XPS studies of the charging effects for the non-conducting rare-earth oxides

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X-ray induced photoemission studies (XPS) of the bulk oxides of the heavy rare-carth elements were made using a Vacuum Generator's ESCA 3 instrument equipped with a X-Y plotter and Mg K_x and Al K_x radiation sources – Charging effects on the core-level binding mergies of Tb₄O₇, Dy₂O₃, Ho₂O₃, Er₂O₃, Tm₂O₃, Yb₂O₃ and Lu₂O₃ are determined. It is found that the charging of the sample is dependent on the sample thickness and also on the radiation source used to probe the sample. The d-arging of the sample is estimated with respect to $4f_{7/2}$ peak for clean gold metal at S3-7 eV. The pxygen 1*S* and carbon 1*S* peaks were also monitored. These results on bulk oxides are compared with the surface oxides reported by the authors elsowhere

1 INTRODUCTION

X-ray induced photoelectron spectroscopy is a poworful tool for studying the binding energies in solids (Siegbahn *et al* 1967). For non-conducting solids, a knowledge of charging effects is necessary for obtaining accurate values of binding energies (Johansson *et al* 1973) Recontly, we reported the core-electron binding energies for the lanthanides (Lang *et al* 1974, 1975, Padalia *et al* (1976, 1977)) In this paper, we present the results of our measurements of the charging effects on the resonance of the term.

2 EXPERIMENTAL

Photoemission spectra of the heavy rare earth oxides were measured using a Vacuum Generators ESCA 3 machine equipped with AlK_{α} and MgK_{α} radiations as exciting sources. Spectroscopically pure oxide powders supplied by Johnson-Mattheys Ltd; England were mounted onto the sample-probe using double-sided scotch-tape. The magnitude of charging was estimated by imbedding a piece of gold wire in the powder and recording the gold 4*f* peaks by X-Yplotter Details of the experimental method and the equipment used in this work were described in our previous communications (Lang *et al*1975a, 1975b, Padalia *et al* (1976, 1977)

B. D. Padalia and W. C. Lang

3. RESULTS AND DISCUSSION

Examples of the 4d photoemission spectra of the sesquioxides of lutetium, ytterbium and holmium are shown in figures 1-3. The binding energies expressed in eV are referred to the measured Fermi level for the corresponding pure metals. The N_{γ} (4f_{7/2}) peak for gold imbedded in terbium oxido powder together with carbon and oxygon 1s signals are presented in figure 4 Photoemission spectra of a blank sellotape and the N_{γ} peak for a piece of gold fixed on this sellotape are given in figure 6. Spectra recorded using AlK_z and MgK_z radiations are labelled with the alphabets a and b, respectively. Charging of the samples are estimated with respect to the 4f_{7/2} peak for clean gold at 83.7 eV. The values of the charging for the heavy rare-earth oxides are given in table 1.

Table 1 Charging of the samples estimated with respect to $4f_{7/2}$ peak for clean gold metal at 83.7 eV

Samples	(cV)	ь (өV)
Lu ₂ O ₃	2.0	2.0
Yb_2O_3	2.1	1.9
Tm ₂ O ₃	2.8	2.0
Er_2O_3	2.1	1.5
Ho_2O_3	2.8	1.1
Dy_2O_3	3.6	1.2
Դb₄Oγ	3.1	1.9
Sellotape (blank)	0.9	1.0

a-AlK_a radiation b-MgK_a radiation.

A scrutiny of the data given in table 1, reveals that the magnitudes of charging, in general are higher for AlK_x than those obtained for MgK_x . This indicates a direct dependence of charging on the penetration power of the radiation used to probe the sample. It is, however not clear why the charging for MgK_x should be more compared to AlK_x in the case of blank sellotape. The measured electron energies for the oxides (figures 1-3) when corrected for the charging effects (table 1), give the actual value of the binding energies. As an example, take the case of lutetium oxide (figure 1). The corrected values of the binding energies for $4d_{5/2}$ and $4d_{3/2}$ are 196.7 eV and 206.4 eV, respectively. The estimated error in the measurement of these energies is found to be ± 0.4 eV. The 4d spectra of the remaining heavy rare earth oxides (figures 2 and 3) are complex because those exhibit multiplet structure (Lang et al 1975).

In our earlier work, it was reported that the evaporated surface oxides of rare earths show no indication of the charging effects (Padalia *et al* (in pross)). It is interesting to note that the binding energies of the $4d_{5/2}$ and $4d_{3/2}$ reported



Fig. 1 4d Photoemission spectrum of lutetium oxide (Lu₂O₃).



Fig. 2. 4d Photoomission spectrum of ytterbium oxide (Yb₂O₃).

for the surface oxide of lutetium are in close agreement with those determined for the bulk oxide of lutetium after applying corrections due to charging. The effect of film thickness on charging were detected by us previously for the evaporated bulk oxide of rare earths. These samples were prepared by



Fig. 3 1d Photoemission spectrum of holmium oxide (Ho₂O₃)



Fig. 4. Carbon and exygen 1s peaks for terbium exide (Tb_4O_7) . $N_7(4f_{7/2})$ peak for gold imbedded in Tb_4O_7 . Curves labelled with a and b corresponding to AlK_a and MgK_a sources, respectively.

evaporating the metal in the presence of oxygon inside a sample preparation chamber maintained at a pressure of 10⁻⁴ mm of Hg – It was found that the magnitude of charging increases with mereasing thickness of the oxide film. In the present case of bulk oxides which were mounted on a selfetape, it is difficult to investigate quantitatively the dependence of charging on the sample thickness. However, the recorded spectra indicate changes in the magnitude of charging with change in the thickness of the sample, using the same radiation as exciting source



Fig. 5 Carbon and oxygon le signals for blank sellotape. N₇ peak for a piece of gold fixed on the sellotape. The curves labelled with a and b corresponding to AlK₂ and MgK₄ sources, respectively.

The charging effect is also displayed by the photoennission speatra of the oxygen and carbon present in the samples. Since carbon and oxygen spectra exhibit the presence of different species, these cannot be taken as standards for calibration. This fact has also been realized by others in recent years. The gold N_7 peak at 83.7 eV binding energy, however continues to be a reliable standard for calibration. This fact is further supported by the present results.

4. CONCLUSIONS

The measured binding energies for the non-conducting materials must be corrected for the charging effects. The charging of the sample is dependent on the radiation source used to probe the sample and also on the sample thickness. The N_7 peak at binding energy 83.7 eV is recommended as a standard for calibration.

B. D. Padalia and W. C. Lang

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