Peridynamics for predicting pit-to-crack transition

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Even though the failure mechanisms in aerospace structures are mainly governed by the incidence of fatigue loading, environmentally assisted fracture related pit-to-crack transition can still occur in many aluminum alloys, stainless steels and high strength low alloy steels. Despite of the relative rarity of this phenomenon, the consequent failures can be extremely destructive and lead to the loss of an aircraft. Prediction of damage evolution starting at corrosion pits acting as precursor to cracking has been hampered by a lack of insight into the process, as well as limitations in visualization and measurement techniques. In this regard, numerical modeling can be beneficial. The current study presents numerical predictions of pit evolution from a flat metal surface to realistic pit morphologies by using a new continuum mechanics formulation, peridynamics. Intergranular/transgranular pit-to-crack transition under different loading conditions and microstructural features are investigated without imposing any limiting assumption on the site of the crack initiation. Based on the numerical results, it can be concluded that microstructure has a significant effect for the prediction of pit-to-crack transition phenomenon.

I. Introduction

Even though the failure mechanisms in aerospace structures are mainly governed by the incidence of fatigue loading, environmentally assisted fracture (EAF) related pit-to-crack transition can still occur in many aluminum alloys, stainless steels and high strength low alloy steels. Moreover, presence of corrosion damage can reduce the fatigue life of the structure [1]. Despite of the relative rarity of this phenomenon, the consequent failures can be extremely destructive and lead to the loss of an aircraft. In February 1992, a military jet crashed in the Dutch city of Hengelo because of a localized corrosion and subsequent EAF of the lever-arm-pin in the jet engine made of austenitic stainless steel [2]. Another relevant incident, which can be attributed to corrosion pitting followed by EAF occurred in February 2007 on the left side of helicopter main landing gear drag beam made of ultrahigh strength steel during landing operations [2].

Prediction of damage evolution starting at corrosion pits acting as precursor to cracking has been hampered by a lack of insight into the process, as well as limitations in visualization and measurement techniques. For long cracks, standards for quantifying environment-assisted crack growth rates can be found [3]. However, in relation to the growth rate of small cracks emerging from corrosion pits, there are no such standards available to guide the measurement process. In this regard, numerical modeling can be beneficial. Nonetheless, it is presently hampered by the popular assumption [3] that the crack has the same depth as the pit at the point of transition and initiates at the pit base, as it has no intrinsic foundation. In fact, novel observations using X-ray tomography of stress-corrosion cracks

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emerging from pits showed that cracks evolved from the pit predominantly at or near the pit mouth and not at the pit base [4]. Moreover, recent finite element analysis of pits in specimens subjected to different static loads revealed that maximum stress and strain can occur at the pit shoulder and in the proximity to the pit mouth [5]. The current study presents numerical predictions of pit evolution from a flat metal surface to realistic pit morphologies by using a new continuum mechanics formulation, peridynamics. Intergranular/transgranular pit-to-crack transition under different loading conditions and microstructural features are investigated without imposing any limiting assumption on the site of the crack initiation.

II. Peridynamics

An alternative to traditional finite element analysis (FEA) is the peridynamic (PD) theory introduced by Silling [6]. FEA is based on classical continuum mechanics which is using partial differential equations to describe the motion of material points. On the other hand, PD is a nonlocal extension of classical continuum mechanics that is based on integro-differential equations involving time and spatial coordinates. This feature allows damage initiation and propagation at multiple sites with arbitrary paths inside the material without resorting to special crack growth criteria. In the PD theory, internal forces are expressed through nonlocal interactions between pairs of material points within a continuous body, and damage is part of the constitutive model.

PD theory was successfully applied for the first time by De Meo et al. [7] to model stress-corrosion cracking (SCC) damage in polycrystalline materials due to hydrogen embrittlement. As an extension of that work, the present study focuses on the transition from corrosion pits to cracks due to EAF. Thus, a new PD algorithm, based on implicit time integration, has been introduced for predicting metal dissolution. This process is modeled using a modified Nernst-Planck equation, where electromigration and diffusion effects are included in an effective diffusion coefficient, whose value is obtained through calibration against experimental data as explained in [8]. The peridynamics version of the governing equation for metal dissolution can be written as:

$$
\dot{C}(\mathbf{x}, t) = \int_{B_d} f_d \left[ C(x', t) - C(x, t) \right] \, \mathrm{d}V_{x'}
$$

(1)

where $\dot{C}(\mathbf{x}, t)$ is the time derivative of metal ion concentration associated with the generic material point $\mathbf{x}$, and $f_d \left[ C(x', t) - C(x, t) \right]$ is called metal dissolution response function which is defined as:

$$
f_d = \frac{d}{|x' - x|}
$$

(2)

in which the peridynamic diffusion bond constant $d$ can be expressed in terms of the effective diffusion coefficient $D_{eff}$ as:

$$
d = \frac{6 \cdot D_{eff}}{\pi \cdot h \cdot \delta^3}
$$

(3)

where $h$ refers to the thickness of the plate and $\delta$ is the PD horizon for a two-dimensional configuration.

III. Numerical results

The PD model of pit-to-crack transition considered in this study is represented by a plate with dimensions of 0.1 mm $\times$ 0.1 mm and a thickness of 1 $\mu$m. The material and environment are austenitic stainless steel grade 304 exposed to 1M NaCl aqueous solution, respectively. For the first phase of the analysis, i.e., pitting evolution, the metal
concentration of $C = 0 \text{ mol/m}^3$ is applied as a boundary condition in the middle of the top surface to a scratch in the oxide film with a length of 6 μm. At the beginning of the simulation, all nodes belonging to the metal have a concentration of metal ions equal to $C_{\text{solid}} = 143000 \text{ mol/m}^3$. As soon as the concentration values drop below the saturation value of the concentration of metal ions in the corrosive solution, $C_{\text{sat}} = 5100 \text{ mol/m}^3$, the related nodes are considered as belonging to the liquid region. The diffusivities of metal ions in the solid and liquid regions are specified as $d_{\text{solid}} = 2 \times 10^{-12} \text{ m}^2/\text{s}$ and $d_{\text{liquid}} = 8.5 \times 10^{-10} \text{ m}^2/\text{s}$, respectively.

The plate is discretized with $100 \times 100$, $200 \times 200$ and $300 \times 300$ PD nodes and the resulting value of grid spacing and horizon radius values are $\Delta = 1 \mu\text{m}$, $0.5 \mu\text{m}$, $0.3333 \mu\text{m}$ and $\delta = 3 \mu\text{m}$, $1.5 \mu\text{m}$, $1 \mu\text{m}$, respectively. The total time is specified as $t = 500 \text{ s}$.

Concerning the second phase of the analysis, i.e., crack propagation, the microstructure of the material is constituted of 50 randomly oriented crystals and the following values of material constants, density and fracture toughness of the material are considered: $c_{11} = 204.6 \text{ GPa}$, $c_{12} = 137.7 \text{ GPa}$, $\rho = 7880 \text{ kg/m}^3$ and $K_{\text{fc}} = 70 \text{ MPa m}^{1/2}$, respectively [9]. A horizontal velocity boundary condition of 0.2 m/s is applied along the left and right edges of the plate. Moreover, the vertical displacement of the nodes in this region is constrained along the vertical direction. The total time is $t = 15 \mu\text{s}$. The time step size has to be small enough to capture the full dynamic characteristic of the problem and different time step sizes, $\Delta t = 0.05 \mu\text{s}$, $0.1 \mu\text{s}$ and $0.5 \mu\text{s}$ are considered to perform a convergence analysis. To model intergranular fracture, only the bonds crossing the grain boundary of the crystals are allowed to break whereas twice stronger grain boundaries with respect to grain interiors are considered for transgranular fracture.
Figure 1. Pit-to-crack transition (100 × 100 particles, \( t = 15 \mu s \), \( dt = \) (a) 0.05 µs, (b) 0.1 µs, (c) 0.5 µs)

Figure 2. Pit-to-crack transition (200 × 200 particles, \( t = 15 \mu s \), \( dt = \) (a) 0.5 µs, (b) 0.1 µs, (c) 0.01 µs)
Figure 3. Pit-to-crack transition (300 × 300 particles, $t = 15$ µs, $dt = 0.1$ µs)

For the intergranular case, based on the results shown in Figures 1-3 for different discretization sizes and time increments, it can be concluded that the crack initiation takes place from the pit base for the randomly generated microstructure.

Figure 4. Pit-to-crack transition (100 × 100 particles, $t = 15$ µs, $dt = (a) 0.05$ µs, (b) 0.1 µs, (c) 0.5 µs)
On the other hand, crack initiation takes place at the left hand side of the pit base when transgranular fracture pattern occurs as shown in Figures 4-6.

IV. Conclusions

In this study, pit-to-crack transition phenomenon is modelled by using a new continuum mechanics formulation, peridynamics. First, pitting evolution is simulated prior to the crack propagation phase. Microstructure is taken into account by randomly generating polycrystalline structure. Both intergranular and transgranular fracture behaviours
are investigated and different crack initiation locations are predicted. Hence, it can be concluded that microstructure has a significant effect for the prediction of pit-to-crack transition phenomenon.

References


