

1 **The role of surface forces in environment-enhanced cracking of brittle solids**

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3 **ABSTRACT**

4 Fracture initiation and propagation in brittle materials is promoted in surface-reactive (sorptive)
5 environments, a phenomenon known as subcritical crack growth (SCG). Laboratory measured crack-
6 propagation velocity vs. stress intensity factor relationships typically exhibit highly nonlinear, multi-
7 stage characteristics that are sensitive to environmental factors such as adsorbate concentration and
8 temperature. For practical purposes, empirical relationships (e.g., a power law) have been used to
9 describe this complex phenomenon. However, how the overall SCG behavior emerges from the
10 underlying fundamental processes near the crack tip, such as the interaction of the crack surfaces
11 separated by only a few nanometers and mass transport within the nano-confined space, is still not well
12 understood. This paper develops a mechanistic, surface-force-based fracture theory (SFFT) which
13 integrates surface force models, fluid transport models, and linear elastic fracture mechanics to
14 quantitatively explain the multi-stage characteristics of SCG in brittle solids. A numerical model is
15 developed based on SFFT and solved through an implicit partitioned scheme for efficiency and
16 modularity. The results are validated by Wiederhorn's data on crack propagation in soda-lime glasses at
17 a wide range of relative humidity levels. We show that, for the first time, the entire range of an SCG
18 curve can be captured by a single physics-based model. The predicted SCG curves reveal that the
19 development of repulsive disjoining pressure behind the crack tip can be responsible for the reduced
20 apparent fracture toughness in a sorptive environment. The shape of the SCG curve, and its changes with
21 respect to the environment, is found to critically depend on the assumed transport models.

22 **Key words:** fracture, kinetics, subcritical crack growth, surface force, sorption.

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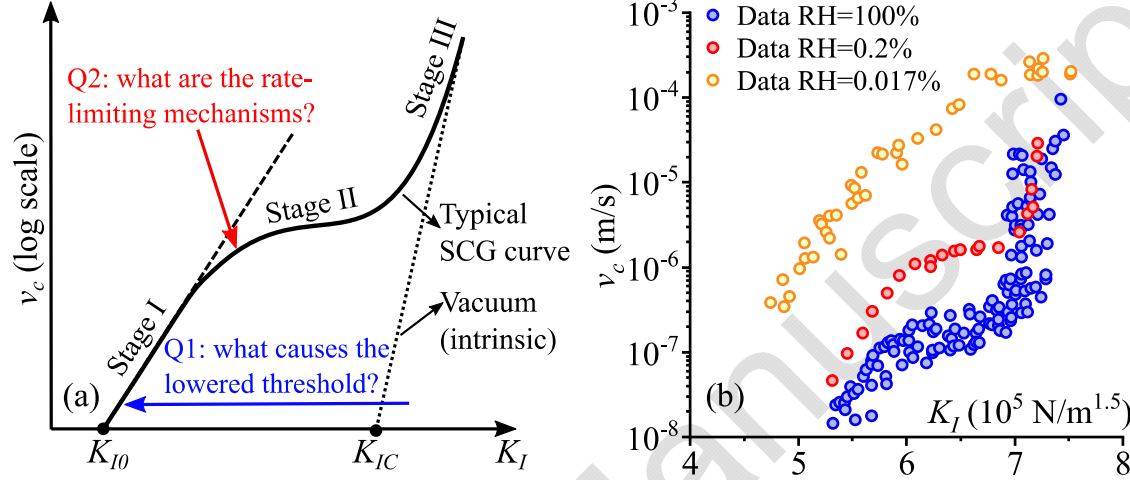
23 **1. BACKGROUND**

24 Subcritical crack growth plays a crucial role in the long-term mechanical performance of natural and
25 engineered materials. In sorptive environments, or as referred to by Rice (1978) the *surface-reactive*
26 environments, stable crack growth with a finite velocity (v_c) can occur at stress intensity factors (K_I)
27 significantly lower than the critical one (K_{IC}) measured under the apparent ‘dry’ or vacuum conditions.
28 This phenomenon is known as subcritical crack growth (SCG) (Atkinson, 1982) or static fatigue
29 (Charles, 1958). SCG controls the rate of important geological processes such as the diagenesis of
30 granular rocks (Chester et al., 2007) and weathering (Eppes and Keanini, 2017). Because it can be
31 viewed as a precursor to abrupt fracturing, understanding and modeling SCG can help improve the
32 prediction of volcanic eruptions (Kilburn and Voight, 1998) and fault failure (Lennartz-Sassinek et al.,
33 2014). SCG may also contribute to the slow degradation of engineering materials (such as concrete,
34 glass, metal, sand, rock), which is important in civil and mechanical engineering practice (Freiman et al.,
35 2009; Karimpour and Lade, 2010; Le et al., 2009; Simmons et al., 1978; Zhang and Buscarnera, 2018).

36 The key challenge of quantitative modeling of SCG is rooted in its nonlinear multistage behavior
37 resulted from complex physiochemical interactions at various length scales. SCG typically exhibits three
38 distinct stages in the $v_c - K_I$ space (Fig. 1a). At relatively low K_I (stage I), v_c increases rapidly with
39 K_I due to solid-environment interactions; at intermediate K_I (stage II), v_c is believed to be controlled
40 by the transport of the active species; stage II continues until a sudden and rapid increase of v_c (stage III)
41 as K_I exceeds the intrinsic fracture threshold (K_{IC}) of the material in vacuum (Freiman et al., 2009;
42 Wiederhorn, 1967). The stage I and II curves strongly depend on environmental conditions such as
43 temperature and reactant (adsorbate) concentration in the environment (Fig. 1b) (Crichton et al., 1999;
44 Wiederhorn et al., 1980). In this context, our SCG model aims to address two key questions: (Q1) how

45 do solid and environment interact to cause the reduced fracture toughness (from K_{IC} to K_{I0} in Fig. 1a)
 46 and the associated exponential kinetics in stage I? and (Q2) what transport mechanism (or combination
 47 of mechanisms) is at work near the crack tip to effectively limit the crack velocity in stage II?

48



49
 50 Fig. 1. (a) Typical SCG curve and the two fundamental questions. (b) SCG data of soda-lime glass at different relative
 51 humidity, RH (Wiederhorn, 1967; Wiederhorn et al., 1980).

52 Regarding the first question, many hypotheses have been proposed for different material-
 53 environment systems. Energy-based approaches explain the reduction of apparent fracture toughness
 54 through the decreased surface energy caused by the adsorption of fluid species on the crack surfaces
 55 (Rice, 1978; Wan et al., 1990). In contrast, atomistic approaches focus on crack-tip chemical reactions
 56 such as hydrolysis of stretched crack-tip bonds, dissolution and transport of crack tip material, and ion
 57 exchange with the crack tip (Atkinson, 1984; Michalske and Freiman, 1982). Concurrent with these
 58 mechanisms are the crack kinetic models by making analogy between crack propagation and reaction
 59 kinetics (Bazant and Planas, 1997; Lawn, 1975). Alternatively, empirical $v_c - K_I$ relationships such as
 60 power law are used as a descriptive tool without considering the underlying mechanisms driving the
 61 crack propagation (Bazant and Planas, 1997; Charles, 1958).

62 For the second question, some mechanisms such as Fickian diffusion (Wiederhorn, 1967), Knudsen
 63 diffusion (Lawn, 1974), and surface diffusion (Crichton et al., 1999) have been proposed to govern the

64 fluid invasion along a crack. However, the temperature-dependence of SCG data observed in phosphate
65 laser glasses suggests that none of these mechanisms alone can fully describe the stage II behavior
66 (Crichton et al., 1999). Mass transport in nano-confined spaces is still an active area of research (Choi et
67 al., 2001; Cihan et al., 2019) and is poorly understood at this moment for the near-crack-tip region. For
68 these reasons, to this day, the modeling of stage II and the transition from stages I to II remains highly
69 simplistic and phenomenological. Frequently, the stage II and III behaviors are neglected altogether, and
70 only power-law relationships are fitted to the $v_c - K_I$ data obtained in stage I, across a range of stresses
71 and environmental conditions (Brantut et al., 2013; Eppes et al., 2018; Nara et al., 2012; Olson, 1993).

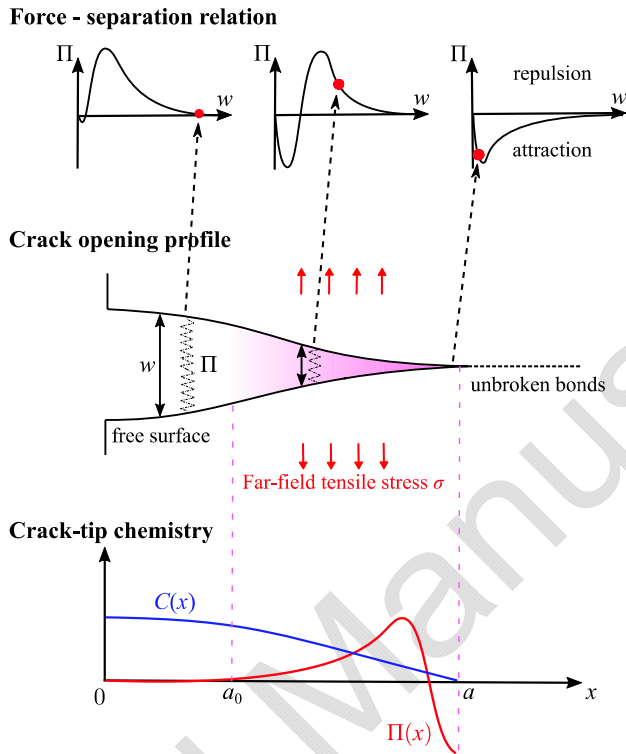
72 Recently, atomic force microscope (AFM) and surface force apparatus (SFA) experiments revealed
73 that repulsive forces (e.g., electrical double layer, hydration, crystallization) can become dominant
74 between mineral surfaces at nanometer separations in the presence of sorptive species (Dziadkowiec et
75 al., 2018; Røyne et al., 2015). Specifically, SFA measurements and molecular dynamic simulations (MD)
76 suggest that the nano-confinement of solutions between calcite surfaces can exert repulsion over
77 hundreds of nanometers of separation (Diao and Espinosa-Marzal, 2016; Dziadkowiec et al., 2019).
78 AFM measurements confirm that forces of similar range exist between glass surfaces in aqueous
79 solutions (Acuña and Toledo, 2008), and their magnitudes depend on the chemical composition of the
80 environmental fluid (Adler et al., 2001). This inspires an alternative explanation of the complex SCG
81 behavior: the repulsive forces along the crack walls can possibly facilitate the initiation and propagation
82 of cracks in sorptive environments, resulting in the characteristic trend of an SCG curve in Fig. 1.
83 Although the potential of surface forces in affecting solid fracture was debated in the glass community a
84 few decades ago (Lawn, 1985; Wiederhorn and Fuller Jr, 1989), it has been largely neglected since then,
85 without much progress.

87 **2. A SURFACE-FORCE BASED FRACTURE THEORY**

88 Provided the renewed evidence of repulsive surface forces induced by surface sorption, we propose
89 an SCG theory to assess a mechanistic, surface-force-based answer to both questions. The key strategy
90 is illustrated in Fig. 2 and explained in the following. In this model, surface forces along crack surfaces
91 are accounted for via a force-separation relation. For a homogenous material in perfect vacuum, $\Pi(w)$
92 is defined by the force required to overcome the solid-solid intermolecular forces in separating two
93 surfaces and is the same everywhere along the crack. The interacting force at different location x along
94 the crack surface can be read from this $\Pi(w)$ curve given the local crack separation $w(x)$. The
95 resistance against crack opening provided by the attractive surface force is macroscopically manifested
96 as the fracture toughness K_{IC} in classical fracture mechanics (Meng and Thouless, 2019). This part is
97 similar to the well-established cohesive crack theory (CCT), where the cohesive forces in the fracture
98 process zone (FPZ) counterbalance the far-field tensile stress to give a net-zero stress intensity factor at
99 the crack tip (Barenblatt, 1962).

100 Now consider that the crack is invaded by sorptive species, resulting in a non-uniform distribution of
101 species concentration $C = C(x)$ along the crack as shown at the bottom of Fig. 2. Intrusion of the
102 sorptive molecules can shift the local $\Pi(w)$ towards the repulsive side (Eskandari-Ghadi and Zhang,
103 2022), the magnitude of which depends on the local chemistry $C(x)$. The surface force at different
104 locations therefore starts to travel along different $\Pi(w)$ curves as opposed to the vacuum case. The
105 $C(x)$ profile is dictated by the species transportation along the crack and is thus coupled with the crack
106 opening profile $w(x)$, the ambient concentration C_0 , and the crack velocity v_c . The proposed theory can
107 potentially explain the first question (Q1) through the reduced attraction or the development of repulsion

108 due to the presence of sorptive species. The second question (Q2) can be also directly addressed through
 109 the coupling between fracture propagation and species transport in cracks.



110
 111 Fig. 2. The key elements of the proposed surface-force based fracture theory.

112 The proposed theory operates at an intermediate scale which is larger than the atomistic theories
 113 without relying on the dynamics of the crack-tip bond but is smaller than the macroscopic energy
 114 approaches by requiring the full resolution of crack opening profile and stress fields near the crack tip. It
 115 also differs from the cohesive crack theory in that, the traction curve in CCT is a constitutive relation
 116 commonly fitted to macroscale fracture testing data, involving only attractive force, and does not vary
 117 along the crack. In contrast, the proposed approach is rooted in surface physics, involves intermediate-
 118 scale force-separation relationships which are either directly measurable by SFA experiments or
 119 theoretically derivable based on intermolecular potentials, and can be strongly repulsive and vary with
 120 the local environment. We therefore refer to the proposed theoretical framework as the Surface-Force
 121 based Fracture Theory (SFFT) to distinguish from the previous works.

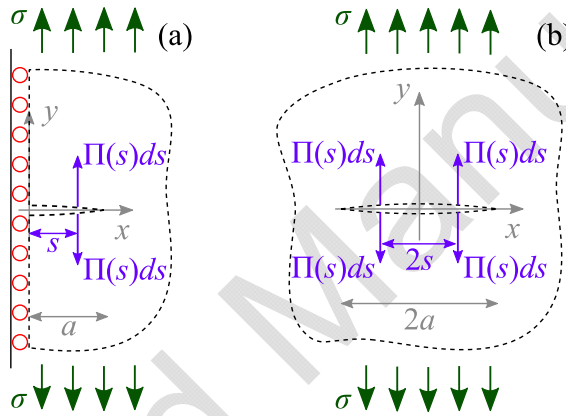
122 The remainder of this paper shall tackle two specific goals: (1) to formalize the mathematical
123 structure of SFFT for mode-I fractures and (2) to numerically implement an SFFT model and test its
124 ability in capturing typical SCG behaviors. For demonstration, the model will be specialized for the
125 glass-in-vapor experiments of Wiederhorn (1967) as our first step towards advancing our general
126 understanding of environment-enhanced cracking of brittle materials. We start by laying out the stress
127 intensity, opening profile, and propagation kinetics of cracks with considering both far-field stress and
128 distributed stresses along the crack surface (Section 3). We then formulate simplified analytical models
129 for surface force under variable environment (Section 4) and species transport along a moving non-
130 parallel slit (i.e., a propagating crack) (Section 5). We numerically solve the system via a partitioned
131 implicit scheme to ensure a robust and efficient coupling between the aforementioned processes (Section
132 6). Finally, the model performance is assessed using the SCG data of soda-lime glass at different
133 humidity levels as presented in Fig. 1b (Section 7). The main conclusions and possible future extensions
134 of this work are discussed at the end (Section 8).

135 **3. FRACTURE MODELING CONSIDERING SURFACE FORCES**

136 **3.1. Fracture mechanics analysis**

137 We adopt the approach of Lawn (1985) by directly acknowledging the disjoining pressure (Π) in
138 the LEFM analysis. The *disjoining pressure* Π is here defined as the net surface force between two
139 solid surfaces normalized by the surface area (Clarke et al., 1986). It lumps the contributions of all solid-
140 fluid interactions together without discerning their physiochemical origins and takes a positive value
141 when the net force is repulsive (or disjoining) and negative when cohesive (or joining). Such distributed
142 pressure along the crack surface can alter the stress field, characterized by the *effective* stress intensity
143 factor (K_{le}), in the vicinity of the crack tip. K_{le} should be distinguished from the applied stress intensity
144 (K_{la}) which solely accounts for the external forces or far-field stresses on the cracked solid.

145 Let us consider a 2D (plane-strain) edge crack of finite length a (Fig. 3a) in an elastic half-space
 146 subject to far-field tensile stress and disjoining pressure $\Pi(x)$ over the crack surface. The boundary
 147 conditions are selected such that the problem is equivalent to an infinite body with a line crack (Irwin,
 148 1957) (Fig. 3b). This allows one to take advantage of the readily available fundamental solutions to
 149 calculate the mode-I crack opening and stress intensity under various loading combinations (Tada et al.,
 150 2000). The dependency of $\Pi(x)$ on fluid chemistry and species transportation will be addressed in
 151 section 4.



152
 153 Fig. 3. Schematic of the model geometry and loading scenarios. (a) A finite edge crack in semi-infinite elastic domain. (b) A
 154 finite center crack in elastic infinite domain. The boundary conditions in (a) guarantee equivalent stress-deformation solution
 155 as (b).

156 LEFM allows one to write the effective stress intensity factor in an additive fashion

157

$$K_{le} = \underbrace{k_{I\sigma}\sigma}_{K_{la}} + \underbrace{\int_{s=0}^{s=a} k_{I\Pi}(s)\Pi(w(s); s)ds}_{K_{I\Pi}} \quad (1)$$

158 where $k_{I\sigma}$ is the stress intensity factor caused by unit far-field tensile stress; σ is the magnitude of the
 159 tensile stress; $k_{I\Pi}(s)$ is the stress intensity factor caused by unit repulsive force on the crack surface at
 160 location s ; $w(x)$ is the crack opening at location x ; $\Pi(w)$ describes a disjoining pressure vs.
 161 separation relation which will be given by the surface force model (see Section 4); K_{la} is the applied

162 stress intensity factor; K_{II} is the stress intensity contribution by the disjoining pressure and takes a
 163 negative value when the net effect of Π is cohesive.

164 For a given σ , the distribution $\Pi(x) = \Pi(w(x); x)$ is not known a priori. Π depends on the surface
 165 separation while its change in turn alters the crack opening profile. The expression of $w(x)$ is given
 166 below based on LEFM (Bazant and Planas, 1997):

$$167 \quad w(x) = c_\sigma(x)\sigma + \int_{s=0}^{s=a} c_\Pi(x, s)\Pi(w(s); s)ds \quad (2)$$

168 where $c_\sigma(x)$ is the crack opening at location x along the crack caused by unit far-field tensile stress;
 169 $c_\Pi(x, s)$ is the crack opening at the same location caused by a unit repulsive force at location s .

170 For the geometry sketched in Fig. 3a and b, $k_{I\sigma}$ and $c_\sigma(x)$ are given by Tada et al. (2000)

$$171 \quad k_{I\sigma} = \sqrt{\pi a} \quad \text{and} \quad c_\sigma(x) = \frac{4(1-\nu^2)}{E} \sqrt{a^2 - x^2} \quad (3)$$

172 Similar fundamental solutions can be found for unit surface force applied at s :

$$173 \quad c_\Pi(x, s) = \frac{8(1-\nu^2)}{\pi E} \begin{cases} \tanh^{-1} \sqrt{\frac{a^2 - s^2}{a^2 - x^2}} & ; \quad x < s \\ \coth^{-1} \sqrt{\frac{a^2 - s^2}{a^2 - x^2}} & ; \quad x > s \end{cases} \quad (4)$$

$$174 \quad k_{II}(s) = 2\sqrt{\frac{a}{\pi}} \frac{1}{\sqrt{a^2 - s^2}} \quad (5)$$

175 It should be noted that Eqns. (4) and (5) are singular at $x = s$. This introduces difficulties to the
 176 numerical integrations of Eqns. (1) and (2). We have circumvented this difficulty by means of numerical
 177 approximation which allows the usage of non-singular analytical solutions (see Appendix A for details).

178 The non-linear coupling between $w(x)$ and $\Pi(w)$ can be assembled as a system of two algebraic
 179 equations, referred to as the $\Pi - w$ system hereafter, to be solved simultaneously (see Appendix A). The
 180 solution to the $\Pi - w$ system provides a crack opening profile $w(x)$ which will be substituted into Eq.
 181 (1) in conjunction with the $\Pi(w)$ model (see Section 4) to obtain K_{le} .

182 **3.2. Crack propagation kinetics**

183 The rate at which crack propagates can be studied in the framework of reaction rate theory (Charles,
 184 1962). Griffith's criterion suggests that non-dissipative crack propagation occurs once $(\mathcal{G} - 2\gamma) > 0$,
 185 where \mathcal{G} is known as the energy release rate and γ is the surface tension of the newly created crack
 186 faces. By analyzing the statistics of bond breakage and healing under the energy difference $(\mathcal{G} - 2\gamma)$ in a
 187 way similar to a chemical reaction driven by chemical potential differences (Lawn, 1975), one can arrive
 188 at the following kinetic equation

$$189 v_c = v_0 \exp\left(-\frac{Q_0}{RT}\right) \sinh\left(\frac{\tilde{\kappa}_0(\mathcal{G} - 2\gamma)}{RT}\right) \quad (6)$$

190 where $\tilde{\kappa}_0$ is a proportionality factor; Q_0 is the activation energy for bond breaking; v_0 contains
 191 information of the solid lattice vibration frequency and lattice length-scale (Meng and Thouless, 2019).
 192 In Eq. (6), the “driving force” for crack propagation is $(\mathcal{G} - 2\gamma)$. Bazant and Planas (1997) have shown
 193 that stress intensity factor can be also taken as the driving force to replace $(\mathcal{G} - 2\gamma)$ in Eq. (6). Inspired
 194 by this, we propose a slightly different equation for crack growth in the presence of disjoining pressure

$$195 v_c = v_0 \exp\left(-\frac{Q_0}{RT}\right) \sinh\left(\frac{\kappa_0 K_{le}}{RT}\right) \quad (7)$$

196 where κ_0 is a proportionality factor related to $\tilde{\kappa}_0$. The decomposition of K_{le} to K_{la} and K_{Π} (i.e., Eq.
 197 (1)) reveals some connections between Eqns. (6) and (7). Firstly, the celebrated Irwin equation directly

198 links K_{Ia} and K_{II} , respectively, to \mathcal{G} and 2γ at equilibrium. $K_{Ie} = 0$ implies that $\mathcal{G} = 2\gamma$ and $v_c = 0$,
199 agreeing that a non-propagating crack attains zero stress intensity and thus finite tensile stress at the tip
200 (Barenblatt, 1962). Secondly, the generation of repulsive disjoining pressure in sorptive environment
201 will reduce K_{II} (Eq. (1)), thus promoting crack propagation under the same applied K_{Ia} . This is
202 equivalent to the surface energy reduction mechanism discussed by Rice (1978). We further argue that
203 Eq. (7) has the unique advantage of directly incorporating the nonuniform reduction of surface energy
204 along the crack (via K_{II}) caused by transient processes such as fluid transport, the inclusion of which in
205 energy-based descriptions is not straightforward.

206 4. SURFACE FORCE MODEL

207 The magnitudes of surface forces depend on the crack aperture w (Bazant and Planas, 1997) as well
208 as the local environment C (Israelachvili, 2011). The net effect of these forces over unit surface area
209 can be collectively encapsulated in the disjoining pressure $\Pi(w, C)$, the modeling of which is essential
210 for the proposed fracture theory. Before tackling this task, we should clarify the ambiguity related to the
211 definition of “crack tip”.

212 From an atomistic perspective, the crack tip may be defined at the location where the solid chemical
213 bonds are broken. Thermodynamically, one may define the crack tip as the point behind which the
214 separation of opposite surfaces is irreversible and involves energy dissipation, and in front of which the
215 separation occurs reversibly or elastically. Indeed, the location of the “crack tip” behind and ahead of
216 which different inelastic processes may be at work is only a matter of perspective and convenience (Sills
217 and Thouless, 2015). From a continuum point of view, undamaged material obeys elastic behavior, and
218 the damaged material in the fracture process zone (FPZ) obeys a softening stress-strain relation (Bažant
219 and Oh, 1983). The point of divergence between continuum and localized crack happens at the

220 maximum value of traction on the force-separation curve. This motivates us to define the crack tip as the
221 point of maximum traction between two surfaces, and the corresponding crack separation at this
222 reference point is taken as $w = 0$ (See Fig. 4).

223 The consideration of purely cohesive interactions (traction) over several micrometers behind crack
224 tip leads to the well-known cohesive crack theory. It has been successfully applied, for example, to
225 remove the stress singularity predicted by LEFM at the crack tip (Barenblatt, 1962), model the kinetics
226 of crack growth (Meng and Thouless, 2019), and capture crack fatigue (Allegri, 2020; Nguyen et al.,
227 2001). It also hosts various material-specific fracture models by treating the traction-separation curve as
228 a constitutive relation (Hillerborg et al., 1976; MAI and Lawn, 1987; Needleman, 1990; Tijssens et al.,
229 2000; Tvergaard and Hutchinson, 1992). However, the true surface forces originated from the physical
230 separation (<100 nm) of solid surfaces (electrostatic, van der Waals, etc.) and environmental attack
231 (surface hydration, electrical double layer, etc.) has been mostly neglected in classical fracture analysis.
232 The goal here is to substantialize a $\Pi(w, C)$ model to capture these nanometer-scale interactions which
233 can be attractive or repulsive.

234 Generally, the disjoining pressure between two solid surfaces in arbitrary environment can be
235 decomposed into two contributions:

236
$$\Pi(w, C) = \Pi_0(w) + \Delta\Pi(w, C) \quad (8)$$

237 where C stands for the concentration of the sorptive species in between surfaces. C could be
238 generalized to a vector $\mathbf{C} = [C_1, C_2, \dots, C_k]$ should multiple sorptive species present in the environment.
239 The first component, Π_0 , accounts for the intrinsic surface forces between two solid surfaces in vacuum;
240 the second term, $\Delta\Pi$, collects the change of disjoining pressure due to solid-species interaction (Adapa
241 and Malani, 2021; Dziadkowiec et al., 2018; Røyne et al., 2015).

242 4.1. Surface force in vacuum $\Pi_0(w)$

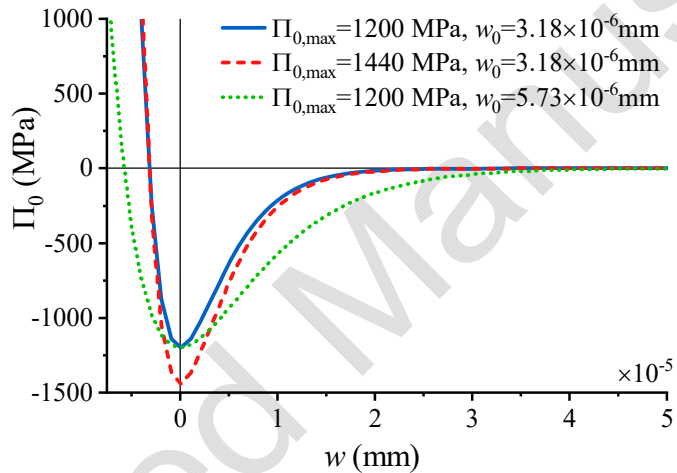
243 $\Pi_0(w)$ may be constructed through smooth interpolations of experimental data provided by
 244 SFA/AFM measurements, or rigorously derived with the knowledge of the total intermolecular pair
 245 potentials of the solid (Israelachvili, 2011). Here we opt for a well-behaved and versatile analytical
 246 expression to represent the in-vacuum surface forces of generic materials. Simple analytical models also
 247 benefit the numerical implementation in terms of both convergence and efficiency. A two-parameter
 248 equation is adopted for this purpose:

$$249 \quad \Pi_0 = -\Pi_{0,\max} \frac{w+w_0}{w_0} \exp\left(1 - \frac{w+w_0}{w_0}\right) \quad (9)$$

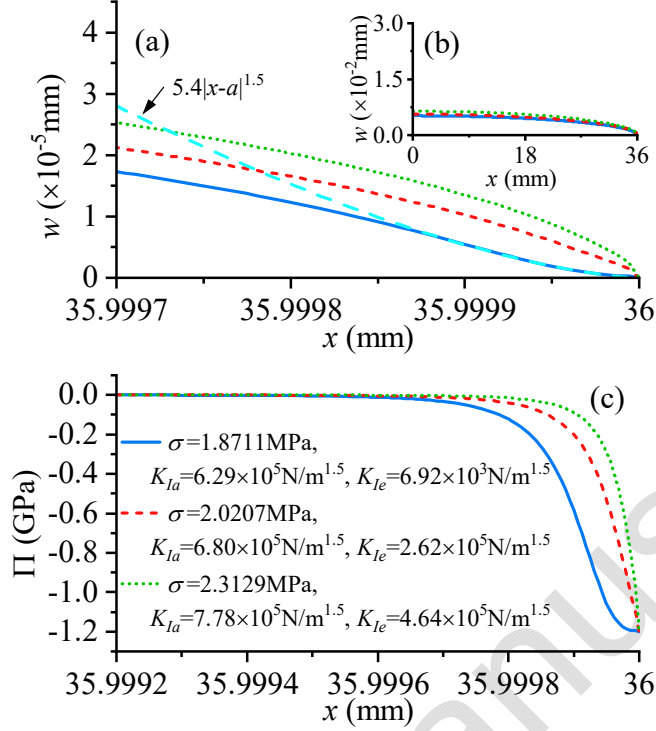
250 where Π_0 attains maximum attraction $\Pi_{0,\max}$ at $w=0$ and decays exponentially for $w>0$, the rate of
 251 which is controlled by w_0 . Eq. (9) also contains an exponentially increasing repulsive branch for $w<0$,
 252 which physically corresponds to the steric repulsion when molecules are brought too close together
 253 (Israelachvili, 2011). This feature is also expected to be numerically beneficial by serving as a penalty
 254 contact to avoid the overlap of two surfaces when they come to contact, in cases where crack healing
 255 may occur (i.e., $K_{le} < 0$). Eq. (9) is visualized in Fig. 4 for three sets of parameters.

256 The solution of Eqns. (2) and (9) together allows us to study crack opening at different levels of
 257 applied far-field stress in vacuum. Using the numerical method detailed later in Section 6 and
 258 parameters in Table 1, the w and Π_0 profiles for a crack of length 36mm under increasing stresses are
 259 plotted in Fig. 5. When the stress is low, the in-vacuum surface force contribution of stress intensity
 260 K_{Π} can effectively balance out the applied stress intensity K_{la} , thus gives a low K_{le} value and a near-
 261 equilibrium crack per Eq. (7). This results in a smoothly converging crack tip profile (the solid blue
 262 curve in Fig. 5a). This is consistent with Barenblatt (1962) who analytically showed that the crack

263 opening profile in the immediate vicinity of the crack tip becomes proportional to the 1.5 power of the
 264 distance from the crack tip when $K_{Ie} \rightarrow 0$ (the dashed cyan curve). As the applied force increases, the
 265 crack is perturbed further away from equilibrium, and its tip geometry becomes blunt and approaches an
 266 elliptical shape. Besides the near-tip region, the overall crack opening profile (Fig. 5b) does not vary
 267 significantly for the range of applied σ , except for a proportional widening of the crack with increasing
 268 σ . Fig. 5c shows that the resultant Π_0 increasingly concentrates at the crack tip as the applied stress
 269 increases and the crack aperture widens up.



270
 271 Fig. 4. The in-vacuum surface force described by Eq. (9). $\Pi_{0,\max}$ controls the magnitude of Π_0 while w_0 controls its spread
 272 along w .

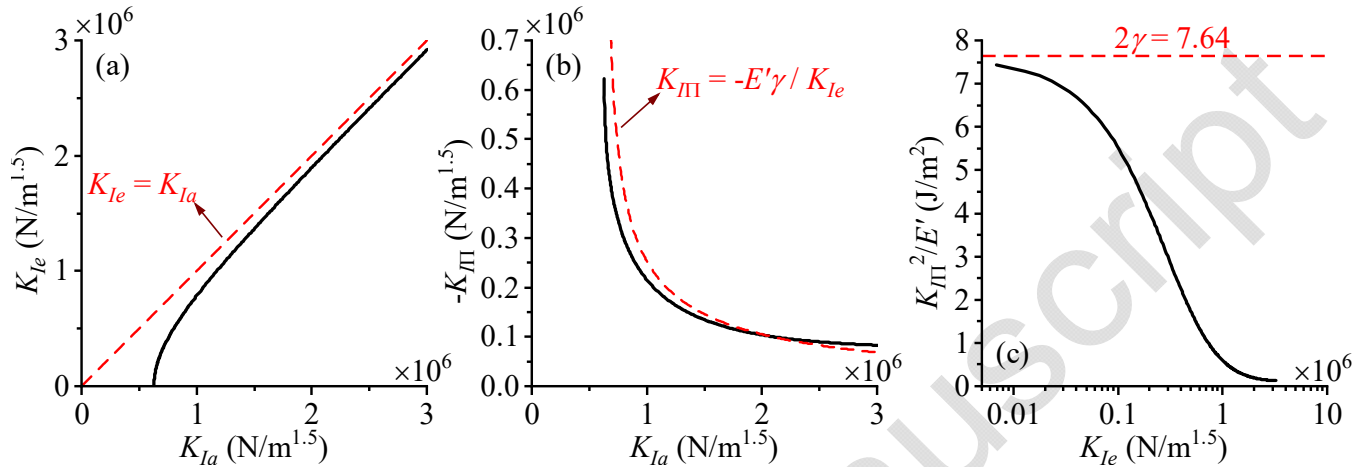


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Fig. 5. The computed (a) crack tip opening profile, (b) overall crack opening profile, and (c) disjoining pressure distribution in vacuum for progressively increasing far-field tensile stress. The model parameters used here are listed in Table 1.

276 Figures 6a and 6b plot the computed K_{Ie} and K_{II} against a wide range of applied K_{Ia} using the
277 same parameters as Fig. 5. As expected, the presence of cohesive forces results in a reduced K_{Ie}
278 compared to K_{Ia} . Increasing K_{Ia} results in a decrease of K_{II} (Fig. 6b), and thus K_{Ie} asymptotes
279 towards K_{Ia} (Fig. 6a). Lawn (1985) analytically derived an expression of K_{II} as $K_{II} = -E'\gamma / K_{Ie}$,
280 where $E' = E / (1 - \nu^2)$ for plane-strain, by neglecting the dependence of crack opening profile on Π_0 .
281 For the selected parameters, this approximation is surprisingly accurate for large K_{Ia} values but deviates
282 from our exact solution of the full $\Pi - w$ system at lower K_{Ia} levels (Fig. 6b). This implies that the
283 consideration of the two-way coupling between Π and w is essential for studying the onset of SCG
284 which always occurs at low K_{Ia} values. Fig. 6c shows that K_{II}^2 / E' asymptotes to 2γ as equilibrium
285 ($K_{Ie} = 0$) is approached. This further supports our arguments that Eq. (7) can be viewed as the

286 mechanistic counterpart to Eq. (6), and the Irwin's equation respectively links K_{Ia} and K_{II} to \mathcal{G} and
 287 2γ at equilibrium.



288 Fig. 6. Solutions of the Π - w system in terms of (a) K_{Ie} versus K_{Ia} to compare with the prediction by Lawn (1985);
 289 (b) $-K_{II}$ versus K_{Ia} to compare with the approximate analysis by Lawn (1985); and K_{II}^2 / E' versus K_{Ie} to study the
 290 application of Irwin's equation in linking K_{II}^2 / E' and 2γ at equilibrium. E' is the elastic constant and equal to $E / (1 - \nu^2)$
 291 for plane strain condition. Parameters used are summarized in Table 1,
 292

293 4.2. Induced disjoining pressure $\Delta\Pi$

294 The magnitude of $\Delta\Pi$ depends on both the surface separation w as well as the local concentration
 295 of sorptive species C (Adler et al., 2001; Eskandari-Ghadi and Zhang, 2021). To construct a $\Delta\Pi$ model,
 296 let us focus the discussion on Wiederhorn (1967)'s glass-in-vapor system from hereon. Particularly, we
 297 consider a single mode-I crack propagation in soda-lime glass in gaseous nitrogen at 25° with variable
 298 relative humidity. The relevant chemistry along the crack can be straightforwardly characterized by the
 299 partial pressure of the water vapor p . In this setting, the environment-induced disjoining pressure can
 300 be expressed by $\Delta\Pi(w, p)$. Note that we did not consider $\Delta\Pi$ as a function of time in this first
 301 treatment. In other words, we have assumed that the adsorption kinetics is much faster than the rate of
 302 subcritical crack propagation ($< 10^{-4}$ m/s), and the only rate-limiting step in generating the repulsive
 303 disjoining pressure $\Delta\Pi$ is the transport of water vapor along the crack, as described by the transport
 304 models detailed in the next section.

305 Physically, many molecular-scale mechanisms can lead to the generation of $\Delta\Pi$ at various vapor
 306 pressures (Clarke et al., 1986). It is believed that these mechanisms mainly have origins in Van der
 307 Waals, electric double layer, and hydration forces, with the hydration force being the dominant
 308 contribution (Dziadkowiec, 2019). The exhaustive discussion of each of them is beyond the scope of this
 309 work. A smooth 4-parameter model is instead adopted for the environment-induced disjoining pressure:

310
$$\Delta\Pi(w, p) = f_{\Delta\Pi}(w)g_{\Delta\Pi}(p) \quad (10)$$

311 where

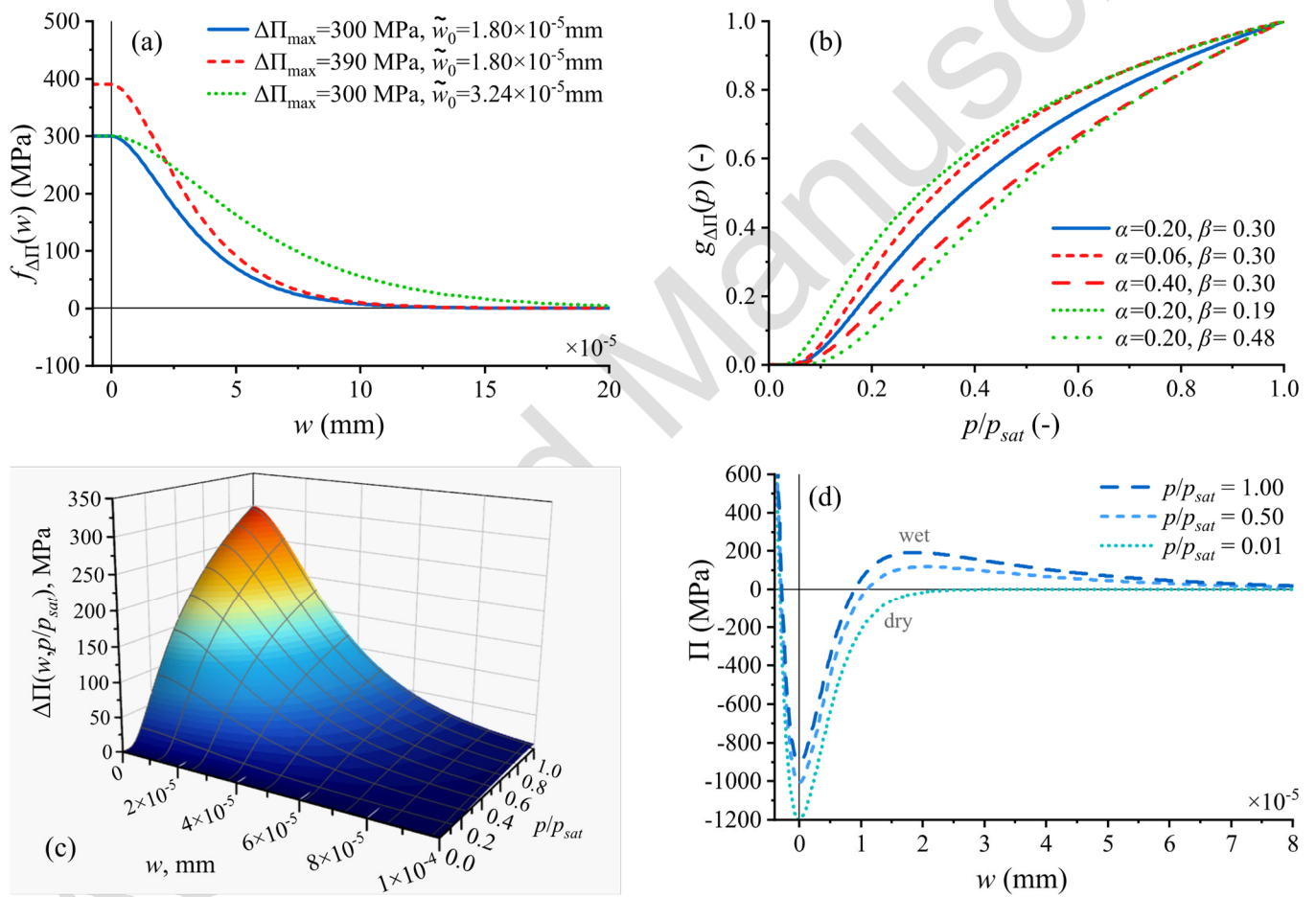
312
$$f_{\Delta\Pi}(w) = \begin{cases} \Delta\Pi_{\max} \frac{w + \tilde{w}_0}{\tilde{w}_0} \exp\left(1 - \frac{w + \tilde{w}_0}{\tilde{w}_0}\right) & ; w \geq 0 \\ f_{\Delta\Pi}(0) & ; w < 0 \end{cases} \quad (11)$$

313 captures the dependency of $\Delta\Pi$ on w by taking similar form as Eq. (9); and

314
$$g_{\Delta\Pi}(p) = \left(\frac{p}{p_{sat}}\right)^\alpha \exp\left(\beta\left(1 - \frac{p_{sat}}{p}\right)\right) \quad (12)$$

315 characterizes the dependency of $\Delta\Pi$ on the vapor pressure p . $\Delta\Pi_{\max}$, \tilde{w}_0 , α , and β are model
 316 parameters; p_{sat} is the saturation vapor pressure. The $f_{\Delta\Pi}$ expression exhibits an overall exponential
 317 decay, which is consistent with expressions adopted for hydration forces (Dziadkowiec et al., 2018;
 318 Israelachvili, 2011). The expression of g is constructed to 1) monotonically increase with increasing
 319 water vapor pressure p , 2) smoothly increase from zero, and 3) contain parameters that control the
 320 pressure and the steepness of the steepest increase. The value of $f_{\Delta\Pi}$ in the range of $w < 0$ is assigned to
 321 ensure the overall Π model is smooth at $w=0$. Eqns. (10), (11), (12) and the total Π from Eq. (8) are
 322 visualized in Fig. 7 for different levels of p / p_{sat} and values of model parameters. Fig. 7a shows that
 323 parameters $\Delta\Pi_{\max}$ and \tilde{w}_0 , respectively and independently, control the spread of $\Delta\Pi$ over separation

324 and its magnitude. Fig. 7b shows that the overall curvature of $g_{\Delta\Pi}$ decreases with increasing α and β .
 325 α also controls the curvature at very small p values, while the effect of β is less concerned with small
 326 p . The resultant $\Delta\Pi$ in Fig. 7c resembles the disjoining pressure sketched in Ash et al. (1973) and the
 327 one derived by Eskandari-Ghadi and Zhang (2021) based on the BDDT isotherm (Brunauer et al., 1940).
 328 The total Π in Fig. 7d can be compared with the conceptual sketch by Røyne et al. (2015) for calcite
 329 surfaces.



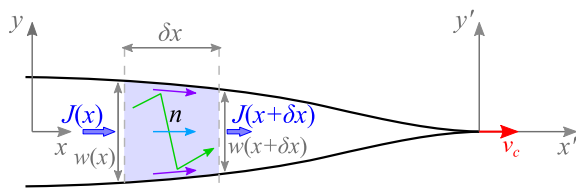
330
 331 Fig. 7. Environment-induced disjoining pressure as a function of (a) separation and (b) relative partial pressure;
 332 (c) the $\Delta\Pi$ surface in the $w-p$ space with $\Delta\Pi_{\max} = 300\text{MPa}$, $\tilde{w}_0 = 1.8 \times 10^{-5}\text{mm}$, $\alpha = 0.2$, and $\beta = 0.3$;
 333 and (d) the total disjoining pressure Eq. (8) with $\Pi_{0,\max} = 1200\text{MPa}$ and $w_0 = 3.18 \times 10^{-6}\text{mm}$.
 334
 335
 336
 337
 338

339 **5. MASS TRANSPORT ALONG THE CRACK**

340 The effect of sorption on crack growth is contingent upon the accessibility of sorptive species near
 341 the crack tip. This is controlled by the species transport along the crack path and is expected to trail
 342 behind as the crack propagates faster. The spatiotemporal evolution of species concentration in turn
 343 impacts the resultant disjoining pressure (Fig. 2) and further the K_{la} and the crack velocity, thus forming
 344 a two-way coupling. Resolving species transport is therefore essential to the proposed SFFT model for
 345 capturing the full SCG behavior. Below we develop a generic transport model to capture the most basic
 346 features of gas flow in cracks. A one-dimensional treatment is adopted, assuming that gas migrates
 347 towards the crack tip along the crack path (x -direction in Fig. 3a). The steady-state solution with respect
 348 to the crack tip of the transport model will be sought, considering crack propagation velocities in stages I
 349 and II are typically slow (of order 10^{-4} m/s or less). In other words, we examine the steady-state crack
 350 propagation scenarios at each K_{la} value during an SCG test without considering the short transient
 351 stages between K_{la} increments.

352 Consider a small segment δx along the crack with respect to the $x-y$ coordinate fixed in space
 353 (Fig. 8), the number of moles of sorptive species must be conserved:

354
$$\delta n(x,t)Lw(x)\delta x = J(x,t)Lw(x)\delta t - J(x+\delta x,t)Lw(x+\delta x)\delta t \quad (13)$$



355
 356 Fig. 8. Schematic of gas transport along propagating crack. The light blue, purple, and green arrows represent, respectively,
 357 bulk gas (viscous) flow, surface diffusion, and Knudsen flow as examples of the transport mechanisms.

358 where L is the depth of the crack in the z direction; n is the molar concentration with a unit of mole
 359 per volume; J is the molar flux with a unit of mole per unit area per unit time; and w is the crack

360 opening treated here as a known. By Taylor expansion of $J(x+\delta x)$ and $w(x+\delta x)$, and taking the limit
 361 of $\delta x \rightarrow 0$ and $\delta t \rightarrow 0$, Eq. (13) can be rewritten as

362

$$\frac{\partial n(x,t)}{\partial t} = -\frac{1}{w(x,t)} \frac{\partial}{\partial x} (w(x,t)J(x,t)) \quad (14)$$

363 For an ever-expanding domain (i.e., a moving crack tip), “steady state” only makes sense for an
 364 observer that moves together with the crack tip, i.e., the near-tip species concentration profile stops
 365 evolving for that observer. Let us therefore define a new coordinate system $x'-y'$ that moves according
 366 to $x' = x - v_c t$ and $y' = y$ (Fig. 8). It follows $n_t(x,t) = n_t(x',t) - v_c n_{x'}(x',t)$ and
 367 $(w(x)J(x,t))_x = (w(x)J(x,t))_{x'}$, where $(\cdot)_t = \partial(\cdot) / \partial t$, $(\cdot)_x = \partial(\cdot) / \partial x$, and $(\cdot)_{x'} = \partial(\cdot) / \partial x'$. Thus the molar
 368 balance in Eq. (14) can be written in the $x'-y'$ coordinate as

369

$$\frac{\partial n(x',t)}{\partial t} = v_c \frac{\partial n(x',t)}{\partial x'} - \frac{1}{w(x',t)} \frac{\partial}{\partial x'} (w(x',t)J(x',t)) \quad (15)$$

370 Examining the steady-state condition $\partial(\cdot) / \partial t = 0$ of Eq. (15) gives the following ordinary differential
 371 equation:

372

$$v_c \frac{\partial n}{\partial x'} = \frac{1}{w} \frac{\partial}{\partial x'} (wJ) \quad (16)$$

373 It relates the steady-state profiles $n(x')$, $w(x')$, and $J(x')$ at given boundary conditions. Without losing
 374 generality, the species transport may be depicted by

375

$$J(x) = -D \frac{\partial n}{\partial x} \quad (17)$$

376 where D is a transport coefficient. At chemical equilibrium, the partial pressure (p), concentration (n),
 377 and chemical potential (μ) of the sorptive species are one-to-one related through the equation of state.
 378 Thus, D can be re-written in terms of permeability for viscous and Knudsen flow or diffusivity for

379 molecular and surface diffusion. This preserves the generality of the model and enables more detailed
380 examinations of the various transport mechanisms in future studies. Substitution of Eq. (17) in Eq. (16)
381 gives

382

$$v_c \frac{\partial p}{\partial x'} = -\frac{1}{w} \frac{\partial}{\partial x'} (wD \frac{\partial p}{\partial x'}) \quad (18)$$

383 where the ideal gas law ($p = nRT$) and isothermal process ($dT = 0$) are assumed. The steady-state
384 pressure distribution $p(x')$ can be obtained by numerically solving Eq. (18) together with the $\Pi - w$
385 system introduced in the previous section. This is however numerically taxing considering solving the
386 $\Pi - w$ system already requires an iterative scheme. Alternatively, an explicit analytical expression of
387 $p(x')$ can be obtained by assuming $\partial w / \partial x' \approx 0$ and constant D in Eq. (18). This compromise
388 significantly eases the global solution scheme and permits quantitative studies of the first-order
389 behaviors of the proposed SFFT model. With these simplifying assumptions, Eq. (18) can be
390 immediately integrated under the boundary conditions of $p = p_0$ at the crack mouth ($x' = -a$) and
391 $J = v_c n$ at crack tip ($x' = 0$) to give:

392

$$p(x') = p_0 \exp\left(-\frac{v_c}{D}(x' + a)\right) \quad (19)$$

393 where p_0 is the partial gas pressure in the environment outside the crack.

394 In reality, it is expected that gas transport from crack mouth to tip involves multiple mechanisms
395 each of which control the transport rate at different crack width (Fig. 9a). For example, bulk gas flow
396 may be dominant at large crack openings (from M to A in Fig. 9b) because of the w^2 – dependency of
397 its diffusivity or permeability D (Wu et al., 2015). As the crack tip is approached and the crack opening
398 narrows, viscous flow rapidly diminishes, and a secondary transport mechanism gains dominance to
399 permit molecular diffusion in nanometer spaces (from B to T in Fig. 9b). This specific transport

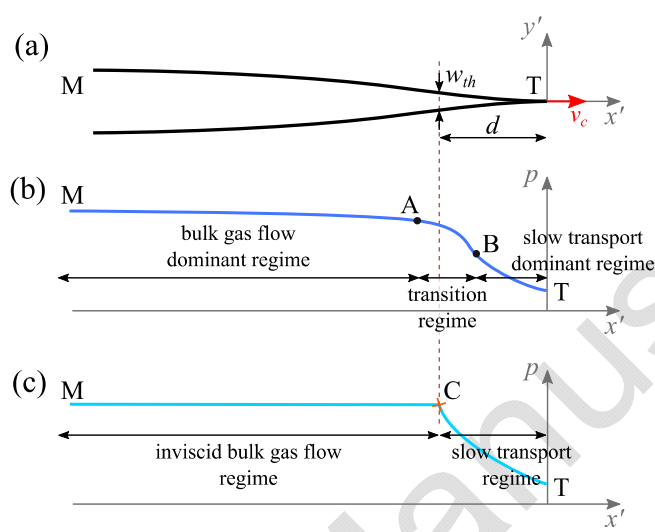
400 mechanism near crack tip is currently not well understood. Lawn (1974) argues that molecular collisions
 401 with the crack walls, known as Knudsen diffusion, is the mechanism of near-tip transport. Wiederhorn
 402 (1967) postulated that there exists a near-tip region, the so called “boundary layer”, that is always
 403 inaccessible to bulk gas flow. Water molecules can only migrate through the gas mixture in this region
 404 via Fickian diffusion at the same rate that they are chemically consumed at the crack tip. Contrarily,
 405 experimental data from glass SCG at different temperatures by Crichton et al. (1999) does not support
 406 the dominance of Fickian diffusion nor Knudsen diffusion in the boundary layer by comparing the
 407 anticipated versus the observed temperature-dependence of the stage-II crack velocity. They suggested
 408 surface diffusion is a plausible mechanism to explain the temperature-dependence, but at the same time
 409 also suggested that more complex mechanisms or their combinations could be responsible. An ongoing
 410 research effort at the Lawrence Berkeley National Laboratory is aimed at providing better understanding
 411 of these complex transport mechanisms in glass cracks at nano-scale.

412 Given the lack of a comprehensive transport model that captures the various mechanisms along the
 413 crack, we opt for a simplified, segmented, analytical steady-state pressure profile to account for the
 414 multi-regime transport of species in complementary to the single-regime model in Eq. (19). Particularly,
 415 species transport is assumed to take form of rapid bulk flow for crack width $w > w_{th}$ (from M to C in Fig.
 416 9c) where the pressure drop is negligible and slow diffusion for $w_{th} < w$ (from C to T in Fig. 9c) where
 417 the pressure profile is similarly described by Eq. (19):

$$418 \quad \begin{cases} p(x') = p_0 & ; \quad \forall x' \mid w(x') > w_{th} \\ p(x') = p_0 \exp\left(-\frac{v_c}{D}(x' + d)\right) & ; \quad \forall x' \mid w(x') \leq w_{th} \end{cases} \quad (20)$$

419 where w_{th} is a threshold crack width to be discussed later; d is defined such that $w(x') > w_{th}$ for all
 420 $x' < -d$. This dual-regime model (Fig. 9c) is conceptually similar to Wiederhorn’s boundary layer

421 model (Wiederhorn, 1967). In essence, the only difference between the single-regime flow Eq. (19) and
 422 the dual-regime flow Eq. (20) is that d is fixed at $d = a$ in the former, but becomes a variable related to
 423 w_{th} in the latter.

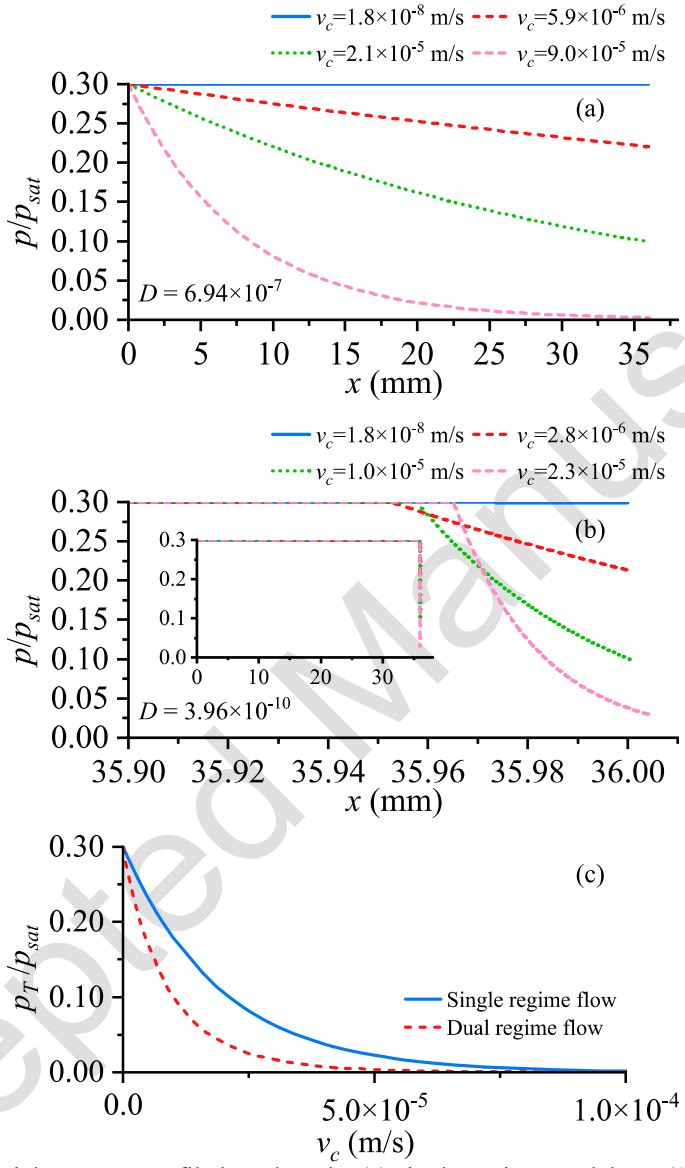


424
 425 Fig. 9. Schematic of the near tip crack profile (a), the realistic pressure profile in nano-spaces (b) inspired by Cihan et al.
 426 (2021), and the approximated dual-regime pressure profile (c).

427 The threshold crack width w_{th} may be selected based on the Knudsen number, $\text{Kn} = \lambda / w$, where λ
 428 is the mean free path of the gas (Lawn, 1974). $\text{Kn} \ll 1$ suggests transport dominated by viscous flow,
 429 while $\text{Kn} \gg 1$ suggests transport by slower diffusion mechanisms (Choi et al., 2001). Therefore, it is
 430 reasonable to define $w_{th} = \lambda / \text{Kn}_0$ where Kn_0 marks the transition point taken to be $1/3$ here, well
 431 within the range of $0.01 < \text{Kn}_0 < 1$ recommended by Lawn (1974). The experiments of glass SCG by
 432 Wiederhorn (1967) is performed in a nitrogen-water vapor environment at room temperature and
 433 atmospheric pressure, based on which the mean free path can be approximated as $\lambda \approx 7 \times 10^{-8} \text{ m}$ (Bird et
 434 al., 2006; Chapman and Cowling, 1990; Jennings, 1988).

435 Fig. 10a and b plot the steady-state vapor pressure profiles at different crack velocity for crack
 436 length $a = 36 \text{ mm}$ and ambient humidity $\text{RH}_0 = p_0 / p_{sat} = 0.3$. The dual-regime flow based on the
 437 current estimation of w_{th} predicts that the slow molecular diffusion only gets activated extremely close

438 (tens of μm) to the crack tip. Both models predict exponential decay of crack-tip vapor concentration
 439 with increasing velocity (Fig. 10c).



440
 441 Fig. 10. Steady-state vapor partial pressure profile based on the (a) single-regime model Eq. (19), and (b) dual-regime model
 442 Eq. (20) at different crack velocities. (c) Crack-tip pressure exponentially decreases as crack velocity increases. The model
 443 parameters used are those of Table 3.

444 6. SOLUTION SCHEME

445 Our SFFT model specialized for glass SCG is now complete with fracture mechanics Eqns. (1) and
 446 (2), crack kinetics Eq. (7), surface force model Eqns. (9) and (10), and transport model Eq. (19) or (20).

447 This section seeks the solution of the system subject to stress, $\sigma(t)$, over domain $x \in [0, a]$, and at each
448 point in time t_k . Over each time increment $(t_k, t_k + \Delta t_k)$, the crack length is updated based on

449
$$a(t_k + \Delta t_k) = a(t_k) + v_c(t_k) \Delta t_k \quad (21)$$

450 where Δt_k is initially set to 0.05 sec. It is reduced if the relative error at t_k is small and increased if the
451 relative error is large, within the range of 0.02 sec – 1 sec.

452 For spatial discretization, nodes are distributed along the crack at the mid-plane, permitted by the
453 one-dimensional nature of the problem. The crack is divided by N nodal points located at (x_1, x_2, \dots, x_N)
454 that are densely distributed (\sim every 6 nm) near the crack tip and coarsely distributed (\sim every 70 μm)
455 close to the crack mouth. This is because of the nature of surface forces: they take large magnitudes and
456 vary dramatically at short separations, then diminish quickly as the separation increases, making the
457 system extremely non-linear near the crack tip. With this discretization, the unknown vectors \mathbf{p} , \mathbf{w} , and
458 $\mathbf{\Pi}$ are defined as $p_i = p(x_i)$, $w_i = w(x_i)$, and $\Pi_i = \Pi(x_i)$, respectively.

459 For the modularity of the numerical scheme, the equations are grouped in three subsystems: **S1**) the
460 fluid transport subsystem, Eq. (19) or (20); **S2**) the fracture mechanics $\Pi - w$ subsystem, Eqns. (1), (2),
461 (9), and (10); and **S3**) the crack propagation subsystem, Eq. (7). They are then iteratively coupled at the
462 global level to search for a solution that simultaneously satisfies all the underlying equations.

463 For S1, the transport models in Eqns. (19) and (20) provide explicit solution at all points. In discrete
464 form, the nodal pressure described by these models are, respectively,

465
$$p_i = p_0 \exp\left(-\frac{v_c}{D} x_i\right) \quad (22)$$

466 and

467

468

$$p_i = \begin{cases} p_0 & ; \quad \forall x_i \mid w(x_i) > w_{th} \\ p_0 \exp\left(-\frac{v_c}{D}(x_i - a + d)\right) & ; \quad \forall x_i \mid w(x_i) \leq w_{th} \end{cases} \quad (23)$$

469 x_i is the coordinate of node i in the coordinate system set at the crack mouth, consistent with that used
 470 for the fracture mechanics solution (Fig. 3). For S2, an iterative method is required to solve the $\Pi - w$
 471 equations at fixed p profiles. Consistent Π and w profiles together with the resultant K_{le} are obtained
 472 by following the solution method of Planas and Elices (1991), while taking advantage of the analytical
 473 solutions made available by Tada et al. (2000) (Appendix A). For S3, Eq. (7) simply inputs an effective
 474 stress intensity and outputs the resultant crack velocity.

475 The global coupling of S1, S2, and S3 is achieved through a partitioned Newton-Raphson method.
 476 This method follows the multi-level Newton method of Kim et al. (2003). At local level, S1, S2, and S3
 477 are solved separately and independently. It is important to distinguish the global unknowns as the
 478 guessed inputs and the response of each subsystem to its pertaining guessed inputs as the local outputs.
 479 The global unknowns are defined here as the partial pressure of the adsorptive fluid, \mathbf{p} ; the crack
 480 opening, \mathbf{w} ; the effective stress intensity factor at the crack tip, K_{le} ; and the crack velocity, v_c . We
 481 shall then denote the solution to the fluid profile in response to guess values of crack opening and crack
 482 velocity by $\hat{\mathbf{p}}(\mathbf{w}, v_c)$, solved by S1; the crack opening and effective stress intensity in response to guess
 483 values of fluid state profile by $\hat{\mathbf{w}}(\mathbf{p})$ and $\hat{K}_{le}(\mathbf{p})$, solved by S2; and the crack propagation velocity in
 484 response to a guess value of stress intensity factor by $\hat{v}_c(K_{le})$, solved by S3. The coupling is achieved
 485 by directing the global residual to zero in an iterative manner.

486 The vector of global unknowns is constructed as

487

$$\mathbf{u} = \begin{bmatrix} \mathbf{p} \\ \mathbf{w} \\ K_{le} \\ v_c \end{bmatrix} \quad (24)$$

488

The global residual is defined as

489

$$\mathbf{R} = \begin{bmatrix} \mathbf{p} - \hat{\mathbf{p}}(\mathbf{w}, v_c) \\ \mathbf{w} - \hat{\mathbf{w}}(\mathbf{p}) \\ K_{le} - \hat{K}_{le}(\mathbf{p}) \\ v_c - \hat{v}_c(K_{le}) \end{bmatrix} \quad (25)$$

490 where \mathbf{p} , \mathbf{w} , K_{le} , and v_c are the values of the unknowns at the current iteration. The global solution
 491 that simultaneously satisfies S1, S2, and S3 can be found by iteratively marching towards $\mathbf{R} = \mathbf{0}$ via the
 492 Newton-Raphson method. In each iteration, the correction to \mathbf{u} , $\delta\mathbf{u}$, can be obtained from
 493 $\delta\mathbf{u} = -\mathbf{J}^{-1}(\mathbf{u}) \cdot \mathbf{R}(\mathbf{u})$ where \mathbf{J} is the global Jacobian matrix defined as:

494

$$\mathbf{J} = \frac{\partial \mathbf{R}}{\partial \mathbf{u}} = \begin{bmatrix} \frac{\partial(\mathbf{p} - \hat{\mathbf{p}}(\mathbf{w}, v_c))}{\partial \mathbf{p}} & \frac{\partial(\mathbf{p} - \hat{\mathbf{p}}(\mathbf{w}, v_c))}{\partial \mathbf{w}} & \frac{\partial(\mathbf{p} - \hat{\mathbf{p}}(\mathbf{w}, v_c))}{\partial K_{le}} & \frac{\partial(\mathbf{p} - \hat{\mathbf{p}}(\mathbf{w}, v_c))}{\partial v_c} \\ \frac{\partial(\mathbf{w} - \hat{\mathbf{w}}(\mathbf{p}))}{\partial \mathbf{p}} & \frac{\partial(\mathbf{w} - \hat{\mathbf{w}}(\mathbf{p}))}{\partial \mathbf{w}} & \frac{\partial(\mathbf{w} - \hat{\mathbf{w}}(\mathbf{p}))}{\partial K_{le}} & \frac{\partial(\mathbf{w} - \hat{\mathbf{w}}(\mathbf{p}))}{\partial v_c} \\ \frac{\partial(K_{le} - \hat{K}_{le}(\mathbf{p}))}{\partial \mathbf{p}} & \frac{\partial(K_{le} - \hat{K}_{le}(\mathbf{p}))}{\partial \mathbf{w}} & \frac{\partial(K_{le} - \hat{K}_{le}(\mathbf{p}))}{\partial K_{le}} & \frac{\partial(K_{le} - \hat{K}_{le}(\mathbf{p}))}{\partial v_c} \\ \frac{\partial(v_c - \hat{v}_c(K_{le}))}{\partial \mathbf{p}} & \frac{\partial(v_c - \hat{v}_c(K_{le}))}{\partial \mathbf{w}} & \frac{\partial(v_c - \hat{v}_c(K_{le}))}{\partial K_{le}} & \frac{\partial(v_c - \hat{v}_c(K_{le}))}{\partial v_c} \end{bmatrix} \quad (26)$$

495

Upon simplification, one has

496
$$\mathbf{J} = \begin{bmatrix} \mathbf{I} & -\frac{\partial \hat{\mathbf{p}}}{\partial \mathbf{w}} & \mathbf{0} & -\frac{\partial \hat{\mathbf{p}}}{\partial v_c} \\ -\frac{\partial \hat{\mathbf{w}}}{\partial \mathbf{p}} & \mathbf{I} & \mathbf{0} & \mathbf{0} \\ -\frac{\partial \hat{K}_{le}}{\partial \mathbf{p}} & \mathbf{0} & 1 & 0 \\ \mathbf{0} & \mathbf{0} & -\frac{\partial \hat{v}_c}{\partial K_{le}} & 1 \end{bmatrix} \quad (27)$$

497 A schematic of this partitioned Newton-Raphson scheme is presented in Fig. 11 for clarity.

498 The derivatives in Eq. (27) can be computed from Eq. (7), Eqns. (A.6) and (A.7) of Appendix A, and
 499 Eq. (22) or (23) depending on the choice of the transport model:

500
$$\frac{\partial \hat{v}_c}{\partial K_{le}} = \frac{v_0 \kappa_0}{RT} \exp\left(-\frac{Q_0}{RT}\right) \cosh\left(\frac{\kappa_0 K_{le}}{RT}\right) \quad (28)$$

501
$$\frac{\partial \hat{w}_i}{\partial p_j} = c_{\Pi,ik} \frac{\partial \hat{\Pi}_k}{\partial p_j} \approx c_{\Pi,ik} \frac{\partial \Pi_k}{\partial p_j} \quad (29)$$

502
$$\frac{\partial \hat{K}_{le}}{\partial p_j} = k_{\Pi,k} \frac{\partial \hat{\Pi}_k}{\partial p_j} \approx k_{\Pi,k} \frac{\partial \Pi_k}{\partial p_j} \quad (30)$$

503
$$\frac{\partial \hat{p}_i}{\partial w_j} = \frac{\partial \hat{p}_i}{\partial d} \frac{\partial d}{\partial w_j} \approx 0 \quad (31)$$

504
$$\frac{\partial \hat{p}_i}{\partial v_c} = -\frac{x_i}{D} p_0 \exp\left(-\frac{v_c}{D} x_i\right) \quad (32)$$

505 for single-regime flow, and

506

$$\frac{\partial \hat{p}_i}{\partial v_c} = \begin{cases} 0 & ; \quad \forall x \mid w(x) > w_{th} \\ -\frac{x_i - a + d}{D} p_0 \exp\left(-\frac{v_c}{D}(x_i - a + d)\right) & ; \quad \forall x \mid w(x) \leq w_{th} \end{cases} \quad (33)$$

507 for dual-regime flow.

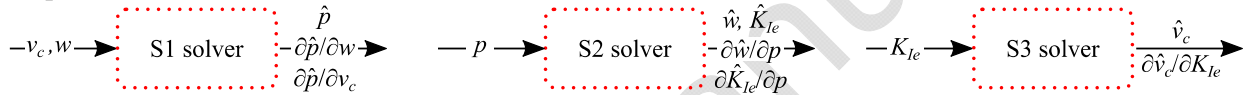
508 The errors are defined as

509

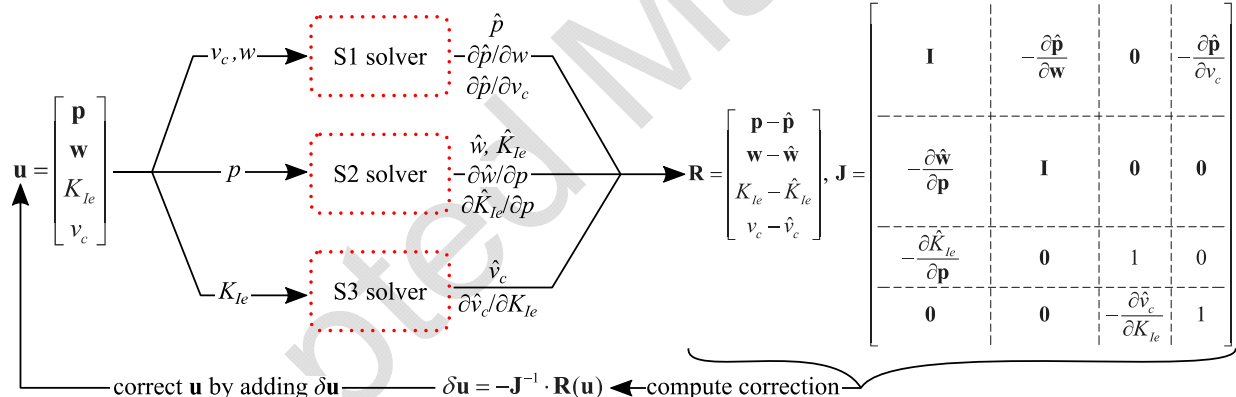
$$\varepsilon_p = \left| \frac{\delta p_i}{p_i} \right|, \quad \varepsilon_w = \left| \frac{\delta w_i}{w_i} \right|, \quad \varepsilon_{K_{le}} = \left| \frac{\delta K_{le}}{K_{le}} \right|, \text{ and } \varepsilon_{v_c} = \left| \frac{\delta v_c}{v_c} \right| \quad (34)$$

510 The iterations continue until the maximum of the four errors falls within a tolerance of 10^{-5} .

(a) Independent solvers:



(b) Global Residual Solver (partitioned Newton-Raphson iterations):



511
512 Fig. 11. A schematic of (a) the inputs and outputs of the S1, S2, and S3 solvers; and (b) the partitioned Newton-Raphson
513 scheme to seek solutions satisfying all subsystems.

514 Note that the length of the slow transport regime d varies continuously as the crack opens or closes,
515 while the numerical representation of the crack opening w is discrete and does not allow smooth
516 variation of d . Therefore, a linear interpolation for the value of $w(x)$ between two nodes is adopted to
517 allow smooth variation of d . Consequently, the derivative $\partial d / \partial w$ cannot be effectively computed
518 without explicit expression of d . This term is expected to be small anyway given the initial guess for w
519 is close to the solution. The derivative in Eq. (31) is thus approximated by zero. In addition, the $\partial \hat{\Pi} / \partial p$

520 term in Eq. (29) must be approximated since the $\Pi - w$ system is implicitly solved without being
521 written in terms of \mathbf{p} . Two versions of the implementation were tested, one with $\partial\hat{\Pi}/\partial\mathbf{p} \approx (\partial\hat{\Pi}/\partial\mathbf{p})_{\hat{w}}$
522 and $\partial\hat{w}/\partial\mathbf{p}$ given by Eq. (29), and the other with $\partial\hat{w}/\partial\mathbf{p} \approx \mathbf{0}$. The second approximation yielded better
523 convergence and was adopted in the final implementation. Finally, a proper initial guess is required to
524 start the search for the solution at each time t_{k+1} due to the highly nonlinear nature of the problem. A
525 robust initial guess algorithm is described in Appendix B.

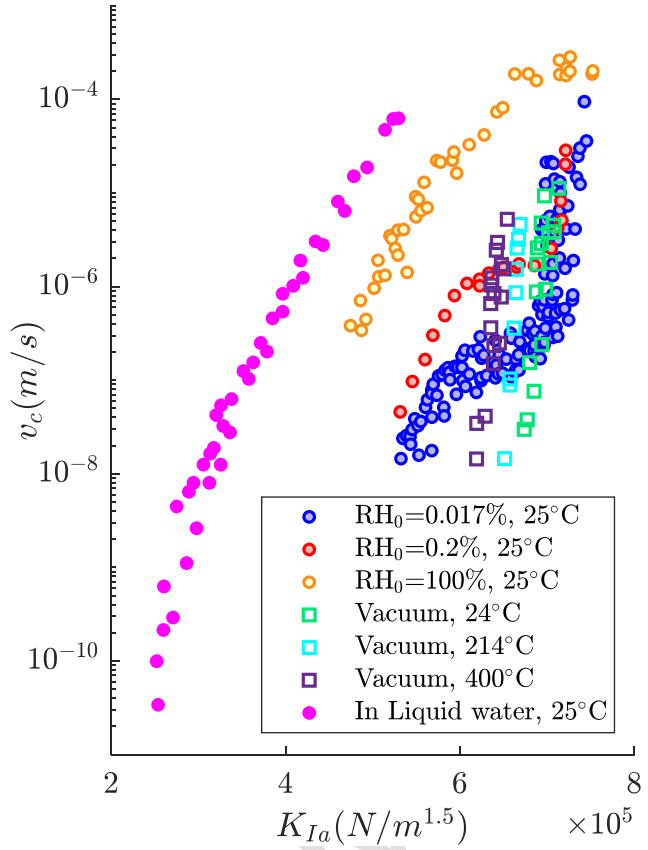
526 7. PERFORMANCE OF THE FRAMEWORK

527 This section compares the SFFT prediction against Wiederhorn's SCG data (Fig. 12) to validate the
528 underlying hypotheses depicted in Fig. 1.

529 7.1. Model calibration

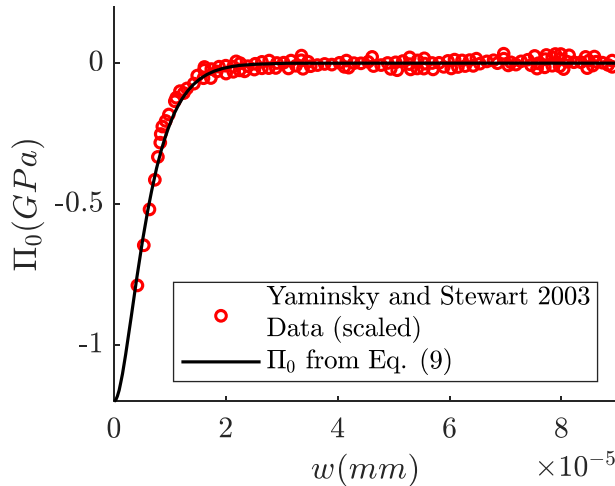
530 We shall calibrate the least coupled parts of the model first. The intrinsic material properties
531 including the elastic parameters, in-vacuum surface force, and the reaction kinetic parameters are
532 therefore calibrated first here. These parameters are expected to control the in-vacuum stage-III SCG
533 response.

534 The standard Young's modulus of soda-lime glass is 70 GPa (572-1, 2012) while the reported ones
535 in literatures vary between 50 GPa and 85 GPa (Meyland et al., 2021). A Poisson ratio in the range of
536 0.2 ~ 0.3 has been reported in some studies (Li and Wei, 2020; Meyland et al., 2021; Xu et al., 2017).
537 Within this ballpark, the selected elastic parameters are shown in Table 1.



538
 539 Fig. 12. Data for crack propagation in soda-lime glass at 25°C and different relative humidity (Wiederhorn, 1967; Wiederhorn
 540 et al., 1980), open circles; at different temperatures at 24°C, 214°C, and 400°C in vacuum (Wiederhorn, 1974), squares; and in
 541 liquid water (Wiederhorn and Bolz, 1970), filled circles.

542 Two types of data are adopted to calibrate the Π_0 parameters. The first data is surface energy for
 543 soda-lime glass measured by Wiederhorn (1969) at near room temperature in an environment with
 544 background nitrogen gas (inert) as $\gamma = 3.82 \text{ J/m}^2$. This measurement in inert gas can estimate the area
 545 underneath the $\Pi_0 - w$ curve, in absence of sorptive species (Gdoutos, 2020). For the selected Π_0
 546 expression Eq. (9), this area is $\gamma = \Pi_{0,\max} w_0$. The second data is the SFA measurements between a glass
 547 sphere in air (Yaminsky and Stewart, 2003), scaled by a factor of 912.6 such that the area underneath
 548 this data matches the surface energy measurement. Parameters $\Pi_{0,\max}$ and w_0 are then selected (Table 1)
 549 to match this data (Fig. 13).



550
 551 Fig. 13. Calibrated in-vacuum surface force ($\Pi_{0,\max} = 1200\text{MPa}$ and $w_0 = 3.1833 \times 10^{-6}\text{ mm}$) vs. SFA measurements. A
 552 constant multiplying factor is applied to the SFA data to roughly match the area under this curve with the surface energy
 553 measurement of $\gamma = 3.82\text{J/m}^2$. The Π_0 data is converted from F/R data multiplied by 912.6.

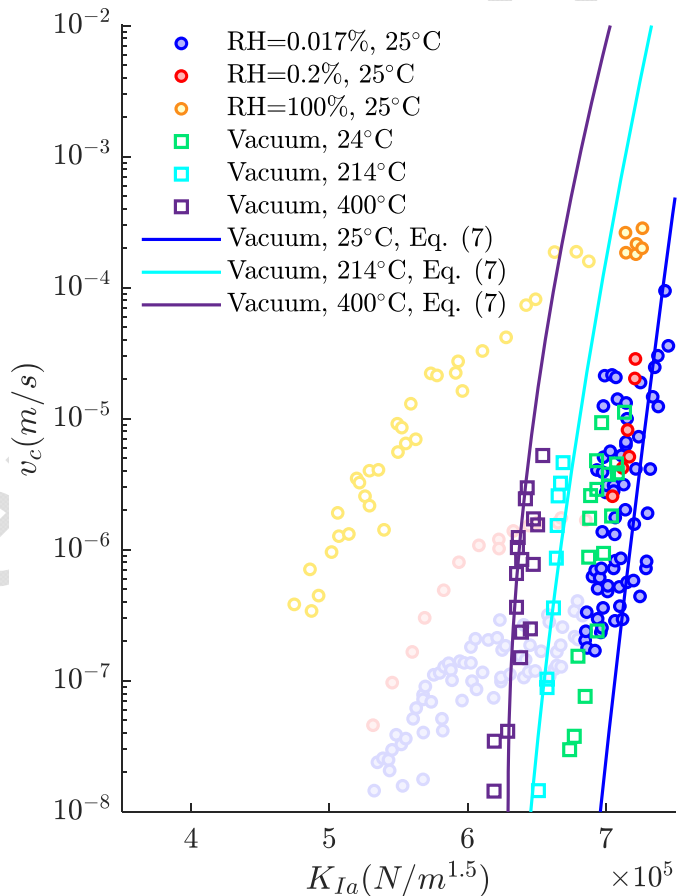
554 The reaction kinetic parameters are calibrated against the in-vacuum $v_c - K_{Ia}$ data. At relatively
 555 large K_{Ia} values, the $K_{Ie} - K_{Ia}$ relation becomes linear (see Fig. 6) and so does the $\log(v_c) - K_{Ie}$ relation
 556 (Eq. (7)). Combining the two gives a linear one-to-one correspondence between K_{Ia} and v_c at high K_{Ia}
 557 levels. Note that there are infinite combinations of v_0 , Q_0 , and κ_0 to match a linear set of $\log(v_c) - K_{Ia}$
 558 data. Such arbitrariness can be removed by considering the temperature dependence of the crack kinetics.
 559 Specifically, the same set of v_0 , Q_0 , and κ_0 should capture the in-vacuum $v_c - K_{Ia}$ curves obtained at
 560 different temperatures, assuming the Arrhenius term in Eq. (7) outweighs the temperature dependence of
 561 the elastic parameters and in-vacuum surface forces. Fig. 14 plots soda-lime glass $v_c - K_{Ia}$ data obtained
 562 at 24°C, 214°C, and 400°C (Wiederhorn, 1974) based on which the kinetic parameters v_0 , Q_0 , and κ_0 are
 563 calibrated (Table 1). Their values are the same orders of magnitude as those presented in Wiederhorn
 564 (1974).

565

566

Table 1. Model parameters that control stage III SCG behavior

Parameter	Unit	Values
Elasticity		
E	[MPa]	50000
ν	[-]	0.2
In-vacuum surface force		
w_0	[mm]	3.1833×10^{-6}
$\Pi_{0,\max}$	[MPa]	1200
Crack propagation reaction		
v_0	[mm/s]	3.303×10^5
Q_0	[J/mol]	1.31×10^5
κ_0	[m ^{2.5} /mol]	0.24

Fig. 14. Calibration of reaction rate parameters v_0 , Q_0 , and κ_0 against SCG data at three temperatures, 25°C, 214°C, and 400°C in vacuum. The irrelevant data are faded for clarity.

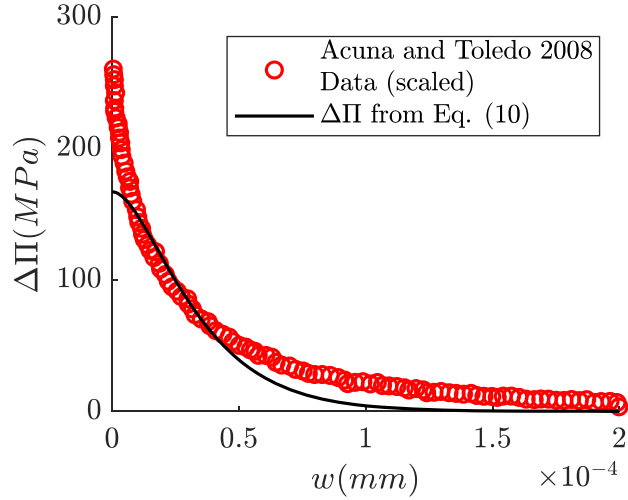
574 The model is now fully calibrated for the in-vacuum condition. We now move on to determine
575 parameters that control the environmental dependency, specifically the environment-induced disjoining
576 pressure $\Delta\Pi$ and the pressure profile $p(x)$. The former is expected to affect the strength reduction in
577 stage I, and the latter limits the rate of crack propagation in stage-II SCG.

578 In Eqns. (10), (11), and (12), parameters $\Delta\Pi_{\max}$ and \tilde{w}_0 control the dependence of $\Delta\Pi$ on the crack
579 opening and parameters α and β independently control the dependence of $\Delta\Pi$ on the partial pressure
580 of the adsorbate. AFM measurements in glass-liquid water system (Acuña and Toledo, 2008) provides a
581 benchmark for the shape of the $\Delta\Pi - w$ relation at $p = p_{sat}$ which allows the determination of \tilde{w}_0 (Fig.
582 15). The area underneath $\Delta\Pi$ reflects the change of surface energy in presence of water vapor, meaning
583 that $\Delta\Pi_{\max}$ can be calibrated based on the reduction of apparent fracture toughness from $\text{RH}_0 = 0\%$ to
584 100% (Fig. 16). Finally, parameters α and β in Eq. (12) are tuned to capture the strength reductions at
585 $\text{RH}_0 = 0.017\%, 0.2\%, 1\%, 10\%,$ and 30% (Fig. 16). These parameters are summarized in Table 2.

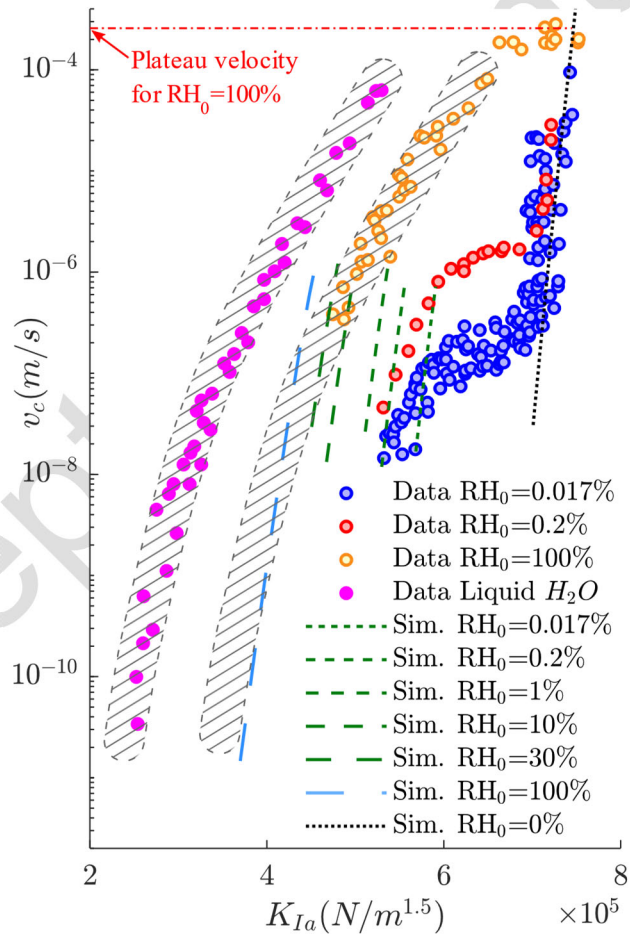
586 Table 2. Parameters that control stage I SCG in complement to Table 1.

Parameter	Unit	Values
Environment-induced disjoining pressure		
\tilde{w}_0	[mm]	1.8×10^{-5}
$\Delta\Pi_{0,\max}$	[MPa]	167
α	[\cdot]	0.067
β	[\cdot]	10^{-5}

587



588
589 Fig. 15. Calibrated $\Delta\Pi$ with parameters $\tilde{w}_0 = 1.8 \times 10^{-5}$ mm and $\Delta\Pi_{\max} = 167$ MPa at $p = p_{\text{sat}}$ based on the F/R
590 measurements from AFM (Acuña and Toledo, 2008) scaled by 120.5. The main purpose here is to match the shape via
591 adjusting \tilde{w}_0 . $\Delta\Pi_{\max}$ is determined in Fig. 16.



592
593 Fig. 16. Measured vs. simulated strength reduction at $\text{RH}_0 = 100\%$ with parameters $\Delta\Pi_{\max} = 167$ MPa and
594 $\tilde{w}_0 = 1.8 \times 10^{-5}$ mm (light blue coarse-dashed line). The strength reductions for intermediate RH values are captured by
595 selecting $\alpha = 0.067$ and $\beta = 10^{-5}$ (green dashed lines). The cross-hatching is meant as a visual guide for the extension of the

596 $\text{RH}_0 = 100\%$ data based on the data in liquid water. The reason for this extension is to obtain a visual of the $\text{RH}_0 = 100\%$
 597 curve at lower velocities for the purpose of calibration.

598 On the transport side, for both $p(x)$ profiles depicted by Eq. (19) and Eq. (20), D is selected such
 599 that the adsorbate relative pressure at the crack tip drops to an arbitrary small value (here taken as
 600 2×10^{-6}) at the stage-II plateau velocity for the $\text{RH}_0 = 100\%$ case ($v_c = 2.5 \times 10^{-4} \text{ m/s}$ at
 601 $K_{la} = 7.44 \times 10^5 \text{ N/m}^{1.5}$ from Fig. 16). The near-tip region length d for the dual-regime flow model is
 602 estimated from the in-vacuum crack opening profile at $K_{la} = 7.44 \times 10^5 \text{ N/m}^{1.5}$. This gives
 603 $d \approx 2.13 \times 10^{-5} \text{ m}$; i.e., $w < w_{th} = \lambda / \text{Kn}_0$ for $a - 2.13 \times 10^{-5} < x$. The values of D for the two cases are
 604 given in Table 3.

605 Table 3. Parameters that control stage II SCG in complement to Table 1 and Table 2.

Parameter	Unit	Values
Single-regime flow		
D	$[\text{m}^2/\text{s}]$	6.94×10^{-7}
Dual-regime flow		
D	$[\text{m}^2/\text{s}]$	3.96×10^{-10}

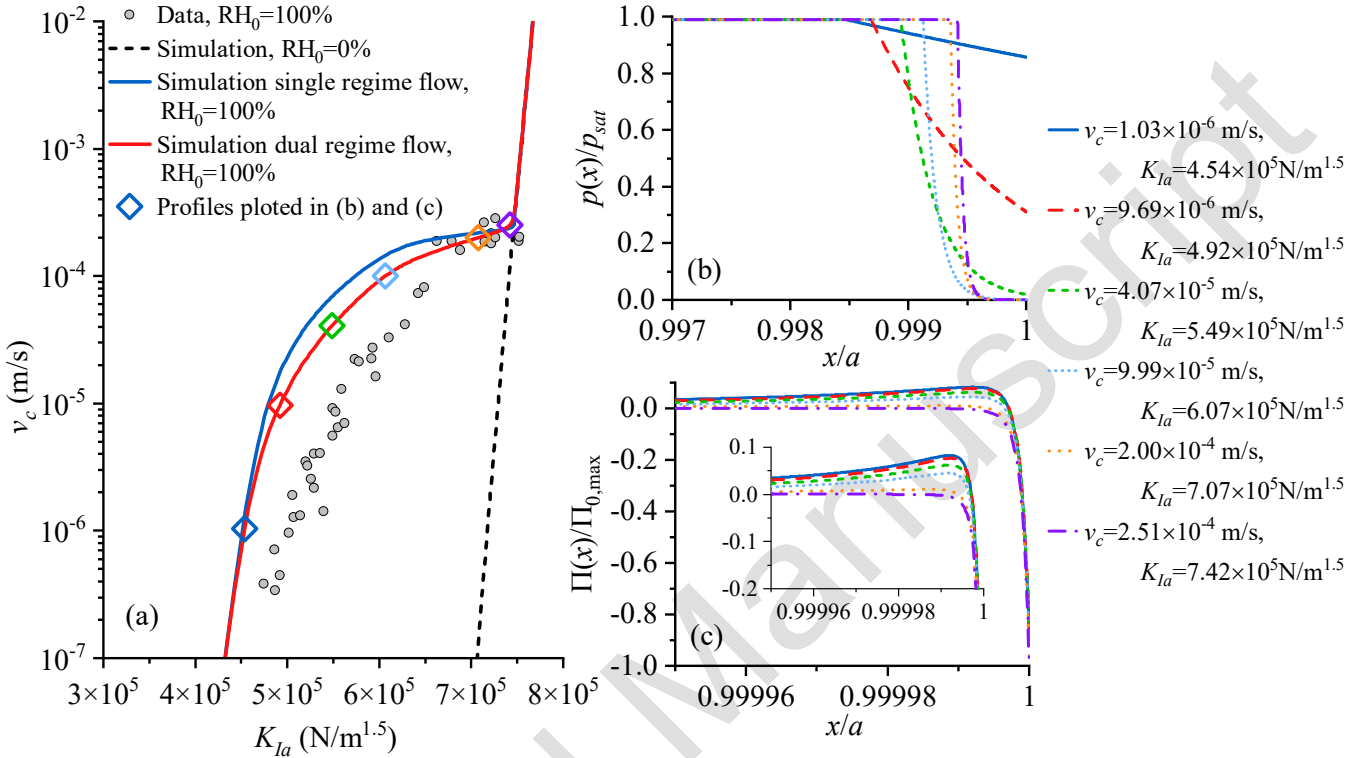
606 These calibrated D values can be quantitatively compared with some theoretical estimates. In a study
 607 of gas transport in shale nanopores, Wu et al. (2015) predicted an equivalent Darcian permeability k (as
 608 in $v_{fluid} = -(k / \mu) \partial p / \partial x$) of order 10^{-17} m^2 for the case of surface diffusion. Conversion of k to D can
 609 be made by rewriting $J(x) = -D \partial n / \partial x$ as $v_{fluid} = J / n = -(D / p) \partial p / \partial x$, giving $D = kp / \mu$. A
 610 viscosity of order $10^{-5} \text{ Pa} \cdot \text{s}$ is assumed (Tsilingiris, 2008; Wu et al., 2015), and a pressure of order
 611 $< 10^3 \text{ Pa}$ can be assumed considering $O(p_{sat}) = 10^3 \text{ Pa}$ for water vapor. This gives .
 612 $O(D_{\text{surf. diff.}}) < 10^{-9} \text{ m}^2/\text{s}$. for surface diffusion, comparable with the calibrated value of D for dual-
 613 regime flow. Similar comparison can be made in the case of Knudsen diffusion (Wu et al., 2015). On the
 614 other hand, considering viscous flow in a slit with aperture w , one can derive $D_{\text{bulk flow}} = RTw^2 / 12\mu V_m$

615 for ideal gas. At room temperature, the molar volume of ideal gas is $V_m = 2.45 \times 10^{-2} \text{ m}^3/\text{mol}$. Fig. 5
616 suggests that the crack opening varies from order 10^{-5} m to 10^{-8} m from the crack mouth to where
617 surface forces are observed. Combining the above gives $O(D_{\text{bulk flow}}) = 10^{-2} \sim 10^{-8} \text{ m}^2/\text{s}$. Note that the
618 estimated $D_{\text{bulk flow}}$ is several orders larger than our calibrated D values (Table 3) and those estimated
619 based on Wu et al. (2015). This justifies our assumption that the pressure drop is negligible in the bulk
620 flow regime up to the transition point (Fig. 9c).

621 **7.2. Model prediction**

622 Using the calibrated parameters in Tables 1, 2, and 3, the SCG curves predicted by the SFFT model
623 for single-regime and dual-regime flow at $\text{RH}_0 = 100\%$ are presented together with experimental data in
624 Fig. 17a. It is exciting to see that, for the first time, the key features of all three stages of SCG and the
625 smooth transition between them are directly predicted by a single mechanistic model. Comparing the
626 two computed SCG curves suggests that the choice of transport model directly impacts the stages I and
627 II SCG behavior. This cannot be observed in existing phenomenological SCG models, since the shape of
628 the whole SCG curve is imposed rather than predicted in these approaches (Charles, 1962; Lawn, 1975).
629 Fig. 17b presents the predicted vapor pressure profiles at different stages of SCG for the dual-regime
630 model. As K_{la} increases, the crack opening widens, and the inviscid regime moves closer to the crack
631 tip. However, the simultaneous increase in crack velocity outpaces the crack widening, which results in
632 an overall decrease of fluid concentration at the crack tip. Fig. 17c shows that the disjoining pressure is
633 extremely concentrated near the crack tip and is highly sensitive to the local vapor pressure and thus the
634 crack velocity. This suggests that the introduction of slight repulsive disjoining pressure in the near-tip
635 region can cause significant reduction of the apparent fracture toughness for brittle solids. The profiles

636 plotted in Fig. 17b and c confirm the common speculation that fluid transport is indeed the rate-limiting
 637 mechanism for the stage II of SCG.



638
 639 Fig. 17. (a) Measured vs. predicted SCG curves at $RH_0 = 100\%$ based on single-regime flow and dual-regime flow.
 640 (b) Pressure profiles at different stages along the SCG curve depicted by the dual-regime flow model.
 641 (c) Disjoining pressure distribution at different stages along the SCG curve.

642 Recalling that the SFFT model was calibrated only for the apparent toughness (where the SCG curve
 643 intersects with the x -axis) and the plateau velocity (where stages II and III intersect) at $RH_0 = 100\%$,
 644 everything in between should be regarded as true predictions. It is observed that the current model
 645 generally overestimates the crack velocity in stages I and II, with the dual regime flow model making
 646 slightly better prediction. This is because the experimental data suggests the stage-I slope is lower
 647 compared to the in-vacuum portion, while the current model predicts the same slopes for stages I and III.
 648 We may remediate this through the reaction rate parameters or the fluid transport parameters. Fig. 18a
 649 demonstrates that the decrease of κ_0 or Q_0 can reduce the slope of SCG curve in the $\log(v_c) - K_{Ia}$ plane.
 650 This implies that the lowered stage-I slope can be accounted for by introducing some environmental

dependency on the kinetic parameters. A rigorous proposal of $\kappa_0(p)$ or $Q_0(p)$ requires detailed study of how the energy barrier for bond rupture at the very crack tip lowers under environmental attack, which is beyond the scope of this paper. Alternatively, an apparent slope reduction can be readily observed by cropping out the very low velocity range of the SCG plots (Fig. 18b).

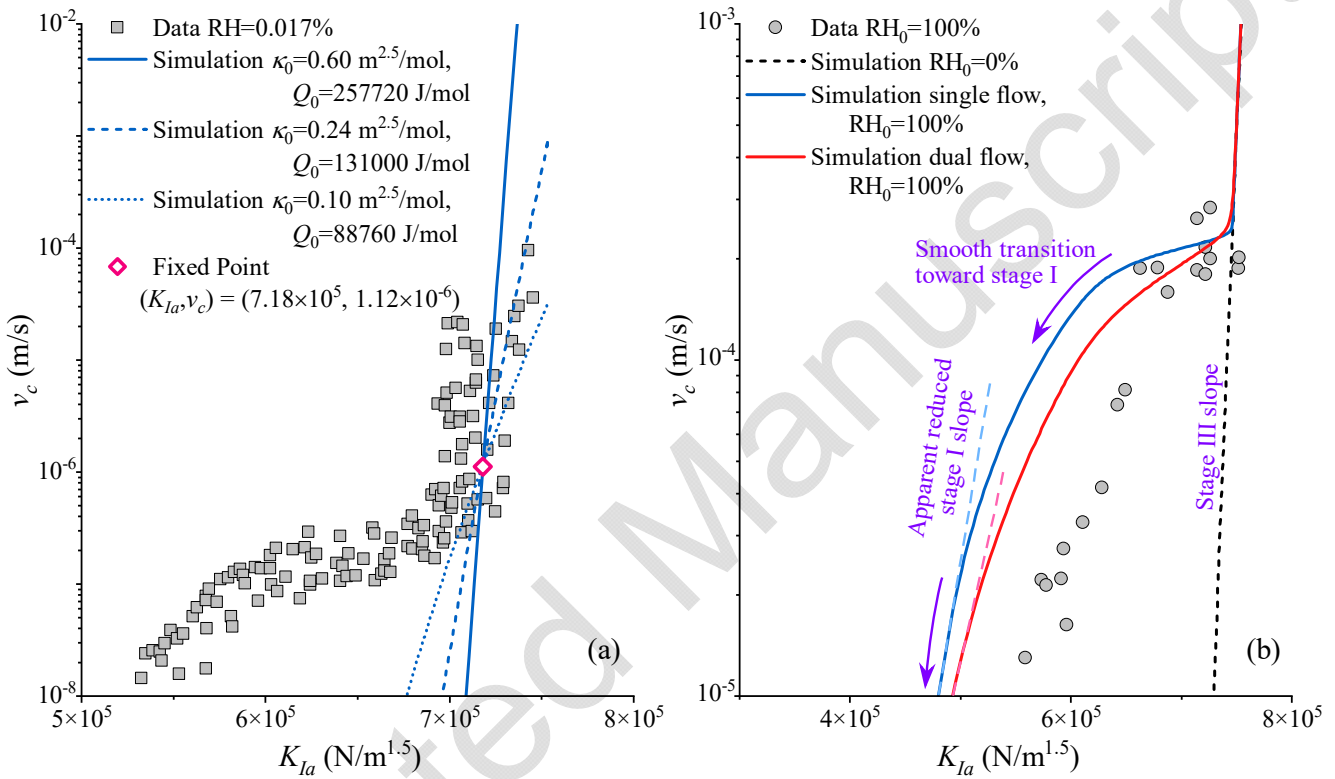
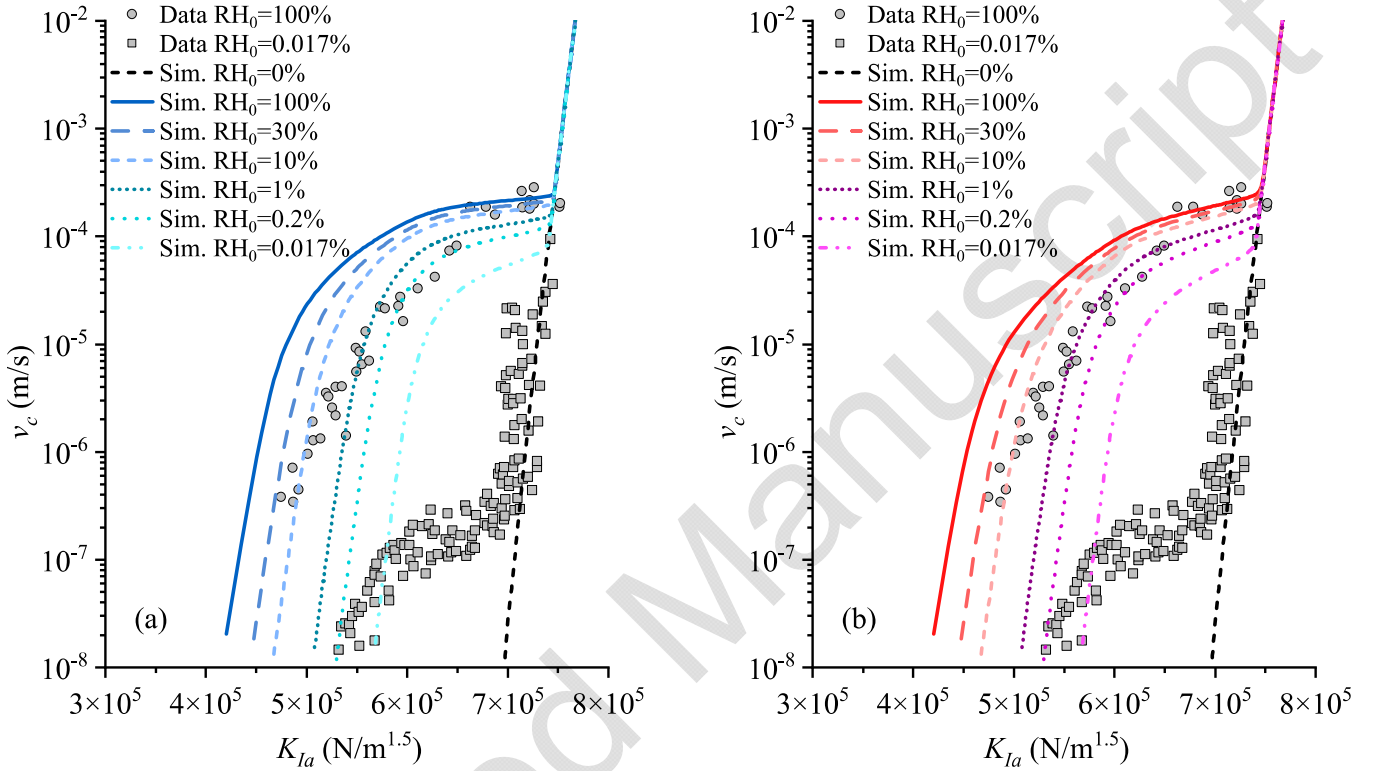


Fig. 18. (a) Variation of SCG slope about a fixed point ($K_{la} = 7.18 \times 10^5 \text{ N/m}^{1.5}$ and $v_c = 1.12 \times 10^{-6} \text{ m/s}$) controlled by parameters κ_0 and Q_0 : $\kappa_0 = 0.60 \text{ m}^{2.5}/\text{mol}$ and $Q_0 = 257720 \text{ J/mol}$ for Simulation A; $\kappa_0 = 0.24 \text{ m}^{2.5}/\text{mol}$ and $Q_0 = 131000 \text{ J/mol}$ for Simulation B; and $\kappa_0 = 0.10 \text{ m}^{2.5}/\text{mol}$ and $Q_0 = 88760 \text{ J/mol}$ for Simulation C. (b) Zoomed-in presentation of Fig. 17 giving an impression of reduced stage-I slope.

This implies that a rate-limiting mechanism that is activated over a wide range of crack velocities can cause the reduced slope in stage I. Perhaps a mechanism with weaker dependence on v_c compared to the current transport models (e.g., a different near-tip diffusion mechanism, or sorption kinetics) can potentially close this gap. In partial support of this hypothesis, the experimental SCG data for crack propagation in liquid water in Fig. 16 which extends to much lower crack propagation velocities shows a slope close to the in-vacuum one. This speculation challenges the commonly accepted picture that SCG

666 in stage I is free from any rate-limiting influences. More investigations are required in this regard to
 667 clarify the origin of environment-dependent slope in stage I SCG.

668 Fig. 19 presents the predicted SCG curves over a range of relative humidity from 0% to 100%.



669
 670 Fig. 19. Measured vs. predicted SCG curves at different RH_0 values based on (a) single regime flow
 671 models.

672 It is observed that both flow models significantly underestimate the variation of plateau velocities at
 673 different humidity levels. Based on our previous observation that the shape of SCG critically depends on
 674 the transport model, we suspect that this underprediction is related to the insensitivity of the vapor
 675 pressure profile $p(x)$ to the RH_0 . Physically, the threshold crack width w_{th} characterizing the point of
 676 transition from bulk flow to molecular diffusion (point C in Fig. 9c) should not be a constant but rather
 677 vary with respect to the boundary conditions (i.e., the ambient humidity). This possibility is explored in
 678 the next subsection.

679

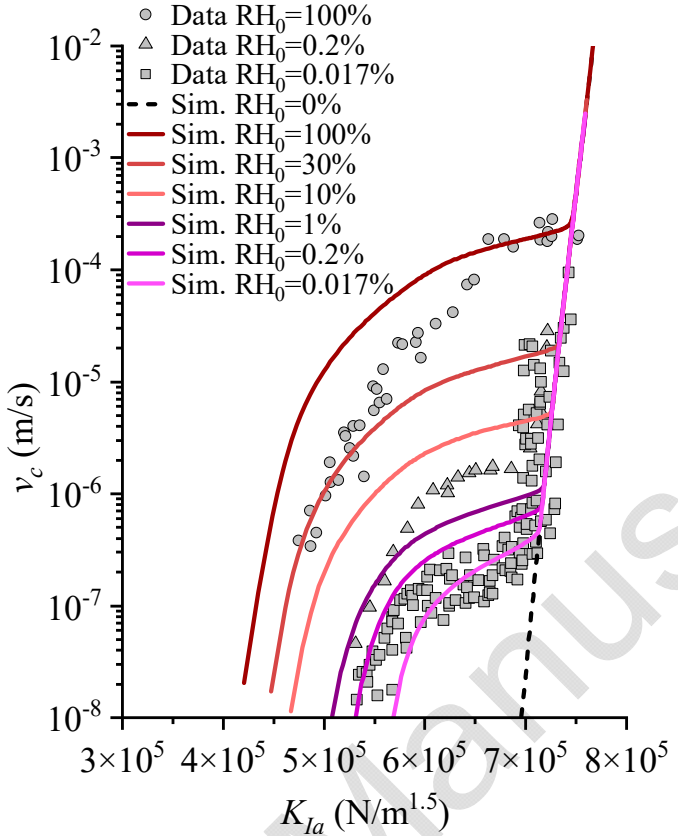
680 7.3. Effect of variable w_{th}

681 Intuitively, lower RH means less water molecules in the environment, and they can be more easily
 682 captured by the crack walls to form the first few adsorbed layers. This promotes the molecular diffusion
 683 of water in adsorbed states rather than free bulk gas flow, thus extending the length of the “boundary
 684 layer” d in Fig. 9a. w_{th} is therefore proposed to be inverse proportional to RH_0 . A simple proposition
 685 to account for this effect is:

$$686 \quad w_{th} = \frac{\lambda / Kn_0}{1 + h_0(RH_0 - 1)} \quad (35)$$

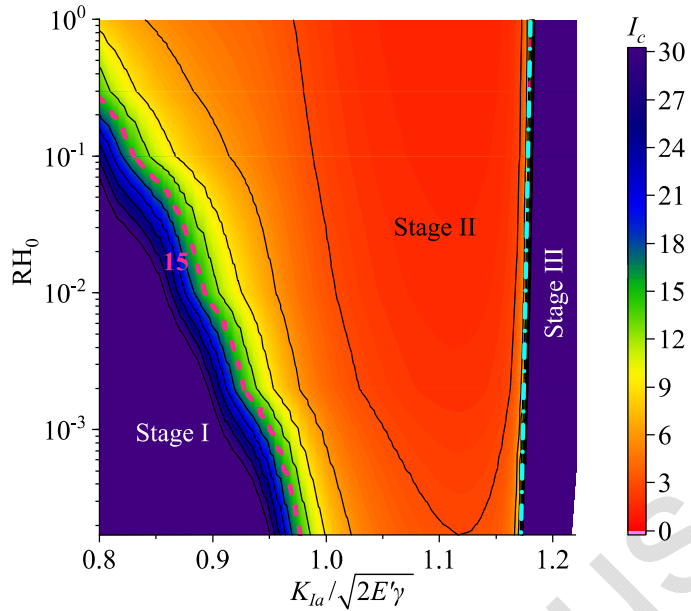
687 where $h_0 < 1$ is a parameter to control how rapid w_{th} varies with RH_0 such that $w_{th}(RH_0 = 1) = \lambda / Kn_0$.
 688 Here, a value of $h_0 = 0.919$ is found to match the plateau velocity for case of $RH_0 = 0.017\%$. The
 689 predicted SCG curves using the updated w_{th} , Eq. (35), for all other RH_0 values are presented in Fig. 20.
 690 We can see that the SFFT model is now capable of quantitatively capturing the shifting of the plateau
 691 velocity and the apparent fracture toughness under humidity changes. Again, this observation highlights
 692 the importance of detailing the transport of sorptive species in the near-tip regime in order to correctly
 693 predict the SCG behavior of brittle solids under various ambient conditions.

694 A subcritical cracking “stage diagram” can be produced (Fig. 21) in the normalized
 695 $\log(v_c / v_0) - (K_{Ia} / \sqrt{2E'\gamma})$ space based on the slope of the predicted SCG curves in Fig. 20. The slope
 696 in the normalized space is defined as $I_c = d \log(v_c / v_0) / d(K_{Ia} / \sqrt{2E'\gamma})$, which is analogous to the
 697 *corrosion index* in the Charles’ law (Atkinson, 1982).



698
699 Fig. 20. Measured vs. predicted SCG curves at different RH_0 values based on dual-regime flow and RH_0 -dependent
700 w_h (Eq. (35) with $h_0 = 0.919$).

701 It is apparent in Fig. 21 that stages I and III are represented by regions with larger I_c , while a drop of I_c
702 to lower values indicates stage II. Transition from stage II to III occurs abruptly (cyan dash-dot line),
703 while the transition from stage I to II happens gradually with a much smoother and wider variation of I_c
704 values in between. A value of $I_c = 15$ is selected to roughly represent this transition (pink dashed line).
705 Fig. 21 predicts that increased ambient relative humidity expands the stage-II zone, indicating that
706 species transport becomes more dominating in shaping the SCG curve over a wider K_{Ia} range at high
707 RH levels. If an SCG curve is plotted in a limited range of K_{Ia} and thus v_c , one would have concluded
708 that the apparent corrosion index (i.e., the slope) is reduced as a function of the ambient humidity. This
709 reiterates one of our earlier speculations that the reduced slopes in stage I SCG could be just a visual
710 impression of an extended stage II zone due to the slow transport of environmental agents.



711
712
713

Fig. 21. A dimensionless stage diagram for SCG. Color indicates the slopes of normalized v_c / v_0 versus $K_{Ia} / \sqrt{2E'\gamma}$ curves from Fig. 20, based on which the three SCG stages can be clearly identified.

714 **8. CONCLUSION**

715 This study develops a surface-force based fracture theory (SFFT) to rationalize and predict the
 716 subcritical cracking of brittle solids in sorptive environments. The theory highlights the fundamental role
 717 of surface forces which present at a few tens of nanometer surface-surface separations on the apparent
 718 fracture toughness, its environmental dependency, and the crack growth kinetics of SCG. Specifically,
 719 analytical LEFM solutions are adopted to consider the repulsive disjoining force induced by sorption
 720 behind the crack tip. The competition between the material's intrinsic cohesive forces and environment-
 721 induced disjoining forces is manifested as the *effective stress intensity factor* operating at the crack tip
 722 and serves as the driving force for crack propagation. This quantitatively explains the environmentally
 723 induced strength reduction of brittle solids. The characteristic shape of the SCG curve is further
 724 impacted by the rate-limiting effect of species transport in the crack. Simplified analytical transport
 725 models are proposed to qualitatively represent the single-regime and dual-regime flows. Finally, the
 726 model is validated against the experimental data on soda-lime glass in water vapor studied by
 727 Wiederhorn (1967). The main findings of this study are highlighted below:

728 1. The development of repulsive disjoining pressure behind the crack tip in sorptive environment
729 can cause the reduction of apparent fracture toughness.

730 2. Species transport along the crack is the rate-controlling mechanism governing the plateau
731 velocity and the stage-II behavior of SCG.

732 3. The effect of transport can go beyond stage II and shape the characteristics of stage-I SCG curve,
733 for example, by reducing the apparent slope in stage I.

734 4. The SCG stage diagram generated by SFFT simulations allows one to visualize the conditions
735 for each stage of SCG and the characteristics of transition between stages.

736 The theory and its numerical solution scheme are set up in such a way that more complex and
737 advanced model components can be readily incorporated for extensions to other brittle materials. Further
738 improvements can be contemplated by (1) incorporating more realistic fluid transport models that
739 automatically capture the smooth transition between different transport mechanisms as the crack
740 aperture varies; (2) considering the underlying sources of surface forces to motivate a physics-based
741 disjoining pressure model; and (3) extending to different systems such as the SCG of quartz or calcite in
742 aqueous solutions for geoscience and geo-engineering applications. Some fundamental aspects require
743 further investigations to enrich the calibration and validation of SFFT. For example, experimental
744 studies and molecular dynamic simulations are needed to quantify disjoining pressure between freshly
745 exposed solid surfaces in sorptive environment and to better understand species transport in nano-
746 confined spaces.

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910

911 APPENDIX A. NUMERICAL SOLUTION TO THE Π - w SYSTEM

912 The solution to the Π - w system involves finding a consistent set of $\Pi(x)$ and $w(x)$ that satisfy
 913 two equations of the general forms $w(x) = w(x, \Pi(x))$ and $\Pi(x) = \Pi(x, w(x))$. Eq. (2) is an example of
 914 the former, and Eqns. (9) and (10) take the form of the latter.

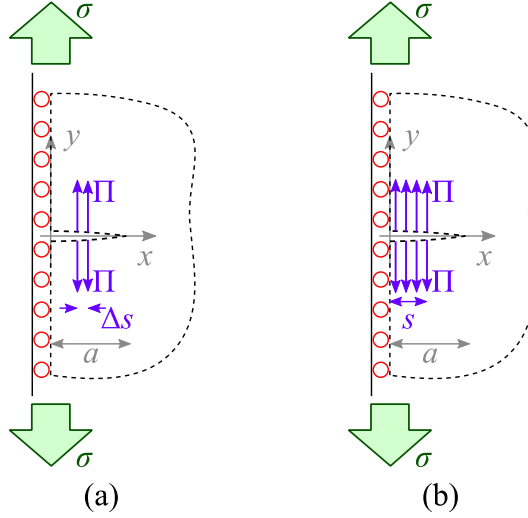
915 To write Eq. (2) in discrete form, an approximation for the integral on the right-hand side is required.
 916 Such approximation is commonly performed by adopting Gauss quadrature methods. However, due to
 917 the singularity of the fundamental solution, Eq. (4), at $x = s$, Gauss quadrature approximation
 918 encountered numerical convergence issues. Here an alternative approximation is adopted. Refer to the
 919 spatial discretization in Section 6. First, the disjoining pressure over the range of x_i and x_{i+1} , $\Pi_i^{elem}(x)$,
 920 is approximated with a constant value, $\Pi_i^{elem} = (\Pi(x_i) + \Pi(x_{i+1})) / 2 = (\Pi_i + \Pi_{i+1}) / 2$. With this, Eqns. (1)
 921 and (2) can be written at each node as

$$922 w(x_i) = c_\sigma(x_i)\sigma + \sum_{j=1}^{N-1} \Pi_j^{elem} \underbrace{\int_{s=s_j}^{s=s_{j+1}} c_\Pi(x_i, s) ds}_{c_\Pi^{elem}} \text{ and } K_{le} = k_{I\sigma}\sigma + \sum_{j=1}^{N-1} \Pi_j^{elem} \underbrace{\int_{s=s_j}^{s=s_{j+1}} k_{I\Pi}(s) ds}_{k_{I\Pi}^{elem}} \quad (A.1)$$

923 The fundamental functions $c_\Pi(x, s)$ and $k_{I\Pi}(s)$ are now the crack opening profile and the stress
 924 intensity factor for uniformly distributed stress over range (s_j, s_{j+1}) in Fig. A.1a, which are non-singular.
 925 For unit repulsive stress ($\Pi = 1$) applied over length s in Fig. A.1b, Tada (1972) provides

$$926 \int_0^s c_\Pi(x, s) ds = \frac{8(1-\nu^2)}{\pi E} \left\{ \begin{array}{l} \sqrt{a^2 - x^2} \sin^{-1} \frac{s}{a} + \frac{1}{2} |x - s| \cosh^{-1} \frac{a^2 - sx}{a|x - s|} \\ + \frac{1}{2} |x + s| \cosh^{-1} \frac{a^2 + sx}{a|x + s|} \end{array} \right\} \quad (A.2)$$

$$927 \int_0^s k_{I\Pi}(s) ds = 2\sqrt{\frac{a}{\pi}} \sin^{-1} \frac{s}{a} \quad (A.3)$$



928
929 Fig. A.1 Geometry considered in (a) this paper in deriving the fundamental solutions and (b) Tada (1972).

930 The fundamental solutions for uniform stress over an arbitrary segment (s_j, s_{j+1}) in the integrals of

931 Eq. (A.1) can be directly expressed in terms of Eqns. (A.2) and (A.3), as

932
$$\int_{s=s_j}^{s=s_{j+1}} c_{\Pi}(x_i, s) ds = \int_0^{s_{j+1}} c_{\Pi}(x_i, s) ds - \int_0^{s_j} c_{\Pi}(x_i, s) ds \quad \text{and} \quad \int_{s=s_j}^{s=s_{j+1}} k_{I\Pi}(s) ds = \int_0^{s_{j+1}} k_{I\Pi}(s) ds - \int_0^{s_j} k_{I\Pi}(s) ds .$$

933 $c_{\sigma}(x_i)$ and $k_{I\sigma}$ are given by Eq. (3). Eq. (A.1) can be written in a fully discretized form

934
$$w_i = c_{\sigma,i} \sigma + \sum_{j=1}^{N-1} c_{\Pi,ij}^{elem} \Pi_j^{elem} \quad \text{and} \quad K_{Ie} = k_{I\sigma} \sigma + \sum_{j=1}^{N-1} k_{I\Pi,j}^{elem} \Pi_j^{elem} \quad (A.4)$$

935 where $c_{\sigma,i} = c_{\sigma}(x_i) \cdot \Pi_i^{elem} = (\Pi_i + \Pi_{i+1})/2$ can also be written as a tensor transformation, $\Pi^{elem} = \mathbf{A}_{\Pi} \cdot \Pi$,

936 where

937
$$\mathbf{A}_{\Pi} = \begin{bmatrix} \frac{1}{2} & \frac{1}{2} & 0 & 0 & 0 & 0 \\ 0 & \frac{1}{2} & \frac{1}{2} & 0 & \dots & 0 & 0 \\ 0 & 0 & \frac{1}{2} & \frac{1}{2} & 0 & 0 & 0 \\ \vdots & & & \ddots & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & \frac{1}{2} & \frac{1}{2} \end{bmatrix}_{(N-1) \times N} \quad (A.5)$$

938 Finally, Eq. (A.4) can be written in matrix-vector notation as:

939
$$\mathbf{w} = \mathbf{c}_{\sigma} \sigma + \mathbf{c}_{\Pi} \cdot \Pi \quad (A.6)$$

940

$$K_{Ie} = k_{I\sigma} \sigma + \mathbf{k}_{I\Pi} \cdot \mathbf{\Pi} \quad (\text{A.7})$$

941 where $\mathbf{c}_\Pi = \mathbf{c}_\Pi^{elem} \cdot \mathbf{A}_\Pi$ and $\mathbf{k}_{I\Pi} = (\mathbf{k}_{I\Pi}^{elem})^T \cdot \mathbf{A}_\Pi$. Substituting the relation $\Pi = \Pi(w, p)$ into Eq. (A.6)
 942 yields a nonlinear algebraic equation that can be solved iteratively. The solution is sought here by
 943 rewriting Eq. (A.6) in the same way as Planas and Elices (1991):

$$\mathbf{\Pi}(\mathbf{w}, \mathbf{p}) - \mathbf{M} \cdot \mathbf{w} + \mathbf{L} \sigma = \mathbf{0} \quad (\text{A.8})$$

945 where \mathbf{p} is the nodal vector of gas partial pressure assumed to be known and constant in this local solver,
 946 $\mathbf{M} = \mathbf{c}_\Pi^{-1}$, and $\mathbf{L} = \mathbf{c}_\Pi^{-1} \cdot \mathbf{c}_\sigma$. A Newton-Raphson iteration method is implemented to seek the solution w_i
 947 at each node. Starting with an initial guess, \mathbf{w}_0 , the correction $\mathbf{w}_{k+1} = \mathbf{w}_k + \delta\mathbf{w}_k$ at iteration k is made
 948 where $\delta\mathbf{w}_k = -(\mathbf{J}^M)^{-1} \cdot \mathbf{R}^M$. \mathbf{R}^M and \mathbf{J}^M are computed as

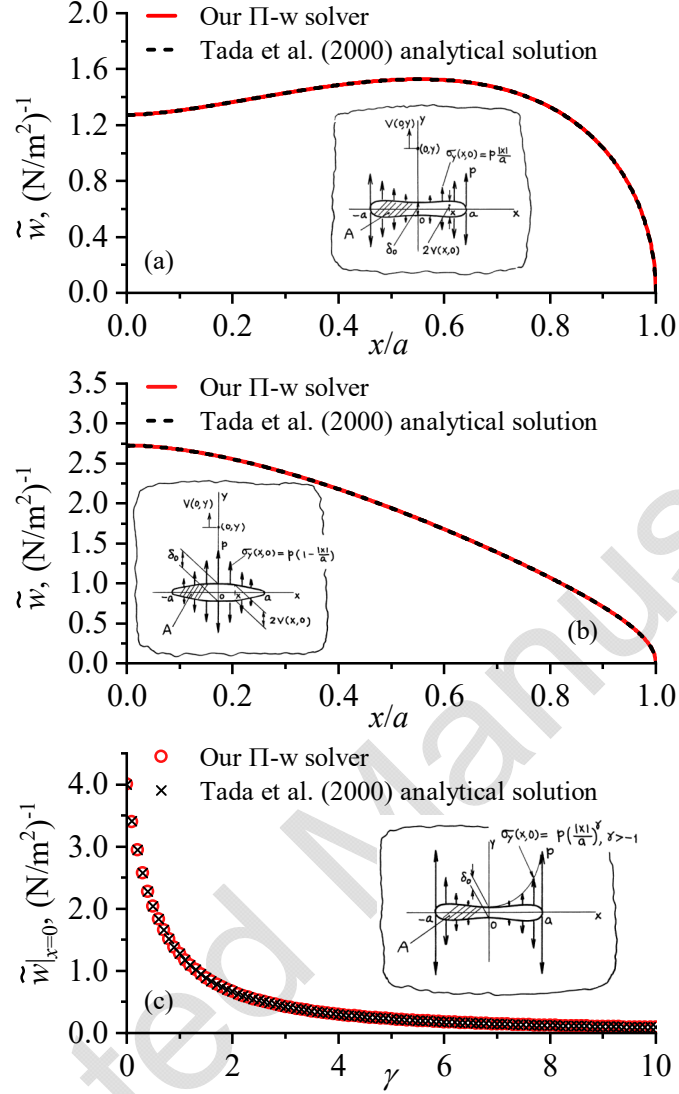
$$949 \quad R_i^M = \Pi(w_i, p_i) - M_{ij} w_j + L_i \sigma \quad \text{and} \quad J_{ij}^M = \frac{\partial \Pi}{\partial w} \Big|_{w_i, p_i} \delta_{ij} - M_{ij} \quad (\text{A.9})$$

950 with $\mathbf{w} = \mathbf{w}_k$. The search for solution continues until the error defined as

$$951 \quad \varepsilon_M = \sqrt{\sum_1^{N-1} (R_i^M)^2 \Delta x_i} \quad (\text{A.10})$$

952 is reduced below a tolerance (here taken as 5×10^{-4}). Once Π and w profiles are obtained, Eq. (A.7)
 953 computes K_{Ie} to conclude the $\Pi - w$ system solver.

954 Our $\Pi - w$ system solver is verified by comparing the computed crack opening profile with the
 955 analytical crack opening solutions of Tada et al. (2000) subjected to different surface force distributions
 956 (Fig. A.2).



957

958

Fig. A.2 Verification of the normalized crack opening profile, $\tilde{w} = wE / (1 - \nu^2)a$, with exact solutions (Tada et al., 2000) for surface force distributions (a) linearly increasing toward the crack tip; (b) linearly decreasing toward the crack tip; and (c) nonlinearly increasing toward the crack tip. Figure insets are courtesy of Tada et al. (2000).

959

960

961

962 APPENDIX B. INITIAL GUESS FOR $T > 0$ AND $T = 0$

963

964

965

The current implementation is stress-controlled (i.e., σ and a vary between timesteps). The crack

velocity is small and the timesteps are close, thus the variation of a over timesteps is very small.

However, a small variation in σ can create a large change in K_{Ie} which exponentially updates v_c and

966 subsequently $p(x)$. It is thus apparent that using the solution at timestep t_k as the initial guess for
 967 timestep t_{k+1} can involve large errors and prevent convergence.

968 We can obtain an initial guess for timestep t_{k+1} by a linear extrapolation. The rate of change of the
 969 solution vector, \mathbf{u} , over time Δt is linearized based on a small variation of σ over a small time $\theta\Delta t$. as

$$970 \quad \frac{\Delta \mathbf{u}}{\Delta t} \approx \frac{\mathbf{u}|_{\sigma(t_k+\theta\Delta t)} - \mathbf{u}|_{\sigma(t_k)}}{\theta\Delta t} \quad (B.1)$$

971 The value of θ is set to 10^{-2} . Since the variation of σ over the time interval $\theta\Delta t$ is now small, the
 972 solution at time t_k works as a good initial guess for the solution at time $t_k + \theta\Delta t$. Thus, $\mathbf{u}|_{\sigma(t_k+\theta\Delta t)}$ is
 973 found using the same partitioned Newton-Raphson solver with the initial guess of $\mathbf{u}|_{\sigma(t_k)}$. The sole
 974 purpose of $\mathbf{u}|_{\sigma(t_k+\theta\Delta t)}$ is to assist with finding a reasonable initial guess for $\mathbf{u}|_{\sigma(t_k+\Delta t)}$, it can thus be
 975 calculated with a more relaxed tolerance to reduce simulation time. Then, the initial guess for time t_{k+1}
 976 is computed via linear extrapolation:

$$977 \quad \mathbf{u}|_{\sigma(t_k+\Delta t)} \approx \mathbf{u}|_{\sigma(t_k)} + \frac{\Delta \mathbf{u}}{\Delta t} \Delta t \quad (B.2)$$

978 While the procedure described here involves an additional set of iterations in search of $\mathbf{u}|_{\sigma(t_k+\theta\Delta t)}$, the
 979 amount of computation required for those iterations is far less than that required for finding the solution
 980 to $\mathbf{u}|_{\sigma(t_k+\Delta t)}$ using the initial guess $\mathbf{u}|_{\sigma(t_k)}$ even if the latter converges.

981 The initial guess for $t = 0$ can be obtained by starting simulations with a small σ so that v_c can be
 982 predicted a few orders of magnitude smaller than d / D . This way, the fluid profile is known, $p(x) = p_0$.
 983 Thus, the $\Pi - w$ system can be solved alone for K_{le} . One must then verify that this K_{le} results in v_c of
 984 similar magnitude as the predicted one.