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Correlation between the fatigue crack growth response and the surface reaction kinetics has been made, and is consistent with a transport-limited model for crack growth. The results also suggest that enhancement of fatigue crack growth by water vapor in the aluminum alloys occurs through a "hydrogen embrittlement" mechanism.

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FRACTURE MECHANICS AND SURFACE CHEMISTRY STUDIES OF FATIGUE CRACK GROWTH IN AN ALUMINUM ALLOY

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by

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ABSTRACT

Fracture mechanics and surface chemistry studies were carried out to develop further understanding of the influence of water vapor on fatigue crack growth in aluminum alloys. The room temperature fatigue crack growth response was determined for 2219-T851 aluminum alloy exposed to water vapor at pressures from 1 Pa to 30 Pa over a range of stress intensity factors (K). Data were also obtained in vacuum (at < 0.50 μ Pa), and dehumidified argon. The test results showed that, at a frequency of 5 Hz, the rate of crack growth is essentially unaffected by water vapor until a threshold pressure is reached. Above this threshold, the rates increased, reaching a maximum within one order of magnitude increase in vapor pressure. This maximum crack growth rate is equal to that obtained in air (40 to 60 pct relative humidity), distilled water and 3.5 pct NaCl solution on the same material. Parallel studies of the reactions of water vapor with fresh alloy surfaces (produced either by <u>in situ</u> impact fracture or by ion etching) were made by Auger electron spectroscopy (AES) and x-ray photoelectron spectroscopy (XPS). The extent of surface reaction was monitored by changes in the oxygen AES and XPS signals

Correlation between the fatigue crack growth response and the surface reaction kinetics has been made, and is consistent with a transport-limited model for crack growth. The results also suggest that enhancement of fatigue crack growth by water vapor in the aluminum alloys occurs through a "hydrogen embrittlement" mechanism.

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INTRODUCTION

The influences of cyclic load frequency and gas pressure on environment assisted fatigue crack growth in high-strength alloys has been under study for many years [1-14]. A complete quantitative explanation of this effect and of the differences in response for different alloy-environment combinations is not yet available. To assist in understanding the influences of these variables, coordinated fracture mechanics and surface chemistry studies have been undertaken to identify the rate controlling processes for crack growth in highstrength steels and to assess the roles of these processes in determining environment assisted fatigue crack growth response [15-17]. In the present study, the role of water vapor-metal reaction in determining the crack growth response of aluminum alloys was examined.

Previous studies [6,7] have shown that the rate of fatigue crack growth in aluminum alloys exposed to water vapor (in the absence of capillary condensation at the crack tip) was significantly affected by pressure and frequency. Bradshaw and Wheeler showed that the rate of fatigue crack growth at a given K level appeared to depend on the product of water vapor pressure and cyclic load period (1/frequency), and suggested that the observed frequency effect resulted from the time available for the reaction of water vapor with the newly created crack surfaces [6]. In distilled water, on the other hand, aluminum alloys exhibited little or no effect of frequency [1,8]. Comparable effects of frequency have been reported for fatigue crack growth in high-strength steels, at K levels below K_{ISCC} (the stress corrosion cracking threshold). Barsom [11] showed that the rate of fatigue crack growth in a 12Ni-5Cr-3Mo steel, tested in 3.5 pct NaCl solution at room temperature, depended on cyclic load frequency, with higher growth rates at the lower frequencies. The influence of frequency has also been reported by Gallagher [12] for a HY-80 steel tested in 3.5 pct NaCl solution, by Miller <u>et al</u>. [13] for a low-alloy steel tested in distilled water, and by Hutin [14] for another low-alloy steel tested in water vapor (without capillary condensation at the crack tip). The frequency effects in steels, however, are observed at lower frequencies, and at higher water vapor pressures (and in aqueous environments) as compared to those in the aluminum alloys. Hudak and Wei [9] suggested that these differences in behavior between steels and aluminum alloys may be attributed to differences in the reactivity of these alloys to water/water vapor. Unfortunately, however, this suggestion could not be assessed at the time because of the lack of relevant surface reaction data on these materials, and because the processes controlling crack growth were not well identified and understood.

Recent fracture mechanics and surface chemistry based studies [15,16] showed that sustained-load crack growth in a low-alloy, high-strength steel, in water/water vapor, is controlled by a slow step in the reaction of water with the steel. This reaction step is associated with the nucleation and growth of oxide on the crack surface and the presumed concomitant production of hydrogen. The enhancement of crack growth is attributed to embrittlement by this hydrogen [16]. Extension of these results to the study of environment assisted fatigue crack growth in this steel by water vapor showed that both steady-state and nonsteady-state crack growth response can be adequately explained in relation to the kinetics of water vapor-steel surface reactions [17]. Environment enhancement of fatigue crack growth (at K levels below K_{ISCC}) appeared to be proportional to the extent of surface reaction during one load cycle, and to depend on exposure (pressure x time).

To further assess the role of surface reactions in determining environment assisted fatigue crack growth response and to follow up on the suggestion of Bradshaw and Wheeler [6], coordinated fracture mechanics and surface chemistry studies have been carried out on an Al-Cu alloy (2219-T851 aluminum alloy).

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Results from these studies are summarized and discussed in terms of a model for crack growth [18].

MATERIAL AND EXPERIMENTAL WORK

Material, Specimen and K Calibration

A 1.65 cm thick plate $\frac{1}{}$ of 2219-T851 aluminum alloy was used in this study. Nominal tensile properties of this plate are as follows: yield strength = 358 MPa and tensile strength = 455 MPa.

Wedge-opening-load (WOL) specimens, with half-height to width ratio (H/W) of 0.486, were selected for use. The specimens were oriented in the longitudinal (LT) orientation. An initial (or crack starter) notch, 1.96 cm in length, was introduced into each specimen by electro-discharge machining (EDM). Each specimen was precracked in fatigue, while exposed to the test environment, through a decreasing sequence of loads that terminated at the desired load level (or initial K) for the actual experiment. The precracking procedure provided a fatigue crack of about 0.33 cm in length from the starter notch, corresponding to a crack length of about 2.3 cm at the start of each experiment. This precracking procedure ensured that the subsequent fatigue crack growth would be through material that had not been altered by the notch preparation procedure, and would be unaffected by the starter notch geometry.

Stress intensity factor, K, for the WOL specimen was computed from Eq. 1 [19,20]:

 $K = \frac{P}{BW} \sqrt{a} [30.96 - 195.8 (a/W) + 730.6 (a/W)^{2} - 1186.3 (a/W)^{3} + 754.6 (a/W)^{4}]$ (1)

1/Plate furnished by Air Force Flight Dynamics Laboratory, WPAB, OH.

P = applied load; B = specimen thickness; W = specimen width; and a = crack length. Both specimen width and crack length were measured from the line of loading.

Test Environment

Crack growth experiments were carried out in dehumidified argon, vacuum, and water vapor at room temperature. For tests in dehumified argon, the environment was maintained around the crack by flowing argon (purified by a suitable purification system) through chambers clamped to the faces of the specimen [21]. Tests in vacuum and in water vapor were carried out inside a commercial ultrahigh vacuum chamber that had been modified to provide mechanical force feedthroughs. Tests in vacuum were made at pressures below 0.50 μ Pa. Water vapor was obtained by backfilling the chamber from a high-purity source following bake-out of the system. Pressure was monitored by a capacitance manometer, and purity was checked with the aid of a quadrupole residual gas analyzer.

Experimental Procedures

The fatigue crack growth experiments were carried out under constant amplitude loading in a closed-loop electrohydraulic testing machine operated in load control at a load ratio (R) of 0.05. Load control was estimated to be better than \pm 1 pct. For the tests in argon, a frequency of 20 Hz was used. Tests in vacuum and in water vapor were carried out at 5 Hz.

Crack Monitoring System

An ac electrical potential system was used for monitoring crack growth [22-24]. For the specimen geometry, analytical relationships between crack length and electrical potential were not available, and calibration had to be established experimentally. Calibration was accomplished either by making simultaneous visual and electrical potential measurements of crack length on specimens fatigued in air, or by comparing electrical potential measurements

against fatigue markings on fracture surfaces produced by intentionally introduced high-load excursions (overloads) during fatigue in air [24].

Calibration results, given in terms of crack length (a) versus potential difference $(V-V_n)^{\dagger}$, can be represented by Eq. (2) [24].

$$a = 1.96 + 0.403 (V - V_{n})$$
 (a in cm)

$$a = 0.77 + 0.159 (V - V_)$$
 (a in in.) (2)

 $V = V(a) = potential corresponding to crack length a, and <math>V_r = reference poten$ tial corresponding to the initial notch. Both V and V_r are given in microvolts.Accuracy of crack length measurement with the ac system was estimated to bebetter than 1 pct, for crack lengths from about 2 to 4.8 cm (0.8 to 1.85 in.).The resolution was better than 0.005 cm (0.002 in.) based on 12.5 nV resolutionin electrical potential.

Fractography

Characterization of the morphology of fracture surfaces was made with the aid of scanning electron microscopy. Entire broken halves of the specimens were placed inside the microscope for examination. Specimen tilt was about an axis parallel to the direction of crack growth.

Surface Reactions

Auger electron spectroscopy (AES) and x-ray photoelectron spectroscopy (XPS)

[†]The electrical potential method provides measurements of crack length averaged through the thickness, while the visual method gives measurements of the crack length at the specimen surface only. Crack length measurements made by these two methods would differ because of crack front curvature. The discrepancy was significant for thick specimens. Corrections for crack front curvature were made by measuring average crack lengths from the fatigue markings (introduced during the calibration tests by changing the load amplitude) after specimen fracture. The average crack length was computed on the basis of five measurements - one at each specimen surface, one along each of the quarterthickness planes and one along the mid-thickness plane. The "corrected" crack lengths are used in deriving Eq. (2) [24].

were used to determine the kinetics of the reactions of water vapor with 2219-T851 aluminum alloy surfaces [25]. For AES analyses, clean surfaces were produced by <u>in situ</u> impact fracture of notched round specimens. A polished and ion-etched alloy surface was used for the XPS analyses. Exposures of these clean surfaces to water vapor were achieved by back-filling the spectrometer chamber from a high purity source. High purity water vapor was obtained by alternately freezing and thawing triply distilled water, and pumping away the dissolved gases.

For these surface studies, background pressure in the spectrometer chamber was typically of the order of 6.7 x 10^{-2} µPa (5 x 10^{-10} torr), as measured on an ionization gage. The reactant gases were admitted into the chamber to pressures below 133 µPa (1 x 10^{-6} torr). The chamber was re-evacuated to below 0.67 µPa (5 x 10^{-9} torr) prior to each analysis. At these low pressures (< 133 µPa), reactions induced by species created by the ionization gage filament were minimal.

A different area of the impact fracture sample surface was analyzed by AES for each exposure, using a scanning Auger microprobe (SAM). For the XPS analyses, a freshly ion etched surface was used for each exposure. These procedures were followed to eliminate effects induced by the incident beam during previous measurements.

RESULTS

Fatigue Crack Growth Response

Room temperature fatigue crack growth data, obtained from tests in water vapor at pressures from 1 to 26.6 Pa $(7.5 \times 10^{-3} \text{ to } 0.2 \text{ torr})$, are shown as a function of stress intensity factor range (ΔK) in Figure 1, along with data obtained from tests in dehumidified argon. The data at 26.6 Pa are comparable to those obtained in air (40 to 60 pct relative humidity), distilled water and

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3.5 pct NaCl solution [24]. The results in dehumidified argon correspond to data in vacuum at less than 0.50 μ Pa (~ 4 x 10⁻⁹ torr). These data are also shown in Figure 2 as a function of water vapor pressure at three ΔK levels. The error bands represent 95 pct confidence intervals computed from the residual standard deviations in each set of data [26,27]. The results show that, at a frequency of 5 Hz, the rate of crack growth is essentially unaffected by water vapor until a threshold pressure is reached. The rates then increased and reached a maximum within one order of magnitude increase in vapor pressure from this threshold. The maximum rate is equal to that obtained in air, distilled water and 3.5 pct NaCl solution (at 20 Hz) [24]. The transition range, in terms of pressure/frequency, is comparable to that reported by Bradshaw and Wheeler on another aluminum alloy [6].

Fractography

A photomacrograph of the fracture surface of a specimen tested in water vapor at 4.66 Pa (0.035 torr) is shown in Figure 3. This water vapor pressure corresponded to a position within the transition region in fatigue crack growth response (Figure 2). Principal interest here is in terms of the general features of the fracture surface morphology rather than with the details of the fracture processes. Examination of the photomacrograph indicates that the fracture surface appearance in the mid-thickness region differs considerably from that in the near-surface region. Scanning electron microfractographs taken from the same specimen indeed show differences in fracture surface morphology between the two regions, Figure 4. The fracture surface morphology may be compared with those observed on specimens tested in dehumidified argon and in water vapor at a pressure above the transition range, Figure 5. It is seen that the fracture morphology in the mid-thickness region (Figure 4a) is comparable to that associated with crack growth in dehumidified argon, an inert

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environment, (Figure 5a), whereas the morphology in the near-surface region corresponds to that for full environmental effect (compare Figures 4b and 5b). This observation suggests that the effective water vapor pressure in the midthickness region of the crack may be substantially below that in the surrounding environment. The implication of this observation will be considered in the correlation of crack growth and surface reaction data.

Surface Reaction Kinetics

The results from AES and XPS studies of the reactions of water vapor and oxygen with clean surfaces of 2219-T851 aluminum alloy are presented in a companion paper [25]. A brief summary of those findings that bear directly on fatigue crack growth in water vapor is given here.

Changes in the normalized oxygen Auger (510 eV) signal as a function of exposure to water vapor are shown in Figure 6. Normalization is based on the average value of oxygen Auger (510 eV) signals from specimens exposed to water vapor for 6.65×10^{-2} to 1.33 Pa-s (5×10^{-4} to 10^{-2} torr-s). Comparable results were obtained from the XPS studies. The results show that the initial rate of reaction with water vapor is rapid and reaches "saturation" following about 2.7 x 10^{-3} Pa-s (2×10^{-5} torr-s) exposure; that is, the extent of reactions with water vapor is limited. XPS results indicate that the reactions are associated with the formation of an oxide or a hydrated oxide layer. The limited reactions with water vapor are consistent with previous results on a high-strength (AISI 4340) steel [16]. The rate of reaction with the aluminum alloy, however, is 8 to 9 orders of magnitude faster than the corresponding rate (associated with the slow, second step) of reaction with AISI 4340 steel. The surface reaction results will be considered in relation to environment assisted fatigue crack growth in the following discussion.

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DISCUSSION

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A comparison between the extent of surface reaction (Figure 6) and the fatigue crack growth response (Figure 2) may be made if one assumes that the enhancement of crack growth is proportional to the extent of surface reaction during one fatigue loading cycle [17,18], and if pressure/2 x frequency (p/2f) is used as an approximate measure of exposure*. This comparison shows that a factor of 10^3 adjustment in exposure (pt or p/2f) will have to be made to obtain a correlation. The need for this adjustment suggests that the rate of reaction at the crack tip is limited by the rate of transport of water vapor to that region. This possibility is also suggested by the differences in fracture surface morphology between the surface and mid-thickness regions of specimens tested within the transition region (<u>i.e.</u>, between 0.13 and 1.3 Pa-s or 10^{-3} and 10^{-2} torr-s), and by the similarity between the observed morphology and the corresponding morphology for specimens tested in argon and at higher water vapor pressures (see Figures 3 to 5).

A transport limited model has been developed [18]. In this model, transport of gaseous environments to the crack tip is assumed to be by Knudsen flow [28]. The governing differential equations for flow and surface reactions are as follows [18]:

$$\frac{dp}{dt} = -\frac{SN_{0}^{r}RT}{V} \frac{d\theta}{dt} + \frac{F}{V}(p_{0}-p)$$
(3)

 $\frac{d\theta}{dt} = k_c p(1-\theta)$ (4)

*It is assumed that the crack growth increment is introduced "abruptly" at the maximum load point in a given fatigue loading cycle. Thus, the newly produced crack surface is exposed to the environment only during one-half (unloading half) of the fatigue cycle.

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The terms in the equations are as follows:

- p = pressure of gas at the crack tip
- \dot{p}_{0} = pressure of gas in the surrounding environment
- R = gas constant
- S = area of newly created crack surface per cycle = α (2B Δa), where Δa = crack growth increment per cycle, B = specimen thickness, and α = empirical constant for surface roughness and crack geometry [18]
- T = absolute temperature
- V = volume associated with the crack growth increment per cycle, <u>i.e.</u>, with S
- N_o = density of surface sites
- θ = fractional surface coverage or extent of reaction of surface per unit area 2

 $F = 8.72 \times 10^4 \beta \frac{\sigma_{ys}^2}{E^2} B\ell \left(\frac{T}{M}\right)^{1/2} \text{ (in cm}^3/\text{s)} = \text{Knudsen flow parameter} \\ \text{that depends on dimension and shape of the capillary, molecular} \\ \text{weight (M) of the gas and temperature (T). The specific form} \\ \text{of this expression reflects an attempt to account for constriction} \\ \text{in flow by the real crack, where } \ell \text{ is an arbitrary distance (of} \\ \text{the order } 10^{-4} \text{ cm} \text{) from the crack tip used in defining a crack} \\ \text{opening and } \beta \text{ is an empirical quantity to be determined from the} \\ \text{crack growth data [18,28].} \end{cases}$

k = reaction rate constant

From Equation (3), it can be seen that the rate of change of pressure at the crack tip depends on the decrease in pressure produced by reaction of the environment with the newly created crack surface and on the increase in pressure from the influx of gas from the external environment. Equation (4) incorporates the

assumption that the surface reaction was first-order in relation to available surface sites [18,25,27]

Equations (3) and (4) are nonlinear, coupled equations which can be solved, at least in principle, for specific cases. Numerical solutions for these equations have been obtained using reasonable, assumed values for S and F and measured value for k [25]. These numerical results are illustrated in Figures 7 and 8. Figure 7 was derived by assuming the initial pressure to be zero at the crack tip, and shows that the percent attenuation in pressure is essentially independent of the external pressure. For Figure 8, the initial pressure at the crack tip was assumed to be equal to the external pressure. It is seen that the pressure decreased rapidly and reached an attenuated value that is essentially the same as the previous case. In either case, the attenuated pressure was more than two orders of magnitude lower than the external pressure, and remained nearly constant as long as the crack surface remained active. The surface coverage (or extent of surface reaction) increased almost linearly with time; the rate of increase being proportional to the local pressure at the crack tip. This transport-limited change in surface coverage would provide much better agreement with the fatigue data.

To provide for a more formal comparison, Equations (3) and (4) may be solved approximately. By combining Equations (3) and (4), one can obtain an expression for the pressure at the crack tip.

$$p = \frac{p_0 - \frac{Vdp}{Fdt}}{\frac{SN_0RTk_c}{F}(1-\theta) + 1}$$
(5)

It has been shown that the term (V/F)(dp/dt) may be neglected in comparison with p_o during the reaction, except at the very beginning of the reaction (or for $\theta \approx 0$) [18]. Using this simplifying assumption, Equation (4) may be rewritten as

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$$\frac{1\theta}{1t} = k_c p(1-\theta) = \frac{k_c p_0(1-\theta)}{\frac{SN_0 RT k_c}{F}(1-\theta) + 1}$$
(6)

Integration of Equation (6) gives the surface coverage θ as a function of pt

$$\frac{SN_{o}RT}{F} \theta - \frac{1}{k_{c}} en(1-\theta) = p_{o}t$$
(7)

For this 2219-T851 aluminum alloy tested in water vapor, SN_0RT/F is estimated to be of the order 1 Pa-s and k_c is of the order $10^3 (Pa-s)^{-1}$ [25]. In other words, SN_0RTk_c/F is of the order 10^3 . Thus, the reactions at the crack tip are limited by flow and θ may be described adequately by the linear term alone up to $\theta \approx 0.99$ for this case, that is

$$\theta \approx \frac{F}{SN_{o}RT} P_{o}t$$
 (8)

If t is taken to be equal to $\tau/2$ or 1/2f, then

$$\theta \approx 4.36 \times 10^4 \frac{\beta}{\alpha} \frac{\sigma_{ys}^2 \ell}{N_o RTE^2} \left(\frac{T}{M}\right)^{1/2} \frac{\cdot p_o}{2f\Delta a}$$
 (9)

In other words, the extent of reaction, as measured by θ , during a cycle is proportional to p_0 , and is inversely proportional to frequency and the extent of crack growth (Δa) during that cycle, up to $\theta \approx 1$. With further increase in pressure or decrease in frequency, θ should remain at 1, <u>i.e.</u>, at saturation coverage. It has been suggested that the rate of environment assisted fatigue crack growth, (da/dN)_e, may be considered to be the sum of two components -the rate of growth in an inert reference environment or "pure fatigue", (da/dN)_r, and the environmental or "corrosion fati_b" component, (da/dN)_{cf} [29]. The growth increment per cycle (Δa) is therefore identified with (da/dN)_e (1). The environmental component is considered to be a function of the amount of hydrogen that is produced (or extent of reaction) during one cycle, and should be proportional to $\Delta a \cdot \theta$, and the number of hydrogen atoms produced per water molecule reaction. Thus, one can expect $(da/dN)_{cf}$ to vary according to Equation (10) for $\theta < 1.0$.

$$\left(\frac{da}{dN}\right)_{cf} = \left(\frac{da}{dN}\right)_{e} - \left(\frac{da}{dN}\right)_{r} \approx 4.36 \times 10^{4} \frac{\beta}{\alpha} \frac{\sigma_{ys}^{2}}{N_{o}RTE^{2}} \left(\frac{T}{M}\right)^{1/2} \frac{p_{o}}{2f}$$
(10)

For combinations of p_0 and f values that would yield $\theta \neq 1$, $(da/dN)_{cf}$ and $(da/dN)_{a}$ would represent maximum enhancement that can be achieved.

Comparison of Equation (10) with fatigue crack growth data is shown in Figure 9. Data at three different ΔK levels are used, and are normalized with respect to the average of (da/dN) rf values for water vapor at 6.9 and 26.6 Pa, air, distilled water and 3.5 pct NaCl solution at each AK level. These average values are used to represent the "saturation" values of (da/dN) cf. Because $(da/dN)_{cf}$ represents the difference between $(da/dN)_{e}$ and $(da/dN)_{r}$, errors tend to be very large, particularly at the lower water vapor pressures. Estimates of error were computed from the residual standard deviations in each set of data, and are shown in Figure 9. The data indicate the desired linear dependence on p_0 or $p_0/2f$ below "saturation", and suggest that the pressure needed to produce "saturation" is essentially independent of ΔK . The latter observation suggests that the characteristic length & in Equation (10) depends on ΔK . If one assumes & to be equal to the growth increment per cycle at "saturation", then the data can be matched to Equation (10) to determine the empirical constants β/α . Alternatively, β/α may be determined from the observed pressure at the onset of saturation response and Equation (9), i.e., for $\theta \approx 1$, using the same assumption. The value of β/α was found to be approximately equal to 1.5. The fitted curves, based on Equation (10), are shown in both Figures 2 and 9, and show good agreement with the data.

If one takes $\alpha = 2$ as a reasonable estimate of the contributions of crack shape and surface roughness to the crack area, then $\beta = 3$. If one now assigns

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β to the constricted flow length, this length is estimated to be approximately equal to $2r_y$ (where $r_y = \frac{1}{2\pi}(K/\sigma_{ys})^2$ is the formally calculated plastic zone correction factor), or equal to about 0.07 cm at $K_{max} = 16.5$ MPa-m^{1/2}. The estimated effective crack opening at this K_{max} is about 4 x 10⁻⁵ cm. Alternatively, one may assign β to the effective crack opening, thus providing a constricted flow length of about 0.2 cm and an effective opening of 7 x 10⁻⁵ cm. These quantities appear to be quite reasonable in either case. Thus, a satisfactory fracture mechanics and surface chemistry explanation of the observed fatigue crack growth response has been achieved.

Examination of Equations (5) and (6) shows that if $SN_0RTk_c/F<<1$, that is for low k_c or "slow" reaction kinetics, the pressure at the crack tip becomes equal to the external pressure following a fast initial build up, and surface reaction is now limited by the surface reaction kinetics. This latter case is consistent with previous observations on AISI 4340 steel in water vapor [17].[•] The observed difference in fatigue crack growth response and a factor of 10^8 to 10^9 difference in reaction rate constants for aluminum alloys and steel with water vapor [16,25], are consistent with these observations.

The fact that no further increase in the rate of fatigue crack growth for this aluminum alloy was observed in the aqueous environments over that in water vapor at "saturation" is consistent with the observation that the aluminumwater reaction is limited [25]. It also suggests that enhancement of crack growth results from hydrogen embrittlement <u>vis-a-vis</u> active path dissolution. The precise mechanics for this hydrogen embrittlement remains to be identified. The fact that significant environment enhancement of crack growth can occur in fatigue in aluminum alloys, as opposed to sustained-load crack growth, resides with the efficiency of the fatigue process in creating "fresh" surfaces to react with the environment.

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SUMMARY

Coordinated fracture mechanics and surface chemistry studies were carried out to develop further understanding of the influence of water vapor on fatigue crack growth in aluminum alloys. The results confirm the earlier suggestion by Bradshaw and Wheeler that the enhancement of crack growth is a function of water vapor pressure and the time available for reaction, that is, of exposure (pressure x time). The enhancement is shown to be dependent on the rate of transport of water vapor to the crack tip, as well as on the surface reaction kinetics. The results also suggest that hydrogen produced by the surface reactions is responsible for the enhancement in growth, <u>vis-a-vis</u> an active path dissolution mechanism. The observed fatigue crack growth response is adequately described in terms of a transport-limited model for fatigue crack growth.

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FIGURE CAPTIONS

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- Figure 1: Influence of water vapor pressure on the kinetics of fatigue crack growth in 2219-T851 aluminum alloy at room temperature.
- Figure 2: Influence of water vapor pressure (or ^{pressure}/_{2xfrequency}) on fatigue crack growth rates in 2219-T851 aluminum alloy at room temperature. Solid lines represent model predictions.
- Figure 3: Photo-macrograph of fatigue fracture surface of 2219-T851 aluminum alloy specimen tested in water vapor (4.66 Pa) at room temperature. $(\Delta K = 16.5 \text{ MPa-m}^{1/2}, R = 0.05 \text{ and } f = 5 \text{ Hz}).$
- Figure 4: SEM microfractographs taken from the mid-thickness region (center) and the near-surface region (edge) of the specimen shown in Figure 3 showing differences in fracture surface morphology.
- Figure 5: SEM microfractographs of specimens tested in argon and in water vapor at 26.6 Pa (full environmental effect) showing similar differences in fracture surface morphology as seen in Figure 4.
- Figure 6: Kinetics of reactions of water vapor with 2219-T851 aluminum alloy at room temperature [25].
- Figure 7: Variation of water vapor pressure and surface coverage at the crack tip as a function of time, assuming that the initial pressure at the crack tip is zero ($p_i = 0$) [18].
- Figure 8: Variation of water vapor pressure and surface coverage at the crack tip as a function of time, assuming that the initial pressure at the crack tip is equal to the external pressure $(p_i = p_0)$ [18].
- Figure 9: Comparison of normalized (corrosion) fatigue crack growth rates with model predictions for pressure and frequency dependence.





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Figure 7: Variation of water vapor pressure and surface coverage at the crack tip as a function of time, assuming that the initial pressure at the crack tip is zero ($p_1 = 0$) [18].



Figure 8: Variation of water vapor pressure and surface coverage at the crack tip as a function of time, assuming that the initial pressure at the crack tip is equal to the external pressure $(p_i = p_0)$ [18].



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