Chemomechanical weakening of muscovite quantified with in situ liquid nanoindentation

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ABSTRACT

Chemomechanical weakening of layered phyllosilicate muscovite mica was studied as a function of chemical environment via in situ liquid-nanoindentation under four conditions (dry, deionized water, and two NaCl solutions of different pH). While traditional indentation analyses of layered materials with extreme mechanical anisotropy have been limited due to displacement bursts (pop-ins), here the bursts were used as proxies for delamination, fracture, and spalling events. Since displacement bursts during an indent represent a stochastic process, 120 indents were conducted for each condition to generate statistically significant amounts of data. In total, over 9000 bursts were assessed using a load–displacement threshold criterion, classifying this as a high-throughput nanoscale fracture technique. For each burst, initiation load, initiation displacement, plastic zone volume at initiation, and energy dissipation were analyzed. A power-law relationship between the burst load and burst energy was noted which separated the bursts into two continuous distributions: (1) bursts due only to the mechanics of the indent and (2) bursts due to both the mechanics of the indent and the environment. By using a cumulative probability distribution, it was found that the NaCl solutions decreased the minimum plastic zone volume necessary to initiate a displacement burst by an order of magnitude relative to the dry condition. Finally, the underlying mechanisms explaining the trends in initiation volume as a function of environment were discussed, with a focus on the chemomechanical degradation processes via chemical attack and cation exchange.
Introduction

Chemical reactions at crack tips and on material surfaces can alter observed mechanical properties from the molecular- to continuum-scales. This recognition has led to new technologies, such as chemical–mechanical polishing [1] and hydraulic fracturing [2]. At the nano to microscales, chemical effects can define material degradation, and manifest as stress corrosion cracking [3] and liquid–metal embrittlement [4]. At the kilometer-scales, chemically assisted slow cracks drive the long-term consolidation processes, formation of fracture networks, and permeability changes in subsurface reservoirs [5, 6].

Chemical effects on fracture have been recognized for over a century [7, 8]. Rehbinder [8] showed that quartz and fluorite crystals fractured more easily in aqueous surfactant solutions. Similar to the Griffith criterion [7], Rehbinder suggested that surface adsorbates can decrease the work of formation for new surfaces during fracture, and therefore that the fracture toughness of a material can be dependent on the chemical environment [9]. Despite the maturity of this research field, current theoretical frameworks fail to predict how the energy release rate, crack growth rate, and geometry of subcritical cracks change with changing chemical environment and material properties. This is because there are a multitude of potential chemical mechanisms that can influence the effective fracture toughness, e.g., adsorption reactions [8, 10–12], redox reactions [13], and chemical attack by a reactive species [14]. Anisotropy, characteristic of layered minerals and sedimentary rock formations, presents an additional challenge in predicting crack growth [15, 16]. Here, we focus on the chemical effects on fracture in the layered mineral muscovite—an abundant member from the mica group, and one of the primary components of rock formations.

The mica mineral group is important in subsurface engineering applications, including resource extraction, carbon sequestration, and nuclear waste storage, because it is susceptible to chemomechanical weakening due to its layered, anisotropic structure and has also been shown to control the strengths and mechanical anisotropies of laminated (layered) rocks such as shale due to its relative weakness in comparison to other silicates [17]. Moreover, mica group minerals are abundant, composing approximately 13% of the upper continental crust [18]. Muscovite is in the dioctahedral subgroup of mica with an ideal chemical formula of \( \text{KAl}_2(\text{Si}_3\text{Al})\text{O}_{10}(\text{OH},\text{F})_2 \) and is the most common of the 55 mica polytypes [19]. Muscovite is composed of TOT (tetrahedral–octahedral–tetrahedral) layered structures, where the ‘T’ layers are silicon oxide that sandwich the ‘O’ layer of aluminum oxide (Fig. 1, structure file from [20]). The ‘T’ layers have random substitutions of Al for Si, while the individual TOT structures are bonded together by potassium cations. TOT layers are nanometer-thick, crystalline sheets that facilitate easy basal slip and perfect basal cleavage on the (001) planes.

The layered structure of muscovite, as depicted in Fig. 1, leads to highly anisotropic mechanical behavior that is dictated by a number of deformation mechanisms depending on the crystallographic orientation of loading. Mares and Kronenberg [17] conducted uniaxial compression of macroscopic, single-crystal muscovite at 400 °C with a confining pressure of 200 MPa at a strain rate of 10^{-5} and observed dislocation glide and low-angle kink boundary formation when loaded 45° to the (001) basal plane, kink band formation when loaded parallel to the basal plane, and fracture when loaded perpendicular to the basal plane. Additionally, the

Figure 1 Laminated structure of muscovite mineral visualized via the Vesta 3.3.9 software. Partial occupancies are due to isomorphous substitutions. Gray: Al, Red: O, Blue: Si, Purple: K, Yellow: Na.
authors found that muscovite could withstand an order of magnitude larger stresses when loaded perpendicular to the basal plane compared to 45° loading. However, despite the progress at the macroscale, microscale, and nanoscale testing of mica and other similarly layered materials has been limited due to this extreme anisotropy [21]. When conducting nanoindentation of mica, traditional analysis techniques are difficult to apply due to the large number of displacement bursts, also known as pop-ins, that occur. And while in situ liquid-nanoindentation was first conducted over 20 years ago [22], it has not been widely employed to analyze this material system due to the difficulties in quantifying results.

In this paper, in situ liquid nanoindentation was used to study the chemomechanical weakening of muscovite, with displacement bursts used as a proxy for kinking, delamination, fracture, and spalling events. Four environmental conditions were explored: (1) dry, (2) deionized (DI) water, (3) 0.1 M NaCl aqueous solution (pH 9.8), and (4) 0.1 M NaCl aqueous solution (pH 12.0). The pH values were chosen because the silicate framework is susceptible to chemical attack by hydroxyl ions (OH\(^-\)) [14]. The measured dissolution rates for muscovite are on the order of 1.20 × 10\(^{-16}\) mol cm\(^{-2}\) s\(^{-1}\) at pH 9.5–4.45 × 10\(^{-16}\) mol cm\(^{-2}\) s\(^{-1}\) at pH 11.8 and at 70 °C [25]. A new crystal was used for each of the three liquid conditions. The samples were stored in an N\(_2\) box and nanoindentation was conducted within 12 h after removal from the box.

Nanoindentation was performed on a Hysitron TribolIndenter 900 with an in situ liquid Berkovich-geometry indenter tip. Prior to nanoindentation, the tip area function and load frame compliance were calibrated over the entire load range of the instrument using fused silica as the reference material. For the Berkovich tip, the tip area function \(A(h)\) was defined as

\[
A(h) = C_0 h^2 + C_1 h^3 + C_2 h^{1/2} + C_3 h^{1/4} + C_4 h^{1/8} + C_5 h^{1/16}
\]

where \(h\) is the contact depth and \(C_0\) through \(C_5\) are coefficients related to tip shape. Assuming a reduced modulus \(E_r = 69.6\) GPa for fused silica, \(C_0\)–\(C_5\) were found through fits to the calibration data. Following calibration, the measurements on muscovite consisted of loading to a maximum load of 10 mN at an indentation strain rate of 0.1 [26], with a 5 s hold at maximum load followed by a 5 s unload. For dry nanoindentation, the relative humidity in the laboratory never exceeded 30%. A total of 120 indents (three 5 × 8 grids) were conducted, with each grid point spaced 60 μm apart. For nanoindentation in DI water and NaCl aqueous solutions, approximately 25 mL of liquid was dispensed into the 9 cm diameter petri dish in order to completely cover the sample to a depth of approximately 3.5 mm. The tip was brought into contact with the sample and retracted 50 μm so that 99% of the shaft was submerged in the

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**Experimental methods**

High-quality synthetic muscovite mica (VI) was purchased from MTI Corporation with lateral dimensions of 1.5 cm\(^2\) and a thickness of 270 μm. The muscovite mica was adhered to the bottom of a petri dish via Loctite 409 Superbonder epoxy from Henkel, such that the epoxy was completely contained under the sample. Four environmental conditions were explored: (1) dry, (2) DI water, (3) 0.1 M NaCl aqueous solution (pH 9.8) and (4) 0.1 M NaCl aqueous solution (pH 12.0). The pH values were chosen due to both the mechanics of the indent and the environment. Finally, the plastic zone volumes \(V_{pl}\), initiation displacement \(h_{in}\), and energy dissipation \(E_{in}\) were assessed using a \(d(P)/d(h) = 0.3 \mu\text{N} / \text{nm}^2\) metric. A power-law relation between the burst load and energy was noted which separated the bursts into two continuous distributions: (1) bursts due only to the mechanics of the indent and (2) bursts due to both the mechanics of the indent and the environment. Finally, the plastic zone volumes \(V_{pl}\) necessary to initiate a burst were calculated assuming a hemispherical region under the tip; decreases in the minimum volume necessary for a displacement burst as a function of environment were justified through chemical attack of hydroxyl ion OH\(^-\) on Si—O bonds and ion exchange of K\(^+\) for Na\(^+\) at the surface defect sites. The mechanistic underpinnings of the property changes were then explored, with a focus on the consequences of environmental corrections to the fracture properties and power-law scaling in the energy data.
liquid as it would be during indentation. Since the liquid affects the perceived load on the tip and the amount of force necessary to move the tip in the z-axis [27], the z-axis calibration was completed in the liquid. At this point, the liquid nanoindentation method was identical to the standard, dry indentation process. All indentation arrays were centered 4 mm away from the edge of the sample and were completed within 12 h after the muscovite was immersed in the liquid. While each condition was indented 120 times, several indents did not run as planned due to an inability to locate the surface. Therefore, the number of indents analyzed for each condition were 115 for dry, 119 for DI water, 118 for NaCl (pH 9.8), and 117 for NaCl (pH 12.0). For lower load (2 mN) indents, subsequent scanning probe microscopy (SPM) images were captured at a normal load (2 mN) indents, subsequent scanning probe microscopy (SPM) images were captured at a normal load of 2 mN and a scan rate of 5 μm/s. 

Indentation of a layered material such as mica is dominated by displacement bursts to the extent that it is difficult to reproduce a specific load–displacement response. This can be seen in Fig. 2a through d. Figure 2a shows 115 indents into the dry (001) muscovite surface. Visual inspection of Fig. 2a shows that the majority of individual indents experience at least one large displacement burst at loads < 5 mN. Inspection of all indents in Fig. 2b through d showed that the response of mica in liquid produced displacement bursts at lower loads and displacements when compared to the dry case. The majority of indents demonstrated large displacement bursts at < 3 mN during indentation in DI water, < 1.9 mN during indentation in the 0.1 M NaCl pH 9.8 aqueous solution, and < 1.3 mN in the 0.1 M NaCl pH 12.0 aqueous solution. This data have been decomposed into burst-only data (Fig. 2e through h) and burst-free data (Fig. 2i through l). The burst-free data collapses to a relatively repeatable load–displacement response for each environmental condition. The influence of the liquid can be observed qualitatively in Fig. 2b through d which shows the P–h response of...
all analyzed indents for each condition. Figure 2b through d also suggests that both $E_r$ (the initial elastic unloading slope) and $H$ ($10 \text{ mN load divided by a larger contact area}$) decrease during indentation in liquid.

The reduced modulus and hardness of each indent was calculated using Eqs. (2–4), and the resulting means and standard deviations for each environmental condition are summarized in Table 1. From the table, it is clear that $E_r$ and $H$ decreased in the following order: dry, DI water, 0.1 M NaCl pH 9.8, and 0.1 M NaCl pH 12. The reduced modulus and hardness decreased by over 40% when comparing 0.1 M NaCl pH 12 to dry indentation. This was mainly due to an increase in frequency of displacement bursts for muscovite in liquid and a decrease of the muscovite stiffness due to intercalation of the liquid between the sheets once surface fractures occurred. Interestingly, $E_r$ and $H$ decreased at a constant ratio, such that $H/E_r$ remained relatively constant at 0.062 ± 0.004.

Table 1 Reduced modulus $E_r$ and hardness $H$ values (means and standard deviations) for each of the four environmental conditions

| Condition          | $E_r$ (GPa) | $H$ (GPa) | $H/E_r$ |
|--------------------|------------|-----------|---------|
| Dry                | 55.1 ± 5.8 | 3.53 ± 0.77 | 0.064   |
| DI water           | 52.8 ± 6.5 | 3.12 ± 0.51 | 0.061   |
| 0.1 M NaCl pH 9.8  | 39.9 ± 9.7 | 2.27 ± 0.47 | 0.058   |
| 0.1 M NaCl pH 12   | 31.0 ± 8.7 | 2.05 ± 0.48 | 0.066   |

$E_r$ and $H$ decreased in the following order: dry, DI water, 0.1 M NaCl pH 9.8, and 0.1 M NaCl pH 12. $H/E_r$ remained relatively constant at 0.062 ± 0.004.

Energy dissipation

The energy dissipated by an individual displacement burst can be calculated as the area under the load-
displacement curve from the beginning to the end of the displacement burst, in other words the burst length multiplied by its corresponding load generally produces an energy on the order of pJ ($J = N \times m$). In order to address which deformation modes the displacement bursts represent, it was necessary to analyze indents that generate only one or two displacement bursts. Figure 3a depicts three shallow indents with a maximum load of 2 mN. Subsequent scanning probe microscopy (SPM) imaging of each indent revealed that unique deformation modes were responsible for each of the displacement bursts. The SPM image of the indent in Fig. 3b indicated delamination between the muscovite basal layers labeled B1, which produced a step-height of $\approx 1.6$ nm as shown by the difference in the green and red markers on the associated line scan; $h_{b1}$, $P_{b1}$, and $E_{b1}$ for this displacement burst were found to be 67.3 nm, 576 mN, and 2.1 pJ, respectively, as shown in Table 2. The SPM image of the indent in Fig. 3c showed radial fracture originating at the contact point between the sample and the apex of the indenter tip, where the two major bursts labeled C1 and C2 expended 7.7 pJ and 11.2 pJ, respectively. Event C2 occurred on unloading; crack growth in low-toughness materials during unloading of an indentation is common [29]. The SPM image of the indent in Fig. 3d showed spalling of the muscovite to a depth of 35 nm, where the two major bursts labeled D1 and D2 (D2 occurred during the hold period) expended 12.3 pJ and 20.8 pJ, respectively. In this latter case, the line scan shows that the base of the spalled material lies along a single basal layer, indicating that the base was formed through a delamination event. In all, the results in Table 2 are in agreement with prior work that suggested smaller displacement bursts represent delamination [30] or kinking of the mica layers [31, 32], whereas larger bursts represent fracture or spallation [30, 33].

Displacement bursts during mechanical deformation are a stochastic process [34–36]. Therefore, large numbers of bursts were necessary to generate enough statistics in order to properly characterize them. Muscovite in the dry condition was indented 115 times generating a total of 1467 displacement bursts for an average of 12.8 bursts per indent. The same sample was submerged in DI water and indented 119 times causing a total of 2178 displacement bursts for an average of 18.3 bursts per indent. A second

Table 2

| Burst | $h_{b}$ (nm) | $P_{b}$ (mN) | $E_{b}$ (pJ) | Deformation mode |
|-------|--------------|--------------|--------------|-----------------|
| B1    | 67.3         | 576          | 2.1          | Delamination    |
| C1    | 160.1        | 1884         | 7.7          | Fracture        |
| C2    | 150.4        | 1477         | 11.2         | Fracture        |
| D1    | 98.6         | 951          | 12.3         | Fracture/spall  |
| D2    | 185.6        | 1998         | 20.8         | Fracture/spall  |

A burst energy of 2.1 pJ was linked to interlayer delamination, while fracture occurred at energies greater than 7 pJ.

![Figure 3](image_url)

**Figure 3**

(a) Nanoindentation of mica to a maximum load of 2 mN. Each indent exhibits at least one displacement burst, and the five major bursts are also detailed in Table 2. **b–d** Subsequent SPM imaging of each indent revealed a unique deformation mode. **b** The red indent indicated delamination between the muscovite basal layers labeled B1 and a step-height of 1.6 nm. **c** The green indent showed radial fracture originating at the contact point and two major bursts labeled C1 and C2. **d** The blue indent showed spalling and two major bursts labeled D1 and D2.
muscovite sample submerged in a 0.1 M NaCl solution with a pH of 9.8 was indented 118 times that produced 2844 bursts (average of 24.1 bursts per indent), and a third sample submerged in a 0.1 M NaCl solution with a pH of 12.0 was indented 117 times generating 2942 bursts (average of 25.1 bursts per indent). Analysis of all conditions totaled 469 indents accounting for 9431 displacement bursts; displacement bursts during unloading were not considered in this analysis. The energy expended during each displacement burst was then calculated using the area under the $P-h$ curve. Figure 4(a–d) summarizes the resulting $E_b$ as a function of $P_b$ data for all four conditions on log–log plots. From the data, a clear trend was observed that follows a power-law relationship of

$$E_b = 4.5 \times 10^{-5} P_b^{0.6}$$

This power-law expression was used to separate the displacement bursts into two groups. Bursts falling below the curve are graphed in black, while bursts falling above the curve are graphed in red. Figure 4e–h shows the bursts graphed as a function of indentation load and displacement in order to give context to the energy-load plots.

Histograms of the energy expended for the displacement burst subsets both above and below the power-law expression are presented in Fig. 5. In order to determine if they are generated from the same or different continuous distributions, the burst subsets were analyzed using a two-sample Kolmogorov–Smirnov test [37]. A comparison matrix of the asymptotic p-values from the test is shown in Table 3, where italic indicates that the null hypothesis was not rejected ($p$ value $> 0.05$, such that the two histograms are assumed to come from the same distribution), while bold indicates that the null hypothesis was rejected ($p$ value $< 0.05$, such that the histograms are assumed to come from different distributions). As shown in Table 3, histograms of the bursts below the power-law relationship are from the same distribution for each of the four conditions. This implies bursts below Eq. (7) are independent of the environment, and only depend on the mechanics of the indent. In other words, this subset of displacement bursts must be spatially isolated from the environment and occur deep within the plastic zone of the indent. In contrast, displacement bursts above Eq. (7) come from different distributions, showing that environment does play a role in the generation of these events. This subset of bursts was exposed to the environment, be it air or liquid. The bursts were generated by a combination of both the mechanics of the indent and the environment. Equation 7 is therefore physically meaningful even though it is empirical. For bursts above the power law, liquid...
increases the likelihood of bursts with $E_b$ in the range of 1–100 pJ when compared to the dry condition. As a point of reference, an $E_b$ of 2.1 pJ was associated with interlayer delamination, while fracture occurred at $E_b$ greater than 7 pJ, as shown in Table 2.

### Plastic zone volume

The plastic zone volume $V_{pl}$ for each displacement burst was calculated with Eqs. (5) and (6), using the $h_b$ values for each burst and the $H/E_r$ ratios in Table 1. The burst energy as a function of plastic zone volume for all bursts is shown in Fig. 6a. To reduce data, the first displacement burst for each indent with an energy $E_b$ greater than the specified value was identified. The volume $V_{pl}$ for that burst was tabulated for each indent to produce a cumulative distribution function (CDF) for each condition. As an example, the CDF for the first burst from each indent with an expended energy above 5 pJ is shown both for bursts below the power law in Fig. 6b, and for bursts above the power law in Fig. 6c. This was repeated for each condition over an energy range from 0.1 to 100 pJ in 0.1 pJ increments.

The mean value of the plastic zone volume of each CDF is graphed as a function of displacement burst energy in Fig. 7, with the displacement bursts below the power law in Fig. 7a and the displacement bursts above the power law in Fig. 7b. Figure 7b shows that the minimum threshold volumes for environmentally assisted fracture decreased with the addition of liquid. In more detail, the threshold volumes for dry muscovite, DI water, and the two 0.1 M NaCl solutions were 0.45 $\mu$m$^3$, 0.15 $\mu$m$^3$, and 0.04 $\mu$m$^3$, respectively. Thus, the NaCl solutions decreased the minimum volume for environmentally assisted fracture by an order of magnitude compared to dry mica. For all cases, bursts with volumes below these threshold values were due entirely to the mechanics
Figure 6  a $E_b$ as a function of $V_{pl}$ for all displacement bursts. b–c The first displacement burst for each indent with an $E_b$ greater than 5 pJ was graphed as a CDF for bursts b below the power-law relationship and c above the power-law relationship.

Figure 7  $E_b$ as a function of $V_{pl}$ for bursts a below the power-law relationship and b above the power-law relationship. In a, the fracture events occurred internally and were not exposed to the environment. In b, fracture events were exposed to the environment via fractures that intersected with the sample surface. The threshold volumes for dry mica, DI water, and the two 0.1 M NaCl solutions were 0.45 $\mu$m$^3$, 0.15 $\mu$m$^3$, and 0.04 $\mu$m$^3$, respectively.

of indentation and were not influenced by the environment. These initial events can be seen in Fig. 7a below the environmental threshold $V_{pl}$ for each condition. For confirmation, see the similarity in deformation response for loads below 1 mN in Fig. 2b and Fig. 4e–h. These bursts depicted the first kinking and delamination events and were internal to the structure. However, once surface fractures and spalling occurred, liquid intercalated between the mica layers, producing bursts from both environmental assistance and indentation mechanics.

Discussion

Two points from the earlier results required further examination: (1) the presence of mechanically dominated and environmentally assisted $E_b$ distributions and (2) the decrease in the minimum $V_{pl}$ for environmentally assisted fracture by an order of magnitude relative to dry mica. On point (1), the fracture bursts generated by “sharp” indenter tips are governed by the underlying elastic–plastic stress fields, and as such, are found to involve the hardness-to-modulus ($H/E$) ratio and fracture toughness [38]. For a material in an inert environment, it was illustrated that the length of an indentation crack $c_0$ in an unrelaxed residual stress field is given by [39]

$$c_0 = \left( \frac{\chi_0}{T_0} \right)^{2/3} p^{2/3}$$

where $T_0$ is the toughness and $\chi_0$ is a dimensionless factor based on the geometries of subsurface cracks and stress fields (both in inert conditions). In previous work, the dimensionless factor was found to be $\chi_0 = 0.016(H/E)^{-1/2}$; the initial term in this empirical relationship was determined by averaging data for “well-behaved” ceramic materials [39]. For a material in air or liquid, it was later shown that the length of an indentation crack $c_{env}$ in a relaxed residual stress field is given by [40]

$$c_{env} = \left( \frac{\chi_{env}}{T_{env}} \right)^{2/3} p^{2/3}$$

where $T_{env}$ is the toughness and $\chi_{env}$ is the dimensionless factor in environmental conditions. The effects of the environmental conditions on the geometries of subsurface cracks and stress fields can be quantified by combining Eqs. (8) and (9) to yield [40]

$$c_{env} = \left( \frac{\chi_0 - \Delta \chi_{env}}{T_0 - \Delta T_{env}} \right)^{2/3} p^{2/3}$$

where $\Delta \chi_{env} = \chi_0 - \chi_{env}$ and $\Delta T_{env} = T_0 - T_{env}$ are environmental corrections that are both positive. Previous data on a range of ceramics via inert [41] and reactive [42] indentation strength measurements
suggest that $\Delta T_{\text{env}} \approx 0.2c_0$ and $\Delta T_{\text{env}} \approx 0.5T_{\text{env}}$ consistent with observations [39] that show $c_{\text{env}} \approx 2c_0$ [40]. However, the exact details of the $c_{\text{env}}/c_0$ ratio are known to be dependent on test conditions (e.g., indentation load [43] and moisture content [44]). Despite the dependencies, it is clear that the differences in the overall $E_b$ distributions (Fig. 4) were from changes in the environmental conditions ($\Delta T_{\text{env}}$ and $\Delta T_{\text{env}}$), and not due to deviations in the inert fracture properties ($\chi_0$ and $T_{\text{env}}$, where $\chi_0 \sim (H/E)^{-1/2}$ and $H/E$ is constant for all conditions). This idea is further supported by Fig. 5 and Table 3, where the $E_b$ distributions below Eq. (7) are statistically similar (and governed by the mechanics of the indent) and the $E_b$ distributions above Eq. (7) are statistically different (and dictated by the mechanics of the indent and the environment).

On point (2), the two NaCl solutions exhibited the same threshold volumes for environmentally assisted fracture, but different fracture efficiencies for energies ranging from $1.5 \, \text{pJ} < E_b < 14 \, \text{pJ}$; the pH 12 solution was more efficient in producing bursts in that range. This is due to the increase in pH, or activity of the hydroxyl ions ($\text{OH}^-$). This increase in efficiency can be explained with two plausible chemical mechanisms: chemical attack by the hydroxyl ions on the Si–O bonds (like that for quartz) [14] and ion exchange, where Na$^+$ in solution replaces K$^+$ in muscovite at the frayed surface edges and possibly within surface defect sites, inducing lattice strain due to size mismatch [45]. For burst energies from $14 \, \text{pJ} < E_b < 50 \, \text{pJ}$, muscovite submerged in DI water had the smallest average plastic zone size, while dry muscovite had the smallest plastic zone size for energies $> 50 \, \text{pJ}$. The large number of bursts generated by the liquid environment decreased the stored elastic energy in the plastic zone to a level that made it difficult for the average contact stress to build to a high enough value to generate large bursts. This is similar to plate tectonics, as it is difficult for an active fault with numerous small earthquakes to store up enough energy to produce a catastrophic earthquake, whereas a less-active fault can fail catastrophically [46]. Interestingly, earthquakes are a standard example of a self-organized critical (SOC) system, a dynamic system that organizes itself to a critical point irrespective of the initial state [47, 48]. At this critical point, a model SOC system exhibits power-law scaling of event sizes, event durations, and in some cases, the $1/f$ scaling of the power spectra [49]. As such, the self-similar or scale-free behavior is described via a scaling exponent $\alpha$, enabling features at one scale to depict features at another scale across two or more orders of magnitude. Several small-scale mechanical phenomena demonstrate SOC behavior, including microfracture in disordered materials [50], shear-induced rearrangements in two-dimensional foams [51], dislocation motion in Ni [35] and Cu microcrystals [52], and structural transitions in self-assembled monolayers [53]. Two common themes were clear from these systems: the extracted values for $\alpha$ (1.3–1.8) were in agreement with those from theoretical models (1.2–2.0) [54] and the subsequent scaling behavior was used to link the nanoscale events to macroscale phenomena. Curiously, the power-law trends detected in Fig. 4 ($\alpha = 1.6$) and Fig. 5 ($\alpha = 1.2$, trendline not shown) fell within these ranges, signifying that chemomechanical weakening of muscovite could represent an SOC system. If so, the bursts in liquid nanoindentation can potentially be used as proxies for delamination, fracture, and spalling events in macroscale layered rocks such as shale, making this new approach a useful tool the development of effective subsurface engineering strategies.

### Conclusions

In summary, in situ liquid nanoindentation was used to characterize chemomechanical weakening of muscovite in four different environmental conditions. From the $P$–$h$ traces, it was found that $E_r$ and $H$ decreased by over 40% when comparing 0.1 M NaCl pH 12 to dry indentation, mainly due to an increase in frequency of displacement bursts and a decrease of the mica stiffness. However, $E_r$ and $H$ decreased at a constant ratio, such that $H/E_r$ remained invariant with environment. The $P$–$h$ traces also demonstrated a large number of displacement bursts. Subsequent SPM imaging of selected low-load indents revealed that bursts were linked to unique deformation modes (e.g., delamination, radial fracture, and spalling) at $E_b$ from 2.1 to 20.8 pJ. The $E_b$ data from all bursts exhibited a power-law trendline as a function of $P_{\text{env}}$; bursts below the trendline originated from a continuous $E_b$ distribution and were dictated solely by the mechanics of the indent, whereas bursts above the trendline came from disparate $E_b$ continuous distributions and were controlled by both mechanics and
environment. The minimum $V_{pl}$ for environmentally assisted fracture decreased by an order of magnitude for the NaCl solutions relative to the dry case. Moreover, an increase in pH led to an increase in fracture efficiency in the low $E_b$ regime, presumably due to an increase in OH$^-$ chemical activity and attack by the hydroxyl ions on the silicate framework and, potentially, an increase in lattice strain at the frayed surface edges and within surface defect sites due to K$^+$ for Na$^+$ cation exchange. Future work will look to quantify the environmental corrections to fracture ($\Delta V_{env}$ and $\Delta T_{env}$) using measurements in inert conditions and to explore the possibility of scale-free behavior through a mean-field theory model.

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