equatorial edge and to two other oxygen atoms not in

![Fig. 1. Structure of Pb₃O₄ projected on the ab plane. The small circles represent the Pb(IV) atoms, and the hatched circles without and with dots Pb(II) atoms at \(z = 0\) and \(z = \frac{1}{2}\), respectively. The large open circles are apical oxygen atoms at \(z = \frac{1}{4}\) and \(z = \frac{3}{4}\) and the double circles are equatorial oxygen atoms at \(z = 0\) and \(z = \frac{1}{2}\).](image)

## Table 1. Interatomic distances (in nm) in Pb₃O₄, β-PbO₂ and β-PbO

<table>
<thead>
<tr>
<th>Distance type</th>
<th>Pb₃O₄</th>
<th>β-PbO₂</th>
<th>β-PbO</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pb(IV)–O</td>
<td>0.220 (x4)</td>
<td>0.2169 (x4)</td>
<td></td>
</tr>
<tr>
<td>Pb(IV)</td>
<td>0.213 (x2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>average Pb(IV)–O</td>
<td>0.218</td>
<td>0.2162</td>
<td></td>
</tr>
<tr>
<td>Pb(IV)–Pb(IV)</td>
<td>0.3282</td>
<td>0.3387</td>
<td></td>
</tr>
<tr>
<td>O–O</td>
<td>0.288</td>
<td>0.2711</td>
<td>0.2944</td>
</tr>
<tr>
<td>Pb(II)–O</td>
<td>0.2215 (x2)</td>
<td>0.2221</td>
<td></td>
</tr>
<tr>
<td>Pb(II)</td>
<td>0.2337</td>
<td>0.2249</td>
<td></td>
</tr>
<tr>
<td>average Pb(II)–O</td>
<td>0.2374</td>
<td>0.2358</td>
<td></td>
</tr>
<tr>
<td>Pb(II)–Pb(II)</td>
<td>0.379</td>
<td>0.3536</td>
<td></td>
</tr>
</tbody>
</table>
The electronic structure of Pb$_2$O$_4$

Table 2. Input parameters for the band-structure calculation of Pb$_3$O$_4$

<table>
<thead>
<tr>
<th>Atom</th>
<th>Start configuration</th>
<th>$R_{WS}$ (pm)</th>
<th>Atomic position</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pb(IV)</td>
<td>[Xe]4f$^{14}$5d$^{10}$6s$^2$6p$^2$</td>
<td>152.4</td>
<td>4d</td>
</tr>
<tr>
<td>Pb(II)</td>
<td>[Xe]4f$^{14}$5d$^{10}$6s$^2$6p$^2$</td>
<td>158.7</td>
<td>8h ($x = 0.140$, $y = 0.163$)</td>
</tr>
<tr>
<td>O1</td>
<td>[He]2s$^2$2p$^6$</td>
<td>105.3</td>
<td>8g ($x = 0.671$)</td>
</tr>
<tr>
<td>O2</td>
<td>[He]2s$^2$2p$^6$</td>
<td>104.2</td>
<td>8h ($x = 0.096$, $y = 0.637$)</td>
</tr>
<tr>
<td>ES1</td>
<td>1s$^2$2p$^6$</td>
<td>152.4</td>
<td>4b</td>
</tr>
<tr>
<td>ES2</td>
<td>1s$^2$2p$^6$</td>
<td>174.4</td>
<td>8g ($x = 0.181$)</td>
</tr>
<tr>
<td>ES3</td>
<td>1s$^2$2p$^6$</td>
<td>153.9</td>
<td>8h ($x = 0.663$, $y = 0.626$)</td>
</tr>
</tbody>
</table>

Equatorial edges. These latter apical oxygen atoms are in adjacent octahedra of the same chain. The equatorial oxygen forms part of a neighbouring chain and the Pb(II) atoms form the border of large channels parallel to the c-axis.

In Table 1 the interatomic distances in Pb$_3$O$_4$ are compared with the distances in $\beta$-PbO [22] and $\beta$-PbO$_2$ [23]. The average Pb-O distances for Pb(II) and Pb(IV) in Pb$_3$O$_4$ are only 1% larger than in PbO and PbO$_2$, respectively. However, the distortions of the Pb(IV)O$_6$ octahedra and the tetragonal Pb(II)O$_4$ pyramids in Pb$_3$O$_4$ is much larger than in the simple oxides.

3. BAND STRUCTURE CALCULATIONS

Self consistent band structure calculations were performed using the augmented spherical wave (ASW) method [24]. Exchange and correlation were treated within the local density approximation [25]. Scalar relativistic effects were included as described by Methfessel and Kübler [26]. To take into account the spin orbit interaction, an extra term $\lambda L S$ was incorporated in the Hamiltonian.

In the ASW method the crystal is subdivided into Wigner–Seitz spheres surrounding the atoms. Within each sphere the potential is taken spherically symmetric; this so-called muffin–tin approximation has proven to give reliable results for close-packed structures. Due to the presence of the asymmetrically coordinated Pb(II) atoms and the presence of large channels, the structure is far from being close packed. A better approximation of the true crystal potential of more open structures can be obtained by filling the unoccupied space with so-called empty spheres (ES). For the calculations on Pb$_3$O$_4$, three types of ES were included. The positions and Wigner–Seitz radii ($R_{WS}$) of the ES are listed in Table 2, together with the Wigner–Seitz radii of the atoms. The empty sphere at the 4b position, ES1, is situated in the middle of the channels and the second type of empty sphere at the 8g position is located inside the PbO$_4$ pyramids. The Pb(IV) atoms are only surrounded by two empty spheres with a distance Pb(IV)-ES2 = 0.225 nm (2x). Each Pb(II) is coordinated by seven empty spheres at the following distances: Pb-ES1 = 0.2505 nm (2x), Pb-ES2 = 0.2283 nm (2x) and Pb-ES3 = 0.2690, 0.2554 and 0.2543 nm. This large coordination of Pb(II) by empty spheres will have implications when the charges on the atoms are considered; a large part of electronic charge in the empty spheres should be assigned to Pb(II) atoms.

For the ASW calculations we used a basis set consisting of 6s, 6p and 6d functions for Pb and 2s and 2p functions for O and 1s and 2p functions for the empty spheres. Pb 5f and O 3d functions were included in the internal summation of the three center contributions to the matrix elements, which can be interpreted as taking these functions as a perturbation. The choice of basis functions implies that the effect of hybridization of Pb 5d with O 2s is not taken into account. This hybridization is quite strong because the orbital energies of Pb 5d and O 2s are nearly the same, but the effect of Pb 5d/O 2s hybridization on the valence and conduction bands and the chemical bonding is small [27].

The Brillouin zone of the tetragonal room temperature phase of Pb$_3$O$_4$ is shown in Fig. 2. The calculated energy bands along symmetry lines are presented in Fig. 3; we have used the symmetry notations of Miller

![Fig. 2. Brillouin zone of tetragonal Pb$_3$O$_4$.](image-url)
Fig. 3. Energy bands of Pb₃O₄ calculated with the ASW method, including spin-orbit interaction.

Fig. 4. Total and partial density of states of Pb₃O₄. Units: total DOS: states/(Ry.unit cell); partial DOS: states/(Ry.atom).

and Love [28]. Figure 4 shows the total and partial density of states and in both Figs 3 and 4, the top of the valence band has been taken as the energy zero. The charges in the muffin–tin spheres and the orbital configuration of the atoms and empty spheres are listed in Table 3. Generally, not too much significance should be attributed to the charges calculated in this way, because the values of the charge within the Wigner–Seitz sphere depend strongly on the values used for the Wigner–Seitz radii and on the presence of empty spheres. The charge within the Wigner–Seitz sphere cannot be identified with the ionic charge. This is clearly demonstrated by the remarkable fact that the positive charge in the Wigner–Seitz sphere of Pb(II) is larger than that of Pb(IV). The reason is that a considerable amount of electronic charge, located in the empty spheres surrounding Pb(II), should be assigned to the atomic charge of Pb(II).

The valence band region consists of 56 energy bands and has a width of 9.0 eV. The maximum of the valence bands is at M, but this maximum is only 0.06 eV higher in energy than the maximum at Γ. The lowest four bands, ranging from −8 eV to the bottom of the valence bands, mainly consist of 6s states of Pb(IV) mixed with O 2p states (Fig. 4). The next eight bands between −6 and −8 eV have mainly Pb(II) 6s character with some hybridization with O 2p states. Four of these bands show a large dispersion of about 2 eV in the Γ–Z direction. The next 36 bands between −6 and −2 eV are the (nearly non-bonding) O 2p states, mixed slightly with Pb(IV) 6p states; these bands have a small dispersion throughout the whole Brillouin zone. The top eight bands, with a total width of 2.3 eV, are predominantly Pb(II) 6s in character, mixed with O 2p states. The hybridization of Pb(IV) 6s and O 2p orbitals is very strong, resulting in a distribution of Pb(IV) states over two energy bands at −9 eV and +2 eV, i.e. separated by 11 eV. Such a very strong Pb(IV) 6s – O 2p hybridization is also present in β-PbO₂ [29] and BaPbO₃ [30]. Hybridization of Pb(II) 6s with O 2p orbitals is also strong and gives rise to bands between −8 and −5 eV and −2 and 0 eV.

The bottom of the conduction band is at Γ. According to our calculations Pb₃O₄ is a semiconductor with an indirect energy gap of 1.1 eV. The
calculated direct energy gap is at $r$, with a value of 1.2 eV. The lowest conduction band consists of mainly Pb(IV) 6s character, with 0 2p states. This conduction band is narrow, and has a total width of 1.3 eV.

The large contribution of Pb(IV) 6p orbitals to the charge density of the valence bands (see Fig. 4 and Table 3) indicates that the bonds between Pb(IV) and oxygen are strongly covalent. The divalent lead atoms and oxygen atoms form bonds which are more ionic in nature.

4. DISCUSSION

The only electronic structure calculation for Pb$_3$O$_4$ reported in literature, was performed by Evarestov and Veryazov [31], using the semi-empirical LUC-CNDO method. These authors give only the total density of states and the eigenvalues at $r$ and their results differ strongly from our calculations. The only electronic structure calculation for Pb$_3$O$_4$ using an electrostatic model. The Pb(II) atoms are highly polarizable and the asymmetric coordination by oxygen ions induces a large dipole moment on Pb(II).

According to our calculations, Pb$_3$O$_4$ is a semiconductor with a band gap of 1.1 eV. This value is smaller than the experimentally observed band gap of 2.1 - 2.2 eV. This is a common defect of many electronic structure calculations using the LD approximation.

The top of the valence band of Pb$_3$O$_4$ consists mainly of Pb(II)(6s) states, the bottom of the conduction band of Pb(IV)(6s) states. Therefore, conduction in p-type semiconducting Pb$_3$O$_4$ will be due to holes in the Pb(II)6s band, which corresponds to Pb$^{2+}$ ions on the Pb(II) sites. Conduction in n-type Pb$_3$O$_4$ is due to electrons in the Pb(IV) 6s band, corresponding to Pb$^{3+}$ ions on the Pb(IV) sites. Thus optical excitation of an electron from the top of the valence band to the bottom of the conduction band corresponds to an inter-valence charge transfer transition Pb$^{3+}$ + Pb$^{2+}$. (We remark that this is a strongly simplified picture because of the strong hybridization of Pb(6s) and O(2p) states.)