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Memorandum of Project MICHIGAN

EFFECT OF CHEMISORBED OXYGEN ON PHOTOVOLTAIC AND PHOTOCONDUCTIVE PROCESSES IN RUTILE

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INFRARED LABORATORY
Institute of Science and Technology
THE UNIVERSITY OF MICHIGAN

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PREFACE

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The emphasis of the Project is upon research in imaging radar, MTI radar, infrared, radio location, image processing, and special investigations. Particular attention is given to all-weather, long-range, high-resolution sensory and location techniques.

Project MICHIGAN was established by the U. S. Army Signal Corps at The University of Michigan in 1953 and has received continuing support from the U. S. Army. The Project constitutes a major portion of the diversified program of research conducted by the Institute of Science and Technology in order to make available to government and industry the resources of The University of Michigan and to broaden the educational opportunities for students in the scientific and engineering disciplines.

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Progress and results described in reports are continually reassessed by Project MICHIGAN. Comments and suggestions from readers are invited.

Robert L. Hess
Director
Project MICHIGAN
SYMBOLS

t = time
k = conductance
Z = excitation rate
D = desorption rate of physically adsorbed atoms from surface to ambient pressures
b = a constant, dependent on the barrier layer formed by adsorbed atoms
p = the partial pressure of gas in equilibrium with the physically adsorbed atoms
C, A = constants
EFFECT OF CHEMISORBED OXYGEN ON PHOTOVOLTAIC AND PHOTOCONDUCTIVE PROCESSES IN RUTILE

ABSTRACT

It has been found that the presence of adsorbed oxygen increases both the response and the response time for the photovoltaic effect in rutile.

The model for the effect of chemisorbed oxygen on the photoconductivity has been modified to take into consideration the effect of oxygen pressure. The conductance of rutile has been measured as a function of oxygen pressure. The results agree with the modified model.

The increase in photovoltaic response and response time with exposure to oxygen has been explained in terms of the barrier layer formed by chemisorbed oxygen.

1 INTRODUCTION

During investigation of photovoltaic effects in rutile [1], it was noted that both the response time and the magnitude of response of a rutile photovoltaic device were affected by exposing the device to the atmosphere.

Figure 1 shows the configuration of the device. It was fabricated as follows. A polished single-crystal disk of rutile was reduced in a hydrogen environment at 600°C for two hours. The surface was then reoxidized by heating in air for ten minutes at approximately 600°C. A semitransparent silver electrode was evaporated onto one face, and a thick indium electrode was evaporated onto the opposite face.

2 EFFECT OF OXYGEN ON PHOTOVOLTAIC CHARACTERISTICS

The rise and decay of the photovoltage, as a result of pulsed ultraviolet radiation incident on the sample, was observed with an oscilloscope. Figure 2(a) shows the rise and decay characteristics of a sample which had been exposed to laboratory conditions for several days. Figure 2(b) illustrates the change in these characteristics as a result of pumping on the sample.
FIGURE 1. CONFIGURATION OF PHOTOVOLTAIC DEVICE

FIGURE 2. EFFECT OF OXYGEN ON PHOTOVOLTAIC CHARACTERISTICS.
(a) In air. (b) In Vacuum.
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continuously for five hours at $10^{-4}$ mm Hg. The response time decreased from 1 msec to
1/2 msec, while the amplitude of the photosignal decreased by a factor of two. When the sam-
ple was exposed to air again, the response time and photosignal returned to their original
values. This procedure was repeated several times with the same results; hence, the process
appears to be reversible.

It was also observed that the rate at which the characteristics changed in vacuum depend-
ted to some extent on whether or not the sample was irradiated with ultraviolet light.

### 3 PHOTOCONDUCTIVITY AND CHEMISORPTION

Miller [2] has discussed the effects of chemisorption on the performance of photocon-
ductive devices. Melnick [3] and Morrison [4] have proposed a model for chemisorption on
semiconductor surfaces during exposure to radiation. The model applies, in particular, to
the interaction of oxygen with an n-type semiconductor such as rutile. They suggest that an
oxygen atom is physically adsorbed on the surface of the semiconductor. There it combines
with an electron from the valence band, thus becoming chemisorbed. Upon irradiation, holes
and electrons are produced near the surface. The holes combine with a chemisorbed atom
which is changed, at least transiently, to a physically adsorbed atom. The physically adsorbed
atoms may then either combine with an electron or diffuse from the surface into the ambient
gas. These phenomena cause a change in the free-carrier concentration, and hence a change
in the conductance of the semiconductor.

The differential equation proposed by Melnick for the change of conductance on exposure
to both oxygen and radiation is:

$$\frac{dk}{dt} = Z - D \exp(bk) + D$$  \hspace{1cm} (1)

where $t =$ time

$k =$ conductance

$Z =$ excitation rate

$D =$ desorption rate of physically adsorbed atoms from surface to ambient pressures

$b =$ a constant dependent on the barrier layer formed by adsorbed atoms

Equation 1 was used to derive the following equation for the photoconductive decay.
\[ \Delta k = b^{-1} \ln (tbZ + 1) \]  

Melnick [3], Medvid [5], and Elovich [6] have found that the photoconductive decay of zinc oxide follows Equation 2.

Another approach to this model is as follows. At equilibrium, \( \frac{dk}{dt} = 0 \). Therefore, at equilibrium, Equation 1 becomes

\[ k = b^{-1} \ln[(Z + D)D^{-1}] \]  

If one assumes the excitation rate \( Z \) to be large compared to the equilibrium desorption rate \( D \), Equation 3 becomes

\[ k = b^{-1} \ln ZD^{-1} \]  

**RESULTS**

In order to introduce the pressure as a variable, one may assume that the desorption rate is dependent on \( p \), the partial pressure of gas in equilibrium with the physically adsorbed atoms. The Langmuir dependency

\[ D = CAp/(1 + Ap) \]  

may be used where \( C \) and \( A \) are constants. Combining Equations 4 and 5, one obtains

\[ k = -b^{-1} \ln p + b^{-1} \ln[Z(1 + Ap)/AC] \]  

For many systems \( A \) is of the order of \( 10^{-4} \) mm\(^{-1} \). Therefore, at low pressures the second term in Equation 6 is a constant, and it may be constant up to several atmospheres.

A single crystal of rutile was cut and polished into a parallelepiped of dimensions 1.6 x 3.6 x 15 mm. Indium electrodes were evaporated onto the ends of the sample. The sample was placed in a dewar and irradiated continuously with ultraviolet light. The conductance at equilibrium was measured as a function of oxygen pressure. Figure 3 shows the results of these measurements. The conductance as a function of pressure follows Equation 6 over 5 orders of magnitude of pressure.

The proposed model predicts a slow change in conductance with pressure if the excitation rate is zero, since, without photo excitation, only thermally excited holes are available to
FIGURE 3. EFFECT OF OXYGEN ON PHOTOCONDUCTIVITY
convert the chemisorbed atoms to physically adsorbed atoms.

It was indeed found that if the sample was kept in the dark there was no change with pressure in the equilibrium conductance over the range of pressures and times used in these experiments.

5 CONCLUSIONS

It appears that oxygen is chemisorbed on the surface of rutile. The observed increase in both the magnitude of the photovoltaic response and the response time in rutile (see Figure 2) may now be explained on the basis of the chemisorption of oxygen on rutile, as discussed above.

According to Hauffle and Pfeiffer [7], chemisorbed oxygen on the surface would produce a barrier layer of the Schottky-Mott type. This barrier should change the magnitude of the barrier formed by the silver-to-rutile contact. The increase in the photovoltaic signal indicates that this barrier is increased by the chemisorption of oxygen.

The chemisorbed oxygen layer would also increase the resistance of the device by reducing the electron concentration. Since it appears that the response time for these photovoltaic devices increases with increasing resistance [1], one would expect an increase in response time with increasing chemisorbed oxygen.

REFERENCES

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