Simulation of wave propagation along fluid-filled cracks using high-order summation-by-parts operators and implicit-explicit time stepping *

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Abstract. We present an efficient, implicit-explicit numerical method for wave propagation in 4 solids containing fluid-filled cracks, motivated by applications in geophysical imaging of fractured 5 oil/gas reservoirs and aquifers, volcanology, and mechanical engineering. We couple the elastic wave 6 equation in the solid to an approximation of the linearized, compressible Navier-Stokes equations in curved and possibly branching cracks. The approximate fluid model, similar to the widely used 8 lubrication model but accounting for fluid inertia and compressibility, exploits the narrowness of the 9 crack relative to wavelengths of interest. The governing equations are spatially discretized using high-order summation-by-parts finite difference operators and the fluid-solid coupling conditions are 11 weakly enforced, leading to a provably stable scheme. 12

13 Stiffness of the semi-discrete equations can arise from the enforcement of coupling conditions, 14fluid compressibility, and diffusion operators required to capture viscous boundary layers near the crack walls. An implicit-explicit Runge-Kutta scheme is used for time stepping and the entire system 15 of equations can be advanced in time with high-order accuracy using the maximum stable time 1617 step determined solely by the standard CFL restriction for wave propagation, irrespective of the 18 crack geometry and fluid viscosity. The fluid approximation leads to a sparse block structure for 19the implicit system, such that the additional computational cost of the fluid is small relative to 20 the explicit elastic update. Convergence tests verify high-order accuracy; additional simulations 21demonstrate applicability of the method to studies of wave propagation in and around branching 22 hydraulic fractures.

Key words. Fluid-filled crack, wave propagation, summation-by-parts, high-order accuracy, implicit-explicit.

25 AMS subject classifications.

1. Introduction. There is considerable interest in wave propagation in solids 26containing fluid-filled cracks. Hydrocarbon reservoirs, enhanced geothermal systems, 27 and groundwater aquifers all feature fractured rock masses saturated in fluid. Frac-2829 tures, or cracks, in these systems are either naturally occurring or created in hydraulic fracturing treatments, and can be as narrow as $\sim 0.1-10$ mm but with lengths exceed-30 ing ~ 100 m. Similarly high-aspect ratio cracks occur at a much larger scale in the 31 form of magma-filled cracks known as dikes and sills, a primary component of active volcanic systems, and water-filled crevasses and basal hydraulic fractures in ice sheets 33 34 and glaciers. Seismic imaging of these systems provides key constraints on the crack geometry and mechanical properties of the fluids and solids. 35

Simulation of wave propagation in and around fluid-filled cracks presents several computational challenges. Many of these arise from the extreme narrowness of the crack relative to wavelengths of interest; the dimensionless ratio of these two length scales is typically $\sim 10^{-3}$ or even less. Direct solution of the elastic wave equation in the solid and linearized compressible Navier-Stokes equation in the fluid, using finite difference, finite element, or discontinuous Galerkin methods, would involve either distorted meshes or very fine grid spacings that might lead to overly restrictive sta-

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bility constraints for explicit time stepping and/or poorly conditioned linear systems 43 44 for implicit time integration of viscous terms. Some studies have taken this direct approach, most commonly by neglecting fluid viscosity and instead using the acoustic 45 wave equation for the fluid [32, 16, 47, 31]. Boundary element and boundary integral 46methods [10, 39, 48] or even hybrid boundary element / finite difference methods [6] 47 overcome many of these issues, but are thus far restricted to inviscid fluids. Viscosity 48 was added recently in two-dimensional finite element models by Frehner and Schmal-49 holz [14], who solved the full linearized Navier-Stokes equation for the fluid using an 50unstructured mesh and a fully implicit Newmark time-stepping scheme. While a fully implicit time-stepping scheme is feasible for two-dimensional problems of moderate size, it likely becomes impractical or at least highly inefficient for three-dimensional 53 54 problems. Nevertheless, their work demonstrates the key role that viscosity plays in damping waves.

Others have taken advantage of the narrowness of the crack by utilizing approxi-56 mate fluid models. In these models, the crack, from the perspective of the solid, is an infinitesimally thin interface. Along this interface, a lower-dimensional set of partial 58 differential equations (PDE) or even local relations between tractions and displace-59 ment discontinuities are used to describe the fluid response. The local relations can 60 be as simple as traction-free interface conditions [31] though more widely adopted is 61 the linear slip model [9]. While local relations can be incorporated into explicit elastic 62 wave propagation codes [9, 47, 31] with relative ease, they fail to capture a fundamen-63 tal type of guided wave that propagates along fluid-filled cracks. This wave, known as 64 65 a Krauklis wave [24, 13, 21], has generated considerable interest in volcanology [13] and the oil and gas industry [21] because Krauklis wave resonance can be used to 66 deduce crack geometry and properties of the fluid within the crack [27, 26]. 67

68 Studies focusing on Krauklis waves have therefore utilized PDE fluid models [8, 69 7], though viscosity is typically neglected or captured by the assumption of fully 70 developed (Poiseuille) flow. However, at the frequencies of interest, viscous dissipation 71 can neither be ignored nor properly described by Poiseuille flow, as it reaches its 72 maximum within boundary layers near the crack walls.

In this work, we present a numerical scheme that combines fully explicit time stepping of the elastic wave equation and a PDE fluid model based on a lubricationtype approximation to the linearized compressible Navier-Stokes equations. We use high-order summation-by-parts (SBP) finite difference operators [25, 42, 35, 44] for spatial discretization. The fluid-solid coupling conditions are weakly enforced using the simultaneous-approximation-term (SAT) penalty technique [5], and geometric complexity is handled with curvilinear, multiblock grids.

We identify several sources of stiffness in the semi-discrete problem, arising from 80 compressibility and viscosity of the fluid. This stiffness is isolated by partitioning the 81 semi-discrete equations, and advancing the partitioned system in time with a high-82 order implicit-explicit (IMEX) Runge-Kutta method [1, 4, 20, 36]. Similar partitioning 83 has been exploited in related fluid-structure interaction simulations [38, 12, 29, 17, 19, 84 46, 15]. A major advantage of our approximate fluid model, over the full linearized 85 86 Navier-Stokes equations, is that the linear system arising in the implicit component of the time-stepping scheme has a sparse block diagonal structure. This substantially 87 88 enhances computational efficiency.

This paper is structured as follows. In Section 2 we describe the overall problem, with focus in 2.1 and 2.2 on the solid and fluid equations. These are combined, in 2.3, through the fluid-solid coupling conditions. These conditions are incorporated into a variational formulation of the continuous problem with a weak enforcement of coupling 93 conditions. We establish well-posedness by deriving an energy estimate. In Section 94 3 we present the semi-discrete approximation and establish stability by deriving a 95 discrete energy estimate. In Section 4 we present the fully discrete approximation 96 by discretizing in time using a high-order IMEX Runge-Kutta method. Section 5 97 demonstrates high-order convergence of the method using the method of manufactured 98 solutions, followed by two application problems illustrating wave propagation in and 99 around a branching fluid-filled crack. In Section 6 we provide a summary of the results 100 and perspectives on future work.

2. Continuous problem. In this section we introduce the governing equations 101 for the solid and fluid, along with conditions for coupling the solid and fluid across the 102moving crack walls. We restrict attention to the two-dimensional problem, as shown 103 in Figure 1. The solid occupies the domain Ω_s and contains a crack, which is treated 104 from the perspective of the solid as an infinitesimally thin interface $\Gamma \subseteq \Omega_s$. The 105crack contains a compressible, viscous fluid defined on the domain Ω_f . Rather than 106 solving the compressible Navier-Stokes equations in their most general form, we seek a 107 108 linearized description of the fluid, assuming small perturbations about a state of rest. Furthermore, we utilize a lubrication-type approximation to take advantage of the 109 110 fact that the crack width is much smaller than wavelengths of interest; however, we retain essential terms in the linearized Navier-Stokes equations that account for fluid 111 compressibility and inertia. Our model generalizes the model of [27] to account for 112 crack curvature and nonplanarity of the crack walls. Similar compressible lubrication 113models are used in engineering, particularly for problems involving gas-filled bearings 114 and in studies of liquid droplet impact on surfaces [45, 41, 3, 18, 2]. 115

2.1. Solid. Assuming linear elastic material response and small strains and rotations, the solid is governed by the elastic wave equation:

118 (1)
$$\rho_s \frac{\partial v}{\partial t} = A_x \frac{\partial \sigma}{\partial x} + A_y \frac{\partial \sigma}{\partial y}$$

119 (2)
$$S\frac{\partial\sigma}{\partial t} = A_x^T\frac{\partial v}{\partial x} + A_y^T\frac{\partial v}{\partial y},$$

121 where

122
123
$$A_x = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 0 & 1 \end{bmatrix}, \ A_y = \begin{bmatrix} 0 & 0 & 1 \\ 0 & 1 & 0 \end{bmatrix},$$

124 $v(x, y, t) = [v_x v_y]^T$ is the particle velocity, $\sigma(x, y, t) = [\sigma_{xx} \sigma_{yy} \sigma_{xy}]^T$ is the stress, ρ_s 125 is the density, and $S = S^T > 0 \in \mathbb{R}^{3 \times 3}$ is the compliance matrix. Note that the x and 126 y subscripts denote the components of the solid velocity and stress and should not be 127 confused with partial derivatives. For an isotropic solid, as used in all simulations in 128 this work, the compliance matrix is

129
130
$$S = \frac{G}{2} \begin{bmatrix} 1 - \nu & 0 & -\nu \\ 0 & 2 & 0 \\ -\nu & 0 & 1 - \nu \end{bmatrix},$$

where G > 0 is the shear modulus and $-1 < \nu < 0.5$ is Poisson's ratio. However, the numerical scheme developed below is applicable to anisotropic linear elastic solids as well.

134 Curvature of the crack and possibly other geometric complexities in the shape 135 of the solid are handled by formulating the elastic wave equation in curvilinear co-136 ordinates. We also utilize a particular splitting of the equations that facilitates the



(b) Fluid grid

FIG. 1. (a) Linear elastic solid Ω_s containing a fluid-filled crack, appearing from the perspective of the solid as an infinitesimally thin interface $\overline{\Gamma}$. \hat{s} and \hat{n} denote unit vectors parallel and normal to Γ , with \hat{n} pointing from the - side to + side of Γ . (b) Zoomed-in view of the fluid domain Ω_f within the crack, along with the mesh used to resolve viscous boundary layers near the crack walls. The arc length along the crack is s and the distance across the crack width, normal to s, is n.

construction of the semi-discrete approximation in a manner that leads to an energy 137estimate and thus stability [33]. Consider the curvilinear coordinate transformation 138 $x = x(q,r), y = y(q,r) \leftrightarrow q = q(x,y), r = r(x,y), \text{ mapping } (x,y) \in \Omega_s \text{ to } (q,r) \in \Omega_s.$ 139We assume a smooth, one-to-one mapping, and define $\tilde{\Omega}_s = [0,1] \times [0,1]$ as the ref-140 erence unit square. Following [11], we transform the elastic wave equation by writing 141142 (1) in conservative form and (2) in non-conservative form [28], which leads to

$$\begin{array}{l} {}^{143}_{144} \quad (3) \qquad \qquad \rho_s J \frac{\partial v}{\partial t} = \frac{\partial}{\partial q} (JA_q \sigma) + \frac{\partial}{\partial r} (JA_r \sigma), \ S \frac{\partial \sigma}{\partial t} = A_q^T \frac{\partial v}{\partial q} + A_r^T \frac{\partial v}{\partial r}, \end{array}$$

145where

146 (4)
$$A_q = \begin{bmatrix} q_x & 0 & q_y \\ 0 & q_y & q_x \end{bmatrix}, \ A_r = \begin{bmatrix} r_x & 0 & r_y \\ 0 & r_y & r_x \end{bmatrix}.$$

In (4), the metric coefficients q_x, q_y, \ldots , are obtained by taking partial derivatives 148149of each coordinate. For example, $q_x = \partial q / \partial x$. The metric coefficients are the only

quantities which use compact derivative notation, and should not be confused with the x and y components of a vector. Furthermore, J > 0 is the Jacobian of the mapping, defined as $J = x_q y_r - y_q x_r$. The metric coefficients satisfy the metric relations $Jq_x = y_r$, $Jr_x = -y_q$, $Jq_y = -x_r$, $Jr_y = x_q$.

The coupling conditions will be stated using the solid fields locally oriented with respect to the curved fluid-solid interface. We therefore define the velocity V and traction T expressed in terms of the normal and tangential components given by the unit normal \hat{n} and unit tangent \hat{s} along Γ (Figure 1a). We have

158 (5)
$$V = [v_n \ v_s]^T = R^T v \text{ and } T = [\sigma_n \ \sigma_s]^T = R^T \frac{A_r}{|\nabla r|} \sigma,$$

where v_n and v_s are the normal and tangential components of the solid particle velocity, respectively, and σ_n and σ_s are the normal and shear components of the solid traction, respectively. To obtain these components, we have introduced the rotation matrix R:

164 (6)
$$R^T = \begin{bmatrix} \hat{n}^T \\ \hat{s}^T \end{bmatrix} = \frac{1}{|\nabla r|} \begin{bmatrix} r_x & r_y \\ r_y & -r_x \end{bmatrix},$$

166 where $|\nabla r| = (r_x^2 + r_y^2)^{1/2}$.

167 **2.2. Fluid.** The fluid is governed by an approximation to the linearized com-168 pressible Navier-Stokes equations. It has density ρ_f , dynamic viscosity μ , and bulk 169 modulus K_f . The fluid equations are stated in a coordinate system (s,n) locally 170 oriented with respect to Γ , for which s is the arc length along Γ and n measures the 171 distance across the width of the crack in the direction normal to s. The upper and 172 lower crack walls are initially located at $n = w_0^{\pm}(s)$ (Figure 1b), but are perturbed to 173 $n = w^{\pm}(s,t)$. The initial width of the crack is defined as $w_0 = w_0^{\pm}(s) - w_0^{-}(s)$.

Following the usual procedure for deriving lubrication-type approximations [41, 3, 18], a scaling analysis of the momentum balance in the *n*-direction establishes uniformity of the fluid pressure across the width of the crack. Conservation of fluid mass, together with a barotropic equation of state, leads to the first governing equation for the fluid. The linearized version of this equation is, on Γ ,

$$\frac{179}{180} \quad (7) \qquad \qquad \frac{w_0}{K_f} \frac{\partial p}{\partial t} + \frac{\partial}{\partial s} (w_0 \bar{u}) = -\left(\frac{\partial w^+}{\partial t} - \frac{\partial w^-}{\partial t}\right),$$

181 for pressure p(s,t) and width-averaged velocity

182 (8)
$$\bar{u}(s,t) = \frac{1}{w_0} \int_{w_0^-}^{w_0^+} u(s,n,t) dn$$

where u = u(s, n, t) is the fluid velocity in the *s*-direction. Equation (7) is derived by integrating the local form of the continuity equation across the crack width, using the kinematic condition to replace the normal component of fluid velocity with the crack opening rate, and linearizing about a state of rest.

188 At this point, the classical lubrication model would neglect inertia by assuming a 189 fully developed Poiseuille flow profile, for which

190 (9)
$$\bar{u} = \frac{w_0^2}{12\mu} \left(-\frac{\partial p}{\partial s}\right) + \frac{v_s^+ - v_s^-}{2}$$

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where $v_s^+ - v_s^-$ is the discontinuity in the tangential component of solid particle velocity 192 across Γ . However, at the frequencies of interest to us, fluid inertia leads to non-193 parabolic velocity profiles with Stokes-type boundary layers adjacent to the crack 194 walls [3]. To obtain a physically relevant \bar{u} we must therefore solve the s-momentum 195balance on the two-dimensional domain Ω_f : 196

197 (10)
$$\rho_f \frac{\partial u}{\partial t} + \frac{\partial p}{\partial s} = \frac{\partial \tau}{\partial n},$$

199where

$$\begin{array}{cc} 200\\ 201 \end{array} \quad (11) \qquad \qquad \tau = \mu \frac{\partial u}{\partial n} \end{array}$$

is the shear stress. Equation (10) retains, on the right-hand side, a single viscous term 202 describing shearing on planes parallel to Γ and diffusive momentum transport across 203 204 these planes. Effects of curvature have been neglected in the momentum balance, 205under the assumption that the radius of curvature of Γ is comparable to or larger than the wavelengths of interest. Note that when inertia is neglected, the solution 206207 to the momentum balance equation (10) with no-slip conditions on the crack walls provides the classical lubrication solution (9). In this classical lubrication limit, the 208 209 associated shear stress on the top and bottom crack walls is

210 (12)
$$\tau^{\pm} = \mp \frac{w_0}{2} \left(-\frac{\partial p}{\partial s}\right) + \frac{\mu(v_s^+ - v_s^-)}{w_0}$$

Note that while the method developed in this paper uses the more general lubrication 212 model that accounts for fluid inertia, it would be a straightforward extension to instead 213 use the classical lubrication model embodied in equations (9) and (12). 214

We apply a coordinate transformation in the s direction for compatibility with 215216 the curvilinear grid used in the solid. We also apply a coordinate transformation in the n direction in the fluid to resolve the boundary layers by clustering grid points 217near the walls. Consider the coordinate transformation $s = s(q), n = n(q, r) \leftrightarrow q =$ 218 q(s), r = r(s, n) that maps $(s, n) \in \Omega_f$ to a reference unit square $\tilde{\Omega}_f = [0, 1] \times [0, 1]$. 219 Note that since s is the arc length of Γ , it can only depend on q. The Jacobian and 220 metric relations become $J = s_q n_r$, $s_r = 0$, $Jq_s = n_r$, $Jr_s = -n_q$, $Jq_n = 0$, $Jr_n = s_q$. 2212.2.2 Transforming (7), (8), (10), and (11) leads to the final governing equations for the 223 fluid:

(13)

$$s_{q} \frac{w_{0}}{K_{f}} \frac{\partial p}{\partial t} + \frac{\partial}{\partial q} (w_{0} \bar{u}) = -s_{q} \left(\frac{\partial w^{+}}{\partial t} - \frac{\partial w^{-}}{\partial t} \right),$$

$$\rho_{f} J \frac{\partial u}{\partial t} + n_{r} \frac{\partial p}{\partial q} = s_{q} \frac{\partial \tau}{\partial r},$$

$$\bar{u}(q, t) = \frac{1}{w_{0}} \int_{r=0}^{r=1} u(q, r, t) n_{r} dr,$$

$$\tau = \mu r_{n} \frac{\partial u}{\partial r}.$$

226 2.3. Fluid-solid coupling conditions and well-posedness. Having made several approximations, we must verify that our problem is well-posed. Well-posedness 227 is established by enforcing the fluid-solid coupling conditions such that the governing 228 229 equations satisfy a mechanical energy balance. In this analysis, we weakly enforce

the coupling conditions. This procedure simplifies the proof of stability in the semi-230 231discrete case, following later.

For simplicity, we consider only the + side of the interface Γ ; the - side is treated 232 in an analogous manner, and boundary conditions on the solid have been discussed 233 extensively in previous work [22, 23, 11]. Since the fluid mass balance is stated on Γ , 234 we only consider the term $\partial w^+/\partial t$ in (13), which will be coupled to the solid on the 235+ side. To simplify the notation in this section, we drop the + superscript. 236

The fluid-solid coupling conditions for a viscous fluid are obtained by balancing 237the tractions across the interface and enforcing the kinematic condition and no-slip 238 condition to ensure that the fluid and solid remain in contact at the crack walls: 239

240 (14)
$$V = [v_n \ v_s]^T = \left[\frac{\partial w}{\partial t} \ u\right]^T \text{ and } T = [\sigma_n \ \sigma_s]^T = [-p \ \tau]^T.$$

The negative sign on fluid pressure arises because pressure is positive in compression, 242 the opposite of the sign convention for solid normal stresses. 243

There are many ways in which the coupling conditions (14) can be enforced. One 244approach is to weakly enforce a common interface velocity $\hat{V} = [\hat{V}_n \ \hat{V}_s]^T$ on the fluid and solid velocities, and a common interface traction $\hat{T} = [\hat{T}_n \ \hat{T}_s]^T$ on the fluid and 245246solid tractions. This procedure uses the fact that the fluid velocities, solid velocities, 247 and tractions are continuous. The interface velocity and traction are determined 248by satisfying the proper mechanical energy balance of the overall problem. In the 249limit when the coupling conditions become strongly enforced, the fluid and solid 250velocities and tractions should be equal to the interface velocity and traction, i.e., $\hat{V} = [v_n \ v_s]^T = [\partial w / \partial t \ u]^T$ and $\hat{T} = [\sigma_n \ \sigma_s]^T = [-p \ \tau]^T$. Since our governing equations are formulated in curvilinear coordinates, we state 251 252

253all integrations with area and length differentials in the curved domain. The rela-254255tionship between the area differential in the curved domain and transformed domain is $d\Omega \leftrightarrow Jdqdr$. Furthermore, the relationship between the line differential for the interface Γ in the curved domain and transformed domain is $ds \leftrightarrow J |\nabla r| dq$. 257

To obtain the mechanical energy balance satisfied by the fluid and solid, we 258consider the following variational form of the solid: 259

$$\int_{\Omega_s} \phi_s^T \rho \frac{\partial v}{\partial t} d\Omega = \int_{\Omega_s} \phi_s^T \frac{1}{J} \frac{\partial}{\partial q} (JA_q \sigma) + \phi_s^T \frac{1}{J} \frac{\partial}{\partial r} (JA_r \sigma) d\Omega + \int_{\Gamma} (R^T \phi_s)^T (T - \hat{T}) ds$$

$$\int_{\Omega_s} \varphi_s^T S \frac{\partial \sigma}{\partial t} d\Omega = \int_{\Omega_s} \varphi_s^T A_q^T \frac{\partial v}{\partial q} + \varphi_s^T A_r^T \frac{\partial v}{\partial r} d\Omega + \int_{\Gamma} \frac{(R^T A_r \varphi_s)^T}{|\nabla r|} (V - \hat{V}) ds.$$

(15)

for smooth, vector-valued test functions $\phi_s, \varphi_s \in L^2(\Omega_s)$. In (15), the integrals along 262the interface Γ are penalty terms, which weakly enforce the coupling conditions. The 263264 rotation matrix R is defined in (6) and arises because the coupling conditions are stated in terms of normal and tangential components. One can derive the penalty 265terms by applying integration by parts to the corresponding volume terms. Another 266 possibility is to introduce an unknown weight Σ in the penalty term and then deter-267mine this weight by satisfying the energy balance [34]. 268

The fluid variational formulation is 269

$$\int_{\Omega_f} & \partial t & \partial q & f & \partial r \\ & -\int_{\Gamma} \varphi_f(\tau - \hat{T}_s) + \mu r_n \frac{\partial \varphi_f}{\partial r} (u - \hat{V}_s) ds,$$

 $\int \varphi_f \rho_f \frac{\partial u}{\partial t} + \varphi_f q_s \frac{\partial p}{\partial t} d\Omega = \int r_n \varphi_f \frac{\partial \tau}{\partial t} d\Omega$

 $\int_{\Gamma} \phi_f \frac{w_0}{K_f} \frac{\partial p}{\partial t} + q_s \phi_f \frac{\partial}{\partial q} (\bar{u}w_0) ds = -\int_{\Gamma} \phi_f \frac{\partial w}{\partial t} ds$

for smooth, scalar test functions $\phi_f \in L^2(\Gamma)$ and $\varphi_f \in L^2(\Omega_f)$. 272

Next, we determine \hat{V} and \hat{T} such that the overall problem satisfies the proper 273mechanical energy balance. The choice of \hat{V} and \hat{T} resulting in well-posedness is 274specified in the following proposition. 275

PROPOSITION 1. The fluid-solid problem (15) and (16) is well-posed and consis-276tent with the coupling conditions (14) if \hat{V} and \hat{T} are chosen as the linear combinations 277

$$\hat{V} = \begin{bmatrix} v_n + \frac{\sigma_n + p}{\alpha_n}, & \frac{\beta_s u + \alpha_s v_s}{\alpha_s + \beta_s} + \frac{\sigma_s - \tau}{\alpha_s + \beta_s} \end{bmatrix}^T \\
\hat{T} = \begin{bmatrix} -p, & \frac{\alpha_s \beta_s}{\alpha_s + \beta_s} (v_s - u) + \frac{\beta_s \sigma_s + \tau \alpha_s}{\alpha_s + \beta_s} \end{bmatrix}^T,$$

for $\{\alpha_n, \alpha_s, \beta_s > 0\} \cup \{\alpha_s = 0, \beta_s > 0\} \cup \{\beta_s = 0, \alpha_s > 0\}.$ 280

Proof. By choosing test functions $\phi_s = v$, $\varphi_s = \sigma$, $\phi_f = p$, $\varphi_f = u$ in (15) and 281(16), combining terms, and integrating by parts, we find 282

$$\frac{dE}{dt} + \Phi = -\int_{\Gamma} \left(v_n \sigma_n + v_s \sigma_s - v_n (\sigma_n - \hat{T}_n) - v_s (\sigma_s - \hat{T}_s) - \sigma_n (v_n - \hat{V}_n) - \sigma_s (v_s - \hat{V}_s) \right) ds$$

283

$$-\int_{\Gamma} p \frac{\partial w}{\partial t} - \tau u + u(\tau - \hat{T}_s) + \tau (u - \hat{V}_s) ds.$$

284

In (1), $\Phi = \int_{\Omega_f} \tau^2 / \mu d\Omega \ge 0$ is the viscous energy dissipation rate and, E is the 285mechanical energy 286

287 (18)
$$E = \frac{1}{2} \int_{\Omega_s} \rho_s v^T v + \sigma^T S \sigma d\Omega + \frac{1}{2} \int_{\Gamma} \frac{w_0}{K_f} p^2 ds + \frac{1}{2} \int_{\Omega_f} \rho_f u^2 d\Omega,$$

where the respective terms are the kinetic and strain energy in the solid, and the 289 elastic and kinetic energy in the fluid. 290

Next, we add and subtract $\hat{T}_n \hat{V}_n$ and $\hat{T}_s \hat{V}_s$ to the right-hand side of (1), which 291 after some algebra leads to 292

$$\frac{dE}{dt} + \Phi = -\int_{\Gamma} \hat{T}_n \hat{V}_n + \hat{T}_s \hat{V}_s - (\sigma_n - \hat{T}_n)(v_n - \hat{V}_n) - (\sigma_s - \hat{T}_s)(v_s - \hat{V}_s)ds$$

$$293 \quad (19) \qquad \qquad -\int_{\Gamma} p \frac{\partial w}{\partial t} - \hat{T}_s \hat{V}_s + (u - \hat{V}_s)(\tau - \hat{T}_s)ds.$$

$$= -(\mathcal{P} + \mathcal{R}),$$

295 where

296 (20)
$$\mathcal{P} = \int_{\Gamma} \hat{T}_n \hat{V}_n + p \frac{\partial w}{\partial t} ds,$$

297 (21)
$$\mathcal{R} = \int_{\Gamma} (u - \hat{V}_s)(\tau - \hat{T}_s) - (\sigma_n - \hat{T}_n)(v_n - \hat{V}_n) - (\sigma_s - \hat{T}_s)(v_s - \hat{V}_s) ds.$$

In (19), we have partitioned the right-hand side into two terms: \mathcal{P} and \mathcal{R} . The first term \mathcal{P} contains the flow of energy from the fluid to the solid and vice versa. When the coupling conditions are enforced, this term must vanish. The second term \mathcal{R} is a residual term arising due to the weak enforcement of the coupling conditions. This term needs to be non-negative and to vanish when the coupling conditions are exactly satisfied. To obtain a well-posed problem, we therefore need to choose \hat{V} and \hat{T} such that $\mathcal{P} = 0$ and $\mathcal{R} \geq 0$. By choosing

$$\hat{T}_n = -p \quad \text{and} \quad \hat{V}_n = \frac{\partial w}{\partial t},$$

308 we obtain $\mathcal{P} = 0$. To bound \mathcal{R} , consider the choice

$$\underset{j \neq 0}{300} \quad (23) \quad \sigma_n - \hat{T}_n = -\alpha_n (v_n - \hat{V}_n), \ \sigma_s - \hat{T}_s = -\alpha_s (v_s - \hat{V}_s), \ \tau - \hat{T}_s = \beta_s (u - \hat{V}_s),$$

for penalty parameters $\alpha_n, \alpha_s, \beta_f \ge 0$. Guidelines for choosing these penalty parameters are given later. Then (21) becomes

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$$\mathcal{R} = \int_{\Gamma} \alpha_n (v_n - \hat{V}_n)^2 + \alpha_n (v_s - \hat{V}_s)^2 + \beta_s (u - \hat{V}_s)^2 \ge 0.$$

By inserting (22) and (23) into (19), we obtain the bound

$$\frac{dE}{dt} = -\Phi - \mathcal{R} \le 0.$$

Note that \mathcal{R} vanishes when the coupling conditions are satisfied exactly, and the energy balance in this limit exactly coincides with the correct mechanical energy balance (i.e., $dE/dt = -\Phi \leq 0$).

When implementing this scheme, we need to determine \hat{V} and \hat{T} . This is done by solving (22) and (23) for \hat{V} and \hat{T} , which yields the stated solution (17). Finally, to show that (17) is consistent with (14), insert (17) into the variational formulations (15) and (16).

3. Semi-discrete approximation. In this section, we utilize the results established in the previous section to construct a stable, semi-discrete approximation. We closely follow the continuous analysis by formulating the semi-discrete approximation in variational form. This will be done by using SBP operators, which are necessary for obtaining a discrete energy estimate and hence a proof of stability.

330 3.1. Definitions. While a multiblock discretization is used for both the fluid and solid domains in realistic applications, we keep the presentation brief by focusing on only one solid and one fluid block. The solid block is located above the crack, as illustrated in Figure 1a. Let the reference domain $\hat{\Omega} = [0, 1] \times [0, 1]$ be discretized by an $(N_q + 1) \times (N_r + 1)$ two-dimensional grid. Furthermore, let the two coordinate directions q and r in the reference domain be discretized by $q_i = i\Delta q$ for $0 \le i \le N_q$, and $r_j = j\Delta r$ for $0 \le j \le N_r$ using grid spacings $\Delta q = 1/N_q$ and $\Delta r = 1/N_r$. For each field, we introduce a grid function $u_{ij}(t) = u(q_i, r_j, t)$, which is stored in a vector u(t) with r being the contiguous direction. The storage order of u_{ij} is, of course, arbitrary, but our particular choice facilitates organization and presentation through use of Kronecker tensor product notation.

Having introduced grids and grid functions, next we define SBP operators. An SBP first derivative difference operator is given in Definition 1; its properties are satisfied by construction.

344 DEFINITION 1. The difference operator D is a summation-by-parts first derivative 345 SBP(2s,s) with interior accuracy 2s and boundary accuracy s with following properties. 346 1. The diagonal matrix H > 0 defines the discrete norm

³⁴⁷
³⁴⁸ (24)
$$\|\phi\|_h^2 = \phi^T H \phi, \ \|\phi\|_h^2 \approx \|\phi\|^2 = \int_0^1 \phi^2 dx,$$

349 for a smooth test function ϕ and a corresponding grid function ϕ . 350 2. The SBP property

$$\frac{351}{352} \qquad (25) \qquad HD + D^T H = B = diag([-1 \ 0 \ \dots \ 1])$$

holds. Here, B is the restriction of ϕ to the right and left boundary:

$$\phi^T B \phi = \phi_N^2 - \phi_0^2.$$

³⁵⁶ For more details concerning accuracy relations, see [42, ?].

357 3.2. Solid. By using the definition of the SBP difference operator, we discretize the variational formulation of the solid (15):

(26)

$$\begin{split} \phi_s^T (I_2 \otimes \rho M_s) \frac{dv}{dt} &= \phi_s^T (I_2 \otimes M_s J^{-1}) \big((I_2 \otimes D_q \otimes I_r) (I_2 \otimes J) A_q \\ &+ (I_2 \otimes I_q \otimes D_r) (I_2 \otimes J) A_r \big) \sigma + (R^T L_s^T \phi_s)^T (I_2 \otimes \bar{M}_s) (T - \hat{T}), \\ \varphi_s^T S (I_3 \otimes M_s) \frac{d\sigma}{dt} &= \varphi_s^T (I_3 \otimes M_s) \big(A_q^T (I_3 \otimes D_q \otimes I_r) + A_r^T (I_3 \otimes I_q \otimes D_r) \big) v \end{split}$$

360

359

+
$$(I_2 \otimes |\nabla r|^{-1} R^T L_s A_r \varphi_s)^T (I_2 \otimes \overline{M}_s) (V - \hat{V}).$$

In (26), all of the material properties, Jacobian, and metric coefficients, evaluated 361 at each grid point, are stored in diagonal matrices. The matrix I_2 is a 2 \times 2 iden-362 363 tity matrix, and \otimes is the Kronecker product. The difference operators D_q and D_r are SBP finite difference operators (see Definition 1). The matrices A_q and A_r are 364 block diagonal matrices containing the metric coefficients (approximated using SBP 365 operators). In the penalty terms, the operator L_s is used to obtain the velocity V 366 and traction T on the interface. For example, we compute V using $V = R^T L_s^T v$, where $L_s = I_2 \otimes I_q \otimes e_0$, and $e_0 = [1 \ 0 \ \dots \ 0]^T$. The rotation matrix R is defined 367 368 using (6). The interface velocity \hat{V} and traction \hat{T} , given in (17), are determined in 369 a similar manner. The mass matrices M_s and \overline{M}_s are diagonal matrices obtained by approximating integrals over Ω_s and along Γ , respectively, using the SBP quadrature 371 rules given in Definition 1. We have $M_s = J(H_q \otimes H_r)$ and $\overline{M}_s = L^T J |\nabla r| L H_q$. 372

Since the variational formulation holds for all non-trivial test functions, we obtain the strong formulation of the semi-discrete approximation of the solid by eliminating the test functions in (26) and inverting the matrices on the left-hand side. Note that the strong form of the equations (26) can be directly advanced in time using explicit time stepping since the mass matrices are diagonal and the inverse of S is known.

3.3. Fluid. Many of the definitions needed to formulate the semi-discrete ap-378 379 proximation of the solid equations are also used to formulate the semi-discrete approximation of the fluid equations. One difference, however, is the appearance of the 380 second derivative operator in the viscous diffusion term. While the second derivative 381 can be constructed by applying the first derivative twice, this procedure leads to a 382 difference operator with sub-optimal stencil width and accuracy. Therefore, in our im-383 plementation, we use a compact second derivative operator with variable coefficients 384 [30]. However, since the presentation and proof of stability become more complicated 385 when using compact operators, in the derivation below we use the first derivative 386 applied twice. 387

The discretization of the weak form of the fluid governing equations (16) is

(27)

388

In (27), the shear stress τ is determined by 391

The width-averaged velocity \bar{u} is computed using the SBP quadrature rule: 394

395 (29)
$$\bar{u} = (w_0 \otimes e_r^T H_r) n_r u \approx \frac{1}{w_0} \int_{r=0}^{r=1} u n_r dr,$$

where $e_r = [1 \ 1 \ \dots \ 1]^T$. We approximate volume integrals over Ω_f and surface 397 integrals along Γ using $M_f = (s_q H_q \otimes H_r) n_r$ and $\bar{M}_f = H_q s_q$, respectively. Since 398 the quadrature rules along the interface on the fluid and solid sides are constructed 399 in the same way, we define $M = M_f = M_s$. Note that the quadrature rule H_r , used 400to calculate \bar{u} , is the same as the one constructing M_f . This is required to obtain an 401 energy balance for the semi-discrete approximation. 402

3.4. Stability. Finally, we show that the semi-discrete approximation is stable 403through the following proposition. 404

PROPOSITION 2. The fluid-solid semi-discrete approximation given by (26) and 405406 (27) is stable.

Proof. The results follow from Proposition 1 and use of the SBP property (25). 407 The energy (18) is approximated as 408

⁴⁰⁹₄₁₀
$$E_h = \frac{1}{2} v^T (I_2 \otimes \rho_s M_s) v + \frac{1}{2} \sigma^T S (I_3 \otimes M_s) \sigma + \frac{1}{2} p^T w_0 K^{-1} \bar{M}_f p + \frac{1}{2} u^T \rho_f M_f u.$$

The semi-discrete approximations (26) and (27) satisfy 411

$$\frac{dE_h}{dt} + \Phi_h = -\alpha_n (v_n - \hat{V}_n)^T \bar{M} (v_n - \hat{V}_n) - \alpha_s (v_s - \hat{V}_s)^T \bar{M} (v_s - \hat{V}_s) - \beta_s (u - \hat{V}_s)^T \bar{M} (u - \hat{V}_s) \le 0.$$

Here, $\Phi_h = \tau^T M_f \tau / \mu \ge 0$ approximates the viscous energy dissipation rate. Since the energy rate of the semi-discrete approximation is non-positive, the numerical solution is bounded, implying stability. The terms arising from the weak enforcement of the coupling conditions yield additional numerical dissipation, vanishing with grid refinement.

4. Fully discrete approximation. Next we turn our attention to time step-419 ping. While the overall problem is dominantly one of wave propagation, there are 420 several sources of stiffness. Our objective here is to advance the solution in time, 421 with high-order accuracy, using a fully explicit method for the elastic wave equation 422 423 (anticipating that this will dominate the computational expense) and with a time step limited only by the usual CFL condition for wave propagation. To overcome stiffness, 424 425we formulate the fully discrete scheme by first partitioning the semi-discrete approximation into stiff and non-stiff parts. The latter accounts for all terms in the governing 426 equations describing wave propagation in the solid and fluid. Then we advance the 427 partitioned system in time using a high-order implicit-explicit (IMEX) Runge-Kutta 428method [1, 4, ?, 20]. The stiff and non-stiff terms are integrated implicitly and ex-429plicitly in time, respectively. 430

431 The semi-discrete approximations (26) and (27) are written in matrix-vector form 432 as

$$\begin{array}{l} 433\\ 434 \end{array} (30) \qquad \frac{dq}{dt} = Wq + Cq + g(t), \ q = \begin{bmatrix} q_f\\ q_s \end{bmatrix}, \ W = \begin{bmatrix} W_f & 0\\ 0 & W_s \end{bmatrix}, \ C = \begin{bmatrix} C_f & C_{fs}\\ C_{sf} & C_s \end{bmatrix}$$

where $q_f = [p^T, u^T]^T$ and $q_s = [v^T, \sigma^T]^T$. In (30), the matrix W holds the difference operators and boundary terms of the fluid and solid, C holds the fluid-solid coupling terms, and q(t) is a forcing function containing external data. We partition (30) into

438 (31)
$$\frac{dq}{dt} = F^{IM}q + F^{EX}q + g(t),$$

440 where

441 (32)
$$F^{IM} = \begin{bmatrix} W_f^{IM} + C_f & C_{fs} \\ 0 & 0 \end{bmatrix}, \ F^{EX} = \begin{bmatrix} W_f^{EX} & 0 \\ C_{sf} & M_s + C_s \end{bmatrix}$$

will be treated implicitly and explicitly, respectively. The partitioning of W_f treats diffusion (contained in W_f^{IM}) implicitly and wave propagation (contained in W_f^{EX}) explicitly. In this work we apply the time integrator ARK4(3)6L[2]SA-ESDIRK (implicit component) and ARK4(3)6L[2]SA-ERK (explicit component) presented in [20]. For future reference, we shall refer to this scheme as ARK4.

448 **4.1. Choice of penalty parameters.** The stiffness of the partitioned, fully 449 discrete scheme (32) is influenced by the penalty parameters α_n , α_s , and β_s appearing 450 in Proposition 1. We explain how to choose the parameters such that the maximum 451 stable time step is set by the usual CFL condition for wave propagation.

To determine the maximum stable time step, we compute the spectral radius of the IMEX stability function given in [4]. This function is the iteration matrix $\widehat{R}(F^{EX}, F^{IM})$ of the fully discrete approximation

$$455 \quad (33) \qquad \qquad q^{k+1} = \widehat{R}q^k,$$

with g(t) = 0 in (32). In (33), q^k denotes the numerical solution at time $t_k = k\Delta t$ for k = 0, 1, 2, ..., and Δt is the time step. If the spectral radius $\rho(\hat{R}) > 1$, then the approximation is not stable. The maximum stable time step is then defined as max Δt s.t. $\rho(\hat{R}) \leq 1$ and \hat{R} is diagonalizable.

461 As in our previous work [22, 23, 11], the solid penalty parameters α_n and α_s are 462 chosen to match the compressional and shear wave impedances, respectively:

$$\alpha_n = Z_p = \rho_s c_p \quad \text{and} \quad \alpha_s = Z_s = \rho_s c_s,$$

where $c_p = \sqrt{M/\rho_s}$ is the compressional wave speed, with $M = 2G(1-\nu)/(1-2\nu)$, and $c_s = \sqrt{G/\rho_s}$ is the shear wave speed. We refer to this choice of penalty parameters as the characteristic choice because α_s and α_n can be obtained by solving the Riemann problem of the elastic wave equation.

469 The fluid penalty parameter β_s is determined by minimizing the spectral radius 470 of the semi-discrete approximation of a one-dimensional model problem describing 471 plane shear waves normally incident on a layer of viscous fluid. Thus, we consider the 472 coupling of the shear wave equation to the diffusion equation in one dimension:

$$\int \phi_s \rho_s \frac{\partial v_x}{\partial t} dy = \int \phi_s \frac{\partial \sigma_{xy}}{\partial y} dy + \left[\phi_s \left(\frac{\alpha_s(\sigma_{xy} - \tau)}{\alpha_s + \beta_s} - \frac{\alpha_s \beta_s}{\alpha_s + \beta_s} (v_x - u) \right) \right]_{y=0^+},$$

$$\int \varphi_s \frac{1}{G} \frac{\partial \sigma_{xy}}{\partial t} dy = \int \varphi_s \frac{\partial v_x}{\partial y} dy + \left[\phi_s \left(\frac{\beta_s(v_x - u)}{\alpha_s + \beta_s} - \frac{\sigma_{xy} - \tau}{\alpha_s + \beta_s} \right) \right]_{y=0^+},$$

$$\int \phi_f \rho_f \frac{\partial u}{\partial t} dn = \int \phi_f \tau dn - \left[\phi_f \left(\frac{\beta_s(\tau - \sigma_{xy})}{\alpha_s + \beta_s} + \frac{\alpha_s \beta_s}{\alpha_s + \beta_s} (u - v_x) \right) + \frac{\partial \phi_f}{\partial n} \left(\frac{\alpha_s(u - v_x)}{\alpha_s + \beta_s} + \frac{\tau - \sigma_{xy}}{\alpha_s + \beta_s} \right) \right]_{n=w_0^+}.$$
473
$$474$$

In (35), the crack is located at y = 0 in the solid. We have weakly enforced the coupling conditions $v_x = u$ and $\sigma_{xy} = \tau$ on the top crack wall ($y = 0^+$ in the solid and $n = w_0^+$ in the fluid) using Proposition 1.

Since the penalty parameters carry units of impedance, a reasonable choice for 478 β_s would be the fluid impedance. For a time-harmonic solution in the boundary layer 479limit the fluid impedance is $Z_f(\omega) = (\mu \rho_f \omega)^{1/2}$. However, since the fluid impedance 480depends on the angular frequency ω , we cannot use it directly. Instead, we estimate 481 it in the following manner. Let ω^* be a frequency of interest. Then, for accuracy, 482 we constrain the fluid and solid grid spacings to be $\Delta x_f = (\mu/\rho_f \omega^*)^{1/2}$ (to resolve 483 the momentum diffusion length at this frequency) and $\Delta x_s = c_s/\omega^*$ (to resolve shear 484 waves), respectively. The impedance parameter can be chosen as $\beta_s = Z_f(\omega^*) =$ 485 $\mu/\Delta x_f$. 486

While $\beta_s = \mu/\Delta x_f$ is a reasonable choice for many problems, it is not always optimal. To demonstrate this, we also investigate two alternative choices that arise in certain limits, specifically when the fluid impedance vanishes ($\beta_s = 0$, as for an inviscid fluid) or when the fluid impedance approaches infinity ($\beta_s \to \infty$). To enforce $\beta_s \to \infty$, we analytically take the limit. Then (35) becomes

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$$\int \phi_s \rho_s \frac{\partial v_x}{\partial t} dy = \int \phi_s \frac{\partial v_x}{\partial y} dy - \left[\phi_s \alpha_s (v_x - u) \right]_{y=0^+},$$

$$\int \varphi_s \frac{1}{G} \frac{\partial \sigma_{xy}}{\partial t} dy = \int \varphi_s \frac{\partial v_x}{\partial y} dy + \left[\varphi_s (v_x - u) \right]_{y=0^+},$$

$$\int \phi_f \rho_f \frac{\partial u}{\partial t} dn = \int \phi_f \tau dn - \left[\phi_f \left((\tau - \sigma_{xy}) + \alpha_s (u - v_x) \right) \right]_{n=w_0^+}.$$

 $\partial \sigma$

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The part of parameter space that we investigate depends on the ratio of the fluid impedance Z_f (in the boundary layer limit) to solid impedance Z_s :

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497
$$\gamma = \frac{Z_f}{Z_s} = \frac{\sqrt{\mu\omega^*\rho_f}}{\rho_s c_s} = \frac{\Delta x_f \rho_f}{\Delta x_s \rho_s}.$$

498 For simplicity, we restrict attention to $\rho_s = \rho_f$ and use the SBP(6,3) operators to discretize (35). For each choice of β_s , we compute the spectral radius $\rho(W+C)$ 499as a function of the impedance ratio γ (Figure 2). Here, the W + C is matrix in 500the semi-discrete approximation of (35), which can be put in the same form as (32). 501Figure 2 shows that $\beta_s = 0$ minimizes $\rho(W + C)$ for $\gamma \ll 1$, whereas for larger values 502 of $\gamma, \beta_s \to \infty$ is the optimal choice. Note that for $\gamma \ll 1$, the spectral radius for 503 $\beta_s = \mu / \Delta x_f$ is nearly identical to that for $\beta_s = 0$. Therefore, in our implementation, 504we never use $\beta_s = \mu/\Delta x_f$, because it is more complicated to implement and shows 505no benefit compared to $\beta_s = 0$ for $\gamma \ll 1$. 506

507 Instead, we propose the following strategy for choosing β_s :

508 (36)
509
$$\beta_s = \begin{cases} 0, & \gamma < \gamma^* \\ \infty, & \gamma > \gamma^* \end{cases}.$$

510 The parameter γ^* is defined as the value of γ at which $\rho(W + C(\beta_s = 0)) = \rho(W + 511 \quad C(\beta_s \to \infty))$, as estimated from Figure 2. For the example shown, $\gamma^* \approx 10^{-1}$.



FIG. 2. Spectral radius $\rho(W+C)$ of the semi-discrete approximation of the problem (35).

512 Note, however, that while this choice ensures that the spectral radius of the semi-513 discrete approximation is minimized, there is no guarantee that the fully discrete 514 approximation is stable for a time step set by the usual CFL condition for wave 515 propagation. For example, the use of energy-conserving coupling conditions, which 516 are traditionally used in many fluid-structure interaction applications, results in loss of stability (see Appendix A for more details). By also analyzing the fully discrete approximation, we have found that when choosing the penalty parameters as (34) and (36), the maximum stable time step remains constant regardless of the amount of stiffness (i.e., the time step is not restricted by the width of the fluid layer or fluid properties). We have therefore achieved our objective of developing a fully discrete scheme that can be advanced with a time step determined only by wave propagation.

523 **5. Numerical Experiments.** In this last part of the paper we investigate the 524 accuracy of our numerical scheme using the method of manufactured solutions and 525 showcase the code capabilities with two application problems featuring a curved, 526 branching crack.

527 **5.1. Manufactured solutions.** We construct a smooth solution and quantify 528 error and convergence rate using the method of manufactured solutions [40]. Param-529 eters are chosen for which the semi-discrete equations are quite stiff; this provides a 530 comprehensive test of the partitioning and IMEX time-stepping procedure.



FIG. 3. MMS verification problem. Two square blocks, Ω_1 (blue) and Ω_2 (red), are joined along the fluid-filled crack Γ , which has the nonplanar geometry shown on the right.

Let the solid domain Ω_s be the rectangle $[0, L] \times [-L, L]$ with a nominally planar crack Γ at y = 0 (see Figure 3). Since the geometry is not curved in this test, we will denote all fields using Cartesian coordinates x, y. The geometry is discretized using two elastic blocks (one on each side of the crack) with $(n + 1) \times (n + 1)$ grid points and a single fluid block of size $(n + 1) \times (m + 1)$, where $n = 12 \times 2^j$, $m = 16 \times 2^j$, and $j = 1, 2, \ldots, 5$. A manufactured solution is constructed by adding forcing functions to the governing equations and boundaries and by exactly satisfying the coupling conditions (14). The manufactured solution in the fluid and the crack geometry are

539 (37)
$$p(x,t) = \sin(kx)\cos(\omega t), \ u(x,y,t) = \sin(kx)\sin(ky)\cos(\omega t) + \sin(kx)\cos(\omega t),$$

Table 1

Fluid and solid properties used in the MMS verification problem. The same fluid properties are used in the branching crack problem.

$$\begin{array}{c|ccccc} n = 24 & n = 48 & n = 96 & n = 192 & n = 384 \\ \hline \rho(M_f)/\rho(M_s) & 2.2 \times 10^4 & 3.0 \times 10^4 & 5.7 \times 10^4 & 1.1 \times 10^5 & 2.3 \times 10^5 \\ \hline & \text{TABLE 2} \end{array}$$

Spectral radius ratio, a measure of stiffness for the MMS verification problem.

542 respectively. We prescribe the motion of the interface using

543 (39)
$$\frac{\partial w^+(x,t)}{\partial t} = \sin(kx)\sin(\omega t), \quad \frac{\partial w^-(x,t)}{\partial t} = -\sin(kx)\sin(\omega t)$$

545 The manufactured solution in the solid is

$$\begin{aligned} v_x(x,y \ge 0^+,t) &= u(x,w_0^+,t)c(ky), \quad v_x(x,y \le 0^-,t) = u(x,w_0^-,t)c(ky), \\ v_y(x,y \ge 0^+,t) &= s(kx)s(\omega t), \quad v_y(x,y \le 0^-,t) = -s(kx)s(\omega t), \end{aligned}$$
546 (40)
$$\begin{aligned} \sigma_{xx}(x,y \ge 0^+,t) &= c(kx)c(ky)c(\omega t), \quad \sigma_{xx}(x,y \le 0^-,t) = c(kx)c(ky)c(\omega t), \\ \sigma_{yy}(x,y \ge 0^+,t) &= -p(x,t)c(ky), \quad \sigma_{yy}(x,y \le 0^-,t) = -p(x,t)c(ky), \end{aligned}$$
547
$$\begin{aligned} \sigma_{xy}(x,y \ge 0^+,t) &= \tau(x,w_0^+,t)c(ky), \quad \sigma_{xy}(x,y \le 0^-,t) = \tau(x,w_0^-,t)c(ky), \end{aligned}$$

where $\tau(x, y, t) = \mu \partial u / \partial y = \mu k s(kx) c(ky) c(\omega t), c(x) = \cos(x), s(x) = \sin(x), k =$ 548 $2\pi/L$ m⁻¹, L = 1 m, and $\omega = 20$ s⁻¹. For the manufactured solution to satisfy the 549 governing equations of the fluid and solid, we need to add forcing functions. These 550forcing functions are obtained by inserting (37)-(40) into the governing equations. To 551conserve space, we have omitted presenting the forcing functions. Initial conditions 552are determined by evaluating (37) and (40) at t = 0. Boundary conditions are enforced 553 by specifying (37) and (40) as data on the incoming characteristic variable. Additional 554parameters are listed in Table 1. After discretizing with SBP-SAT, the semi-discrete 555approximation becomes stiff (Table 2). 556

The numerical error $e^{(n_j)} = u^{(n_j)} - u^*$ is defined as the difference of the numerical solution $u^{(n_j)}$, and the exact solution u^* sampled at the grid points of the j^{th} grid and computed using (37) and (40). The convergence rate is

560
561
$$\operatorname{rate} = \log_2 \left(\frac{\|e^{(n_j)}\|_h}{\|e^{(n_{j+1})}\|_h} \right),$$

where $||e^{(n_j)}||_h$ is the norm of the error, in the energy norm on the j^{th} grid. Time integration is carried out using ARK4 to the final time t = 0.16 s with a time step $\Delta t = h/c_p$, where h is the grid spacing in the solid. We test using the SBP(6,3) operators. Table 3 shows that the scheme is 4^{th} -order accurate, confirming the expected order of accuracy [43].

567 **5.2.** Branching cracks at a material interface. Next we present two ap-568 plication problems featuring a curvilinear crack branching into two additional crack

		n = 24	n = 48	n = 96	n = 192	n = 384
$w_{1} = 1 m$	$\log_{10} \operatorname{error}$	0.04	-1.13	-2.75	-4.27	-5.64
$w_0 = 1 \text{m}$	rate		3.90	5.37	5.05	4.56
$w_{1} = 0.1 m$	$\log_{10} \operatorname{error}$	-0.55	-1.83	-3.13	-4.42	-5.73
$w_0 = 0.1 \text{ m}$	rate		4.27	4.31	4.30	4.35
$w_{\rm c} = 1 \mathrm{am}$	$\log_{10} \operatorname{error}$	-0.63	-1.87	-3.13	-4.42	-5.73
$w_0 = 1 \operatorname{Cm}$	rate		4.11	4.21	4.29	4.35
$w_{\rm e} = 1 \mathrm{mm}$	$\log_{10} \operatorname{error}$	-0.63	-1.87	-3.13	-4.42	-5.73
$w_0 = 1 \min$	rate		4.11	4.21	4.29	4.35
$w_{\rm e} = 0.1 \mathrm{mm}$	$\log_{10} \operatorname{error}$	-0.63	-1.87	-3.13	-4.42	-5.73
$\omega_0 = 0.1$ IIIII	rate		4.12	4.21	4.29	4.34

TABLE 3

Errors and convergence rates for the MMS verification problem.

	$ ho_s$	G	ν	c_p	c_s
Ω_1	2.4 g/cm^3	10 GPa	0.3	$3800 \mathrm{~m/s}$	2000 m/s
Ω_2	2.4 g/cm^3	20 GPa	0.3	$5400 \mathrm{~m/s}$	2800 m/s
	- ,	ΤA	DIE 4		

Solid material properties for Ω_1 and Ω_2 in the branching crack problem. Compressional and shear wave speeds c_p and c_s have been computed from the density ρ_s , shear modulus G, and Poisson's ratio ν .

segments along a material interface. This geometry can arise in many natural and 569 engineered systems (volcanoes, oil and gas reservoirs, and glaciers) due to hydraulic fracturing of material Ω_1 . Continued influx of fluid causes the crack to grow through 571 Ω_1 until it encounters a stiffer material Ω_2 . The crack then branches by exploiting 572joints (pre-existing fractures) along the material interface. Figure 4 shows the setup. 573 The fluid-filled crack is represented by 5 piecewise smooth and connected segments 574 Γ_i (see Appendix 2 for coupling conditions at the crack junction). The main crack is 5 mm wide and the branches are 1 mm wide at the junction and 0.01 mm wide at the crack tips. The fluid and solid material properties are listed in Tables 1 and 4, respectively. 578

The computational domain is discretized using a multiblock grid (Figure 4). Boundary- and interface-conforming structured grids are generated using cubic Bsplines and transfinite interpolation. The Jacobian and metric coefficients for each grid are computed using the SBP(6,3) first derivative operators. While the Jacobian is smooth inside each block, it is discontinuous across the interfaces.

We use the SBP(6,3) finite difference operators and qualitatively assess grid convergence by performing several levels of grid refinement. To advance the solution in time, we use ARK4 with a time step $\Delta t = 0.7 \times h_{min}/c_{max}$, where $c_{max} = c_p^{(2)}$ and h_{min} is the minimum grid spacing in the solid. On the coarse grid, the minimum grid spacing in the solid is $h_{min} = 1.9$ mm and in the fluid (in the *n* direction) it is $h_{min} = 62.5$ nm. With a fully explicit time stepping scheme we estimate that we would need to reduce the time step by at least two orders of magnitude.

591 Below we present results for the two problems. Both have exactly the same 592 geometry, mesh, material properties, and boundary conditions; they differ only in 593 how waves are excited. In both problems, the fluid and solid are initially at rest, 594 except as indicated.



FIG. 4. Geometry of branching crack problem. A fluid-filled crack Γ cuts through the solid Ω_1 (blue) before branching, at the interface with a stiffer solid Ω_2 (red), into two crack segments along the material interface.





FIG. 5. Snapshots in time of Krauklis waves propagating along a fluid-filled crack (solid line); color shows velocity in x direction. The discontinuity in color indicates opening/closing motions of the crack. 241×241 grid points per block.



(c) Cross section 1.

(d) Cross section 2.

FIG. 6. (a) and (b) Snapshots in time of fluid velocity field inside the main crack and branches. (c) and (d) Velocity profiles for the cross sections marked in (b). Note boundary layers and nonmonotonic profiles, both characteristic of oscillatory flows at high frequency. 241 grid points along each crack segment and 241 grid points across the crack width.

5.2.1. Excitation at the crack mouth. In this first problem, waves are excited 595by specifying a pressure boundary condition p(0,t) = q(t) on Γ_1 (the bottom end of 596 the main crack, referred to below as the crack mouth). Excitation at the crack mouth 597preferentially generates Krauklis waves that propagate along the cracks, ultimately 598leading to resonance at specific frequencies determined by the crack geometry. Crack 599mouth excitation can arise from pressure changes transmitted to the crack by an 600 601 unmodeled narrow conduit or pipe, such as a well in hydraulic fracturing operations in an oil or gas reservoir. For more details on this problem class, see [26]. The 602 boundary data is $g(t) = A \sin(\omega t) \exp(-\eta t)$, where A = 100 kPa, $\omega = 1.2 \times 10^5$ s⁻¹, 603 and $\eta = 100 \text{ s}^{-1}$. This function is a chirp with a maximum frequency $f_{max} \approx 2000$ 604 Hz at 1% of peak amplitude. The maximum frequency f_{max} is used to estimate 605 the minimum wavelength λ_{min} that needs to be resolved in the simulation. The 606607 relationship between wavelength λ and frequency f is determined by the dispersion

relation of the Krauklis waves propagating along the crack. For an infinitely long, planar crack filled with an inviscid fluid, we have [24]

$$\lambda = \left(2\pi \frac{Gw_0}{(1-\nu)\rho_f f^2}\right)^{1/3}$$

We can then estimate $\lambda_{min} \sim \lambda(f_{max}) \sim 0.1$ m, suggesting that λ_{min} will be wellresolved on both the coarse and fine grids. To set the grid spacing in the *n*-direction within the fluid, we estimate the ratio of the boundary layer thickness to crack width as $\sqrt{\mu/(\rho_f \omega)} / \max w_0 \sim 10\%$, which should also be well-resolved on both the coarse and fine grids.

The pressure perturbation applied at the crack mouth excites Krauklis waves 617 propagating along the fluid-filled crack (Figure 5). As Krauklis waves propagate along 618 the crack, the crack walls oscillate inward and outward. A pair of counter-propagating 619 620 waves are formed when the waves are partially reflected at the crack tips and the crack junction. Krauklis waves are attenuated primarily by viscous dissipation in the fluid, 621 which in the wider parts of the crack is confined to boundary layers at the walls of 622 the crack (Figure 6). The coarse and fine grid simulations are visually identical (not 623 shown). 624

5.2.2. Excitation in the solid. This second problem, involving excitation in 625 the solid, demonstrates the potential of our method for studying seismic wave scat-626 tering from fluid-filled cracks. Seismic waves in the solid can be excited by ex-627 plosions or other active sources, or by naturally occurring impulsive perturbations 628 such as small earthquakes (i.e., microseismic events). The latter, when the earth-629 quakes are much smaller than modeled wavelengths, can be treated as point mo-630 ment tensor sources. Details on how to discretize the singular source terms with 631 high-order accuracy can be found in [37]. Here, for simplicity, we excite waves by 632 specifying a Gaussian function as the initial condition in the solid: $v_x(x, y, 0) =$ 633 $\exp\left(-\frac{1}{2a^2}(x-x_*)^2-\frac{1}{2a^2}(y-y_*)^2\right)$ mm/s, where $a=1/\sqrt{200}\approx 7.1$ cm and 634

 $(x_*, y_*) = (-1.5, -4)$ m with the origin located at the bottom end of Γ_1 . All of the other solid and fluid fields are initially zero.



FIG. 7. Snapshots in time of seismic wave scattering from a fluid-filled crack (solid line) and material interface (dashed line). 481×481 grid points per block.

The initial disturbance excites both compressional (P) waves and shear (S) waves that scatter from the fluid-filled crack (Figure 7). P-to-S conversion along the crack



FIG. 8. Grid refinement study at t = 5 ms (grid points per block listed listed in sub-captions).

generates shear head waves, which have curved wavefronts due to the curvature of 639 the crack itself. All waves undergo reflections and additional mode conversations 640 upon reaching the branch segments (and material interface). Diffracted waves from 641 642 the crack junction are also evident. Note that in contrast to the previous problem, Krauklis waves are almost absent. This is because excitation in this problem is from 643 a perturbation to particle velocity approximately normal to the main crack and with 644 symmetry across the crack. Opening or closing motions of the crack are therefore 645 646 negligible.

Figure 8 shows a zoomed-in version of the wavefield at the final time (t = 5 ms)at three different grid resolutions. Dispersion errors that are evident on the coarsest mesh vanish with refinement.

To investigate the computational cost of the fluid model, we compare the per-650 formance of simulations with and without the fluid. In the simulation without the 651 fluid, we remove the fluid blocks and instead couple to the neighboring solid blocks 652 653 directly to one another. We measure the time to solution for both simulations after a fixed number of time steps. The test is conducted on the grid with lowest resolution, 654 655 using 121×121 grid points per block in both the solid and fluid (when present). The computational cost of the fluid is only about 8% of the total cost. This cost is slightly 656less than the ratio of the number of fluid to solid blocks used in this test. 657

658 **6.** Conclusions. We have developed a method to simulate wave propagation 659 in and around cracks containing a viscous, compressible fluid. Our method achieves 660 computational efficiency relative to other commonly used methods through two key 661 components.

First, rather than solving the full linearized Navier-Stokes equations for the fluid, we use a lubrication-type approximation of the fluid response. Viscous effects enter only through one-dimensional diffusion operators in the direction spanning the crack width. Even with this approximation, the semi-discrete system of equations can be quite stiff, such that fully explicit time-stepping methods would require several orders of magnitude smaller time steps than the time step required for explicit integration of the elastic wave equation alone.

669 Second, the computational efficiency is enhanced by partitioning the semi-discrete 670 equations in conjunction with an implicit-explicit Runge-Kutta time-stepping method. 671 Specifically, we treat the elastic wave equation in the solid and the wave propagation 672 part of the fluid equations in a fully explicit manner, whereas the viscous (diffusion) 673 term and fluid-solid coupling terms in the fluid are treated in an implicit manner. By 674 enforcing the coupling conditions using characteristic variables, the overall system of equations can be integrated using the maximum stable time step for wave propagation only. For typical fluid and solid properties, this corresponds to the typical CFL-limited time step used for explicit solution of the elastic wave equation.

Although we developed the numerical scheme in the context of high-order finite differences, the fluid model and many of the results related to the coupling formulation and partitioning should be applicable to other provably stable schemes with weakly enforced coupling conditions in SBP-SAT form (e.g., discontinuous Galerkin methods).

Finally, the method was applied to several application problems involving waves 683 in and around fluid-filled cracks. Excitation at the crack mouth generates large am-684 plitude Krauklis waves, and simulations like the ones shown in this work can be used 685 686 to quantify Krauklis wave resonances and their relation to crack geometry [26]. The method can also be used to study scattering of seismic waves by fluid-filled cracks. 687 Obvious applications include seismic imaging of fractured hydrocarbon-bearing reser-688 voirs, crevasse systems in glaciers and ice sheets, and magmatic dike and sill complexes 689 beneath active volcanoes. 690

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Appendix A. Energy conserving penalty parameters. This appendix continues Section 4.1 with a more detailed investigation of how the stability of the fully discrete approximation is influenced by how the coupling conditions are enforced. This is done by further investigation of the one-dimensional model problem (35), but with a different choice of penalty parameters. Specifically, we take $\alpha_s \to \infty$ and $\beta_s = 0$, for which (35) becomes

$$\int \phi_s \rho_s \frac{\partial v_x}{\partial t} dy = \int \phi_s \frac{\partial \sigma_{xy}}{\partial y} dy + \left[\phi_s \left(\sigma_{xy} - \tau \right) \right]_{y=0^+},$$

$$\int \varphi_s \frac{1}{G} \frac{\partial \sigma_{xy}}{\partial t} dy = \int \varphi_s \frac{\partial v_x}{\partial y} dy$$

$$\int \phi_f \rho_f \frac{\partial u}{\partial t} dn = \int \phi_f \tau dn - \left[\mu \frac{\partial \phi_f}{\partial n} \left(u - v_x \right) \right]_{n=w_0^+}.$$

$$(41)$$

The fluid traction is enforced as a Neumann condition on the solid and the solid velocity is enforced as a Dirichlet condition on the fluid. This way of enforcing the coupling conditions is a very common approach in many fluid-structure interaction schemes [19]. A consequence of using (41) is that there is no additional numerical energy dissipation:

709 (42)
$$\frac{1}{2}\frac{d}{dt}\left(\int \rho_s v_x^2 + \frac{1}{G}\sigma_{xy}^2 dy + \int \rho_f u^2 dn\right) = -\int \frac{\tau^2}{\mu} dn.$$

We next investigate the impact of this coupling procedure on the semi-discrete and fully discrete approximations of (41). We write the semi-discrete approximation of (41) in the strong form:

714 (43)
$$\frac{dq}{dt} = (W+C)q,$$

where $q = [v_x^T \sigma_{xy}^T u^T]^T$. We use the partitioning $dq/dt = (F^{EX} + F^{IM})q$, with $F^{EX} = W^{EX} + C^{EX}$ and $F^{IM} = W^{IM} + C^{IM}$. Consider the following choice of the 716 717 partitioning: 718

$$W^{EX} = \begin{bmatrix} 0 & D_y/\rho_s & 0\\ GD_y & 0 & 0\\ 0 & 0 & 0 \end{bmatrix}, \ W^{IM} = \begin{bmatrix} 0 & 0 & 0\\ 0 & 0 & 0\\ 0 & 0 & \mu/\rho_f D_n D_n \end{bmatrix},$$

$$719 \quad (44) \qquad C^{EX} = \begin{bmatrix} 0 & H_y^{-1}L_s L_s^T & -\mu H_y^{-1}L_s L_f^T D_n\\ 0 & 0 & 0\\ 0 & 0 & 0 \end{bmatrix},$$

$$C^{IM} = \frac{\mu}{\rho_f} \begin{bmatrix} 0 & 0 & 0\\ 0 & 0 & 0\\ H_n^{-1} D_n^T L_f L_s^T & 0 & -H_n^{-1} D_n^T L_f L_f^T \end{bmatrix}.$$

$$720$$

7

721 For simplicity, the grid spacing is uniform, implying that there are no metric coeffi-

cients. This partitioning is the same as the one we used before; see (32). The solid is 722 fully explicit and the fluid is fully implicit (because in this one-dimensional problem 723

there is no wave propagation in the fluid). 724



FIG. 9. Spectral radii of the semi-discrete approximation of the problem (41). The spectral radii $\rho(W+C)$ (complete problem) and $\rho(F^{EX})$ (explicit part of the semi-discrete approximation) are shown.

Next, we compute the spectral radius of the semi-discrete approximation while 725varying $\Delta x_f / \Delta x_s$. As in Section 4.1, we use $\rho_s = \rho_f$ and discretize using the SBP(6,3) 726 operators. However, for this new choice of penalty parameters, Figure 9 shows that 727 the spectral radius $\rho(W+C)$ is at about one order of magnitude larger than the 728 spectral radius $\rho(F^{EX})$ (which is determined by the explicit part of the semi-discrete 729 approximation, i.e., by wave propagation). Note that the spectral radius of the explicit 730 part $\rho(F^{EX})$ does not change as $\Delta x_f / \Delta x_s$ is varied, indicating that there is no source 731



FIG. 10. Maximum stable CFL number of the fully discrete scheme approximating the problem (41) with partitioning (44).

of stiffness in the explicit part. However, when we look at the maximum stable time 732step of the fully discrete approximation, we find that it is necessary to decrease the 733time step for stability (Figure 10). The important lesson here is that this choice of the 734 penalty parameters, when used in combination with the partitioning (44), causes the 735scheme to be unstable for sufficiently large $\Delta x_f / \Delta x_s$, unless the time step is decreased. 736 In contrast, the scheme is stable for the penalty parameter choice presented in Section 737 4.1 regardless of the value of $\Delta x_f / \Delta x_s$. 738

Another way to stabilize the scheme (41), without changing the penalty parame-739ters, is to modify the partitioning. This modification should be done such that energy 740 is conserved in the fully discrete approximation as well. Consider the following energy 741 conservative partitioning: 742

743 (45)
$$C^{EX} = \begin{bmatrix} 0 & H \\ 0 \end{bmatrix}$$

743 (45)
$$C^{EX} = \begin{bmatrix} 0 & H_y^{-1}L_sL_s^T & 0\\ 0 & 0 & 0\\ 0 & 0 & 0 \end{bmatrix},$$

744
$$C^{IM} = \begin{bmatrix} 0 & 0 & -\mu/\rho_sH_y^{-1}L_sL_f^TD_r\\ 0 & 0 & 0\\ \mu/\rho_fH_n^{-1}D_n^TL_fL_s^T & 0 & -\mu/\rho_fH_n^{-1}D_n^TL_fL_f^T \end{bmatrix}$$

In this case, one can easily show that each sub-problem satisfies an energy balance, implying $\langle q, F^{EX}q \rangle \leq 0$ and $\langle q, F^{IM}q \rangle \leq 0$, where $\langle u, v \rangle = \sum u_j v_j h$ is the discrete inner-product, i.e., F^{EX} and F^{IM} are both semi-bounded. We again obtain a fully 746 747 748 discrete scheme with the attractive property that the maximum stable time step is set 749 by wave propagation. In practice, however, we do not use this partitioning because 750751 the solid penalty terms are treated implicitly, making it more difficult to implement.

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2. Crack junction coupling conditions. Consider the coupling of three cracks
 at a junction. Prior to weakly enforcing the coupling conditions, the work rate at the
 junction is

884
885

$$\frac{dE}{dt} = -\sum_{i=1}^{3} n^{(i)} w_{0}^{(i)} \hat{p}^{(i)} \hat{\bar{u}}^{(i)} - \mathcal{R}$$

where (i) labels each crack. We use $n^{(i)}$ to keep track of the sign at each end $(n^{(i)} = -1$ and $n^{(i)} = 1$ for the left and right end, respectively). While the crack can be in any direction, the left end is defined at the minimum arc length s along the crack. At the junction, the pressure is continuous: $\hat{p} = \hat{p}^{(1)} = \hat{p}^{(2)} = \hat{p}^{(3)}$. Mass conservation, in the context of our linearized model, requires

891 (46)
$$\sum_{i=1}^{3} n^{(i)} \rho_{f}^{(i)} w_{0}^{(i)} \hat{u}^{(i)} = 0$$

893 To ensure $\mathcal{R} \geq 0$, we need

$$893 \quad (47) \qquad \qquad \alpha^{(i)} n^{(i)} w_0^{(i)} \hat{u}^{(i)} + \hat{p} = \alpha^{(i)} n^{(i)} w_0^{(i)} \bar{v}^{(i)} + p^{(i)}, \text{ for } i = 1, 2, 3.$$

896 Multiplying (47) by $\rho_f^{(i)} \alpha^{(i+1)} \alpha^{(i+2)}$, and cyclically summing over i = 1, 2, 3 results in

897
898
$$\sum_{j=1}^{3} \alpha^{(j)} \sum_{i=1}^{3} n^{(i)} \rho_{f}^{(i)} w_{0}^{(i)} \hat{\bar{u}}^{(i)} + \sum_{i=1}^{3} \rho_{f}^{(i)} w_{0}^{(i)} \sum_{j \neq i} \alpha^{(j)} \hat{p} = \zeta.$$

899 where $\zeta = \sum_{i=1}^{3} \rho_{f}^{(i)} w_{0}^{(i)} p^{(i)} \sum_{j \neq i} \alpha^{(j)} + \sum_{i=1}^{3} w_{0}^{(i)} n^{(i)} v^{(i)} \sum_{j=1}^{3} \alpha^{(j)}$. Finally, using 900 (46) results in

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$$\hat{p} = \frac{\zeta}{\sum_{i=1}^{3} \rho_{f}^{(i)} w_{0}^{(i)} \sum_{j \neq i} \alpha^{(j)}}, \ \hat{\bar{u}}^{(i)} = \bar{\bar{u}}^{(i)} + \frac{p^{(i)} - \hat{p}}{\alpha^{(i)} n^{(i)} w_{0}^{(i)}}.$$