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REPORT NO. NADC-86028-60

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SEMICONDUCTOR/INSULATOR FILMS FOR CORROSION PROTECTION

F. C. Jain
The University of Connecticut
Electrical Engrg. & Computer Science, U-157
Storrs, Connecticut 06268

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OCTOBER 1985

FINAL REPORT
NADC Contract N62269-83-M-3295

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Prepared for
NAVAL AIR DEVELOPMENT CENTER
Warminster, PA 18974-5000

86 0 0 041

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REPORT DOCUMENTATION PAGE

1a REPORT SECURITY CLASSIFICATION Unclassified		1b RESTRICTIVE MARKINGS	
2a SECURITY CLASSIFICATION AUTHORITY		3 DISTRIBUTION AVAILABILITY OF REPORT Approved for Public Release; Distribution is Unlimited	
4 PERFORMING ORGANIZATION REPORT NUMBER(S) NADC-86028-60		5 MONITORING ORGANIZATION REPORT NUMBER(S) N/A	
6a NAME OF PERFORMING ORGANIZATION Naval Air Development Center		6b OFFICE SYMBOL <i>(if applicable)</i>	7a NAME OF MONITORING ORGANIZATION N/A
6c ADDRESS (City, State, and ZIP Code) Warminster, PA 18974		7b ADDRESS (City, State, and ZIP Code) N/A	
8a NAME OF FUNDING SPONSORING ORGANIZATION Naval Air Systems Command		8b OFFICE SYMBOL <i>(if applicable)</i>	9 PROCUREMENT INSTRUMENT IDENTIFICATION NUMBER Contract No. N62269-83-M-3295
8c ADDRESS (City, State, and ZIP Code) Washington, D.C. 20361		10 SOURCE OF FUNDING NUMBERS	
		PROGRAM ELEMENT NO	PROJECT NO
		TASK NO	WORK UNIT ACCESSION NO
11 TITLE (Include Security Classification) Semiconductor/Insulator Films for Corrosion Protection			
12 PERSONAL AUTHOR(S) F.C. Jain			
13a TYPE OF REPORT Final	13b TIME COVERED FROM _____ TO _____	14 DATE OF REPORT (Year, Month, Day) 1985 October	15 PAGE COUNT
16 SUPPLEMENTARY NOTES			
17 COSATI CODES		18 SUBJECT TERMS (Continue on reverse if necessary and identify by block number)	
FIELD	GROUP	SUB GROUP	
19 ABSTRACT (Continue on reverse if necessary and identify by block number)			
<p>- Aluminum surfaces exhibit significantly improved corrosion protection when they are coated with suitable semiconductor/insulator thin films. These coatings, generally realized in Metal-Semiconductor (MS) or Metal-Insulator-Semiconductor (MIS) structural configurations, given rise to an interfacial electric field which acts as an effective built-in electronic barrier. This active barrier significantly impedes the transfer of electrons from the aluminum surface to foreign species which cause oxidation by accepting the electrons. Anodic polarization data on numerous samples fabricated in both MS [e.g. Al-Indium Tin Oxide (ITO)] and MIS [e.g. Al-SiO₂-ITO] configurations have demonstrated the protective nature of the built-in active electronic barrier. In particular, we have observed a rest potential of -1.126 volts (1% NaCl, 2pH solution) for Al-SiO₂-pITO samples in polarization tests. The electronic barrier heights have been shown to increase with: (1) the presence of a thin (20-100 Å) SiO₂ layer at the metal-semiconductor interface; and (2) the energy gap of ITO which depends upon the indium content. A comparison of these results with data obtained on plasma-deposited Al-Si₃N₄ samples is also presented.</p>			
20 DISTRIBUTION AVAILABILITY OF ABSTRACT <input type="checkbox"/> UNCLASSIFIED UNLIMITED <input checked="" type="checkbox"/> SAME AS RPT <input type="checkbox"/> DTIC USERS		21 ABSTRACT SECURITY CLASSIFICATION Unclassified	
22a NAME OF RESPONSIBLE INDIVIDUAL V.S. Agarwala		22b TELEPHONE (Include Area Code)	22c OFFICE SYMBOL

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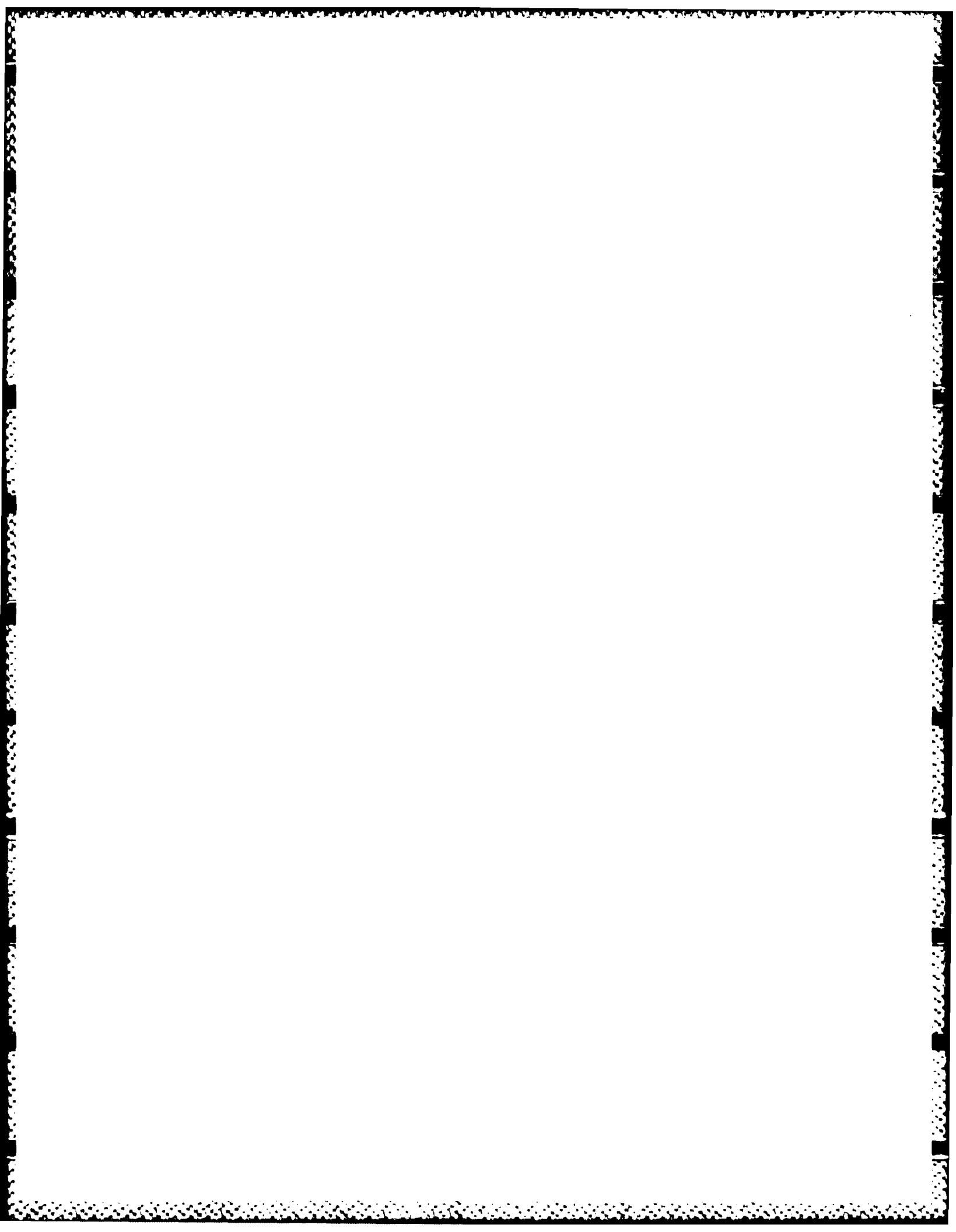


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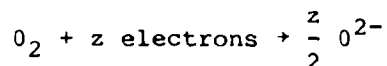
I. INTRODUCTION

The oxidation of a metallic surface involves a net transfer of electrons from the metal to a foreign species such as oxygen, sulphur etc. Oxidation is favored energetically as it results in a decrease in the free energy of the system. The reaction is commonly expressed⁽¹⁾ as:

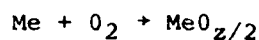
Oxidation:



Reduction:



Net:



A schematic illustration of the oxidation process is shown in Fig. 1. Reactions involving chloride, nitride, and sulphide formations, which generally occur in the environment, can be expressed similarly. Theoretically, the interactions at a solid surface with the species available in a given ambient are determined by its chemical and electronic properties.

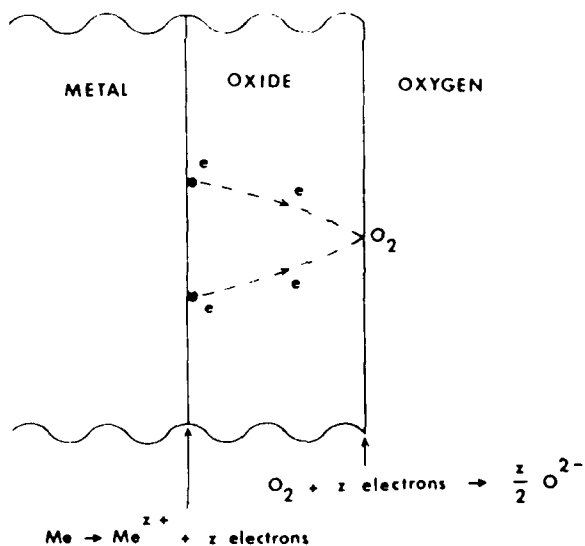


Fig. 1. Electron Transfer During Oxidation
(only the electron transfer is specifically shown).

Foreign species may interact and subsequently be adsorbed at the surface of a solid in a variety of ways. For instance, a species may be physisorbed, chemisorbed (via ionosorption or local chemical bonding), or form a new phase (e.g. oxidation on a metallic surface). Adsorption of oxygen on clean semiconductor and metallic surfaces has been a topic of intense experimental (surface spectroscopies) and theoretical (local density of states LDOS calculations) investigations.⁽²⁻⁶⁾ The chemical and electronic properties of adsorption on clean surfaces are described by atomistic (surface site or surface molecule), and band (surface state or rigid band) models. Morrison⁽⁷⁾ and Schreiffer and Soven⁽⁸⁾ have combined the two approaches by considering the chemical bonding and the energy band interaction. The nature of the chemical bonding of a foreign species to the solid surface determines whether the surface corrodes, acts as a catalyst in a chemical reaction, or remains relatively inert (passivated) in an ambient.

Metallic oxides are generally used as protective coatings to inhibit corrosion. In some cases (e.g. Ni/NiO, Cr/Cr₂O₃) the native oxide does not grow above a thickness value of 10-15 Å at room temperature, regardless of exposure time to the atmospheric conditions. In other cases, the oxidation takes place continually. The corrosion resistance of oxide coatings, grown naturally or intentionally (using one of several methods available), depends on the chemical and electronic structure, the thickness, and the porosity of the oxide layer. SiO₂, NiO, Cr₂O₃ are well known examples of naturally occurring protective thin oxides on Si, Ni, and Cr surfaces, respectively. In addition, growing oxide on Al, Ti, and Ta surfaces using anodic oxidation is a common practice to protect these metals against corrosion.^(1,9-11) The theory of anodic oxidation of metals has been reviewed by Wagner.⁽¹²⁾ Coatings of silicides, aluminides, and nitrides have also been used as protective layers.⁽¹⁾ Schneider, Bauer, and Grunling⁽¹³⁾ have studied the failure mechanisms of Si, Cr, and Ni-Cr-Si coatings for turbine applications. Among these, silicon coatings with native SiO₂ layers have been found to be the most protective against high temperature corrosion and oxidation. However, the Ni-Cr-Si coatings were found to be more desirable due to their ductility, toughness, and reasonable corrosion resistance. The theory of metal oxidation has recently been reviewed by Fromhold.^(14,15)

II. THEORY OF ACTIVE ELECTRONIC BARRIER FORMATION AT METALLIC SURFACES

The ability of an oxide or nitride layer to prevent corrosion is dependent upon its electrical resistivity. The higher the electrical resistance of the oxide layer the more effective it will be in preventing electron transfer. In general, the current flow due to the oxidation of a metal surface will depend on the resistance/impedance of the oxide layer. The higher the resistance, the better will be the protection. The electrical resistance of an oxide layer depends upon many factors including the oxide thickness, as well as the electronic (defect states in the energy gap etc.) and physical (pin holes etc.) qualities of the oxide. Fig. 2 shows the energy band diagram of an oxide with defect states in the energy gap. Fromhold^(14,15) has summarized the mechanisms responsible for electronic and ionic current flow during oxidation in oxides ranging in thickness from 10 Å to 100 microns. This type of protection can be viewed as a passive electronic barrier at the metal surface.

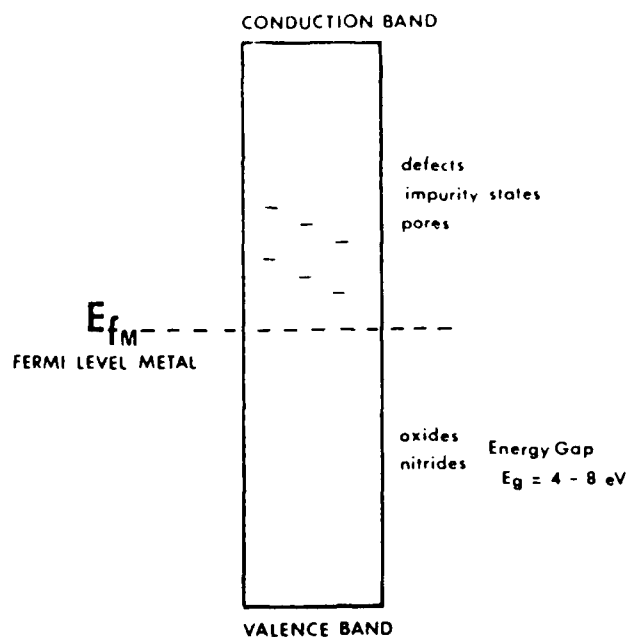


Fig. 2 Electronic Energy Band Diagram of a Metal - Oxide System (passive electronic barrier)

This paper presents a new approach to corrosion prevention. Rather than utilizing a simple passive barrier, we may create conditions at the metal surface such that it hosts a built-in electric field. The presence of such an electric field E , as shown in Fig. 3, will impede the transfer of electrons from the metal surface. Electrons now see a retarding force which prevents their transfer to the oxidizing species. This force can be expressed as $F = -eE$, where e is the magnitude of the electron charge. It should be noted that the interfacial electric field is not due to any externally applied voltage, rather it is built-in at the metal-semiconductor. Metallic surfaces host positive dipole layers when they are interfaced with appropriately doped semiconductors to form metal and metal-semiconductor and metal-oxide (insulator)-semiconductor structures.⁽¹⁶⁾ These interfacial space charge layers result in a built-in electric field and cause bending of the electronic energy bands. The net band bending is defined as the active electronic barrier. This novel approach to corrosion prevention was first proposed by Jain.⁽¹⁷⁾ It may be added that by properly selecting the semiconductor coating material for a metal surface one can realize both the traditional passive as well as the novel active barriers. The formation of an active electronic barrier is described next.

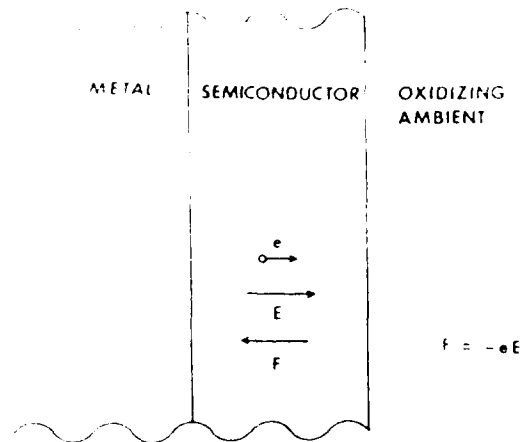


Fig. 3 Electronic Barrier Due to a Built-In Electric Field

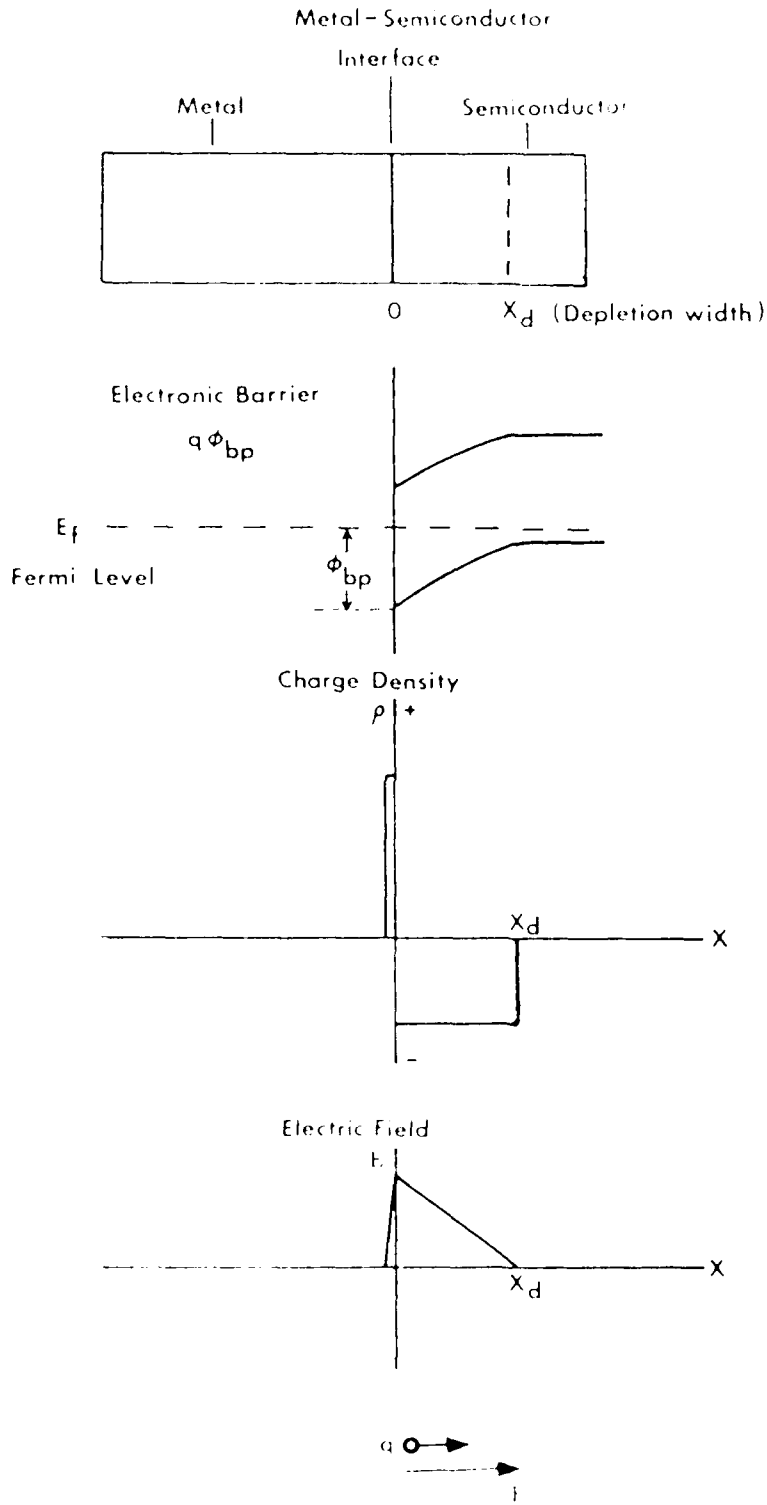
II.1 Metal-Semiconductor Interfaces

The behavior of a metal-semiconductor interface, commonly called a Schottky barrier, is known to depend on the work function of the metal, the conductivity type, the carrier concentration, and the energy gap of the semiconductor. (16,18-20) Fig. 4 shows the energy band diagram, charge distribution, and electric field profile in a metal-p type semiconductor interface. It may be added that n-type semiconductors will also form barriers with metals.

The magnitude of the electronic barrier is dependent upon the choice of metal (i.e. its work function) and the properties of the semiconductor. The presence of impurities, which can act as traps for electrons and holes in a p-type semiconductor layer, are also known to affect the barrier height. For a given metal, the barrier height of a metal-semiconductor interface is dependent upon the doping levels and the energy gap of the semiconductor. In general, wider energy gap semiconductors yield higher barrier. Recently, Brillson(21) and others have investigated the barrier formation in its initial phases using electron loss and other spectroscopic techniques.

II.2. Metal-Oxide (Insulator)-Semiconductor Interfaces

A metallic surface can also develop a positively charged dipole layer and the associated active electronic barrier in a metal-oxide(insulator)-p semiconductor (MOS) configuration. The physics of the barrier formation depends upon the thickness of the oxide. The manifestation of the electronic barrier in thick oxide ($> 150 \text{ \AA}$) systems is similar to those of conventional MOS



Electric field impeding the transfer of electrons

Figure 4. Energy barrier formation at a metal-p semiconductor interface.

devices used in Si integrated circuit technology.^(16,19) Thick oxide MOS structures are not treated in this paper. Structures utilizing oxide (insulator) thickness in the range of (20-100 Å) are generally known as MIS structures. These systems behave similarly to the metal-semiconductor systems described above. However, the barrier has been found to be larger in MIS than in MS configurations.⁽²⁴⁻²⁶⁾ For instance, we have found that the barrier height is increased in Al-nSi and Au-nGaAsP interfaces with the introduction of a thin 20-30 Å SiO₂ layer. Reference is made to Jain and Marciniec,⁽²⁵⁾ and Jain and Nichols⁽²⁶⁾ for detailed conditions of barrier enhancement in Schottky type MOS structures.

The active electronic barrier inhibits the net transfer of electrons from the metal surface to the oxidizing species, resulting in a lower probability of oxidation/corrosion. Additionally, the electronic barrier may help in regions having micropores and pinholes in the semiconductor layer. In these regions we expect a finite electric field (due to field fringing effects) to retard the transfer of electrons.

III. ACTIVE ELECTRONIC SYSTEMS FABRICATED - EXPERIMENTAL

III.1. Selection of Semiconductor/Insulator Coating Systems

The electronic barrier at the aluminum surface is realized by depositing a thin coating of n or p-type indium tin oxide (ITO). ITO films can be grown in n-doped or p-doped form depending on the reactor growth conditions. The indium tin oxide was selected for several reasons including:

- a. adjustable large energy gap (3.5 to 4.0eV for films having higher In/Sn ratio),
- b. potentially controllable n- and p-type doping (in contrast to doped polyacetylene, phthalocynine etc.),
- c. stable and durable film coatings (it has been recently reported that phthalocyanine⁽²⁷⁾ and doped polyacetylene⁽²⁸⁾ are not stable under illuminated and oxidizing environments, respectively),
- d. adaptability to chemical vapor deposition (CVD) and spray deposition techniques.

In addition to the active electronic barrier at the metal interface, the ITO film also acts as a passive barrier due to its high resistivity. The resistivity/passivity of metals and semiconductor has been the topic of a recent conference in France.⁽²⁹⁾

The use of a thin (300-100 Å) SiO₂ insulator layer in a MIS configuration is chosen for two reasons: (1) to enhance the electronic barrier at Al-ITO interfaces; and (2) to protect Al alloys (e.g. 7075-T6) from being attacked by reaction products in the ITO CVD reactor. A final Si₃N₄ coating was used to physically protect the ITO surface against wear and abrasion. It will be shown in the next section that the active electronic barrier is the primary mechanism preventing the corrosion of Al substrates, and that the Si₃N₄ coatings without the ITO film is a poor corrosion inhibitor. The sole purpose of the nitride coating is to protect the surface against wear.

III.2 Structures Fabricated

The semiconductor (ITO) and insulator (SiO_2 , Si_3N_4) films were deposited in a variety of configurations. Fig. 5 shows the metal-semiconductor and metal-thin insulator-semiconductor structures fabricated using the ITO, SiO_2 and Si_3N_4 films. It should be noted that a thin layer of aluminum oxide was grown on all the Al substrates prior to the deposition of other coatings.

The aluminum substrates were prepared from several different grades of aluminum and aluminum alloys. For example, commercial purity as well as high purity aluminum were used for polycrystalline substrates. High purity, single crystal aluminum substrates were also used. In the case of aluminum alloy, the 7075-T6 alloy was used. The ITO semiconductor coatings were deposited in the thickness range of 0.1-0.3 μm . The SiO_2 coating thickness varied between 20-100 Å. The outermost Si_3N_4 coatings were approximately 0.75 μm thick.

III.3. Fabrication Techniques

Aluminum and aluminum alloy sample surfaces were prepared using mechanical, chemimechanical, and chemical polishing techniques. A thin layer of native aluminum oxide was grown in a dry oxygen ambient at the relatively low temperature range of 300-375°C. Aluminum substrates with a thin native oxide were then deposited with either thin SiO_2 or ITO films depending on the structural configurations desired.

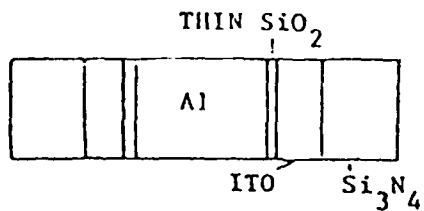
III.3.A. Substrate Preparation

All aluminum substrates (including Al alloy) were mechanically and chemically polished to a mirror-like finish prior to film deposition. The following procedure was used:

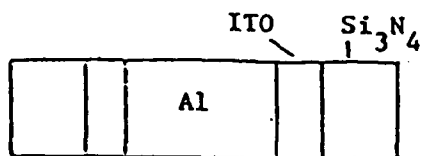
- 1) Boil in Trichloroethylene
- 2) Acid etch in 10% Hydrofluoric acid (5 seconds)
- 3) Mechanical lapping
- 4) Boil in Trichloroethylene
- 5) Mechanical polish with 5 micron alumina polish
- 6) Mechanical polish with 0.3 micron alumina polish
- 7) Mechanical polish with 0.05 micron alumina polish
- 8) Mechanical polish in ethylene glycol
- 9) Wash in distilled water
- 10) Boil in Trichloroethylene
- 11) Chemical microetch in 10% phosphoric acid
- 12) Rinse in distilled water

After polishing, the substrates were placed in an oxide furnace for aluminum oxide growth. The temperature was maintained at 300-375°C in a pure oxygen environment. After 15-20 min., the substrates were removed. At this point, a thin layer of approximately 20 Å of Al_2O_3 covered the aluminum surface.

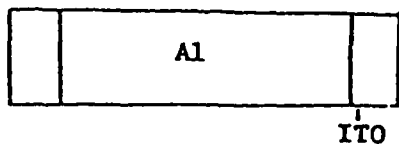
Aluminum substrates with a thin native oxide were then processed for SiO_2 , ITO, and silicon nitride depositions. These growths are described in the next two sections.



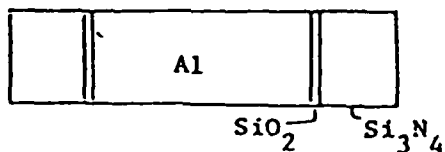
1. Al-Thin SiO₂-ITO-Nitride



2. Al-ITO-Si₃N₄



3. Al-ITO



4. Al-Thin SiO₂-Si₃N₄

FIGURE 5 - Structures fabricated to test corrosion prevention

III.3.B Semiconductor Coatings: Deposition of SnO₂ and ITO Films

The semiconductor films of Indium Tin Oxide (ITO) and Tin Oxide (SnO₂) were deposited by three techniques: 1) chemical vapor deposition (CVD), 2) spray injection chemical vapor deposition, and 3) air spray pyrolysis.

Chemical Vapor Deposition Reactor

Fig. 6a shows the CVD reactor which was utilized in the early phases of the project work. The system features separate feed lines for the reactant chemicals to allow for precise control of the film deposition. Indium chloride, water, and ethanol were mixed to form one reactant solution, while anhydrous tin chloride was used for the second solution. By controlling the flow rates of the two reactant solutions, either SnO₂ or ITO films could be fabricated. It should be noted that the anhydrous SnCl₄ was contained in a methylene chloride base, which initiated a competing reaction in the CVD reactor.

Spray Deposition Reactors

Two types of reactor systems were developed to spray deposit the semiconductor films. The first system utilized a controlled ambient environment for a chemical vapor deposition. Films grown in this reactor were of very high quality and were used for laboratory characterization. The second reactor system was developed to simulate a commercial application process. This system utilized an open air environment with a high pressure spray gun. The films grown in this system were very thick, although of somewhat lower quality.

The spray deposition technique was developed to increase the film uniformity and thickness. In addition, the composition and quality of the deposited films could be easily controlled by changing the makeup of the spray solution. The spray solution contained varying amounts of InCl₃, SnCl₄·5H₂O, CH₃CH₂OH, and H₂O. The amount of indium in the deposited ITO films could be increased or decreased by altering the concentration of InCl₃ in the starting solution. This allowed for an effective means of tailoring the bandgap of the semiconductor film. The films deposited by this method were found to be uniform and semiconducting. The two systems are described in detail below.

Spray Injection Chemical Vapor Deposition Reactor

Figure 6b shows a schematic diagram of this system. This system utilizes a controlled ambient environment in which the temperature is precisely controlled. The starting solution is aspirated through the injector nozzle using a high pressure nitrogen gas stream. The resultant mist is sprayed into the reactor where it is preheated at the nozzle tip and then vaporized in zone I. The aluminum substrate is placed in a quartz holder which is located at the zone I/zone II boundary. The temperature of zone I is typically maintained at 575°C while zone II is at 500°C. This difference creates a temperature gradient which enhances the chemical vapor deposition on the aluminum substrate. The actual substrate temperature is typically 390-430°C. The composition of the ITO films is varied by changing the spray solution makeup and by varying the zone temperature. The

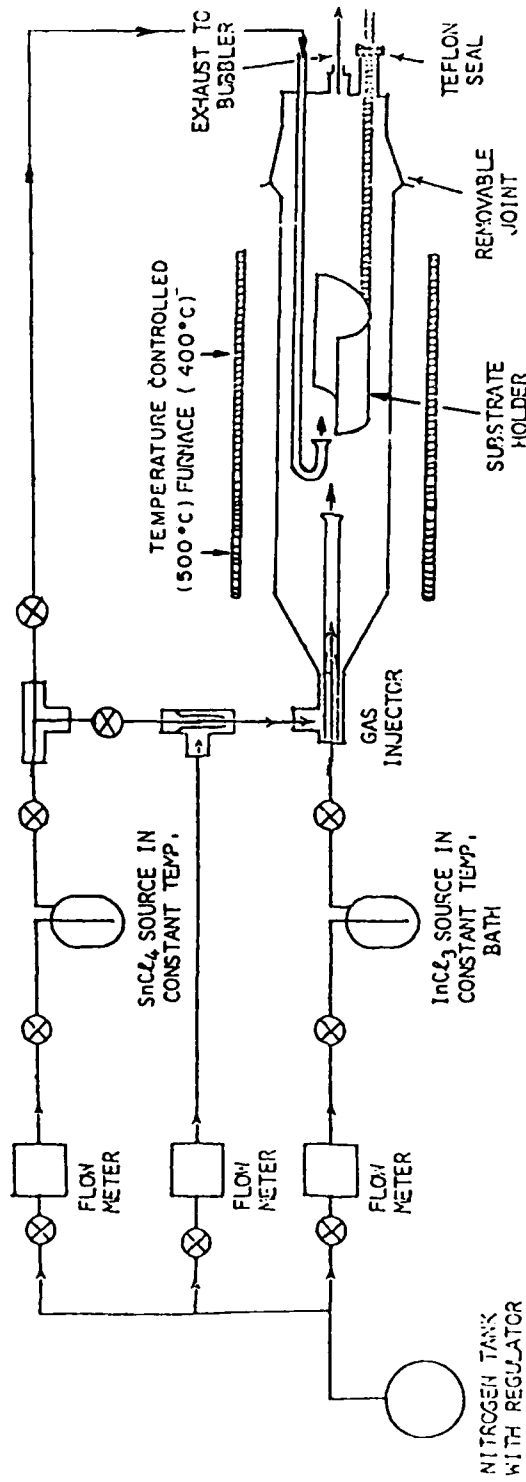


FIGURE 6a - Schematic of Indium-Tin-Oxide (ITO)
Chemical Vapor Deposition reactor

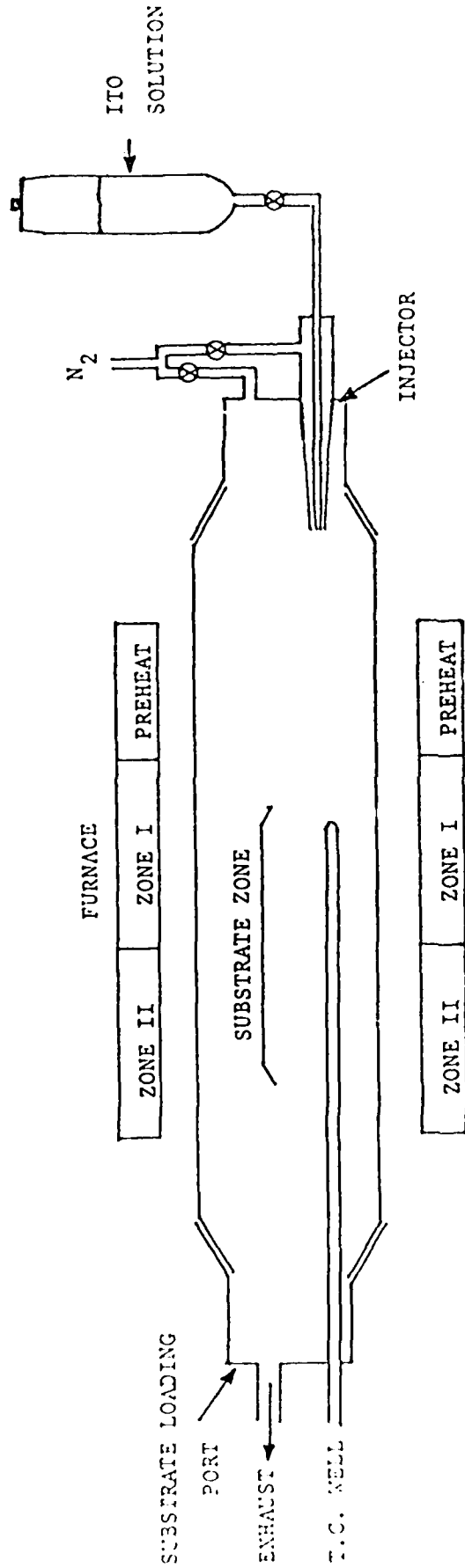


FIGURE 6b - Controlled ambient spray injection CVD ITO reactor

thickness of the film is determined by the quantity of solution injected and the precise location of the substrate.

The films deposited in this reactor were of very high quality. Due to the controlled ambient environment, the number of defects and impurity states in the semiconductor films could be minimized. These films were used for analysis and characterization (i.e.: Auger, SEM analysis, spectrophotometric analysis, and electronic characterization).

Air Spray Reactor System

A second spray deposition system was developed with the goal of simulating a commercial type application of the semiconducting films. This system is shown schematically in figure 7. In this reactor, the spray deposition takes place in an open air environment. The air is filtered through a submicron filter, but is held at room temperature. In order to achieve the high substrate temperature required for the reaction, the substrate is placed on a hot plate at approximately 390-430°C. The starting solution is aspirated with nitrogen and sprayed directly on the aluminum substrate. The height and angle of the spray nozzle is fully adjustable to optimize film thickness and quality.

The films deposited in this "open-air" system were typically very thick and uniform. However, due to the influence of impurities in the ambient, the films did not form an electronic barrier as large as those deposited in the CVD reactor.

ITO and SnO₂ Deposition Reactions

ITO Chemical Reaction

The chemical reaction which occurs in the ITO CVD and ITO spray deposition processes is an oxidation-reduction reaction. Table 1 gives the reaction in detail. The reactants used in the spray deposition solution were 3.9M InCl₃, SnCl₄.5 H₂O (hydrated crystals), CH₃CH₂OH (ethanol), and water. A typical mixture used to yield an indium-rich film of ITO consists of:

0.5 cc 3.9M InCl₃

1.25 g SnCl₄.5 H₂O

7 cc CH₃CH₂OH

3 cc H₂O

The resultant solution is then sprayed through the injection nozzle.

Tin-Oxide Chemical Reaction

In forming tin-oxide, the InCl₃ is deleted from the list of reactants above. A typical solution mix is:

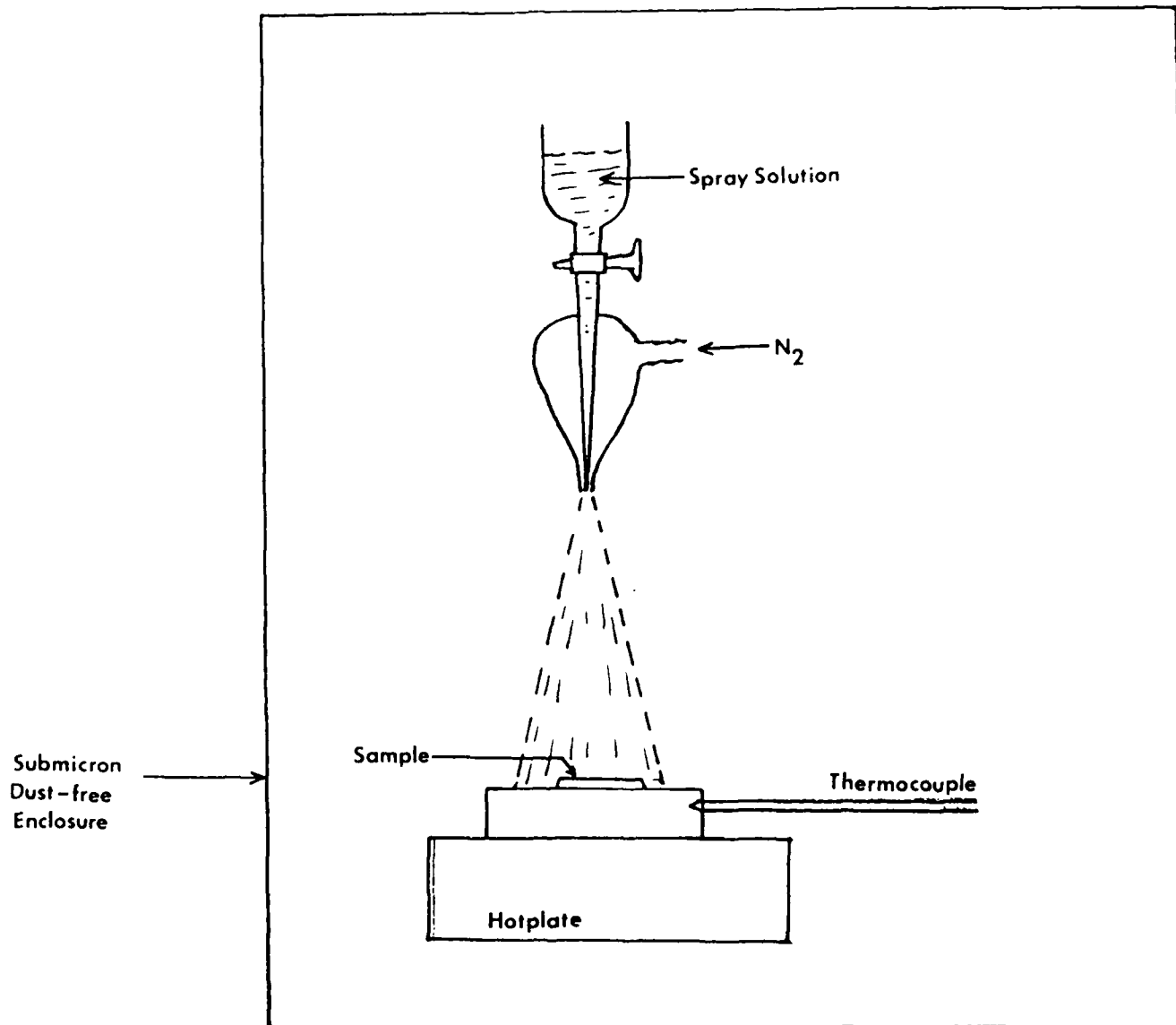
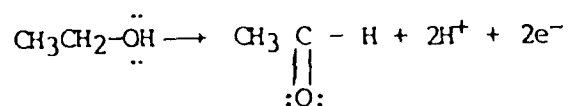
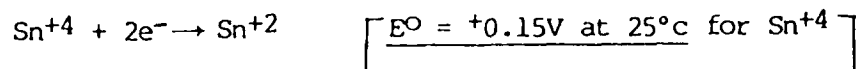
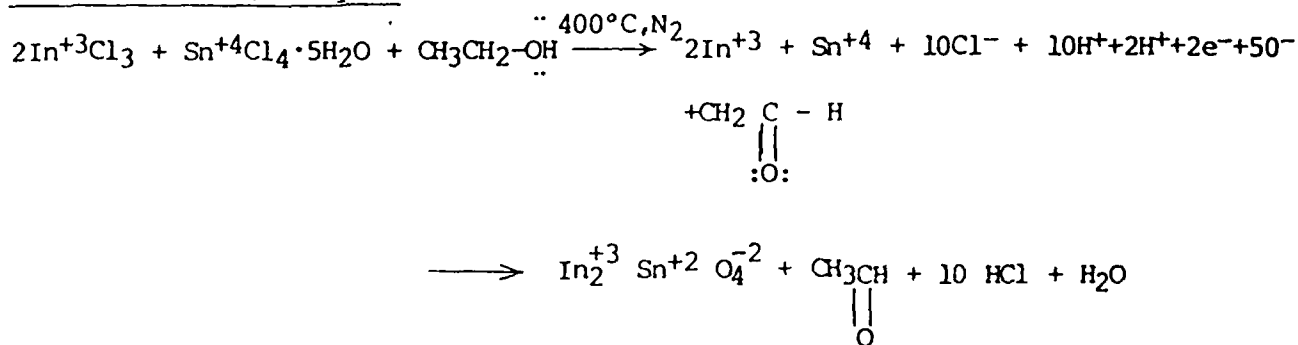
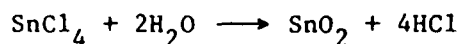
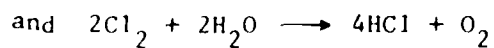
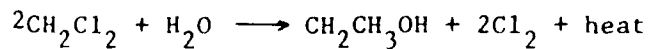


FIGURE 7 - SPRAY PYROLYSIS APPARATUS

TABLE 1 - Chemical reactions

ITO Spray Deposition ReactionOxidationReductionTotal Reaction (Aldehyde)SnO₂ Formation in absence of significant InCl₃Methylene Chloride competing reaction (Ethanol production)

1.25 g SnCl₄ · 5 H₂O

3.0 cc H₂O

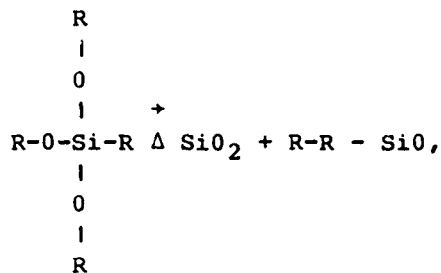
7.0 cc CH₃CH₂OH

Tin-oxide films can be formed at a lower temperature than ITO films. The minimum reaction temperature is approximately 350°C. However, the SnO₂ films have a lower energy bandgap than ITO and do not form as large an electronic barrier. The reaction which occurs in SnO₂ deposition is also shown in Table 1. Also given at the bottom of Table 1 is the competing reaction involving methylene chloride which occurred in the CVD reactor (with separate feed solutions of InCl₃ and anhydrous SnCl₄).

C. Insulator Films

Silicon Dioxide Growth

Fig.8 gives a schematic representation of the SiO₂ reactor. The deposition of SiO₂ involves the pyrolytic decomposition of tetraethylortho-silicate (TEOS) at above 700°C. However, the samples are kept in a lower temperature zone ranging between 450-500°C. The reaction resulting in SiO₂ deposition is as follows.



Where R is the hydrocarbon radical.

IV. RESULTS

IV.1 Characterization of ITO, SnO₂, SiO₂, and Si₃N₄ Coatings

IV.1.A. Physical Characterization

Important parameters in the physical characterization are:

- a. chemical composition
- b. grain size
- c. film thickness

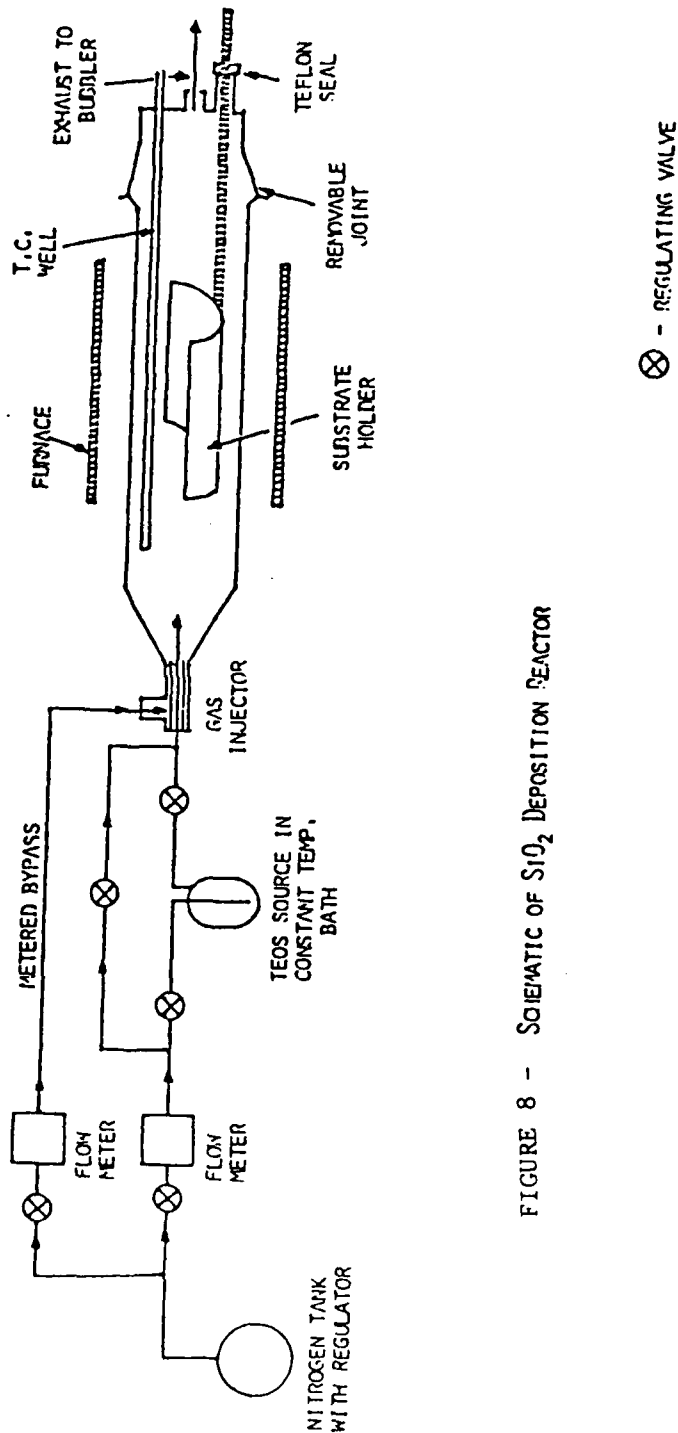


FIGURE 8 - SCHEMATIC OF SiO₂ DEPOSITION REACTOR

The chemical composition is determined using energy and wavelength dispersive techniques such as EDAX and WDAX. These analytical techniques are available as an attachment to a scanning electron microscope. In addition, an Auger depth profiling setup (at the United Technologies Research Center) is used for additional information on the deposited films. The scanning electron and optical microscopes were used to determine the grain size of ITO films.

The WDAX data is shown in Table 2.

Table 2. - Chemical Composition of ITO Films

Sample	Film Structure	Composition	Molar In/Sn ratio	Growth Method
#1	ITO/Si	In. _{.26} Sn. _{.42} O. _{.75}	0.62	Injection CVD
#2	ITO/SiO ₂ /Al	In. _{.003} Sn. _{.14} O. _{.67}	0.002	CVD
#3	ITO/SiO ₂ /Al	In. _{.28} Sn. _{.46} O. _{.74}	0.61	Injection CVD
#4	ITO/SiO ₂ /Al	In. _{.34} Sn. _{.44} O. _{.47}	0.78	Injection CVD

The SEM micrographs of ITO films on Si and Al substrates are shown in Figs. 9(a) and 9(b), respectively. Fig. 9(a) shows an ITO film deposited by the CVD technique. The film surface appears extremely fine grained with little evidence of pinholes. Fig. 9(b) shows a spray injection deposited ITO film. The film structure is coarser and shows some pinholes.

IV. 1.B Optical Characterization of SnO₂ and ITO Coatings

The optical absorption studies on ITO films were done using a Varian spectrophotometer (Model 17D). SnO₂ and ITO films were grown under identical growth conditions (CVD/CVD Spray) on quartz substrates. The quartz samples were located in the same zone as the aluminum samples which were used for corrosion testing. The absorption data results are shown in Figs. 10 and 11.

The bandgap of ITO (In/Sn ratio of 0.62) was found to be approximately 3.5 eV. In contrast, the SnO₂ film showed an energy gap of about 1.5 eV.

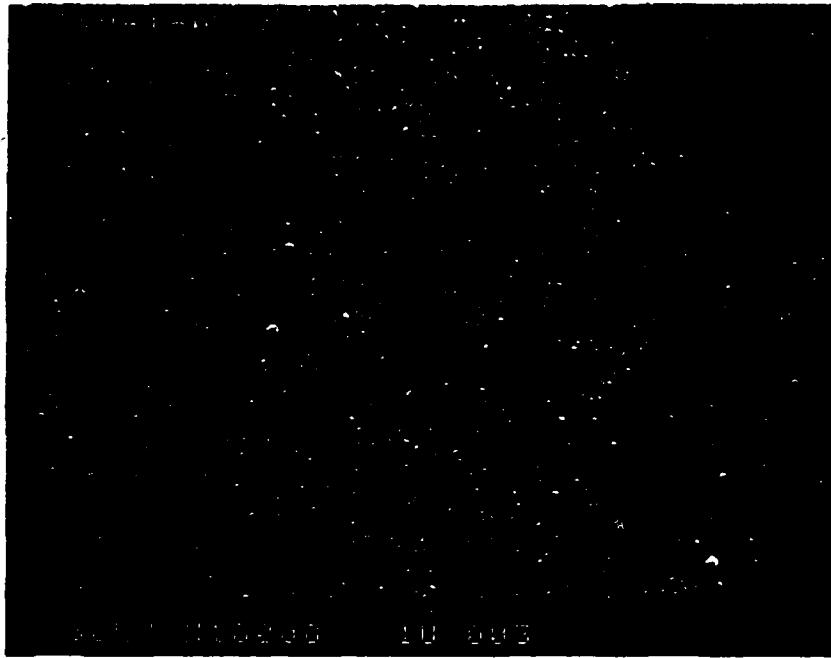


FIGURE 9a - SEM micrograph of CVD deposited ITO film

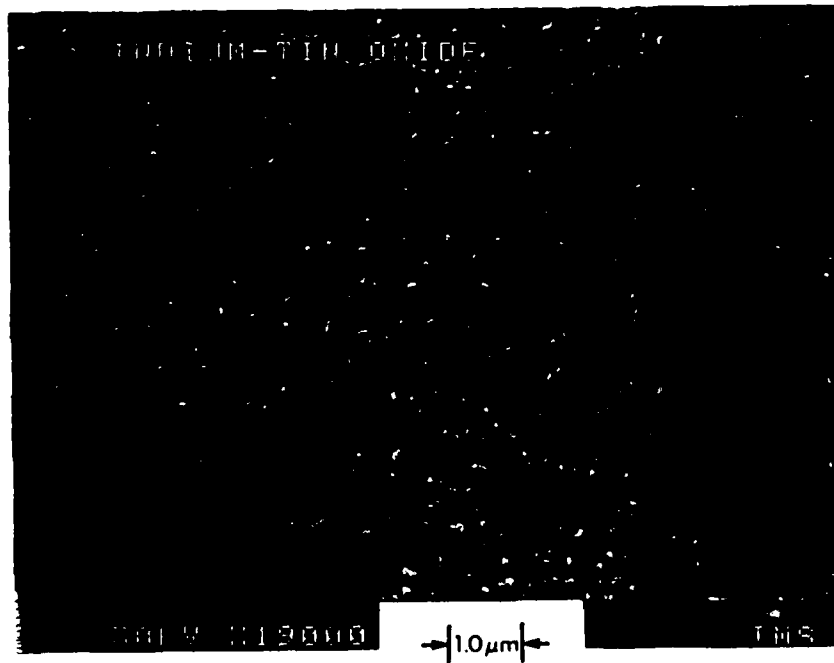


FIGURE 9b - SEM micrograph of spray CVD deposited ITO film

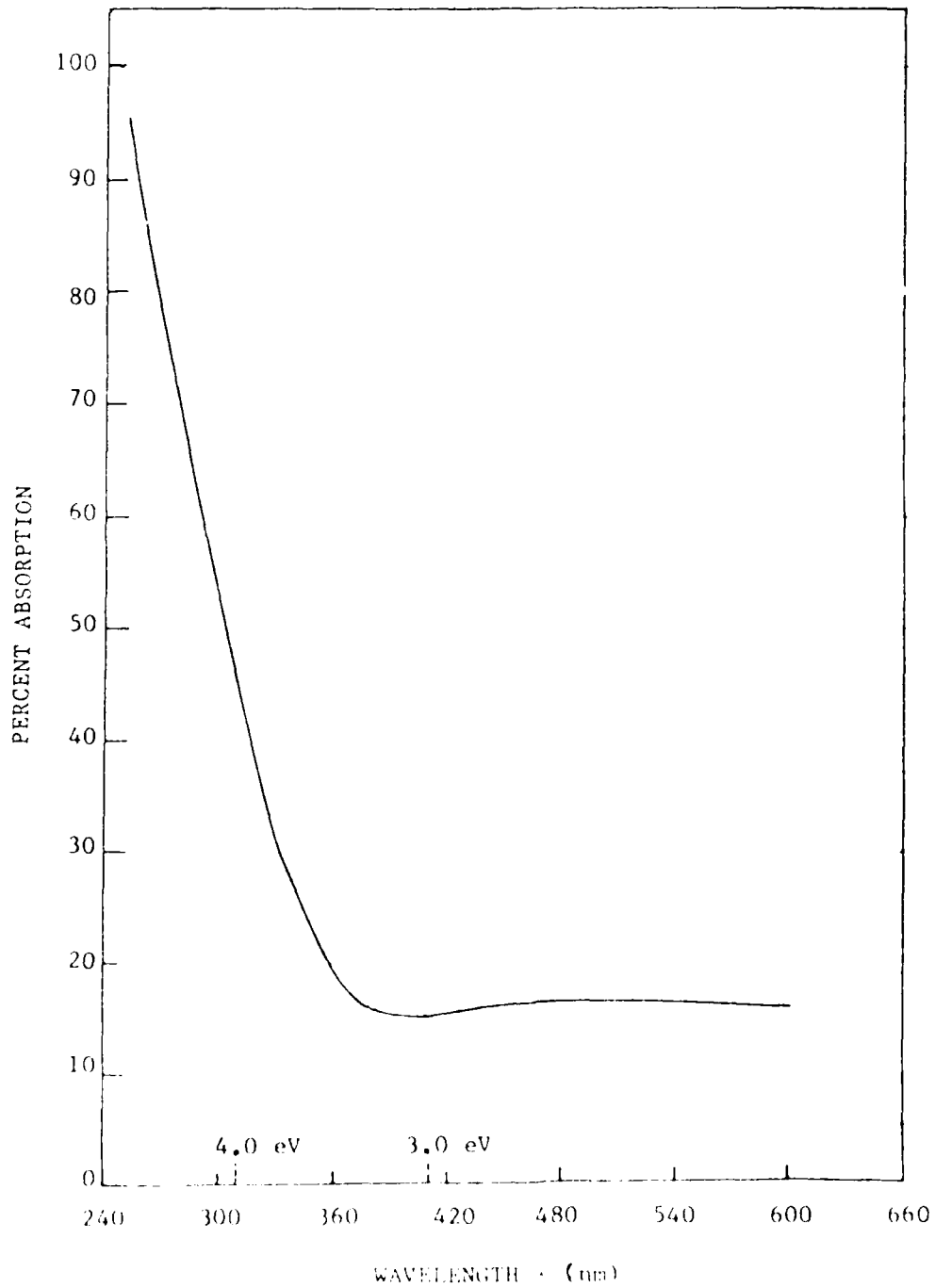


FIGURE 10 - The Absorption Edge of Indium Tin Oxide

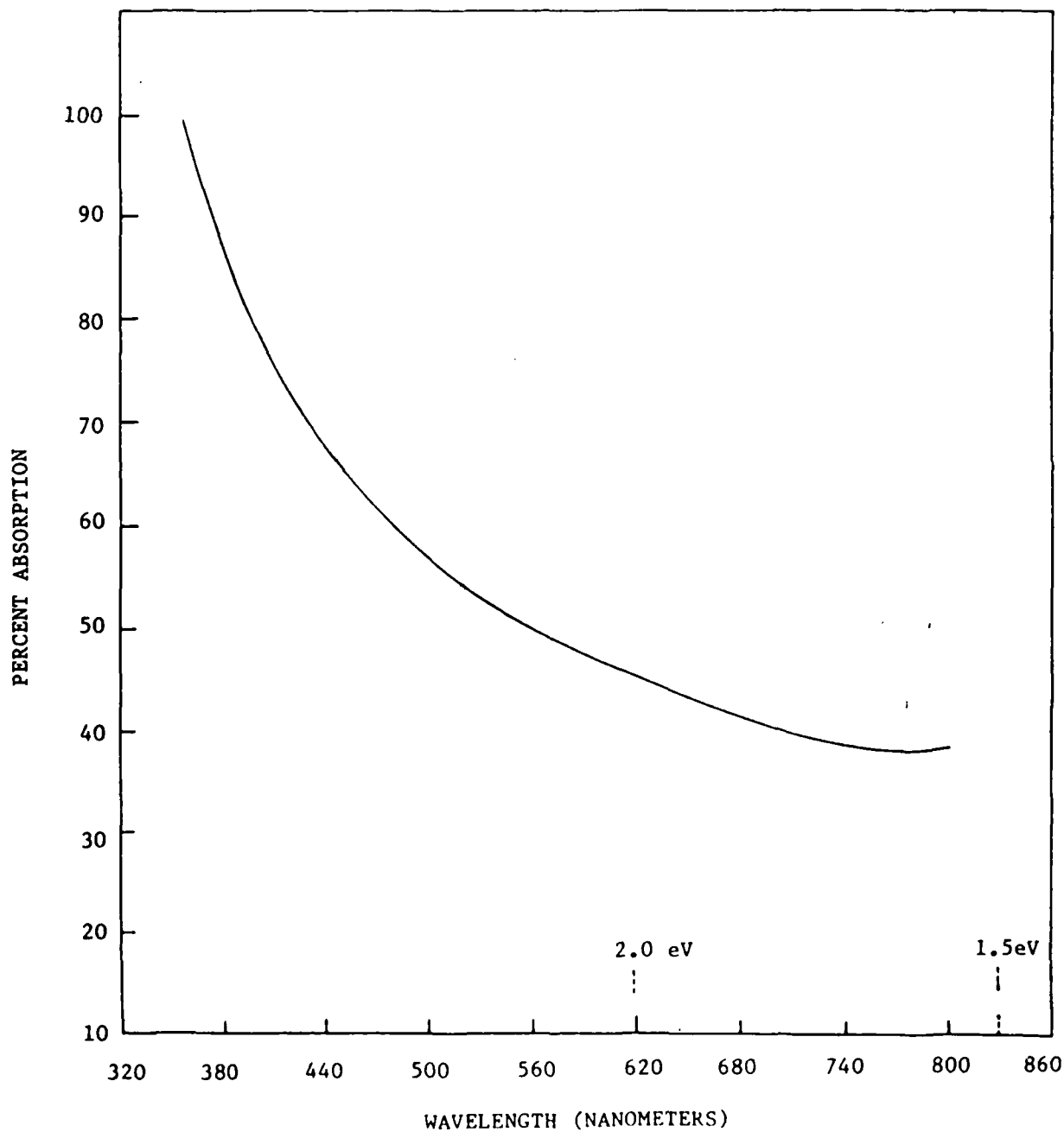


FIGURE 11 - The Absorption Edge of Tin Oxide

The optical data is a clear indication that the films are semiconducting. In addition, it shows that the energy bandgap is significantly larger in ITO.

IV.1.C Electronic Measurements

Resistivity Measurement

A Four-point setup was used to determine the electrical resistivity of the ITO films. The tested ITO films included films grown in Si as well as quartz substrates. This technique does not work on ITO films grown on Al substrates as the aluminum, being a good conductor, interferes with the measurements. Table 3 lists the results of some resistivity measurements.

V-I Characteristics:

Direct evidence of the electronic barrier formation at Al/ITO interfaces was provided by analyzing voltage-current data of ITO/quartz samples.

Fig. 12 shows the arrangement used for voltage-current characteristics. ITO/quartz samples having Al and Ag deposited contacts were used. Here, the aluminum contact simulates the Al substrate (as in the Al/ITO samples used for corrosion testing) and Ag serves as the ohmic contact to the ITO film.

The V-I characteristics of two different (spray CVD) samples having In/Sn ratio of about 0.6 are shown in Figs. 13 and 14, respectively. Fig. 13 shows good rectification behavior ('REVERSE'), characteristic of a good electronic barrier. However, the 'FORWARD' characteristic seems to indicate series resistance (ohmic contact). Fig. 14 exhibits soft 'REVERSE' V-I characteristics, an indication of surface and/or defect states short-circuiting the barrier. The surface states provide an alternate path for current conduction, allowing part of the current to flow around the electronic barrier. Regardless, both samples verify the formation of the electronic barrier at Al/ITO interfaces. The absence of a barrier would result in a straight-line type of V-I plot under both forward and reverse biasing. Fig. 15 shows the V-I characteristics of an ITO/Si sample (prepared using the CVD technique) under dark and illuminated conditions. Computations based on dark voltage-current and photovoltaic measurements yielded an effective barrier height in the range of 0.15 - 0.4 eV.

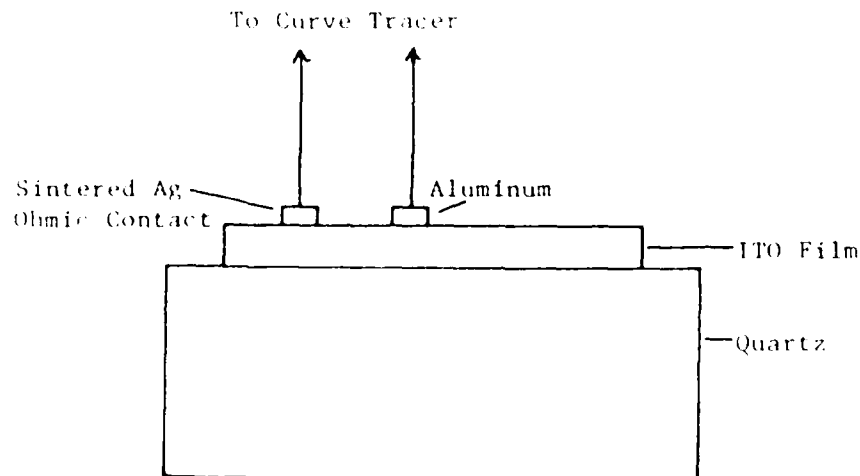
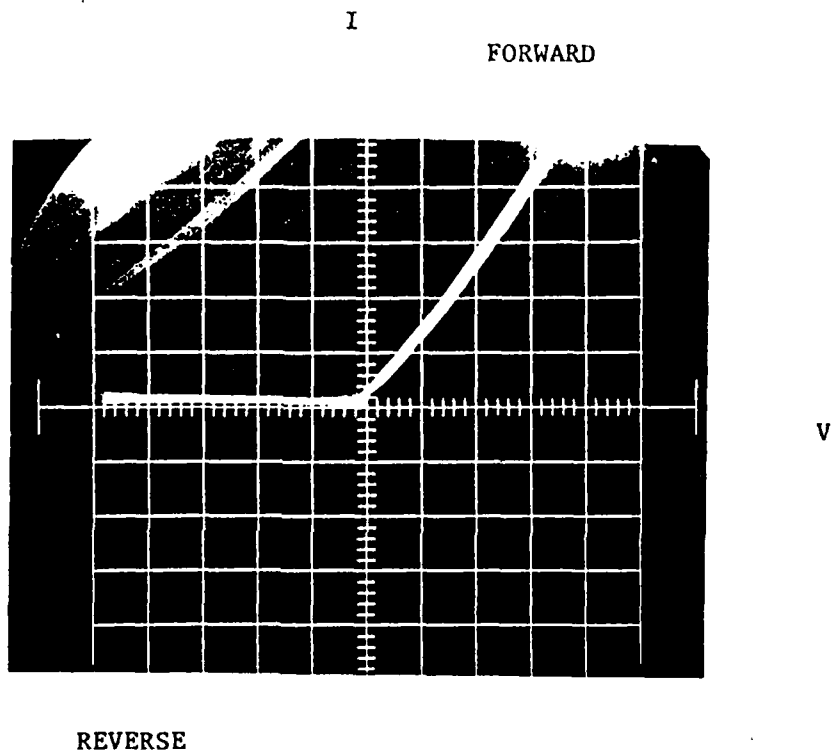


FIGURE 12 - V-I Characteristics Test Setup

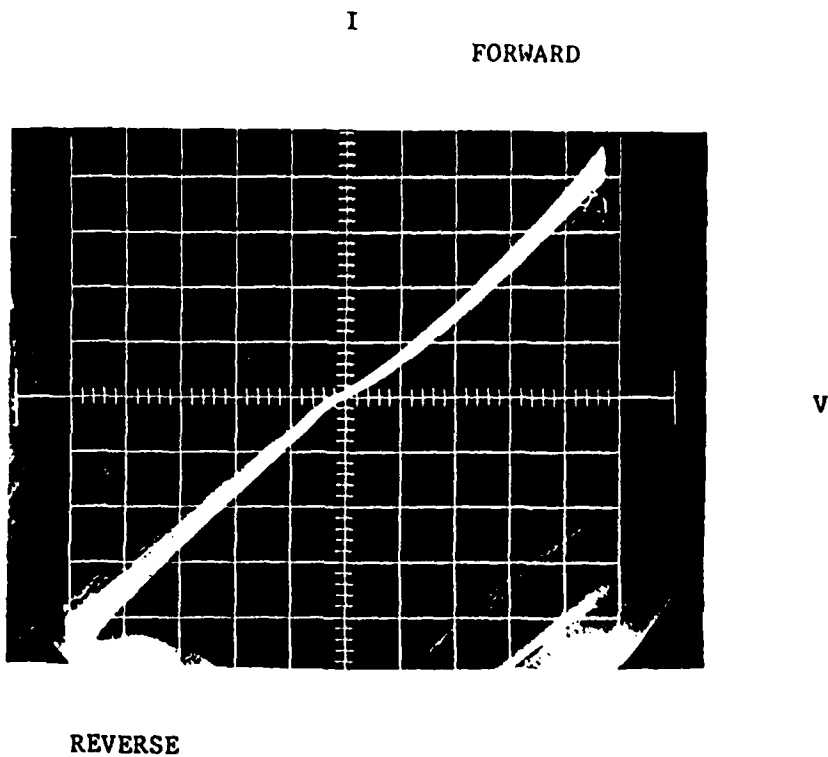
TABLE 3 - Four-Point Probe Resistivity Measurements

<u>Control Sample</u>	<u>Method of Deposition</u>	<u>ITO Conductivity Type</u>	<u>Resistivity Ω-cm</u>	<u>Growth Temp. $^{\circ}$C</u>
ITO/Si	Open Air Spray	n	3.7×10^{-3}	485
ITO/Si	Open Air Spray	p	5.9×10^{-3}	485
ITO/Si	Open Air Spray	n and p	8.5×10^{-3}	485
ITO/Si	CVD Spray	p	5.2×10^{-4}	460
ITO/Quartz	CVD Spray	n	3.6×10^{-4}	465
ITO/Quartz	Open Air Spray	n	5.9×10^{-3}	480



	FORWARD	REVERSE
CURRENT:	10 μ A/DIV	10 μ A/DIV
VOLTAGE:	0.5 V/DIV	0.5 V/DIV

Figure 13 -CURRENT-VOLTAGE CHARACTERISTIC OF AN Al-ITO/QUARTZ SAMPLE



	FORWARD	REVERSE
CURRENT:	5 μ A/DIV	5 μ A/DIV
VOLTAGE:	0.5 V/DIV	0.5 V/DIV

Figure 14 -CURRENT-VOLTAGE CHARACTERISTIC OF AN Al-ITO/QUARTZ SAMPLE

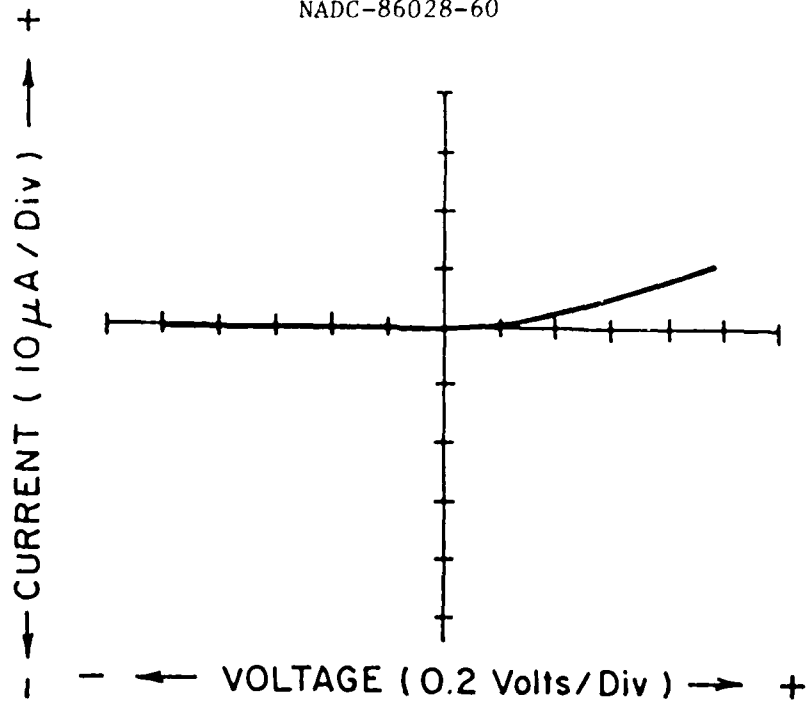


FIGURE 15a - Current-voltage characteristic of CVD Al-ITO sample (dark conditions)

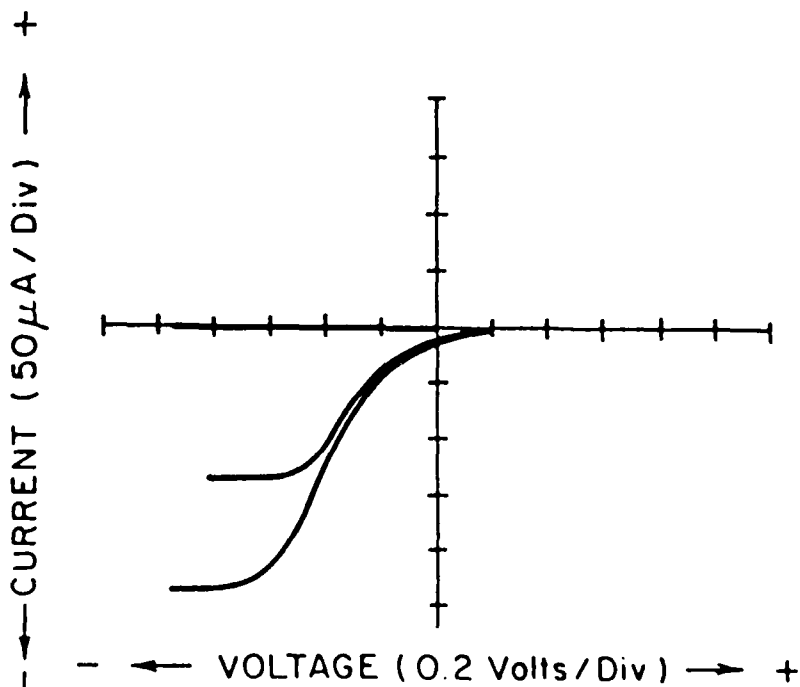


FIGURE 15b - Current-voltage characteristic of a CVD Al-ITO sample under illumination

IV. 2 Corrosion Testing

Fabricated structures were tested using weight-loss, and cathodic and anodic polarization techniques. A comparative study of various structures was conducted. The anodic polarization measurements were made at the NADC facilities using a potentiostatic setup. The details of this method are described elsewhere.⁽³³⁾ Table 4 summarizes some representative samples tested.

TABLE 4 - Anodic Polarization results (1% 2NaCl, 2pH)

Aluminum Type	Structure	Method of Growth	Resp Potential (-) Volt	Polarization results in Figure #
C.P.	Al-SiO ₂ -ITO	CVD Spray	0.980	16
C.P.	Al-SiO ₂ -ITO-Si ₃ N ₄	Air Spray	0.845	16
C.P.	Al	Control Sample	0.72	16
H.P.	Al-Si ₃ N ₄	Plasma	0.7	17
H.P.	Al-SiO ₂ -ITO	CVD Spray	0.66	17
H.P.	Al-SiO ₂ -ITO	Air Spray	0.707	17
Alloy	Al-SiO ₂ -ITO	CVD Spray	0.825	18
Alloy	Al-SiO ₂ -ITO	CVD Spray	0.9	18
C.P.	Al-SiO ₂ -ITO/TO-Si ₃ N ₄	CVD Spray	0.833	19
C.P.	Al-SiO ₂ -ITO/TO-Si ₃ N ₄	CVD Spray	0.965	20

C.P. = Commercial purity;

H.P. = High Purity;

Alloy = 7075-T6

Figs. 16-20 present polarization results on these samples. In all the voltage-current plots presented here, the point at the extreme left (intersecting the voltage axis) represents the rest potential. Fig. 21 shows polarization results of Al samples with low indium content ITO films. The role of the indium content will be discussed in the next section.

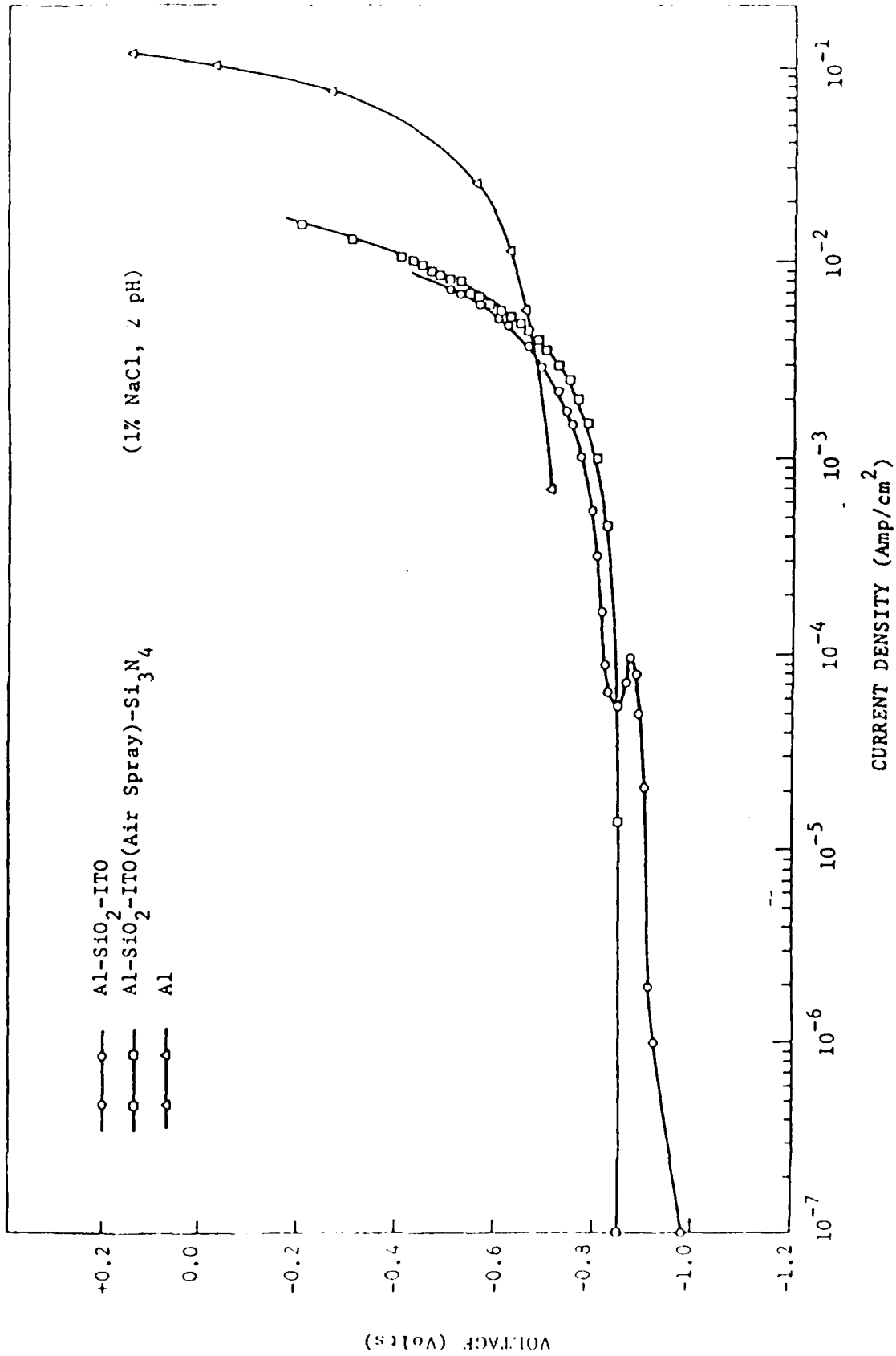


FIGURE 16 - Polarization characteristics of commercial purity aluminum substrates

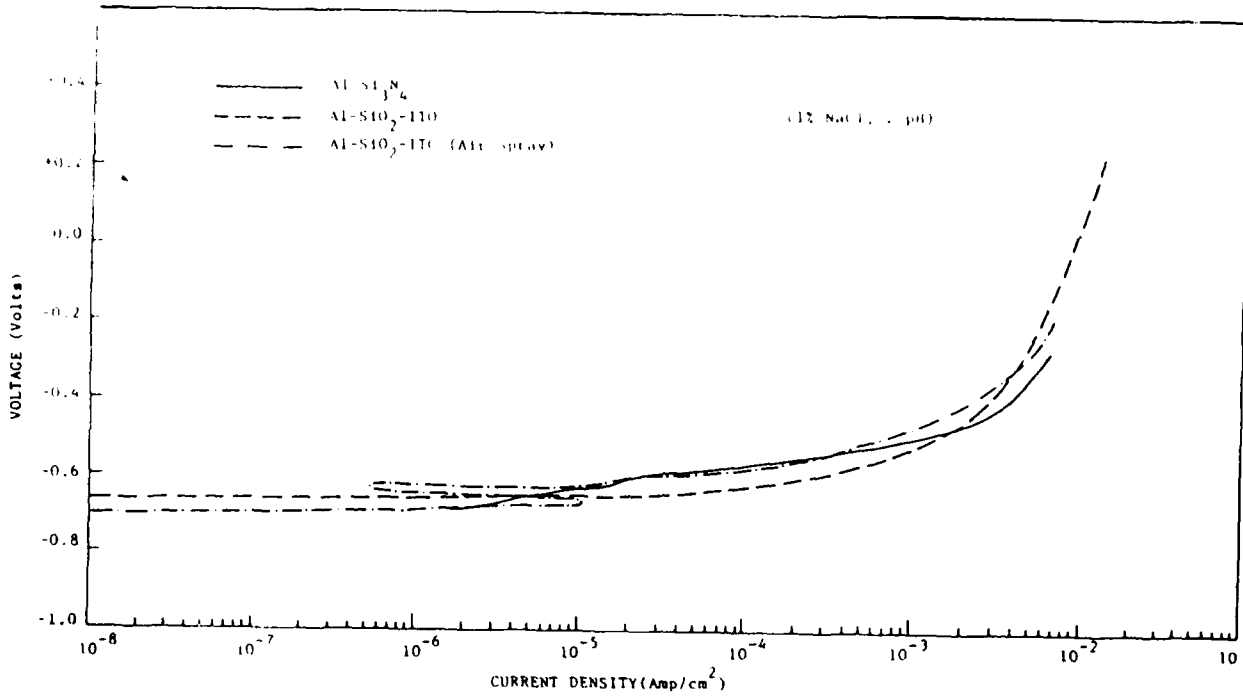


FIGURE 17 - Polarization characteristics of high purity aluminum substrates

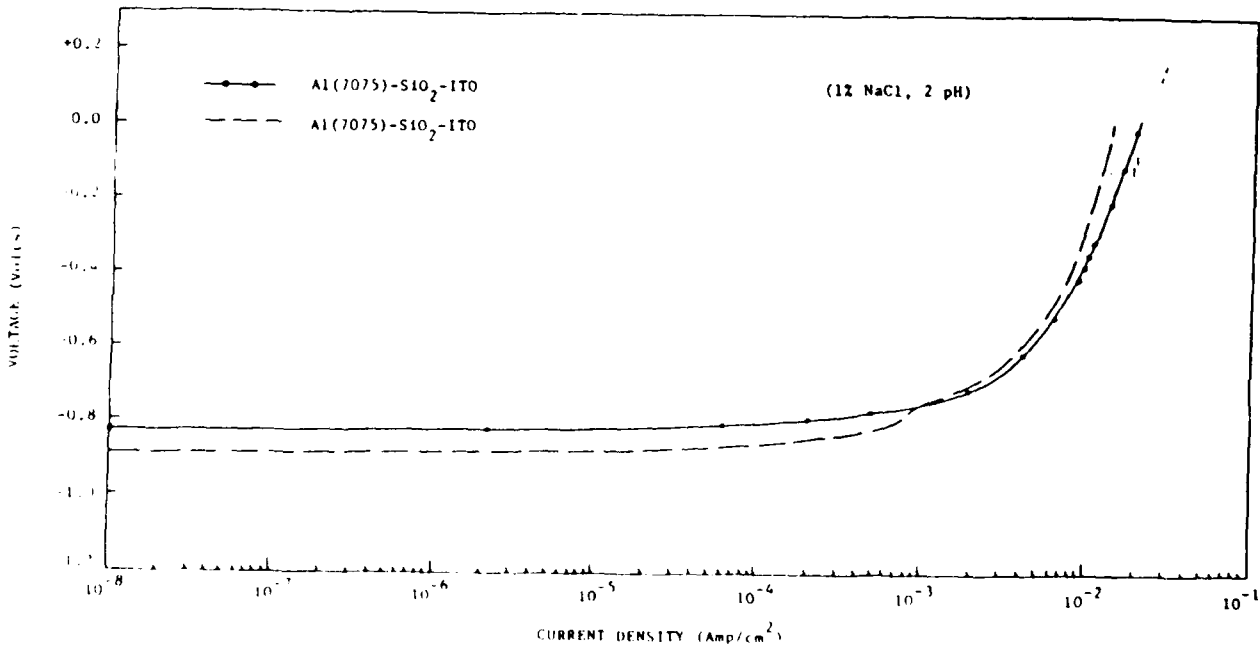


FIGURE 18 - Polarization characteristics of aluminum alloy (#7075-T6) substrates

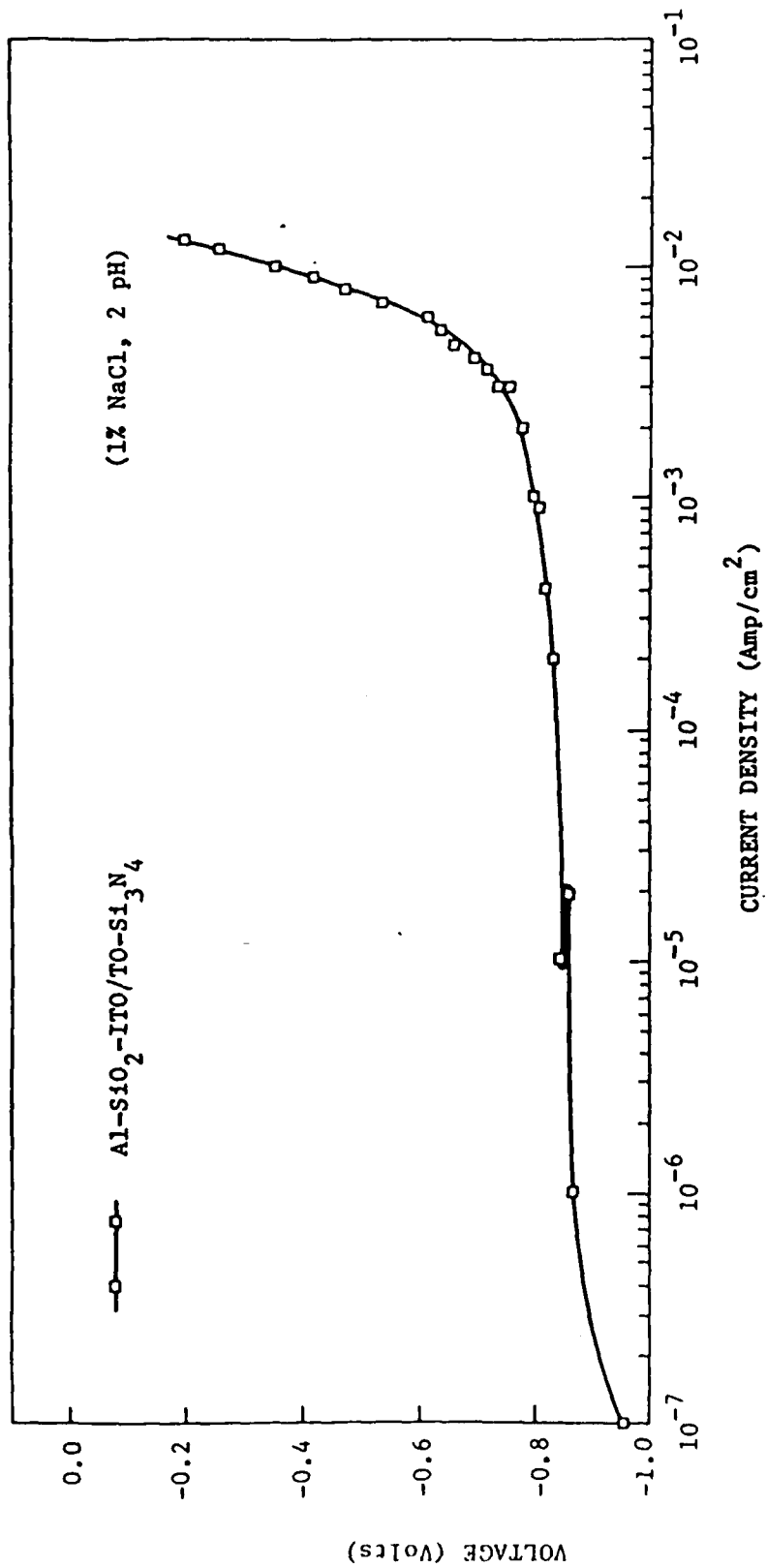


FIGURE 19 - Polarization Characteristic of a Commercial Purity Aluminum Substrate (With Composite ITO/TO Film)

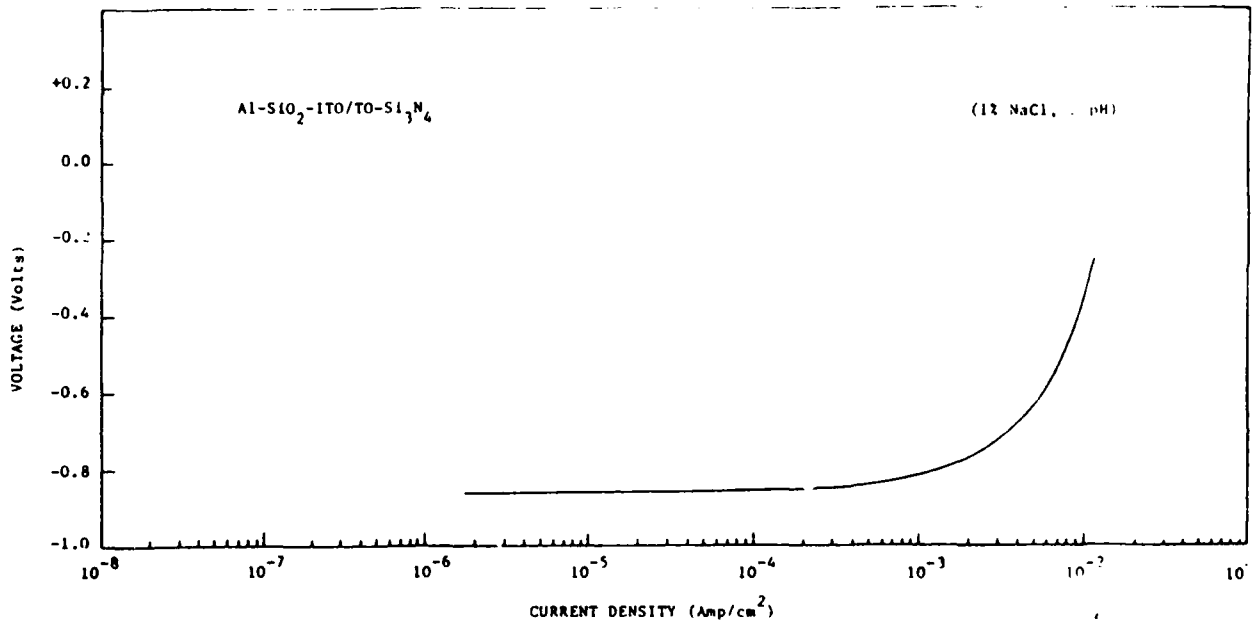


FIGURE 20 - Polarization characteristic of a scratched aluminum substrate (Commercial Purity)

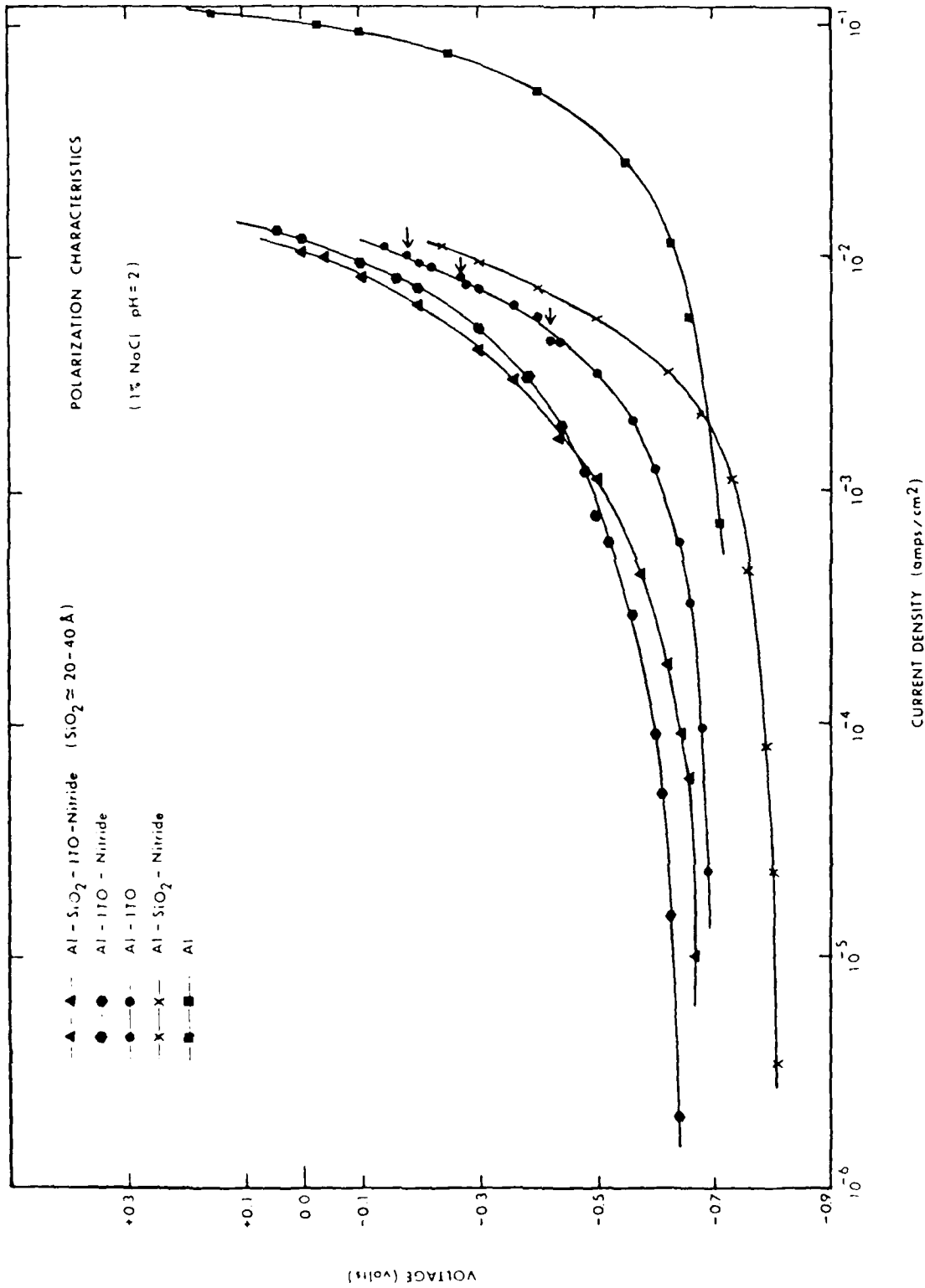


FIGURE 21 - Polarization characteristics of CVD deposited low indium content ITO samples

Weight-loss measurements were performed in our laboratory using 1-3% NaCl (with 2 pH) solutions. The weight loss measurements were conducted on Al-ITO samples coated with SiO₂ or Si₃N₄. The samples showed little or no weight loss. This data is shown in Table 5. High purity aluminum substrates were used for testing. The ITO films used in these samples were of low indium content. These films were grown in a CVD reactor.

TABLE 5.- Weight Loss Measurement on CVD Deposited Samples of Al in 3% NaCl Solution at pH 2

Sample #	Initial Wt gm	Wt after 70 Hrs. gm	Weight loss gm
Al-ITO-NITRIDE	2.08867	2.08863	0.00004
Al-ITO-SiO ₂ (THICK)	0.75436	0.75412	0.00024
Al	0.81532	0.81307	0.00225

[Surface area \approx 5 cm²]

The ITO films deposited by the spray deposition technique having higher indium content showed different weight loss results. Films deposited on quartz and mirror smooth silicon substrates showed negligible weight loss. For example, a typical Si sample (surface area \approx 7.84 cm²) coated with ITO having a total weight of 0.83714 gms showed no measurable weight-loss over a time period of 48 hours in a 1% NaCl solution (with pH=2). Similar results were found for ITO coated quartz samples. However, films deposited on commercial grade aluminum substrates exhibited higher weight losses. The rate of weight loss was small initially but accelerated with time.

V. DISCUSSION

This section interprets the results of the corrosion test data presented in the preceding section. The anodic polarization data provides the most useful means of evaluating the protective nature of the ITO films. The significant increase in the magnitude of the rest potential in the presence of ITO semiconducting films is evident from Figs. 16-21. This increase, in conjunction with the electronic V-I characteristics, confirms the existence of the electronic barrier at the metal-semiconductor interfaces. Samples exhibiting higher magnitudes of rest potential also show reduced corrosion currents, particularly at lower voltages (i.e. within the range of 100 to 200 mV above the rest potential). Fig. 21 shows the importance of ITO as a protective coating in contrast to both SiO₂ and Si₃N₄ coatings. This figure compares unprotected Al, Al-passive films and Al-active barrier films. It should be mentioned that a relatively thick (0.7 μ m) plasma-deposited Si₃N₄ layer is not as protective as the much thinner (0.2 μ m) ITO layer. Additional comparisons of Al-Si₃N₄ films with Al-SiO₂-ITO and Al-SiO₂-ITO-Si₃N₄ films are given in Figs. 16 and 17. Another important feature evident from the comparative data is that the presence of a thin SiO₂ layer (50-100 Å) does indeed further increase the effective barrier height. This effect is evident in Figs. 16-21. The results also indicate that the SiO₂/ITO system is effective in protecting both aluminum (commercial and high purity Figs. 16, 17, 19-21) and aluminum alloy (Fig. 18) substrates.

A comparison of Fig. 21 with Figs. 16-20 illustrates the effect of indium content in the ITO films. The samples of Fig. 21 had lower indium-to-tin ratios ($\text{In/Sn} \approx 0.002$) than those of Figs. 16-21 ($\text{In/Sn} \approx 0.3-0.6$). Clearly the polarization results of Figs. 16-20 show an increased magnitude in the rest potential and lower corrosion currents. This can be explained in terms of an increased electronic barrier at the Al-ITO interface with high In/Sn ratio films. It is well known that the electronic barrier at a metal-semiconductor interface increases with the energy gap of the semiconductor.⁽¹⁶⁾ The ITO films containing more indium have been found to have an increased energy gap. This has been confirmed by the spectrophotometric measurements. The increased rest potential in the anodic polarization data has graphically demonstrated the importance of the semiconductor energy gap.

In addition to ITO coatings, composite coatings of ITO and tin oxide (TO) were also investigated. The motivation for this was twofold: (1) to increase the film thickness in order to minimize the occurrence of pinhole/micropores; and (2) to explore the possibility of a double active electronic barrier. The second barrier would exist at the ITO/TO heterointerface. Figs. 19 and 20 show these results. While these structures show promising results, further work is needed to characterize the properties of the second barrier.

The weight loss data for Al samples coated with ITO show different results for films deposited by CVD and spray deposition techniques. CVD deposited samples, having a low indium content, exhibit very low weight losses, as shown in Table 5. However, ITO films deposited by the injection spray technique show higher weight loss in spite of the higher active electronic barrier. This contradictory behaviour is attributed to the relatively poor surface qualities of the films grown by this method on Al substrates. Surface morphological analysis of weight-loss samples revealed that the weight loss was concentrated at selective sites. It is believed that these areas hosted large pinholes in the ITO films where selective undercutting occurred. Fig. 22 shows photographs of these areas magnified by 420X. The substrates were commercial grade aluminum alloy (#7075-T6) with controlled ambient CVD spray deposited ITO films. The sequence of photographs dramatizes the growth of a pinhole due to undercutting at the pinhole site. It is further believed that the undercutting was reduced, if not eliminated, at small pinhole sites due to electric field fringing. Additional confirmation of this theory is provided by the weight loss results of Si-ITO and quartz-ITO samples. These results show that the spray deposited ITO films are indeed chemically impervious to the test environment.

We can further analyze the polarization data quantitatively. For instance, if we determine the change in current Δi for a small change in applied voltage $\Delta V = (V_{\text{applied}} - V_{\text{Rest Potential}})$, we can compare the effectiveness

of the various coatings. If the change ΔV is less than 25 mV (i.e. $\frac{kT}{q}$; where

k is the Boltzman Constant, q is the magnitude of electron charge, and T the temperature in Kelvin) we can obtain information on the intrinsic corrosion resistance of the film. Reference is made to Fontana and Greene³³ for additional details. Data extracted from the polarization measurements, particularly in the vicinity of rest potential (10 mV-20 mV), shows that the corrosion resistance was significantly greater than 10^8 ohms for the Al-SiO₂-ITO sample of Fig. 16 (the current change was less than 10 nA, and was not measured accurately on the setup used). In fact, it took a change of about

FIGURE 22 - Undercutting of ITO films at large pinhole sites

Pictures taken after exposure to 1% NaCl (2 pH) solution for following durations: a) 15 minutes, b) 45 minutes, c) 75 minutes,

d) 105 minutes and e) 135 minutes. Magnification = 420X ; Scale : 1 cm = 24 μ m



Figure 22a



Figure 22b



Figure 22c



Figure 22d

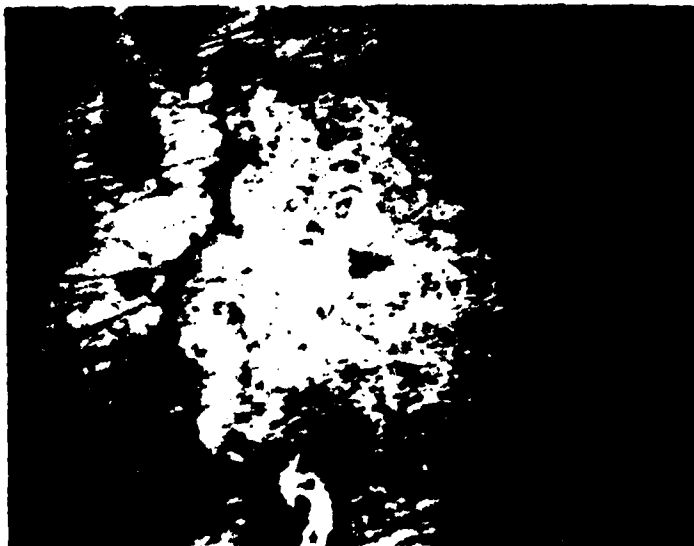


Figure 22e

0.1 volt to see a change of about 1 nA for this sample. Table 6 shows Δi versus ΔV for two samples having different compositions of ITO.

Table 6

Effect of Increased Indium Content in ITO

Sample	Growth Process	$\Delta V = V_A - V_{Rest} $ volts	Δi amps	In/Sn Ratio
Al-SiO ₂ -ITO	Spray Injection CVD ITO	0.06	1×10^{-6}	= 0.6
		0.10	80×10^{-6}	
		0.20	0.78×10^{-3}	
Al-SiO ₂ -ITO-Si ₃ N ₄	CVD ITO	0.6	188×10^{-6}	0.002
		0.10	430×10^{-6}	
		0.20	1.55×10^{-3}	
Al-SiO ₂ -Si ₃ N ₄	Plasma Deposited Nitride	0.06	385×10^{-6}	
		0.10	1185×10^{-6}	
		0.20	3.20×10^{-3}	

The change in current Δi for a given change in voltage ΔV has been found to be greater in the case of air-sprayed ITO in comparison to the CVD spray (see Fig. 16 and 17).

An important feature, observed in samples exhibiting increased rest potential due to good ITO coatings, is the phenomenon of current lowering (passivation) in spite of an increase in the applied voltage. This is observed in Fig. 16 (Al-SiO₂-ITO sample), Fig. 17 (Al-SiO₂-ITO) and Fig. 19 (Al-SiO₂-ITO/TO/Si₃N₄ sample).

VI. CONCLUSIONS

The use of an active electronic barrier in corrosion prevention is a novel concept⁽¹⁷⁾, and the validity of this approach has been demonstrated using Al-ITO and Al-SiO₂-ITO samples. The protective nature of the active electronic barrier in Al-SiO₂-ITO films is clearly evident when the polarization

results of these samples are compared with Al-Si₃N₄ (plasma-deposited) passive structures. The ITO coated samples with a film thickness of only 0.2 microns yielded lower currents than the Si₃N₄ coated (0.75 microns) samples. Weight-loss measurements further verify the concept of an active electronic barrier for use in corrosion prevention.

ITO films containing more indium have been found to have an increased electronic barrier at the Al/ITO interface. This has been shown in both the voltage-current characteristics and the anodic polarization data. The increased rest potential in the anodic polarization data has confirmed the increased active barrier with In rich films. The increase in the indium content has clearly shown a corresponding increase in the energy gap of the ITO films. This has been confirmed by spectrophotometric measurements.

A commercially viable processing system has been simulated by the deposition of ITO films using an injection spray technique. This has been done in a controlled ambient as well as in air. Results of the controlled ambient system have been better than the air spray system, and quite comparable to the chemical vapor deposition technique developed earlier in our laboratory. However, further process optimization is required to obtain high quality ITO films with a minimum number of pinholes.

The concept of active electronic barriers could also be applied to other semiconductor coatings on metals such as silicides, doped polyacetylene, phthalocyanine, chlorophyll, porphyrins, etc.

VII. FUTURE WORK

A. Controlled Doping of ITO

High quality films of ITO and SnO₂ have been grown in both n and p-type dopings. However, the control over the amount of doping, necessary to obtain high resistivity films of n and p-type, has been poor. Current films of each type exhibit low resistivities (10^{-2} to 10^{-3} Ω .cm). These low resistivity films do form an effective electronic barrier at the aluminum surface, however, the corrosion protection can be further enhanced by increasing the film resistivity.

In addition, the controlled doping will allow for the optimization of the active electronic barrier.

B. Growth of Thicker Homogeneous Films

The growth of thicker ITO films (> 1.0μm) is necessary to minimize the occurrence of microscopic pinholes. Extremely uniform films of ITO have been grown on Si wafers using the CVD spray deposition system. However, ITO films grown on aluminum substrates have been of lower quality. It is suspected that it is due to rapid cooling of the aluminum surface during the spray deposition. Additional work is needed in the reactor design to improve film quality and homogeneity.

C. Current Transport Modeling

A theoretical model of the current transport in Al-SiO₂-ITO structures is needed to characterize the effective height of the barrier formed at the interface.

A relationship to predict effective barrier height in terms of the ITO energy gap E_g and the location of defect trap levels is needed. This model will be general enough to be of use for other systems such as Al-polyacetylene, Al-phthalocyanine, etc. The model would also be used to obtain relationships for the corrosion current. In addition, the nature of the ITO-electrolyte interface (as in polarization testing) must be studied. This information would be useful in understanding the effects of the ITO-ambient interface.

D. Investigation of Multiple ITO/TO Coatings

Preliminary investigation suggests that the semiconductor heterointerface formed between ITO and SnO_2 layers results in a second active electronic barrier which enhances the corrosion resistance. Further work is needed to characterize this phenomena.

A C K N O W L E D G E M E N T S

The author wishes to acknowledge J. J. Rosato and K. S. Kalonia for their assistance during the course of this study. Thanks are due to Dr. V. S. Agarwala for helpful discussions and avowed interest in the project. He is grateful to J. G. Bouchard(Sprague Electric Company, Worcester, MA) for the deposition of nitride films.

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