# **General Disclaimer**

# One or more of the Following Statements may affect this Document

- This document has been reproduced from the best copy furnished by the organizational source. It is being released in the interest of making available as much information as possible.
- This document may contain data, which exceeds the sheet parameters. It was furnished in this condition by the organizational source and is the best copy available.
- This document may contain tone-on-tone or color graphs, charts and/or pictures, which have been reproduced in black and white.
- This document is paginated as submitted by the original source.
- Portions of this document are not fully legible due to the historical nature of some of the material. However, it is the best reproduction available from the original submission.

Produced by the NASA Center for Aerospace Information (CASI)

NASA Technical Memorandum 83711

# Surface Effects of Corrosive Media on Hardness, Friction, and Wear of Materials

(NASA-TM-83711) SURFACE EFFECTS OF N85-15896 CORROSIVE MEDIA ON HARDNESS, FRICTION, AND WEAR OF MATERIALS (NASA) 20 p HC A02/MF A01 CSCL 116 Unclas G3/27 13453

Kazuhisa Miyoshi, Donald H. Buckley, George W.P. Rengstorff, and Hiroyuki Ishigaki Lewis Research Center Cleveland, Ohio

NASA



Prepared for the International Conference on Wear of Materials—1985 sponsored by the American Society of Mechanical Engineers and supported by ASTM, AIME, ASM, ACerS, ASLE, and ACOT Vancouver, Eritish Columbia, April 14-18, 1985

## SURFACE EFFECTS OF CORROSIVE MEDIA ON HARDNESS, FRICTION,

AND WEAR OF MATERIALS

Kazuhisa Miyoshi, Donald H. Buckley, George W.P. Rengstorff,\* and Hiroyuki Ishigaki\*\* National Aeronautics and Space Administration Lewis Research Center Clevelang, Ohio 44135

#### ABSTRACT

Hardness, friction, and wear experiments were con-ducted with magnesium oxide exposed to various corrosive media including two acids, a base, and a salt and with elemental iron and nickel exposed to water and various concentrations of NaOH. Chlorides such as MgCl<sub>2</sub> and sodium-containing films were formed on cleaved magnesium oxide surfaces from their interactions with HC1containing solutions. The MgCl<sub>2</sub> films softened the magnesium oxide surfaces and caused high friction and great deformation. Hardness was strongly influenced by the pH value of the HCl-containing solution. The lower the pH, the lower the microhardness. Neither the pH value of nor the immersion time in NaOH-containing, NaCl-containing, and HNO3-containing solutions influ-enced the microhardness of magnesium oxide. NaOH formed a protective and low-friction film on iron surfaces. The film was broken up by sliding action at low concentrations but remained on the surfaces at high concentrations. The coefficient of friction and the wear for iron were low at concentrations of NaOH higher than 0.01 N. An increase in NaOH concentration resulted in a decrease in the concentration of ferric oxide (mainly Fe2O3) on the iron surface. It took less NaOH to form a protective, low-friction film on nickel than on A small amount of NaOH on nickel gave a low iron. coefficient of friction and low wear damage, and the values did not change with NaOH concentration. In water alone the nickel surface sustained considerable wear damage.

#### INTRODUCTION

The presence of surface-active agents on metals and nonmetals can alter the surface activity of these materials as well as their mechanical properties (1,2). Tribological properties such as adhesion, friction, deformation, wear, and lubrication of solid surfaces in contact are extremely dependent on the adsorbed surface-active ions or molecules (i.e., environmental constituents).

Merceletter Astronetering

Ē.

10000

.

ų

It was anticipated that the hardness of solids would be influenced by environment because hardness is one of the measurements used to indicate the extent of surficial plastic deformation. Many investigators have studied the effects of environment on the hardness of a variety of nonmetals such as CaF2, Al2O3, and LiF (2-5). These effects include strengthening by dissolution of the solid surface, the Joffe effect (6); surface hardening, the Roscoe effect (7,8); surface softening, the Rehbinder effect (9); correlation between hardness and the zeta-potential (10,11); and the effect of adsorbed water on indentation creep (5,12).

Since environmental effects on the hardness of nonmetals are so surface dependent, the present authors took a different analytical approach to determining environmental effects on hardness. A surface analytical tool (i.e., x-ray photoelectron spectroscopy (XPS)) was used to study the surficial or outermost atomic layers of solids.

The first objective of the present paper is to examine the surface chemistry, hardness, friction, and wear of a ceramic (single-crystal magnesium oxide) exposed to various corrosive media including two acids, a base, and a salt.

Although corrosion is recognized as an important variable in the friction and wear of metals (13), its role is not well understood. In the sliding, rolling, or rubbing contact of materials the surfaces become strained as a result of the mechanical activity that takes place. A wear surface is different electrochemically from its surroundings. It contains metal that is highly strained and that reaches locally high temperatures at shearing points or asperities (14,15). Electrochemical potentials may be established locally to either impede or enhance corrosion; cyclic stresses may promote stress corrosion and corrosive fatigue (16). Resistance to corrosion is often the result of the

Resistance to corrosion is often the result of the formation of some type of film on the metal. Sliding action can destroy such films - or it can develop better corrosion-resistant films by producing new surfaces. The coefficient of friction is, like corrosion resistance, highly sensitive to surface films.

<sup>\*</sup>Summer Faculty Fellow. \*\*NRC-NASA Research Associate.

The second objective of this paper is to examine the surface chemistry, hardness, friction, and wear of elemental iron and nickel exposed to various concentrations of NaOH and to water, as well as to analyze their surface chemistry by XPS.

#### MATERIALS

The arc-melted, single-crystal magnesium oxide used was 99.99 percent pure as determined from the manufacturer's data. Magnesium oxide was selected as the material to examine for a number of reasons: its slip and fracture behavior are well understood, the Rehbinder effect has been observed with it, fresh, atomically clean surfaces can be prepared by cleavage, and it can be used as a bearing material (17).

The iron used was more than 99.99 percent pure and had been fully annealed. Its hardness was Rockwell B40. The nickel used was electrolytic and had been annealed to a hardness of Rockwell B30.

### **APPARATUS**

The apparatuses used in this investigation have been described previously (18-22). The Vickers microhardness tester was fixed on a vibration-free table and measured indentation hardness in a controlled environment (18,19). Specimens were indented manually with a diamond indenter. The friction and wear apparatuses were basically a pin (rider) on a flat (e.g., Fig. 1). An ultra-high-vacuum system containing the XPS system and an ion gun (18-20) was used for ion sputter-etching.

#### EXPERIMENTAL PROCEDURE

#### Hardness, Friction, and Wear Experiments on Magnesium Oxide

The magnesium oxide bulk crystals were immersed in room-temperature aqueous solutions with various pH values and then cleaved to a specimen size of 5 mm by 5 mm along the (001) surface by using a knife and hammer. The submerged specimens were then mounted on the Vickers microhardness tester and the friction apparatus. Indentations and sliding friction experiments were made on cleaved (001) surfaces in the desired aqueous solution and in water without exposing the specimen surface directly to the air or to any other environment. For comparison tests the magnesium oxide bulk crystals were similarly prepared in laboratory air at room temperature.

The indentation diagonals were parallel to the <001> directions on the (001) surfaces. The time in contact was 20 s. A hemispherical diamond rider (20-µm radius) was made to slide in the <001> directions on the magnesium oxide (001) surfaces under a load of 0.25 N and at a sliding velocity of 10 mm/s.

Table I lists the aqueous solutions used in this investigation and their analyzed pH values. The solutions contained NaOH, NaCl, HNO3, and HCl.

#### Hardness, Friction, and Wear Experiments on Iron and Nickel

Tron and nickel flats were metallographically lapped with 600-grit silicon carbide paper. They were then polished with diamond paste, first 6-um paste and then  $3_{-\mu m}$  paste, or with  $3_{-\mu m}$  paste alone. Final pol-ishing was with  $0.3_{-\mu m}$  alpha alumina on a flooded, flat polishing wheel. The polished specimen was cleaned with water and then with ethanol. The specimens were subse-quently etched until the residual deformed region was removed and the grain structure was visible. For iron the etchant was 5 percent Nital, and for nickel it was 1 part nitric acid to 1 part acetic acid (glacial). The specimen was then lightly polished with the 0.3- $\mu$ m alumina and cleaned with distilled water and ethanol.

情况在中国主义的学习学习

Before the Vickers microhardness measurements drops of the desired solution were placed on the flat metal surface. After 1-hr exposure an indentation was made with a diamond Vickers indenter through the drops. T The metal specimen and the diamond indenter were then cleaned with distilled water and ethanol. The indenta-tion diagonals were measured right after cleaning. The hardnesses of iron and nickel were also measured in laboratory air. All indentations were made at a load of 0.1 N at room temperature. The time in contact was 20 s. Each hardness value is the average of 8 to 12 measurements.

In the sliding friction experiments the specimens were clamped in the vise of the apparatus. Several drops of the desired solution were placed under the rider and on the flat surface to form one large drop covering most of the surface of the flat specimen. The rider, a 6.4-mm-diameter Al203 (sapphire) sphere, was lowered through the drop until a strain gauge indicated that it was just in contact with the metal flat surface. The weight was then loaded onto the rider and sliding begun. The rider and its point of contact with the specimen were always in the interior of a large liquid drop. Furthermore a meniscus formed between the rider and the flat specimen provided a continuous supply of solution to wet the flat specimen in and near the wear track. The rider slid in reciprocating motion, on the iron and nickel flats at 1.5 mm/s, at a load of 2.5 N (250 g), at room temperature. Each experiment lasted 60 min. Thus the rider passed over the track 540 times (270 times each way) during an experiment.

To estimate the wear characteristics of iron or nickel, their wear tracks were scanned by a surface profilometer.

<u>XPS Analysis</u> For XPS analysis (18-20) the specimens were placed in the vacuum chamber, and the system was evacuated and subsequently baked out to 26 nPa ( $2x10^{-10}$  torr). The surfaces were examined with a magnesium ka x ray at a source power of 400 W (10 kV, 40 mA). The analyzer work function was determined by assuming the binding energy for the gold 4f 7/2 peak to be 83.8 eV; that is, the gold 4f 7/2 peak was used as the reference line.

#### RESULTS AND DISCUSSION

Hardness, Friction and Wear of Magnesium Oxide The distribution of dislocation etch pits shown in Fig. 2 is typical for indentations made on magnesium oxide surfaces cleaved in air. Because magnesium oxide is highly ionic, slip occurs not on the close-packed (001) cubic planes but on the (011) cubic diagindentation on the (001) surface (Fig. 2) contains screw dislocations in the <001> directions and edge dislocations in the <O11> directions. The arrays of screw and edge dislocations were 4.9 and 7.7 times wider, respectively, than the average length of the two diagonals of indentation.

The Vickers microhardnesses of magnesium oxide surfaces cleaved in air and in water were essentially the same at 824 and 822, respectively. Indentation experiments were also conducted in corrosive solutions containing NaOH, NaCl, HNO3, and HCl with various pH values. The microhardness as a function of the time that the magnesium oxide crystal was immersed in the corrosive solution is given in Fig. 3. Each hardness value is the average of 10 measurements. There were no changes in microhardness with immersion time for surfaces immersed in solutions containing NaOH (pH = 13.5),

NaCl (pH = 6.7), and HNO<sub>3</sub> (pH = 1.0). The pH value had little effect on microhardness: the difference in microhardnesses of the surfaces immersed in these solutions was relatively small. However, the surfaces immersed in HCl-containing solutions with a pH or 1.0 exhibited both low microhardness and a slight change in microhardness with immersion time. These microhardness data could not be taken beyond an immersion time of 7200 s because the magnesium oxide surfaces were considerably roughened.

To gain an understanding of the surfaces of magnesium oxide in the HCl-containing solution, indentation experiments were conducted in HCl-containing solutions with pH values of 4.2, 2.5, and 1.0. The data (Fig. 4) indicate that the microhardness of a magnesium oxide surface immersed in HCl solutions is influenced by the pH value as well as by the immersion time. The stronger the acid, the lower the microhardness observed. The microhardness of the surface immersed in the HCl solution with a pH of 1.0 for 900 s was almost the same as those of surfaces immersed in solutions with pH values of 4.2 and 2.5 for 3.5x10<sup>5</sup> s. There was a significance in the differences in the

distribution length of edge dislocations for the HCl solution (pH = 1.0), the NaCl solution (pH = 6.7), and the NaOH solution (pH = 13.5). The lengths of the rows of etch pits on the solid surface reflect the extent of deformation. The longer the rows, the greater the deformation and the lower the microhardness. The distribution lengths of the edge dislocations are consist-ent with the microhardness data shown in Fig. 3 (18,19).

XPS spectra of  $Cl_{2p}$ ,  $Na_{1s}$ ,  $Mg_{2p}$ , and  $O_{1s}$ were obtained from narrow scans takenon the magnesium oxide surfaces cleaved in  $10^{-2}$  N aqueous NaCl solutions containing HCl (pH = 1.0) or NaOH (pH = 13.5) before and after argon-ion sputter etching for 30 min (18,19). Typical results are presented in Figs. 5 and 6. Chlorine, sodium, and magnesium peaks for the sur-faces exposed to the HCl solution can be seen (Fig. 5) before and after sputtering. The solium peak was larger after sputtering. Oxygen peaks were also evident on surfaces exposed to the acid solution before and after sputtering (19).

Figure 5 reveals that chemically reacted chlorideand sodium-containing films formed as a result of the HCl solution interacting with the magnesium oxide surface and deeply in the subsurface. Chlorine chemically reacted with the magnesium to form compounds such as  ${\rm MgCl}_2$  in the surficial layer. Even after 30 min of sputtering these chemically reacted films were not removed; that is, the interactions occurred not only on the surface, but also in the subsurface. The chloride films played a role in softening the magnesium oxide surface (Figs. 3 and 4).

A chlorine peak is clearly evident (Fig. 6(a)) in the spectrum of the magnesium oxide surface exposed to the NaOH solution (pH = 13.5) before sputtering, no chlorine is apparent after sputtering. Although sodium peaks appear both before and after sputtering (Fig. 6(b)), the amount of sodium was decreased by sput-tering. No magnesium peak is observable before sputtering, but afterward a magnesium peak does appear in the spectrum.

The magnesium oxide surface cleaved in this alkali solution was covered with chloride- and sodiumcontaining compound films. Its outermost surficial layer consisted of mostly NaOH and NaCl films with no magnesium oxide exposed. However, argon ion sputtering completely removed the chloride films from the surface and partially removed the sodium films, exposing the magnesium oxide. These sodium-containing films played a slight role in softening or hardening the magnesium oxide surface (Fig. 3).

Sliding friction experiments were conducted with a diamond rider having a 20-µm-radius hemispherical tip in contact with magnesium oxide surfaces exposed to an HCl solution (pH = 1.0), an NaOH solution (pH = 13.5), water, and air. Plastic deformation occurred and permanent grooves were formed in the surfaces during sliding. The coefficients of triction and the ground were influenced by the solutions and environments The coefficients of friction and the groove widths (Table II). When magnesium oxide was exposed to the HC1 solution, the MgCl2 (as detected by XPS analysis) formed in its surficial layer caused greater plastic flow and plowing action and resulted in higher friction and greater groove width than for magnesium oxide exposed to the NaOH solution, water, and air. The dif ferences for magnesium oxide exposed to the NaOH solu-The diftion, water, and air were small. Thus, the friction and deformation of magnesium oxide, as well as its hardness are markedly affected by exposure to an HCl solution.

清朝日朝

Hardness, Friction, and Wear of Iron and Nickel The Vickers microhardnesses of Iron and nickel were measured as a function of NaOH concentration. At 0.001 N NaOH the hardness of iron (Fig. 7(a)) was higher than that measured in laboratory air (81.5), increased to a maximum at 0.01 N NaOH, and continuously decreased with increasing NaOH concentration.

The Vickers microhardness of nickel exposed to NaOH solutions was also influenced by the concentration of NaOH. At 0.001 N NaOH the hardness of nickel was almost the same as that measured in laboratory air (88.8). The higher the concentration, the lower the microhardness observed.

Friction and wear experiments were conducted with iron and nickel flats in contact with 6.4-mm-diameter Al<sub>2</sub>O<sub>3</sub> spherical riders in NaOH at a load of 2.5 N for 50 min. There were marked differences in the friction and wear behavior of the two metals. For iron (Fig. 8) the maximum coefficient of friction and wear track width occurred at 0.01 N NaOH. The coefficient of friction continuously decreased with increasing NaOH  $\,$ concentration, but the wear initially decreased and then leveled off at 0.4 N NaOH. On the other hand, the coefficient of friction and wear track width for nickel (Fig. 9) did not change with NaOH concentration over the entire range from 0.001 to 20 N. The coefficient of friction was low under all of these conditions, as is consistent with nickel being highly resistant to corrosion by NaOH.

Because the difference in wear between iron and nickel under corrosion conditions is important, iron and nickel wear surfaces were examined in more detail by using optical and scanning electron microscopy and a surface profilometer (Figs. 10 to 12). At 0.001 N and 0.01 N NaOH overall corrosive attack was low, but con-siderable wear was indicated on the iron. At 0.001 N NaOH the pitting and grain boundary attack inherent in caustic embrittlement were caused by the applied and residual stresses in the wear track. Pits were seen in the optical microscope as well as in the surface profile of the wear track (Fig. 10). Iron exhibited the highest friction and wear at 0.01 N NaOH. Although there was no evidence of deep pits, the wear track surface was severely damaged (Fig. 11).

The friction and wear of iron were high in dilute NaOH but low in concentrated NaOH. In 0.4 N to 20 N NaOH (Fig. 10) the wear surfaces were smooth, as also observed in optical and scanning electron micrographs (20,21). The sliding action left a permanent groove on the wear surface, with deformed iron piled up along the sides of the groove. The amount of iron removed, how-ever, was much less than for dilute NaOH solutions. The The tenacious film formed was not broken up by the sliding action, and this film acted as a lubricant.

3

. وما تخلف المتحت .

Sliding the Al<sub>2</sub>O<sub>3</sub> sphere on a nickel surface in O.1 N NaOH solution (Fig. 12) resulted in a permanent groove on the nickel surface with a considerable amount of deformed nickel piled up along the sides of the groove. The wear track on nickel was rough and irregu-lar, but the surface was not damaged as it was for iron.

Sliding the Al<sub>2</sub>O<sub>3</sub> sphere on a nickel surface in water created a highly torn and distorted surface in the wear track (Fig. I3). Corrosion principles suggest that a tenacious film protects nickel from corrosion by water but that this film might be disrupted in the wear track.

XPS spectra for wear surfaces of iron in 0.001 N, 0.01 N, and 20 N NaOH were compared (Fig. 14). 0xygen bonded directly to iron (as in Fe<sub>2</sub>O<sub>3</sub>) had an oxygen 1s peak at 530 eV (Fig. 14(a)). The height of this peak decreased with increasing NaOH concentration. The oxygen peak at approximately 532 eV is generally attributed to a bond involving hydrogen as well as iron (i.e., Fe-O-H, such as in FeODH). This peak is dominant in 20 N NaOH and gets relatively weaker with decreasing NaOH concenon iron in NaOH solutions (Fig. 14(b)). No sodium peaks were detected in any scans between

1068 and 1082 eV with NaOH concentrations even up to 20 N.

#### CONCLUSIONS

The following conclusions were drawn from the data presented herein on the tribological characteristics of magnesium oxide, iron, and nickel in corrosive media.

#### Magnesium Oxide

1. MgCl2 films are formed on cleaved magnesium oxide surfaces as a result of an HCl-containing solution interacting with the surface. The MgCl<sub>2</sub> films soften the magnesium oxide surfaces and cause higher friction and greater deformation. The pH value and the immersion time in HCl solutions strongly influence microhardness. The smaller the pH value and the longer the immersion time, the lower the microhardness.

2. There is almost no change in microhardness with immersion time on magnesium oxide surfaces immersed in NaCl solution containing NaOH with a pH of 13.5, in water containing NaCl with a pH of 6.7, and in water containing HNO<sub>3</sub> with a pH of 1.0. The difference in hardness between these surfaces is also relatively small.

3. Sodium-containing films are formed on cleaved magnesium oxide surfaces as a result of an NaOHcontaining solution interacting with the surface. oxide surface, and the pH value of the NaOH solutions does not influence its microhardness.

#### Iron and Nickel

1. In low NaOH concentrations, such as 0.01 N, the friction for iron is high and the wear track is torn up and considerably damaged. The hardness, friction, and wear of iron peak at 0.01 N NaOH and decrease with increasing concentrations of NaOH. This decrease is accompanied by a decrease in the concentration of ferric oxides on the iron surface and an increase in a tenacious film composed of complex iron-oxygen compounds.

2. The general conclusion is that NaOH forms a protective, low-friction film on iron that is destroyed by wear at low NaOH concentrations but remains at high concentrations.

3. Nickel behaves differently from iron in that only a little NaOH gives low values of hardness, coefficient of friction, and wear damage.

#### REFERENCES

A SALES AND A SALES AND A SALES

- Miyoshi, K. and Buckley, D.H., "Effect of Oxygen and Nitrogen Interactions on Friction of Single-Crystal Silicon Carbide," NASA TP-1265, Aug. 1978.
   Westwood, A.R.C. and Goldheim, D.L., "Occurrence and Mechanism of Rehbinder Effects in CaF2," Journal of Applied Physics, Vol. 39, No. 7, June 1989 and 2401 2401
- 1968, pp. 3401-3405. Westwood, A.R.C., "The Rehbinder Effect and the Adsorption-Locking of Dislocations in Lithium
- Adsorption-Locking of Dislocations in Lithium Fluoride," <u>Philosophical Magazine</u>, Vol. 7, No. 76, Apr. 1962, pp. 633-649. Gilman, J.J., "The Mechanism of Surface Effects in Crystal Plasticity," <u>Philosophical Magazine</u>, Vol. 6, No. 61, Jan. 1961, pp. 159-161. Westbrook, J.H. and Jorgensen, P.J., "Indentation Creep of Solids," <u>Transactions of the Metallurgical Society of AIME</u>, Vol. 233, No. 2, Feb. 1965, pp. 425-428 425-428.
- Joffe, A.F., <u>The Physics of Crystals</u>, 1st ed., McGraw-Hill, <u>New York</u>, 1928. Roscoe, R., "The Plastic Deformation of Cadmium Single Crystals," <u>Philosophical Magazine</u>, Vol. 21, 7. 1936, pp. 399-406.
- Harper, S. and Cottrell, A.H., "Surface Effects and the Plasticity of Zinc Crystals," <u>Proceedings of</u> the Physical Society, London, Section B, Vol. 63, May 1950, pp. 331-338. Rehbinder, P.A. and Shchukin, E.D., "Surface Phe-8.
- nomena in Solids During Deformation and Fracture Processes," <u>Progress in Surface Science</u>, Vol. 3, Part 2, 1972, pp. 97-188. Macmillan, N.H., Huntington, R.D., and Westwood,
- 10. A.R.C., "Relationship Between c-Potential and Dislocation Mobility," <u>Philosophical Magazine</u> Vol. 28, No. 4, Oct. 1973, pp. 923-931.
- Macmillan, N.H. and Westwood, A.R.C., "Surface Charge-Dependent Mechanical Behavior of Non-Metals, 11. "Surfaces and Interfaces of Glass and Ceramics,
- Plenum, New York, 1974, pp. 493-513. Westbrook, J.H. and Jorgensen, P.J., "Effects of Adsorbed Water on Indentation Hardness Anisotropy in Crystals," Anisotropy in Single-Crystal Refrac-tory Compounds, Vol. 2, Plenum, New York, 1958,
- tory Compounds, Vol. 2, Plenum, New York, 1968, pp. 353-360. Eyre, T.S., "Wear Characteristics of Metals," Source Book on Wear Control Technology, American Society for Metals, Metals Park, OH, 1978, pp. 1-10; Tribology International, Vol. 9, No. 5, Oct. 1976, pp. 203-212. Bowden, F.P. and Tabor, D., "The Friction and Lubrication of Solids, Vol. 2, Clarendon Press, Overand 1964 13.
- Oxford, 1964. Rabinowicz, E., <u>Friction and Wear of Materials</u>.
- 15.
- Wiley, New York, 1965. Staehle, R.W., "Stress Corrosion Cracking (and Corrosion Fatigue)," <u>Materials Science and</u> Engineering, Vol. 25, No. 1/2, Sept./Oct. 1976, pp. 207-215.
- Dufrane, K.F. and Glaeser, W.A., "Study of Rolling-Contact Phenomena in Magnesium Oxide," NASA 17.
- CR-72295, Sept. 1967. Miyoshi, K., Ishigaki, H., and Buckley D.H., "Influence of Mineral Oil and Additives on Micro-hardness and Surface Chemistry of Magnesium Oxide 18.
- (001) Surface," NASA TP-1986, Mar. 1982. Ishigaki, H., Miyoshi, K., and Buckley, D.H., "Influence of Corrosive Solutions on Microhardness 19. and Chemistry of Magnesium Oxide (001) Surfaces," NASA TP-2040, Aug. 1982.

Ishigaki, H., Miyoshi, K., and Buckley, D.H., "Influence of Corrosive Solutions on Microhardness and Chemistry of Magnesium Oxide (OO1) Sur-faces," NASA TP-2040, Aug. 1982.
 Rengstorff, G.W.P., Miyoshi, K., and Buckley, D.H., "Friction and Wear of Iron in Corrosive Media," NASA TP-1985, Mar. 1982.

- 24 1

- Rengstorff, G.W.P., Miyoshi, K., and Buckley, D.H., "Friction and Wear of Iron and Nickel in Sudium Hydroxide Solutions," <u>ASLE Transactions</u>, Vol. 26, No. 4, Oct. 1983, pp. 509-516.
   Ishigaki, H. and Buckley, D.H., "Effects of Envi-ronment on Microhardness of Magnesium Oxide," NASA TP-2002, Apr. 1982.

৵

사망

常

Solution	pH value <sup>3</sup>
HCl added to $10^{-2}$ N aqueous NaCl solution	1.0 2.5 4.2
$10^{-2}$ N aqueous NaCl solution	6.7
NaOH added to $10^{-2}$ N aqueous NaCl solution	10.5 12.3 13.5
HNO3 added to distilled water	1.0

TABLE I. - SOLUTIONS WITH VARIOUS pH VALUES

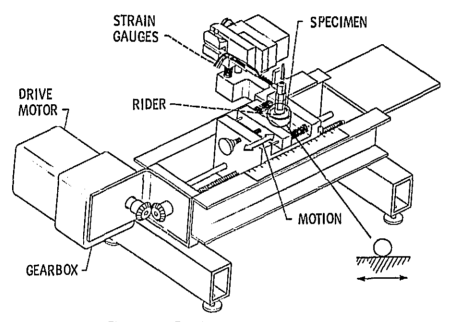
THE REPORT

<sup>a</sup>Potentiometric method was employed for pH determination.

## TABLE II. - COEFFICIENT OF FRICTION AND GROOVE WIDTH FOR MAGNESIUM OXIDE IN SLIDING CONTACT WITH HEMISPHERICAL DIAMOND RIDER IN VARIOUS MEDIA

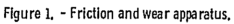
[Single-pass sliding; load, 0.25 N; sliding velocity, 10 mm/min; rider radius, 20 µm.]

Medium and environment	Coefficient of friction	Groove width, <sup>µm</sup>
HCl added to 10-2 N aqueous NaCl solution (pH = 1)	0.23	41
NaOH added to 10 <sup>-2</sup> N aqueous NaCl solution (pH = 13.5)	.11	28
Distilled water	.12	32
Laboratory air	.10	27



.

日本体験



ORIGINAL PAGE IS OF POOR QUALITY

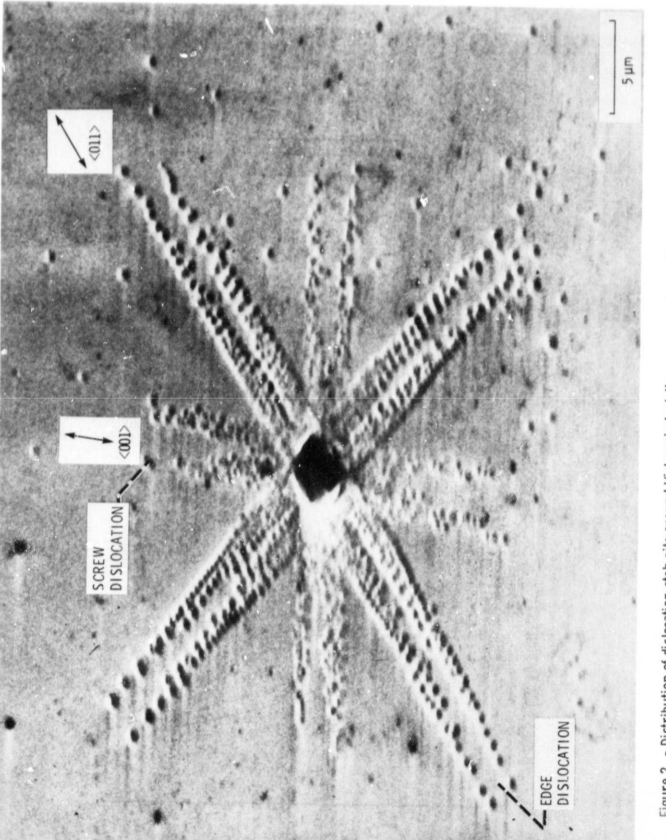
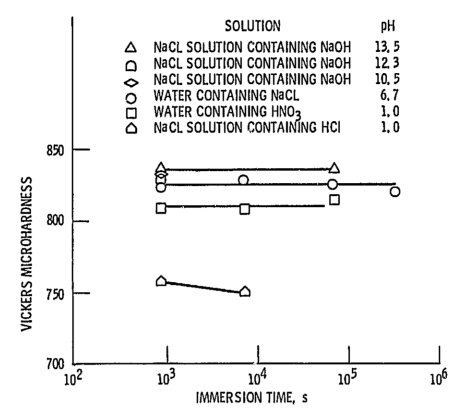
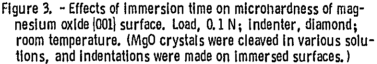
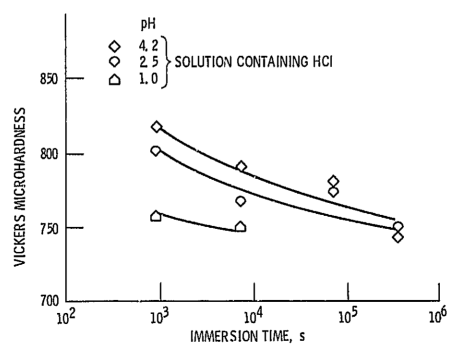
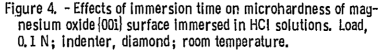


Figure 2. - Distribution of dislocation etch pits around Vickers indentation on magnesium oxide surface. Indepting surface, {001} cleaved in laboratory air; load, 0.1 N; indenter, diamond.

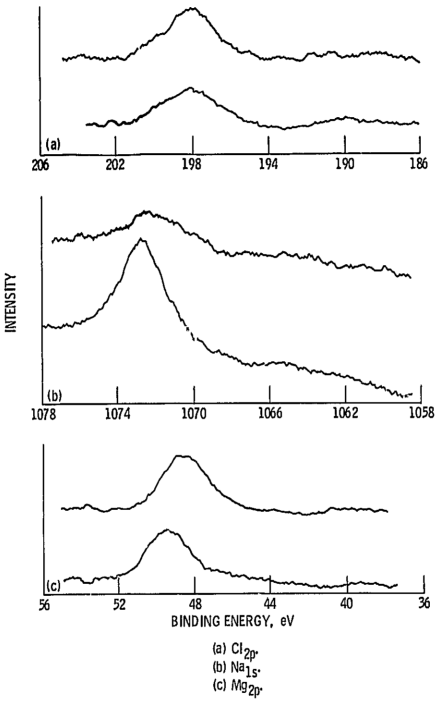






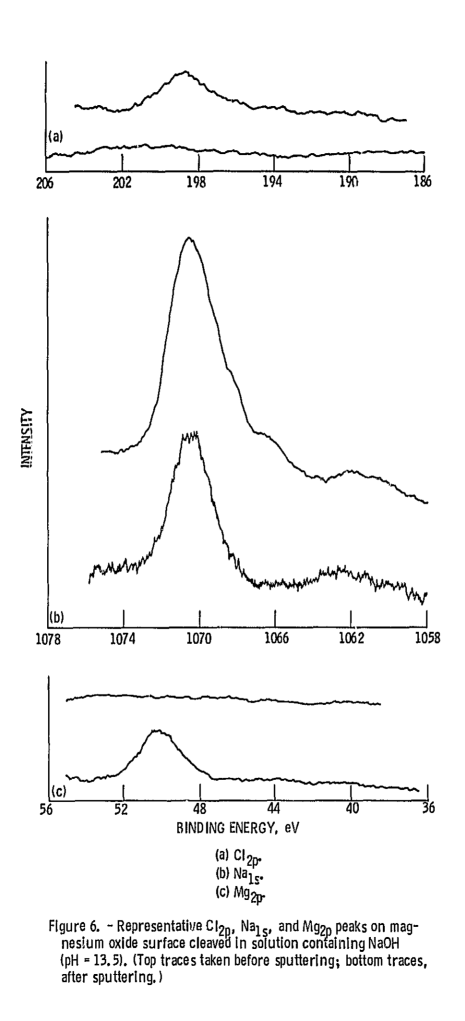


ALC: NOT THE REAL PROPERTY.

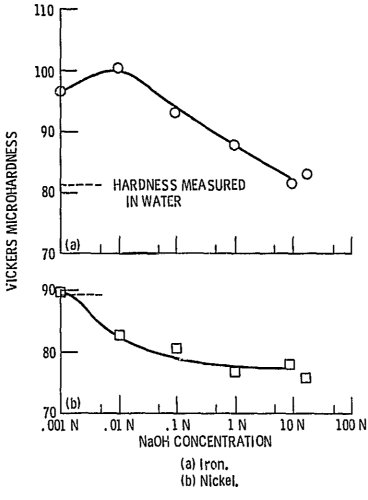


'n.

Figure 5. - Representative Cl<sub>2p</sub>, Na<sub>Is</sub>, and Mg <sub>2p</sub> peaks on magnesium oxide surface cleaved in solution containing HCl (pH = 1). (Top traces taken before sputtering; bottom traces, after sputtering.)



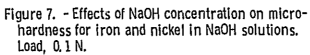




- TV 2\*

2

a 2



\* \*

. 13

214 - F. F

( #

-17

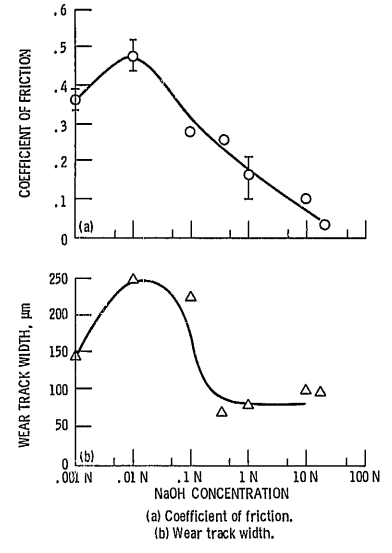
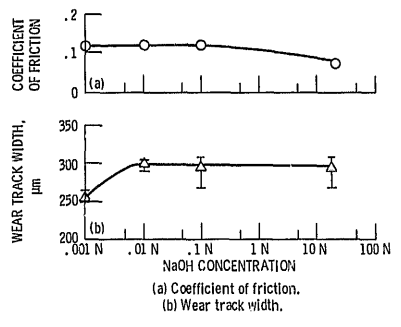
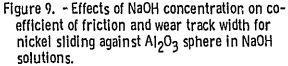
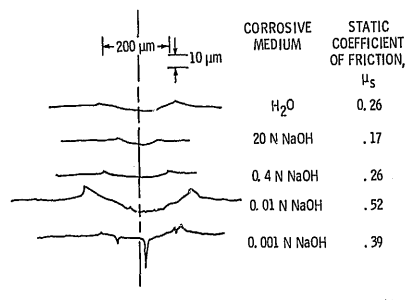


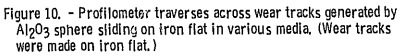
Figure 8. - Effects of NaOH concentration on coefficient of friction and wear track width for iron sliding against Al<sub>2</sub>O<sub>3</sub> sphere in NaOH solutions. zîpi

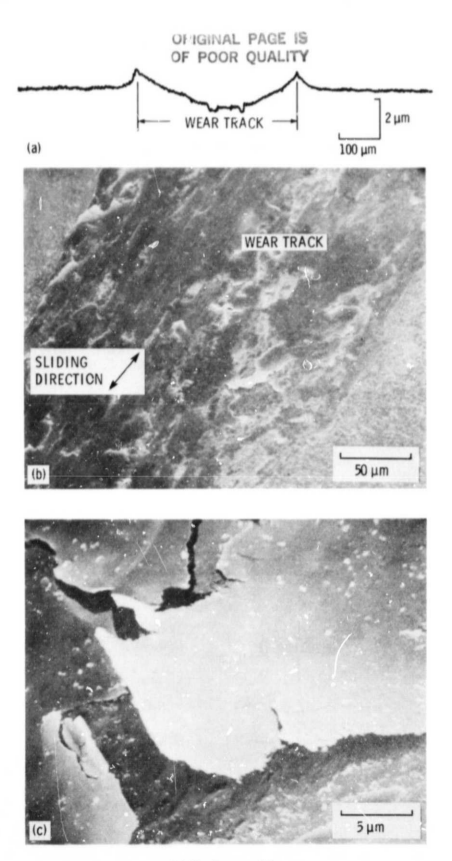


**举注社会的现代** 





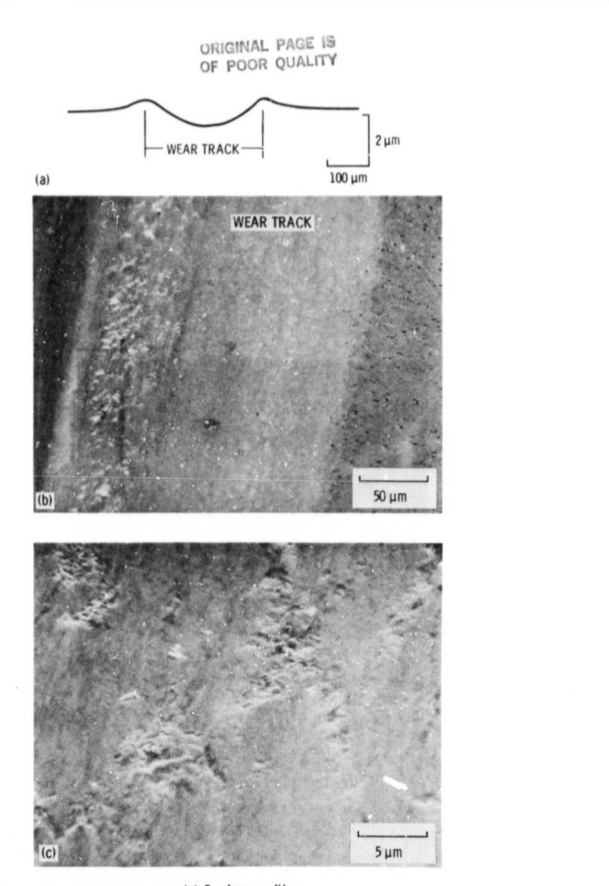




(4

(a) Surface profile.
(b) Wear track (low magnific
(c) Wear damage (high magnifica)

Figure 11. - Surface profile and scanning electron micrographs of wear track on iron surface sliding against  $Al_2O_3$  sphere in 0.01 N NaOH solution.



(a) Surface profile.(b) Wear track (low magnification).(c) Slightly roughened surface (high magnification).

Figure 12. - Surface profile and scanning electron micrographs of wear track on nickel surface sliding against Al<sub>2</sub>O<sub>3</sub> sphere in 0.1 N NaOH solution. ORIGINAL PAGE IS OF POOR QUALITY

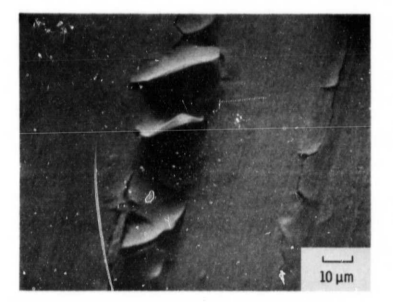


Figure 13. - Scanning electron micrograph of wear track on nickel surface sliding against Al<sub>2</sub>O<sub>3</sub> sphere in water. Static coefficient of friction, 0.28.

