X-Ray Photoelectron Spectroscopy Study on The Double Layer at an Al₂O₃-Al Interface

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X-ray photoelectron spectroscopy study on the electrical double layer at an $\text{Al}_2\text{O}_3$-Al interface

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ABSTRACT

Upon oxidation of a clean Al surface, an electrical double layer (EDL) is formed at the Al-$\text{Al}_2\text{O}_3$ interface. This EDL is investigated using XPS data available in the literature. The EDL strength, measured as a potential difference across the EDL, depends on the Al surface and the oxidation process. The polarity of the EDL is however invariably the same: the $\text{Al}_2\text{O}_3$ side of the Al-$\text{Al}_2\text{O}_3$ interface is always positively charged. The reduction of the Al work function upon oxidation is attributed to this EDL. The asymmetry in the potential barrier shape formed in Al-$\text{Al}_2\text{O}_3$-Al sandwiches is also attributed to a strong EDL at the electrode-film interface and a weaker EDL at the interface between the counter electrode and the film.
I. INTRODUCTION

When a pure aluminum metal is oxidized in dry O₂ at room temperature, an amorphous Al₂O₃ film of about 30 Å is formed on its surface [1]. At the Al-Al₂O₃ interface, as well as at any other phase boundaries, an electrical double layer (EDL) is expected to develop [2]. This EDL produces a potential difference (PD) across the interface and modifies the work function of Al. However, measurement of the Al work function with an oxidized Al surface generally involves not only the Al-Al₂O₃ interface but also the Al₂O₃-vacuum interface. In this paper, we use XPS (X-ray photoelectron spectroscopy) to investigate the PD produced across an Al-Al₂O₃ interface. Our XPS study, unlike the work-function measurements, can separate out the effects of the Al-Al₂O₃ interface from those of the Al₂O₃-vacuum interface.

The Al-Al₂O₃ system is chosen, simply because rather accurate XPS data are available for the aluminum-oxygen compounds [1,4]. We first present our model and later verify it with experimental results available in the literature.

II. THEORETICAL BACKGROUND

A. General features of XPS for oxidized Al

In XPS measurement on an oxidized Al metal, an electron-energy analyzer generally faces the oxide surface as shown in Fig. 1. The oxide (Al₂O₃) film is generally so thin (~10Å) that electrons originating from the metal portion, as well as those from the oxide portion, can reach the analyzer. Consequently, two peaks appear both in the Al 2p photoelectron spectrum and in the Al KLL Auger spectrum. These two peaks arise from Al atoms located in the oxide and metal portions of the sample. One can easily identify the origin (metal or oxide) of these peaks based on the expected chemical shifts for Al (AlO) and Al₂O₃ (Al⁺³); and on the dependence of the relative intensity of the two peaks with change in the Al₂O₃-film thickness. Figure 2 shows two such peaks in the Al 2p photoelectron spectra. The Al⁺³ binding energies in the figure are aligned for comparison.

We denote the electron kinetic energies of the two peaks in the Al KLL Auger spectrum as KE(Auger,AlO) and KE(Auger,Al⁺³) and those in the Al 2p photoelectron spectrum as KE(photo,AlO) and KE(photo,Al⁺³). KE(Auger) denotes the energy of the most intense KLL₂₂₃ transition for an Al atom and hence does not depend on the incident X-ray energy. On the other hand, KE(photo) depends on the incident X-ray energy. However, the binding energies of the Al 2p electrons,

\[ KE(Al^0) = h\nu - KE(photo,Al^0) \]  
\[ KE(Al^{+3}) = h\nu - KE(photo,Al^{+3}) \]

are independent of the incident X-ray energy, h\nu. In short, XPS measurements for an oxidized Al metal provide four physically significant energies: KE(Auger,AlO), KE(Auger,Al⁺³),
BE(Al\(^0\)) and BE(Al\(^{+3}\)). We employ, however, an alternative set of four independent energies defined by

\[
\begin{align*}
E(Al^0) & = KE(Auger,Al^0) + BE(Al^0) \\
E(Al^{+3}) & = KE(Auger,Al^{+3}) + BE(Al^{+3}) \\
E(M/O) & = BE(Al^{+3}) - BE(Al^0) \\
E(O/V) & = BE(Al^{+3}).
\end{align*}
\]

These energy parameters are more useful because, except for 
E(O/V), they eliminate the static-charge referencing problem. 
E(Al\(^{+3}\)) and E(Al\(^0\)) are nothing but the modified Auger parameters (MAP) introduced by Wagner [3] and are independent of the interfacial potentials. E(M/O) depends on only the metal-oxide interfacial potentials, as explained below.

Electrons originating from the metal portion must cross the M/O (metal-oxide) and O/V (oxide-vacuum) interfaces to reach the electron-energy analyzer (see Fig. 1), and consequently experience the potential differences (PD) across these interfaces denoted as PD(M/O) and PD(O/V). These PDs enter in both KE(Auger,Al\(^0\)) and BE(Al\(^0\)), since their electrons originate from the metal portion (Al\(^0\)). However, because of Eq. (1), the PDs in KE(Auger,Al\(^0\)) and BE(Al\(^0\)) enter with opposite signs. Therefore, a sum of these energies, that is E(Al\(^0\)), is independent of both PD(M/O) and PD(O/V).

Electrons originating from the oxide portion (Al\(^{+3}\)), on the other hand, cross only the O/V interface on the way to the electron-energy analyzer and hence experience only PD(O/V).

This PD(O/V) also enters in \(E(\text{Auger},\text{Al}^{+3})\) and BE(Al\(^{+3}\)) with opposite signs, so that E(Al\(^{+3}\)) is similarly independent of both PD(M/O) and PD(O/V).

E(M/O) is defined as an energy difference between BE(Al\(^{+3}\)) and BE(Al\(^0\)). Since PD(O/V) enters in BE(Al\(^{+3}\)) and BE(Al\(^0\)) with the same sign, the PD(O/V) contributions cancel in E(M/O) and consequently E(M/O) does not depend on PD(O/V). In other words, E(M/O) depends only on PD(M/O). The PD(M/O) is expected to be stable and reproducible, while PD(O/V) is harder to obtain reproducibly because of adventitious surface contamination.

B. Theoretical model

Let us first consider an imaginary Al\(_2\)O\(_3\)-Al system. Suppose that a thin γ-Al\(_2\)O\(_3\) film is made separately and then brought into contact with a clean Al surface. In this case, electron migration across the Al\(_2\)O\(_3\)-Al interface will not take place, since the Fermi level of the Al metal is situated inside the forbidden band gap of the insulator Al\(_2\)O\(_3\), far away (more than 2 eV) from the top of the valence band (VB) and the bottom of the conduction band (CD). The electron energy diagram for this Al\(_2\)O\(_3\)-Al system is nothing but a series of energy diagrams for isolated γ-Al\(_2\)O\(_3\) and element Al, where the vacuum level between them is eliminated, as shown in Fig. 3a. Note that, in the figure, the vacuum level (VL) connected to Al\(_2\)O\(_3\) is equal to that connected to Al. We use this imaginary Al\(_2\)O\(_3\)-Al system as a reference system.
Now, we consider a more realistic Al₂O₃-Al system: A thin Al₂O₃ film is grown on a clean Al surface by exposing it to dry O₂ at room temperature. It is shown that the oxide film formed in this way is an amorphous γ-Al₂O₃ and almost homogeneous throughout except at the phase boundaries [5]. The main difference between this Al₂O₃-Al system and the reference system described above is therefore the phase boundaries. It is quite reasonable that the asymmetric forces, which are always present at phase boundaries, are adjusted in the process of the Al₂O₃ film formation. Adjustment at a phase boundary normally results in the formation of EDL at the phase boundary and hence a PD across the phase boundary [2]. Consequently all of the electronic energy levels of Al₂O₃ shift uniformly in comparison with those levels of Al₂O₃ in the reference system (a), as shown in Fig. 3b where the uniform shift is denoted as PD(M/O). The work function of Al metal also changes by PD(M/O) + PD(O/V), where PD(O/V) is due to the EDL at the O/V interface.

XPS measurements on our reference Al₂O₃-Al system would give E(M/O) as BE(γ-Al₂O₃) - BE(element Al), where BE(γ-Al₂O₃) and BE(element Al) are the Al 2p binding energies for the isolated γ-Al₂O₃ compound and isolated elemental Al, respectively. On the other hand, XPS measurements on the Al₂O₃-A sample represented by Fig. 3b will yield the E(M/O) energy parameter to be

\[ E(M/O) = BE(\gamma-Al_2O_3) - BE(element Al) - PD(M/O). \]  

provided that the XPS chemical states of the Al₂O₃ film and the Al substrate are still those of the isolated γ-Al₂O₃ and element Al.

from this equation, we obtain

\[ PD(M/O) = BE(\gamma-Al_2O_3) - BE(element Al) - E(M/O). \]  

The sign of PD(M/O) is chosen to be negative when the electronic levels of Al₂O₃ are lowered relative to the Fermi level of Al. The EDL at the M/O interface shown in Fig. 3b therefore gives a negative PD(M/O) and the EDL at the O/V interface gives a positive PD(O/V).

The interfaces of an Al-Al₂O₃-Al sandwich are closely related to those of the previous two Al₂O₃-Al systems. In a sandwich, an ultra-thin Al₂O₃ film is grown on an aluminum metal (Al') by exposing its surface to O₂ and then another aluminum metal (Al'') is deposited on the already grown Al₂O₃ film. Therefore, the Al'-Al₂O₃ interface will be almost identical to the Al-Al₂O₃ interface represented by Fig. 3b, while the Al₂O₃-Al'' interface will be similar to the Al₂O₃-Al interface of the reference system (Fig. 3a). Figure 4a shows a simplified electron-potential-energy diagram for the Al'-Al₂O₃-Al'' sandwich. In this diagram, the effect of the EDL at the Al₂O₃-Al'' interface is neglected in comparison with that at the Al'-Al₂O₃ interface. All electron energy levels of Al₂O₃ and Al'' and also the vacuum level connected to the Al'' metal are lowered uniformly by PD(M/O) due to the EDL at the Al'-Al₂O₃ interface. When the Al' and Al'' electrodes of the sandwich (Fig. 4a) are short-circuited, electrons flow from Al' to Al''
until the Fermi levels of these two metals coincide with each other. As a result, positive charge is left behind at the Al'-
Al₂O₃ interface and excess electrons are built up on the Al⁺
surface at the Al⁺-Al₂O₃ interface. This pair of positive and
negative charges produces a trapezoidal potential barrier
between the two Al metals, as shown in Fig. 4b. Thus, the
asymmetry of the potential barrier can originate from a strong
double layer at the Al⁺-Al₂O₃ interface. In short, there is a
close connection between the potential barrier shape of the Al-
Al₂O₃-Al sandwich and the EDL strengths at the two phase
boundaries.

III. EXPERIMENTAL RESULTS

A. Modified Auger parameters $E(Al^{+3})$ and $E(Al^{0})$

Wagner conducted a comprehensive survey on Auger and
photoelectron energies published up to 1982 and compiled these
energies and also the modified Auger parameters in Ref. [3].
The claimed accuracy of the data for aluminum-oxygen compounds
is around ± 0.1 eV [4]. Table I lists the modified Auger
parameters (MAP) obtained from Ref. [3,4] for Al metal and
Al₂O₃ oxides. For easier comparison, the MAP minus 1460 eV are
listed.

Oxidized Al, which is denoted as Al₂O₃/Al in Table I,
exhibits two MAPs, since it has two (metal and oxide) phases.
The $E(Al^{0})$ of the oxidized Al agrees with the MAPs of pure Al
(new Table I), indicating that the metal portion of the
oxidized Al was not altered significantly by oxidation as far
as XPS can detect. Further, the $E(Al^{+3})$ of oxidized Al is
equal to the MAP of pure $γ$-Al₂O₃, which differs substantially
from the MAPs of other Al₂O₃ compounds, as seen in Table I.
This confirms that an oxide film on Al is indeed a $γ$-like
alumina as found previously [5]. Furthermore, this agreement
also confirms that $E(Al^{+3})$ and $E(Al^{0})$ are indeed independent of
the interfacial potentials.

B. $E(\text{M/O})$ and $P(\text{M/O})$

Table II lists some of the observed $E(\text{M/O})$, which is
defined as the energy difference between the two Al 2p peaks
(see Eq. 5). Fig. 2 exhibits these two peaks in the Al 2p
spectra for the first and the last data entries in Table II.
We can see clearly the difference. Also included in the figure
is the Al 2p binding energy of the isolated Al metal relative
to that of the isolated $γ$-Al₂O₃ compound. The Al⁺ 3 binding
energies are aligned for comparison. The $E(\text{M/O})$ values appear
to depend on the oxidation conditions. Later we will show that
$E(\text{M/O})$ also depends on the Al surface indices (i.e., surface on
which the Al₂O₃ film grows). For the XPS measurements
referenced in Table II, the Al surfaces were not well
characterized. To separate out these two effects, we need more
systematic measurements with well defined Al surfaces and
oxidation conditions.
The $E(M/O)$ value depends not only on $P(M/O)$ but also on the XPS chemical states of the $Al_2O_3$ film and the Al metal. Changes in the XPS chemical states can be monitored by measuring the MAPs. But unfortunately no such monitoring has been carried out except for the first case in Table II, which shows that the XPS chemical states of the $Al_2O_3$ films and the Al substrate are those of pure $\gamma-Al_2O_3$ and elemental Al as detected by XPS. Furthermore it is known that the anodic films formed on Al are also $\gamma$-like $Al_2O_3$ [5]. Based on these facts, we speculate that the XPS chemical state changes of the $Al_2O_3$ and the Al substrate contribute less than 0.5 eV to the $E(M/O)$ value, which is the largest XPS chemical state change observed among the $Al_2O_3$ compounds (see Table I).

The $PD(M/O)$'s in Table II are calculated from Eq. (8) with $BE(\gamma-Al_2O_3)-BE(\text{element Al})=0.8$ eV, which are evaluated with the data from Ref. [1]. Here we have assumed that the XPS chemical states of the $Al_2O_3$ film and the Al substrate are the same as those of $\gamma-Al_2O_3$ and elemental Al. The error in $PD(M/O)$ due to this assumption could be as large as 0.5 eV as discussed in the previous paragraph. The sign of $PD(M/O)$ is, however, invariably negative for all cases in Table II. This means that the polarity of the EDL is always the polarity shown in Fig. 3b, namely, an excess positive charge on the $Al_2O_3$ surface and the induced counter electron charge on the Al surface.

C. Magnitude of $PD(O/V)$

Now, we attempt to estimate the magnitude of $PD(O/V)$ due to an EDL at the O/V interface. The difference between $BE(\alpha-Al^{+1})$ of the $Al_2O_3$ film and $BE(\gamma-Al_2O_3)$ of the isolated $\gamma$-$Al_2O_3$ should give $PD(O/V)$, provided that the following conditions are met: (a) the static-charge-referencing problem for these two measurements do not introduce errors and (b) not only the XPS chemical state but also the surface conditions of the $Al_2O_3$ are the same as those of the isolated $\gamma-Al_2O_3$.

Assuming that the data cited in Ref. [3] satisfy these conditions, we obtain $PD(O/V)=0.4$ eV. Although the error in $PD(O/V)$ could be as large as ±0.5 eV, we may conclude that the magnitude of $PD(O/V)$ is substantially smaller than that of $PD(M/O)$.

D. Work function change

The work function changes ($\Delta\phi$) of the three Al surfaces ($\{111\}$, $\{110\}$, and $\{100\}$) upon oxidation with a high oxygen exposure (>400 Langmuir) are about -0.2, -0.8, and -1.3 eV respectively [9,10]. In our model, the work function change ($\Delta\phi$) of the oxidized Al is given by

$$\Delta\phi = PD(M/O) + PD(O/V),$$

as can be seen in Fig. 3b. From XPS data, we have deduced a large negative value for $PD(M/O)$ and a small value for $PD(O/V)$. Therefore we expect a negative $\Delta\phi$ for all cases agreeing with the above observations. Conversely, we may state that the work function change is mainly due to the formation of an EDL at the
Al-Al$_2$O$_3$ interface. This statement implies that the EDL strength depends on the Al surface on which the Al$_2$O$_3$ film grows. This is, we think, quite reasonable.

E. Potential barrier shape for Al-Al$_2$O$_3$-Al sandwich

The composition of the Al-Al$_2$O$_3$-Al sandwich suggests a symmetric potential barrier shape. However, experiments consistently show an asymmetric potential barrier of approximately the trapezoidal barrier shape as shown in Fig. 4b [11-13]. The barrier height $\phi_1$ at the Al'-Al$_2$O$_3$ interface is always found to be lower than the barrier height $\phi_2$ at the Al$_2$O$_3$-Al" interface [11-13], as indicated in Fig. 4b. The observed value for $\phi_2-\phi_1$, however, varies from 0.4 eV [11] to 1.9 eV [12] and other measurements [13] fall between these two values. In our model described in Section II.B, we have

$$\phi_2 - \phi_1 = -PD(M/O)$$ (10)

 neglecting the effect of the weaker EDL at the Al$_2$O$_3$-Al" interface. Since PD(M/O) was always found to be negative from the analysis of the XPS data, we expect $\phi_2-\phi_1>0$ for all cases as observed. Conversely, the main cause of the asymmetric barrier shape for Al-Al$_2$O$_3$-Al sandwiches can be attributed to a strong EDL formed at the Al-Al$_2$O$_3$ interface by oxidation.

VI. SUMMARY

XPS data, work function changes, and potential barrier shapes all consistently reveal that a strong EDL is formed at the Al-Al$_2$O$_3$ interface upon oxidation of a clean Al surface. The EDL strength depends on the Al surface indices and the oxidation conditions. The polarity of the EDL is always found as such: a positively charged Al$_2$O$_3$ interface surface and the counter electrons in the Al metal.

In a larger sense, this work has demonstrated for the first time that XPS can be used to investigate the interfacial potential induced between an ultra-thin film and its substrate. The photoelectron-binding-energy (or Auger-line-energy) difference of a selected atom of the film and another selected atom of the substrate directly depends on this interfacial potential. If the modified Auger parameters of the film and the substrate are unchanged, a change in the energy difference equals a change in the interfacial potential. The advantage of utilizing these (Photoelectron or Auger) energy differences is that it is independent of the surface conditions of the thin film.

ACKNOWLEDGMENTS

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REFERENCES


FIGURE CAPTIONS

FIG. 1. Configuration of sample and electron detector for typical XPS measurements on oxidized Al. The electron detector faces the oxidized surface of Al. As a result, electrons originating from the metal portion must cross the two (M/O and O/V) interfaces in order to enter the detector, while those from the oxide portion cross only the O/V interface on their way to the detector.

FIG. 2. Al 2p photoelectron spectra for air-oxidized Al and anodically oxidized Al obtained from Ref. [6]. The Al$^0$ and Al$^{1+}$ peaks have been attributed to electrons originating from the metal (Al$^0$) and the oxide (Al$^{1+}$) portions of the samples. Also included is the Al 2p binding energy of isolated Al metal relative to that of isolated Al$_2$O$_3$(γ), which are taken from Ref. [3]. The Al$^{1+}$ binding energies are aligned for comparison.

FIG. 3. Simplified electron potential energy diagrams for two distinct Al$_2$O$_3$-Al systems: For (a), the Al$_2$O$_3$ film is made separately and then brought into contact with a clean Al surface; the Al$_2$O$_3$ film for (b) is formed on the Al surface by exposing to O$_2$ gas. Only in case (b) does an electrical double layer develop at the phase boundaries, especially strongly at the M/O boundary.
FIG. 4. Simplified electron potential energy diagrams for an Al-Al$_2$O$_3$-Al sandwich. The Al$_2$O$_3$ film is grown on an aluminum substrate Al' by exposing to O$_2$ gas, and then another aluminum Al" film is deposited on the already grown Al$_2$O$_3$ film. The Al' and Al" electrodes of the sandwich is open in (a), but shortcircuited in (b).

TABLE I. The modified Auger parameters (MAP) obtained from Ref. [7] for Al metal and Al$_2$O$_3$ compounds. For easier comparison, the MAP (in eV) minus 1460 eV are listed.

<table>
<thead>
<tr>
<th>Compound</th>
<th>E(Al$^{+3}$)-1460eV</th>
<th>E(Al$^{0}$)-1460eV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al$_2$O$_3$/Al</td>
<td>1.6</td>
<td>6.1</td>
</tr>
<tr>
<td>Al$_2$O$_3$/Al</td>
<td>1.6</td>
<td>6.2</td>
</tr>
<tr>
<td>Al$_2$O$_3$ (γ)</td>
<td>1.6</td>
<td>---</td>
</tr>
<tr>
<td>Al$_2$O$_3$ (α)</td>
<td>2.1</td>
<td>---</td>
</tr>
<tr>
<td>Al$_2$O$_3$ sapphire</td>
<td>2.0</td>
<td>---</td>
</tr>
<tr>
<td>Al$_2$O$_3$ sapphire$^b$</td>
<td>2.0</td>
<td>---</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>1.9</td>
<td>---</td>
</tr>
<tr>
<td>Al</td>
<td>---</td>
<td>6.1</td>
</tr>
<tr>
<td>Al</td>
<td>---</td>
<td>6.1</td>
</tr>
<tr>
<td>Al</td>
<td>---</td>
<td>6.2</td>
</tr>
</tbody>
</table>

$^a$Calculated from the MAP difference and the E(Al$^{+3}$) value.
$^b$Heated in vacuum at 450°C.
TABLE II. Observed $E(M/O)$ values for $\text{Al}_2\text{O}_3$ films grown on an Al surface under various oxidation conditions.

<table>
<thead>
<tr>
<th>Oxidation conditions</th>
<th>$E(M/O)$ [in eV]</th>
<th>$\Delta E(M/O)^a$ [in eV]</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air, at room temperature</td>
<td>2.7</td>
<td>1.9</td>
<td>[6]</td>
</tr>
<tr>
<td>$\text{O}_2$, at room temp.</td>
<td>2.7</td>
<td>1.9</td>
<td>[1]</td>
</tr>
<tr>
<td>$\text{O}_2$, at 700°C for 16 min.</td>
<td>3.2</td>
<td>2.4</td>
<td>[8]</td>
</tr>
<tr>
<td>$\text{O}_2$, at 700°C for 42 min.</td>
<td>3.6</td>
<td>2.8</td>
<td>[8]</td>
</tr>
<tr>
<td>Anodic</td>
<td>3.4</td>
<td>2.6</td>
<td>[7]</td>
</tr>
<tr>
<td>Anodic, in $\text{Na}_2\text{SO}_4$</td>
<td>4.0</td>
<td>3.2</td>
<td>[6]</td>
</tr>
</tbody>
</table>

*a) Calculated from Eq. (8) with $\text{BE}(\gamma-\text{Al}_2\text{O}_3) - \text{BE}(\text{pure Al}) = 0.8$ eV obtained from Ref. [3].
Fig. 2

Fig. 3
Fig. 4

**Fig. 4**

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