Particle velocity based universal algorithm for numerical simulation of hydraulic fractures

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Abstract

In the paper, we propose a new effective mathematical formulation and resulting universal numerical algorithm capable of tackling various HF models in the framework of a unified approach. The presented numerical scheme is not limited to any particular elasticity model or crack propagation regime. Its basic assumptions are: i) proper choice of independent and dependent variables (with the direct utilization of a new one - the reduced particle velocity), ii) tracing the fracture front by use of the Stefan condition (speed equation), which can be integrated in a closed form and sets an explicit relation between the crack propagation speed and the coefficients in the asymptotic expansion of the crack opening, iii) proper regularization techniques, iv) improved temporal approximation, v) modular algorithm architecture. The application of the new dependent variable, the reduced particle velocity, instead of the usual fluid flow rate, facilitates the computation of the crack propagation speed from the local relation based on the speed equation. In this way, we avoid numerical evaluation of the undetermined limit of the product of fracture aperture and pressure gradient at the crack tip (or alternatively the limit resulting from ratio of the fluid flow rate and the crack opening), which always poses a considerable computational challenge. As a result, the position of the crack front is accurately determined from an explicit formula derived from the speed equation. The underlying ideas employed in the algorithm are combined together producing a robust and efficient numerical scheme. Its performance is demonstrated using classical examples of 1D models for hydraulic fracturing: PKN and KGD under various fracture propagation regimes. Solution accuracy is verified against dedicated analytical benchmarks and other solutions available in the literature. Most of the ideas developed here, can be directly extended to more general 2D and 3D cases.

1 Introduction

Hydraulic fracture is a process of a crack propagating in a brittle material, as a result of pressurized liquid injection. It can be observed in many natural phenomena, like magma driven dykes [60], subglacial drainage of water [66] and others [53, 58]. Recently it has been associated mostly with the method of reservoir stimulation used in the oil and gas industry to maximize hydrocarbon extraction. Although this technology can be backdated to the 1930s [28], it has been in the last twenty years that hydrofracturing has become commonplace, for both economic reasons and the technological development itself. Without any doubt, hydrofracturing has revolutionized the exploitation of shale oil and gas and has become a key technology when it comes to extraction from non-conventional reservoirs.
Mathematical modeling of this multiphysics process is a challenging task. The main difficulties associated with this problem are: (a) strong non-linearities resulting from the interaction between the solid and fluid phases, (b) singularities in the physical fields near the fracture front, (c) moving boundaries, (d) degeneration of the governing equations in the near-tip region, (e) pronounced multiscale effects and others. The complexity of the problem enforces many simplifications when trying to conduct the analysis. The first mathematical models were proposed in 1940s and 1950s \[14, 29, 30, 31, 62\]. These studies together with the later works led to the formulation of the basic 1D models of hydraulic fractures: i) the PKN model \[55, 57\], ii) the KGD model \[27, 34\], iii) the radial or penny shaped model \[63\]. Such simplified variants of the problem were used for many years to design the treatments. However, in the 1980s with the increasing number and sizes of treatments, the necessity for more advanced and accurate modeling emerged. The so called pseudo 3D models (P3D) appeared \[19\]. They approximate the behaviour of the planar 3D fractures, including those in the stratified reservoirs, with minimal computational costs. Another attempt at advancing the mathematical modeling of hydraulic fractures was the introduction of the planar 3D models (PL3D) \[4, 13, 67\], in which case the crack footprint and the internal fluid flow are described by the 2D mesh of cells and combined with the full 3D elasticity equations. In this way the fracture aperture is determined as a function of the fluid pressure. In the recent years there have also been attempts to develop full 3D models utilizing various numerical techniques, e.g. finite element method \[12, 62, 40, 68\], boundary element method (or combination of the two mentioned \[11, 70\]), the discrete element methods \[15\] or other techniques \[37, 39\]. A broad review of the topic can be found in \[3\] where it is has been shown that, although immense progress has been made since the pioneering works in the area, there is still a demand for further improvements in efficiency and credibility of computations.

Alongside the development of more sophisticated mathematical models of hydraulic fractures, fundamental research aimed at identifying the basic features inherent to any solution have been carried out. In this way, a deeper insight into the classical models resulted in the formulation of the basic principles for constructing a solution. Then, feedback into numerical simulations enabled an increase in the credibility and efficiency of computations. Special attention has been paid to the tip behaviour of the solution. The early works introducing the correct tip asymptotics can be backdated to the 1980s \(64\) - for the KGD model, \(63\) - for the PKN model. More comprehensive studies on this problem were presented in \[16\] for the zero toughness impermeable case, in \[12\] for the zero toughness leak-off dominated variant. In \[9\], the near tip process in an impermeable elastic medium for the plane strain conditions was enriched by the presence of a lag between the fracture tip and the fluid front. The authors of \[61\] have proposed asymptotic solutions in the case of a penny shaped fracture driven by the Newtonian fluid for both small and large toughness values. The analysis for plain strain and penny shaped fractures propagating in the toughness dominated regime in permeable rock was delivered in \[7\], giving the early and large time asymptotes. Results pertaining to the plain strain fracture driven by the shear-thinning fluids can be found in \[22\].

Simultaneously, recognition of the importance of the near-tip behaviour of solutions has led to classification of the basic fracture propagation modes \[18, 25, 26\]. They have been categorized in the parametric space, confined by four limiting physical regimes: i) leak-off dominated, ii) storage dominated, iii) toughness dominated, iv) viscosity dominated. The hydraulic fracture is considered to evolve in time between these specific modes depending on the injection rate, the rock and fracturing fluid properties. A number of semi-analytical and numerical solutions have been constructed for such asymptotic regimes. In particular, the case of zero toughness impermeable rock was analyzed in \[2, 61\], small toughness zero-leak-off variant in \[21\], large toughness impermeable in \[23\], and finite toughness permeable in \[52\].

All these efforts have underlined the importance of the multiscale character of the problem. It is now well understood that the coupling between non-linear, non-local and history dependent physical fields results in a complex solution structure, where respective physical processes manifest themselves with different intensity at various temporal and spatial scales. It has been proved that the global behaviour of a fluid driven fracture depends critically on features of the local solution in the near-tip region,
which entails respective consequences for the numerical implementation. Furthermore, the nature of the moving boundary, resulting in degeneration of the governing equations and the boundary conditions at the crack tip, make tracing the fracture front an extremely difficult task [56]. All these factors clearly indicate the challenge in understanding the solution structure (especially the tip asymptotics) and its appropriate application in the computational schemes. In the recent studies by Lecampion et al [47] it has been shown that the algorithms which use the appropriate multi-scale hydraulic fracture asymptote in the near tip region provide much better results than those which do not apply it. Moreover, when accounting for the proper tip asymptotics, very good results can be obtained even for coarse meshing.

The analysis given in [38, 45, 51, 69] proves that proper mathematical formulation of the problem of hydraulic fracture facilitates the correct introduction of the basic asymptotic features of the solution to the numerical algorithm. This in turn results in an appreciable improvement to the accuracy and efficiency of computations. A key role in properly formulating the problem is the choice of independent (spatial and temporal meshing) and dependent variables. These modifications enable one to rigorously apply the solution tip asymptotics, and simultaneously to make use of dedicated regularization techniques like the so called \( \varepsilon \)-regularization (see [38, 46]) or operator regularization [69].

In this paper we propose a new effective mathematical approach, from which the resulting universal numerical algorithm is capable of tackling various HF models within the framework of a unified approach. Its basic assumptions are: i) proper choice of independent and dependent variables (with a direct utilization of a new one - the reduced particle velocity), ii) tracing the fracture front by use of the Stefan condition (speed equation) which can be integrated in a closed form to give an explicit relation between the crack propagation speed and the leading coefficients of the crack opening asymptotics, iii) proper regularization techniques, iv) improved numerical approximation of the temporal derivative of the solution, v) modular algorithm architecture.

The application of the new dependent variable, the reduced particle velocity, instead of the usual fluid flow rate, facilitates the computation of the crack propagation speed from the local relation based on the speed equation. In this way we avoid numerical evaluation of the undetermined limit of the product of fracture aperture and pressure gradient at the crack tip (or alternatively the limit resulting from division of the fluid flow rate by the crack opening), which always poses a considerable computational challenge. As a result the position of the crack front can be accurately determined from an explicit formula derived from the speed equation (see e.g [40, 51]), being an evolution of the Stefan condition. With regards to the numerical modeling of hydraulic fractures, this condition was originally introduced by Kemp [33] but later abandoned. Recently, it has been rediscovered by Linkov [44, 45]. Information about the solution tip asymptotics is utilized rigorously in the numerical scheme together with appropriate regularization techniques. Some elements of the employed scheme has already been presented in [38, 51, 69], where a broad discussion on the advantages of application of proper dependent variables and regularization techniques can be found. One of the key points of the developed universal algorithm is utilization of the explicit relation between the crack propagation speed and the leading terms of the crack opening asymptotic expansion in the form (51) – (54). For the PKN model it was found and described in [38, 69], while for the KGD formulation respective results have been recently reported at a number of conferences.

All these devices, accordingly combined together, produce a robust and efficient numerical scheme. Its performance is demonstrated against examples of classical 1D models for hydraulic fracture: PKN and KGD under various fracture propagation regimes. The solution accuracy is verified against dedicated analytical benchmarks and other solutions available in the literature. Most of the ideas developed here, can be directly extended to more general 2D and 3D cases.

The paper is organized as follows. In Section 2 a general mathematical description of the problem is given. Moreover the basic idea of fracture front tracing is explained, also the motivation for and
advantages of the applied approach are presented. We introduce here a dimensionless formulation of the problem, which is henceforth used. Section 3 contains a detailed characterization of the solution tip asymptotics, and its link to the mechanism of crack tip tracing. A universal mathematical description is proposed for different elasticity models and crack propagation regimes. A complete definition of the mechanism of crack front tracing based on the speed equation (Stefan condition) is given. In section 4, a new dependent variable, the reduced particle velocity, is introduced. The governing equations are reformulated in terms of the new variable. Section 5 describes the reduction of the problem to a self-similar version for two different time-dependent functions: the power function and the exponential one. Then the self-similar variant of the universal algorithm is presented. The performance of the algorithm is verified against dedicated analytical benchmarks (detailed description in Appendix) as well as the reference solutions available in the literature. Fully analytical benchmarks for the KGD variant of the problem are introduced here for the first time. In Section 6, the idea of a universal solver is adapted to the transient case where the improved approximation of the temporal derivative is one of the key elements. Extensive accuracy analysis is given. Final discussions and conclusions are presented in Section 7.

2 Problem formulation

2.1 Basic equations for hydrofracturing

Let us consider a rectilinear symmetrical crack of the length $2l\ x \in [-l, l]$. The crack is fully filled by a Newtonian fluid injected at the middle point ($x = 0$) with a known rate $q_0(t)$. As a result, the crack front ($x = \pm l$) moves and the crack length, $l(l(t))$, is a function of time. Below we present a classic set of the governing equations for the 1D formulation of the problem, which can be found in e.g. [2, 20, 35] for various elasticity models. As usual, due to symmetry of the problem, we restrict our analysis to one of the symmetrical parts of the crack $x \in [0, l(t)]$.

The mass conservation principle is expressed by the continuity equation:

$$\frac{\partial w}{\partial t} + \frac{\partial q}{\partial x} + q_l = 0, \quad t > 0, \quad 0 < x < l(t),$$

(1)

The fluid flow inside the fracture is described by the Poiseuille equation which, in case of Newtonian fluid, is written in the following form:

$$q = -\frac{1}{M} w^3 \frac{\partial p}{\partial x}, \quad t > 0, \quad 0 < x < l(t),$$

(2)

In the above equations $w = w(t, x)$ stands for the crack opening, $q = q(t, x)$ is the fluid flow rate, $p = p(t, x)$ ($p = p_f - \sigma_0$, $\sigma_0$ - confining stress) refers to the net fluid pressure. Constant $M$ involved in the Poiseuille equation is computed as $M = 12\mu$, where $\mu$ denotes the dynamic viscosity. Function $q_l = q_l(t, x)$, in the right-hand side of the continuity equation (1), is the volumetric rate of fluid loss due to formation in the direction perpendicular to the crack surfaces per unit length of the fracture. Usually it is assumed (local formulation) to be given [38, 55, 50]. More accurate analysis involves a nonlocal formulation where the mass transfer in the entire external domain should be taken into account [36]. We will comment on the possible behaviour of $q_l$ later on.

The group of fluid equations is to be supplemented by the relation describing the deformation of rock under applied hydraulic pressure. Thus, the net pressure in the fracture, is given by the relationship:

$$p(x) = \mathcal{A} w(x), \quad 0 < x < l(t),$$

(3)

where operator $\mathcal{A}$ refers to the chosen model of elasticity (related to the predefined fracture geometry). In this paper we consider the two most popular linear models (local and nonlocal, respectively):
• the PKN model \[55\]
\[ A_1w = k_1w, \] (4)

• the KGD model \[63\]
\[ A_2w = k_2 \int_0^{l(t)} \frac{\partial w(t,s)}{\partial s} \frac{s}{x^2 - s^2} ds. \] (5)

For the classical PKN model constant \(k_1\) can be found in \[55\] for an elliptical crack of height \(h\), while \(E\) and \(\nu\) are the elasticity modulus and Poisson’s ratio, respectively.\footnote{The multiplier \(k_1\) may depend on \(x\) and/or \(w\) as well: \(k_1 = k_1(x,w)\), constituting the so-called pseudo 3D model (P3D) \[49, 47\]. This case can also be considered in the framework of the presented approach.} The multiplier \(k_2\) in the KGD model follows, for example, from \[54, 63\]:
\[ k_1 = \frac{2}{\pi h} \frac{E}{1 - \nu}, \quad k_2 = \frac{E}{2\pi(1 - \nu^2)}. \] (6)

The inverse operators for (4) – (5) are:
\[ A_1^{-1}p = \frac{1}{k_1}p, \] (7)
\[ A_2^{-1}p = \frac{2}{\pi^2 k_2} \int_0^{l(t)} p(t,s) \ln \frac{\sqrt{l^2(t) - x^2 + \sqrt{l^2(t) - s^2}}}{\sqrt{l^2(t) - x^2 - \sqrt{l^2(t) - s^2}}} ds. \] (8)

Note that the original Cauchy type singular integral in the elasticity equation \[5\] is defined (compare \[63\]) over the entire crack, \(-l(t) < x < l(t)\), and that representation \[5\] is valid only under the assumption
\[ \frac{\partial w}{\partial x}(t,0) = 0. \] (9)

On the other hand, the form of the integral operator \(A_2^{-1}\) from \[8\] guarantees this property if the net pressure is smooth enough (differentiable) near the zero point and the following condition is enforced:
\[ \int_0^{l(t)} \frac{p(t,s)ds}{\sqrt{l^2(t) - s^2}} < \infty, \] (10)
which can be easily checked by direct differentiation. This condition has a clear physical sense. It states that the Stress Intensity Factor (SIF) defined by the integral in \[10\] is finite. Thus, when equation \[8\] instead of \[5\] is utilized in the analysis condition, \(9\) is satisfied automatically. Moreover, using condition \[10\] one can also prove that:
\[ \frac{\partial w}{\partial x}(t,0) = O(x \log x), \quad x \to 0. \] (11)

The foregoing equations should be supplemented with the initial and boundary conditions. Thus, the influx boundary condition and two tip boundary conditions at the crack tip are:
\[ q(t,0) = q_0(t), \] (12)
\[ w(t,l(t)) = 0, \quad q(t,l(t)) = 0. \] (13)

We accept in this paper non-zero initial conditions:
\[ l(0) = l_0, \quad w(0,x) = w_0(x), \quad x \in (0,l_0). \] (14)
Usually, the uniform initial conditions are suggested instead (e.g. [55]):
\[ w(x,0) = 0, \quad l(0) = 0. \] (15)

However, for time dependent problems, the authors tend to replace them with condition (14), where \( l_0 = l_{sf}(t_0) \) and \( w_0(x) = w_{sf}(x,t_0) \) for \( x \leq l_0 \) are taken from the corresponding self-similar solution which neglects the leak-off to the formation (early time asymptote). Note that (14) in its general form accounts for a preexisting hydraulic fracture.

In the case when fracture evolution is analyzed in the framework of Linear Elastic Fracture Mechanics (LEFM), that is for the toughness driven regime for the KGD model, the following propagation condition is imposed:
\[ K_I(t) = K_{IC}, \] (16)
where \( K_{IC} \) is the material toughness [60] while \( K_I \) is the already mentioned SIF computed by the following formula [54] (compare (10)):
\[ K_I = 2\sqrt{\frac{l(t)}{\pi}} \int_0^{l(t)} \frac{p(t,s)ds}{\sqrt{l^2(t) - s^2}}, \] (17)

Note that the tip asymptote for the crack opening is defined in this case as:
\[ w(t,x) \sim \frac{8(1-\nu^2)}{\sqrt{2\pi}} \frac{K_I}{E}\sqrt{l(t)-x}, \quad x \to l(t). \] (18)

Finally, the global fluid balance equation takes the form:
\[ \int_0^{l(t)} [w(t,x) - w_0(x)]dx - \int_0^t q_0(t)dt + \int_0^{l(t)} \int_0^t q_l(t,x)dtdx = 0. \] (19)

Which is usually used to determine the crack length (see e.g. [3]).

The above set of equations and the respective conditions constitute the classical model of hydraulic fracture [3, 46].

In our analysis we will utilize another dependent variable, the average fluid velocity through the fracture cross-sections (called also a particle velocity), \( v \), defined as follows:
\[ v(t,x) = \frac{q(t,x)}{w(t,x)} = -\frac{1}{M} w^2 \frac{\partial p}{\partial x}, \quad 0 < x < l(t). \] (20)

This variable has been frequently mentioned in publications (see for example [26, 23]), but not used directly in computational algorithms. In [49, 46] it was suggested to incorporate it as a dependent variable instead of the net pressure or the fluid flux to improve the algorithm performance. In our analysis we are going to confirm this fact.

Throughout this paper we assume that there are no flow stagnation or inversion points along the fracture, which means that \( v(t,x) \) should be finite and positive.
\[ 0 \leq v(t,x) < \infty, \quad t > 0, \quad 0 < x < l(t). \] (21)

As a result, one can conclude, that the spatial derivative of the net pressure is negative along the whole fracture
\[ \frac{\partial p}{\partial x} < 0, \quad t > 0, \quad 0 < x < l(t), \] (22)

Moreover, taking into account the fact that the crack opening vanishes at the crack tip (compare [13,1]), one can conclude that
\[ \frac{\partial p}{\partial x} \to -\infty, \quad x \to l(t). \] (23)
2.2 Description of the crack front movement

Tracing the fracture front evolution is one of the major challenges when solving the nonlinear problem of hydraulic fracture. In the recent paper by Detournay & Peirce, [19], various approaches to this task have been discussed (we refer a prospective reader to this publication). In the numerical scheme proposed in sections 5-6 below, we will use a strategy for finding the fracture tip different from that advocated in [19]. Indeed, in the recalled paper there is a paragraph devoted to our approach, but is misleading rather than helpful. For this reason and in order to eliminate any further confusions, we would like to clearly identify the key assumptions and steps of our approach.

First let us underline that the standard methods of simulating hydrofracturing mostly employ the crack opening, \( w(t, x) \), and the fluid flow rate, \( q(t, x) \) as the dependent variables. It allows one to directly account for respective boundary conditions at the fracture tip (compare (13)), and has a few other benefits as discussed in \[3\]. However, when determining the crack tip position, this approach causes serious difficulties, which was comprehensively analysed in [19]. In particular, the authors pointed out the major computational problem of evaluation of the fluid front velocity \( x^* = x^* \in \partial \Omega \) from the equation

\[
v^*(t, x^*) = \lim_{x \to x^*} \frac{q(t, x) - w(t, x)}{}
\]

where both dependent variables vanish at the crack tip.

Our approach utilizes another pair of dependent variables: i) the standard one - the crack opening \( w(t, x) \), and ii) the particle velocity (the average through a channel cross section fluid flow velocity), \( v(t, x) \), instead of the fluid flux, \( q(t, x) \). The next crucial assumption used in our paper is a condition that

- **the particle velocity takes a finite value at the crack front**.

This assumption is valid for most hydraulic fracture models, with some reservations in the cases of severe leak-off regimes (e.g. fluid driven regime for the KGD model combined with a Carter law - see e.g. [52]). Note that the assumption has a clear physical motivation as all the basic equations were derived neglecting the inertia effects. This state of affairs has been also confirmed in many experiments [59, 6, 24, 8].

The first consequence of the chosen set of dependent variables is that, when the first of the conditions (13) is satisfied, the second one is fulfilled automatically. Moreover, when solving the problem in such a formulation, one eliminates completely the difficulties mentioned in [19], as well as those related to computing of the crack propagation speed from relation (24).

Naturally, this new approach requires reformulation of all the governing equations in terms of the new dependent variables \( w(t, x) \) and \( v(t, x) \). Then stability and cost effectiveness of the new system is to be confirmed, which is one of the goals of our paper. However, to make this possible, one needs also to answer a crucial question:

- **what is the relationship between the crack propagation speed, \( v_{cr} \), and the finite value of the fluid front velocity, \( v^* \).**

Note that this question is not specific to our approach. It arises in any formulation of the hydraulic fracture problem (what is not always, however, clearly highlighted). As stated in [19], the case where the fluid front coincides with the fracture tip is much more challenging from a computational point of view. In the class of problems with moving boundaries such a condition is usually called the **Stefan condition** [65, 43, 19]. In mathematical modeling of hydraulic fractures it was probably Kemp [33] who first explicitly wrote and directly used this condition in computations. The Stefan condition is usually employed in the analysis of hydrofracturing in its implicit form as a compatibility condition (e.g. [21], [22]) and has been defined in an explicit way for the steady state problems [26], [17].
Kemp’s condition has recently been rediscovered by Linkov [44] and named the speed equation. In our analysis we shall use both names interchangeably. The recalled condition has the following form in the 1D formulation:

\[ \frac{dl}{dt} = v_*(t) \equiv v(t, l(t)). \]  

(25)

Note that the left-hand side of (25) describes the speed of the fracture tip, while the right-hand side refers to the fluid front velocity. Equation (25) is valid under the assumption that the fluid front coincides with the fracture tip. This implies that there is no lag between them. Also the invasive zone ahead of the crack should be excluded from consideration. Note that both of the recalled cases are also tractable (accountable for) within the framework of the presented approach, if the speed equation (25) is supplemented with an additional term which takes into account the respective phenomenon manifesting itself in the near tip region.

As an additional argument to justify the applicability of the speed equation in form (25), one can examine a possible behaviour of the leak-off function \( q_l \) near the crack tip. Note that \( q_l \) in (1) describes the rate of fluid flow from the fracture into the surrounding formation, and generally speaking is not known in advance. To describe this phenomenon properly, one needs to formulate a complete coupled multiphysics problem linking the processes within the fracture to those appearing in the rock formation, where the latter depends on various external conditions (geometry, porosity, permeability, fluid concentration and others). The respective nonlocal formulation complicates enormously the problem under consideration. The usual way to overcome this difficulty is to predefine the function \( q_l = q_l(t, x, w, p) \) (which anyway can be dependent on the processes within the fracture through the crack opening, the net pressure and the properties of the adherent rock). Only in the case of impermeable rock does the problem of hydraulic fracture simplify in this respect and one can put \( q_l = 0 \).

In the local empirical formulation, the leak-off function is assumed to be known and dependent on the solution. We accept in this paper the following behaviour of \( q_l \) near the crack tip:

\[ q_l(t, x) \sim Q(t)(l(t) - x)\eta, \quad \text{as} \quad x \to l(t), \]  

(26)

for some constant \( \eta \geq -1/2 \). Note that \( \eta = -1/2 \) corresponds to the classic empirical Carter law [10].

It is well known (see for example [26, 38, 52, 2, 21, 35]) that the leading term of the crack opening asymptotic expansion in the near-tip region can be expressed as:

\[ w(t, x) \sim w_0(t)(l(t) - x)^\alpha, \quad \text{as} \quad x \to l(t), \]  

(27)

and it does not depend on the value of \( \eta \). Here, the constant \( \alpha \) depends only on the particular elasticity model for the problem and the crack propagation regime (52, 38) (see Table 1). However, further terms of the asymptotics of the crack opening and other dependent variables depend essentially on the behaviour of the leak-off function near the crack tip and play a crucial role in the analysis as they determine the smoothness of the particle velocity function in the vicinity of the fracture front.

Let us then discuss the possible behavior of the function \( q_l \), taking into account one of the basic assumptions used when deriving the lubrication equation. Namely, it is accepted that the fluid flow inside a thin channel is predominantly directed along the channel walls. This, among others, implies that

\[ q_l(t, x) \ll q'_l(t, x), \quad t > 0, \quad 0 < x < l(t), \]  

(28)

at any point along the crack surfaces including the fracture tip. On the other hand, the fluid flux near the crack tip behaves similarly to the that of the crack opening (since the particle velocity is finite at the crack tip). As a result, validity of the following condition should be accepted:

\[ 1 + \eta \geq \alpha. \]  

(29)

\[ \text{This fact was noted by Prof. A. Linkov in a private correspondence with GM.} \]
Throughout this paper we assume that the condition (29) holds true in its stronger variant:

\[ 1 + \eta > \alpha. \] (30)

Note that the condition (30) is equivalent to the speed equation in the form (25), as it follows immediately from the continuity equation (1).

The assumption (30) fails when one considers the toughness driven KGD model (\( \alpha = 1/2 \)) with the classical Carter law (\( \eta = -1/2 \)). In this particular case, an additional term \(-2Q(t)w^{-1}_0(t)\) should be introduced in the right-hand side of (25) to guarantee its validity. Here \( Q(t) \) is a known constant or a functional defined on the solution \( Q(t) = F[w,v](t) \). However, in the case of the fluid driven KGD model (\( \eta = -1/2, \alpha = 2/3 \)), failure to comply with condition (29) results in an infinite value of the crack propagation speed \( 4 \).

The authors believe that the empirical Carter law loses its physical sense at the crack tip. We have proved this earlier in [38] for the PKN model. It turned out that a perturbation of the law in a very small region near the fracture front (at a distance less than \( 10^{-4} \) of the crack length) produces a change in the fracture length, which amounts to a few percent (in other words a change in the law over a distance less than 1 mm would result in the deviation of the crack length greater than 1 m).

In the following we restrict ourselves to the case (26) with assumption (30), and consequently with the speed equation in the form (25). The latter will be used throughout this paper to trace the fracture front. On the other hand, the speed equation serves as the boundary condition at the crack tip for the dependent variable, which is now the particle velocity, \( v \), not the fluid flux as in the standard approach.

The advantages of such an approach have already been shown in [44, 45, 46, 51, 38].

2.3 Problem normalization

Let us normalize the problem by introducing the following dimensionless variables:

\[
\tilde{x} = \frac{x}{l(t)}, \quad \tilde{t} = \frac{t}{t_n}, \quad \tilde{w}(\tilde{t}, \tilde{x}) = \frac{w(t,x)}{l_*}, \quad \tilde{L}(\tilde{t}) = \frac{l(t)}{l_*}, \quad \tilde{q}(\tilde{t}, \tilde{x}) = \frac{t_n}{l_*}q(t,x),
\]

where \( \tilde{x} \in [0, 1] \), and parameter \( m \) takes value either 1 or 2 for the PKN and KGD model, respectively. The value of \( m - 1 \) coincides with the order of homogeneity of the operator \( A_m \). Note that the normalization (31) is not attributed to any particular influx regime, elasticity model or asymptotic behaviour of the solution.

In the normalized variables, the continuity equation (1) takes the form:

\[
\frac{\partial \tilde{w}}{\partial \tilde{t}} - \frac{L'}{L} \frac{\partial \tilde{w}}{\partial \tilde{x}} + \frac{1}{L} \frac{\partial (\tilde{w} \tilde{v})}{\partial \tilde{x}} + \tilde{q} = 0,
\] (32)

where the fluid flow rate was replaced by the product of the crack opening and the particle velocity according to (20).

The normalized particle velocity yields:

\[
\tilde{v} = \frac{\tilde{q}}{\tilde{w}} = -\frac{1}{L} \tilde{w} \tilde{\alpha} \frac{\partial \tilde{p}}{\partial \tilde{x}},
\] (33)

and the speed equation (25) transforms now to:

\[
\frac{dL}{d\tilde{t}} = -\frac{1}{L(\tilde{t})} \left[ \tilde{w} \tilde{\alpha} \frac{\partial \tilde{A} \tilde{w}}{\partial \tilde{x}} \right]_{\tilde{x}=1}.
\] (34)

\footnote{In this case, many authors tend to accept the validity of the Carter law only away from the fracture tip - see e.g. [52].}
The normalized elasticity equation (3) takes the form:

\[ \tilde{p} = \tilde{A}\tilde{w}, \]  

(35)

where an identity operator,

\[ \tilde{A}_1 = \tilde{A}_1^{-1} = I, \]  

(36)
corresponds to the PKN model, while the integral operators for the KGD model are

\[ \tilde{A}_2\tilde{w} = -\frac{1}{L(t)} \int_0^1 \frac{\partial \tilde{w}(\tilde{t}, \eta)}{\partial \eta} \frac{\eta}{\eta^2 - \tilde{x}^2} d\eta, \]  

(37)

and

\[ \tilde{w} = \tilde{A}_2^{-1}\tilde{p} = \frac{2}{\pi^2} L(\tilde{t}) \int_0^1 \tilde{p}(\tilde{t}, \eta) \ln \frac{\sqrt{1 - \tilde{x}^2} + \sqrt{1 - \eta^2}}{\sqrt{1 - \tilde{x}^2} - \sqrt{1 - \eta^2}} d\eta, \]  

(38)

for equations (5) and (8), respectively. In our computations we shall use an alternative form of (38) obtained using integration by parts:

\[ \tilde{w} = -\frac{2}{\pi^2} L(\tilde{t}) \int_0^1 \frac{\partial \tilde{p}(\tilde{t}, \eta)}{\partial \eta} K(\eta, \tilde{x}) d\eta + \frac{4L(\tilde{t})}{\sqrt{\pi}} \tilde{K}_I \sqrt{1 - \tilde{x}^2}, \]  

(39)

where the kernel \( K(\eta, \tilde{x}) \) is:

\[ K(\eta, \tilde{x}) = (\eta - \tilde{x}) \ln \frac{\sqrt{1 - \tilde{x}^2} + \sqrt{1 - \eta^2}}{\sqrt{1 - \tilde{x}^2} - \sqrt{1 - \eta^2}} + \tilde{x} \ln \left( \frac{1 + \eta \tilde{x} + \sqrt{1 - \tilde{x}^2} \sqrt{1 - \eta^2}}{1 + \eta \tilde{x} - \sqrt{1 - \tilde{x}^2} \sqrt{1 - \eta^2}} \right), \]  

(40)

and \( \tilde{K}_I = \tilde{K}_I(1 - \nu^2)/(E\sqrt{l_*}) \) stands for the dimensionless toughness. Consequently, the asymptotic estimate (18) can be now rewritten in the form:

\[ \tilde{w}(\tilde{t}, \tilde{x}) = \frac{8}{\sqrt{2\pi}} \tilde{K}_I \sqrt{L(\tilde{t})} \sqrt{1 - \tilde{x}}, \quad \tilde{x} \to 1. \]  

(41)

From definition (17) one can determine the dimensionless toughness as:

\[ \tilde{K}_I(t) = \frac{\sqrt{L(\tilde{t})}}{\pi^{3/2}} \int_0^1 \frac{\tilde{p}(\tilde{t}, s) ds}{\sqrt{1 - s^2}}, \]  

(42)

The boundary conditions (12) – (13) are converted to:

\[ \tilde{w}(\tilde{t}, 0)\tilde{v}(\tilde{t}, 0) = \tilde{q}_0(\tilde{t}), \quad \tilde{w}(\tilde{t}, 1) = 0. \]  

(43)

The initial conditions are now:

\[ L(0) = \frac{l_0}{l_*}, \quad \tilde{w}(0, \tilde{x}) = \tilde{w}_*(\tilde{x}) \equiv \frac{1}{l_*} w_*(l_0\tilde{x}), \quad \tilde{x} \in [0, 1]. \]  

(44)

The transformation of the global fluid balance equation (19) gives:

\[ \int_0^1 [L(\tilde{t})\tilde{w}(\tilde{t}, \tilde{x}) - L(0)\tilde{w}_*(\tilde{x})] d\tilde{x} - \int_0^f \tilde{q}_0(\tilde{t}) d\tilde{t} + \int_0^f L(\tilde{t}) \int_0^1 \tilde{q}(\tilde{t}, \tilde{x}) d\tilde{x} d\tilde{t} = 0, \]  

(45)

Note that relation (45) is valid under the assumption that the process of hydraulic fracturing is monotonous \( (L'(t) > 0) \).
Remark 1. Note that the boundary conditions
\[ \tilde{q}(\tilde{t}, 1) = 0, \quad \frac{\partial \tilde{w}}{\partial \tilde{x}}(\tilde{t}, 0) = 0, \]
are satisfied automatically and no longer need to be enforced. However, when appropriate, they can be implemented in the code.

For simplicity, from now on, we omit the "\(\sim\)" symbol for all normalized variables and parameters, and consider respective dimensionless values only.

3 Tip asymptotics and the crack propagation speed

In [38, 69] it was demonstrated that, in order to utilize the speed equation (34) efficiently, one needs to properly employ the tip asymptotics. This is the key point of the proposed method: such an approach makes it possible to evaluate the crack propagation speed without the technical difficulties discussed in [19] and uncertainties related to dividing two infinitesimally small values near the crack tip (the fluid flow rate and the crack opening). We shall show that there is a unique relationship between the crack length and the multipliers of one (or two) leading term(s) of the solution’s tip asymptotics. This relationship has a universal form and can be used regardless of the elasticity model or crack propagation regime. The analysis presented below is nothing but a direct extension (for the KGD model) of the approach introduced and verified in [38, 69].

It has been proved that the crack aperture in the vicinity of the fracture tip can be expressed as (see [26, 38] and the references therein, where most of the information can be found for at least the two leading terms):
\[ w(t, x) = w_0(t)(1 - x^m)^{\alpha_0} + w_1(t)(1 - x^m)^{\alpha_1} + O\left((1 - x^m)^{\alpha_2}\ln^\kappa(1 - x^m)\right), \quad x \to 1, \]
where powers \(\alpha_i\) are given in Table 1 for respective elasticity models. In the case of non-local elasticity, it is more convenient to base the asymptotic representation on terms \((1 - x^2)\). For the PKN and the fluid driven KGD models \(\kappa = 0\), while in general for the toughness driven variant of KGD: \(\kappa = 1\).

Note that the asymptotic behaviour (47) guarantees that the condition (21) is satisfied near the crack tip. As a consequence, the asymptotics of particle velocity yields:
\[ v(t, x) = v_0(t) + v_1(t)(1 - x^m)^{\beta_1} + O\left((1 - x^m)^{\beta_2}\right), \quad x \to 1. \]

The values of \(\beta_i\) \((0 < \beta_1 < \beta_2)\) are collected in Table 1 for various models in the case of impermeable rock. Note that \(v_0(t) = v(t, 1)\), is bounded and equal to the crack propagation speed:
\[ v_0(t) = \frac{dL}{dt} = -\frac{1}{L} \left[ w^2 \frac{\partial p}{\partial x} \right]_{x=1} < \infty. \]

It can be checked that, for all considered elasticity equations and fracture propagation regimes, the limiting value of the right-hand side of equation (49) is defined by no more than the first two terms of the asymptotic expansion for the crack opening (47). Indeed, let us adopt the following symbolic representation:
\[ \lim_{x \to 1} w^2 \frac{\partial A}{\partial x} A w = -C_A \frac{\mathcal{L}(w)}{L_{m-1}} < 0, \]

The powers, starting from the second one (the third one in toughness driven KGD model), were taken for the case with leak-off vanishing at the crack tip at least as fast as for the crack opening: \(\eta \geq \alpha_0\) (compare [29]). However, the proposed algorithm is applicable to any other permissible leak-off regime - see [29] and the discussions there. In such a case the table is to be modified, e.g. respective data for the PKN model can be found in [38].
| Elasticity model | m   | κ   | α₀ | α₁ | α₂ | β₁ | β₂ | ζ₀ | ζ₁ | ζ₂ |
|------------------|-----|-----|----|----|----|----|----|----|----|----|
| PKN              | 1   | 0   | 1/3| 4/3| 7/3| 1  | 2  | 1  | 2  | 3  |
| KGD (fluid driven)| 2   | 0   | 2/3| 5/3| 8/3| 1  | 2  | 1  | 2  | 3  |
| KGD (toughness driven) | 2   | 1   | 1/2| 1  | 3/2| 1  | 3/2| 1  | 3/2| 2  |

Table 1: The values of basic constants involved in the asymptotic expansions of $w$, $v$ and $\phi$ for the zero leak-off case.

where $L(w)$ is a functional on the fracture aperture $w$, related to the form of the elasticity operator, while $C_A$ is a known constant. In other words, the formula

$$v_0 = \frac{1}{L^m} C_A L(w),$$

is a universal one (valid for all elasticity formulations) and constitutes a relation between the crack propagation speed and the multiplier(s) of the leading term(s) of the crack opening tip asymptotics.

The values of the constants $C_A$ and the functionals $L(w)$ for respective models are:

- **PKN model**

  $$C_A = \frac{1}{3}, \quad L(w) = w_0^3,$$

- **KGD model - fluid driven regime**

  $$C_A = \frac{4}{9\sqrt{3}} \pi, \quad L(w) = w_0^3,$$

- **KGD model - toughness driven regime**

  $$C_A = 2, \quad L(w) = w_0^2 w_1.$$

Note that, when using the asymptotic expansion based on terms $(1-x)$ only, one needs to modify relations (53–54).

Taking into account that $v_0(t) = L'(t)$, equation (51) can be directly integrated to compute the crack length:

$$L(t) = \left[ L^{m+1}(0) + (m+1)C_A \int_0^t L(w) d\tau \right]^{\frac{1}{m+1}}.$$  

(55)

This universal formula in turn, sets a nonlocal relation between the crack length and the leading term(s) of $w(t,x)$ asymptotic expansion.

However, for the toughness driven regime of KGD model the following condition is satisfied:

$$w_0(t) = \frac{4}{\sqrt{\pi}} K_I \sqrt{L(t)}.$$  

(56)

As a result, a slightly different approach to define the crack length at any time, $t$, can be applied. Namely, by combining (56) with (50), (54) and (49) one obtains an alternative formula:

$$L(t) = \left[ L^2(0) + C_I K_I^2 \int_0^t w_1(\tau) d\tau \right]^{\frac{1}{2}}, \quad C_I = \frac{64}{\pi}.$$  

(57)
Moreover, the formula (56) itself can be directly used to determine the fracture length:

\[ L(t) = \frac{\pi}{16R^2} w_0^2(t). \]  

(58)

Thus, for the toughness driven KGD model it is possible to use different strategies for the computations. Namely, for small toughness is is natural to use the representation (57), for large toughness – (58), while a "mixed" strategy based on two first terms in the asymptotic expansion (47) and formulae (55), (54) can be adopted for the intermediate cases.

**Remark 2.** It follows from the foregoing analysis that numerical evaluation of the leading coefficient (or two first multipliers for the toughness driven KGD model) in the asymptotic representation (47) plays a crucial role in computing the crack length. This will be discussed in details in Section 6.

**Remark 3.** Note that there is another way to determine the crack length by employing the balance condition (45). Indeed, this relation represents the Volterra integral equation of the second kind with respect to the crack length \( L(t) \). Such an equation has always a unique solution and there are effective numerical methods to solve it [48]. The kernel of the equation, which is an indefinite integral itself, should be, generally speaking, modified at every time step and thus this approach is not cost effective. Moreover, in case of impermeable rock \( q_l = 0 \) the integral equation degenerates and the crack length can be found explicitly regardless of the elasticity model and fracture propagation regime.

### 4 Problem reformulation in terms of reduced particle velocity.

In this section we reformulate the problem by replacing the particle velocity with a new dependent variable, the reduced particle velocity, and demonstrate its advantages.

Let us introduce a new dependent variable called, from now on, the reduced particle velocity:

\[ \phi(t, x) = v(t, x) - x v_0(t). \]  

(59)

Its asymptotic behaviour can be described qualitatively as (compare with Table 1):

\[ \phi(t, x) = \phi_0(t)(1 - x^m) \zeta_0 + \phi_1(t)(1 - x^m) \zeta_1 + O((1 - x^m) \zeta_2), \quad x \to 1. \]  

(60)

In numerical implementation \( \phi \) possess all the advantages of the particle velocity, \( v \), allowing simultaneously to set a new boundary condition

\[ \phi(t, 1) = 0, \]  

(61)

instead of \( v(t, 1) = v_0(t) \), with the unknown speed of the crack, \( v_0(t) \). The latter is not easy to be used as it is directly defined by the product of two terms where one tends to zero while the other to infinity (Compare (49), (23) and (43)\(^2\)). Qualitatively the new variable, \( \phi \), exhibits similar asymptotic behaviour to the crack opening - it tends to zero at the fracture tip. Thus, we can define a respective boundary condition for this variable (see (61)).

Combining (59) with (43), and substituting the result into the continuity equation (32), one obtains its modified form:

\[ \frac{\partial w}{\partial t} + \frac{1}{L} \frac{\partial}{\partial x} (w \phi) + \frac{v_0}{L} w + q_l = 0. \]  

(62)

The boundary condition (43) can be now replaced by:

\[ w(t, 0) \phi(t, 0) = q_0(t), \]  

(63)

while the crack tip conditions are (43)\(^2\) and (61).
Note, that the following solvability condition for the equation (62) should be satisfied:

\[
\left. \int_0^1 \frac{\partial w}{\partial t}(t, x) + q_l(t, x) \right|_{\partial_t} + \frac{\partial v_0(t)}{L(t)} \int_0^1 w(t, x) dx = \frac{1}{L(t)} q_0(t). \tag{64}
\]

It constitutes a local equivalent of the global balance condition (45) and, since \(w > 0\) for any \(x \in [0, 1]\), it allows one to uniquely define the value of the crack velocity, \(v_0\), at every time step. As a result, equation (62) always has a unique solution with respect to the reduced particle velocity, \(\phi\).

From (33), the pressure derivative can be expressed as:

\[
\frac{\partial p}{\partial x} = -L w^2 \left( \phi + x v_0 \right). \tag{65}
\]

In the case of PKN model \((p = w)\), equation (65) can be transformed to a functional equation with respect to the crack opening, \(w\), which can be written in a symbolic manner:

\[
w = B_w(\phi, L, w, v_0). \tag{66}
\]

The latter representation corresponds to the proper variable approach discussed in [46, 51] and is usually more effective in computations. In particular, it does not require iterations in numerical computing. However, e.g. in the case of the P3D model, equation (66) takes the form

\[
w(t, x) = -\frac{2}{\pi^2} L^2(t) \int_0^1 \frac{\phi(t, \eta) + \eta v_0(t)}{w^2(t, \eta)} K(\eta, x) d\eta + \frac{4L(t)}{\sqrt{\pi}} K \sqrt{1 - x^2}, \tag{67}
\]

where \(F(w, x)\) is the proportionality coefficient in the elasticity relation, \(p = F(w, x)w\), (see [49, 47]). For this single reason, we stay with both representations (67) and (68). Respective implications for the numerics shall be discussed later on.

In the case of KGD model, equation (65) should be combined with the transformed form of elasticity relation (39) to eliminate the pressure gradient, \(p'_x\), and to find the crack opening, \(w\), again in form the (66)

\[
\frac{\partial p}{\partial x} = -L \int_0^1 \phi(t, \eta) + \eta v_0(t) w^2(t, \eta) d\eta + p_0(t). \tag{67}
\]

It is obvious that the boundary condition (43) holds automatically. It can be proved that for the KGD model, also condition (46) is fulfilled by definition.

As a result, the basic system of equations now utilizes two dependent variables: the crack opening, \(w(t, x)\), and the reduced particle velocity, \(\phi(t, x)\). We will be looking for a solution to the lubrication equation (62), under boundary conditions (43), (61), (63), initial conditions (44) and the speed equation (49) written in one of the transformed forms (55) – (58). Finally, the pressure in the KGD model can be computed by integrating equation (65):

\[
p(t, x) = -L(t) \int_0^\pi \frac{\phi(t, \eta) + \eta v_0(t)}{w^2(t, \eta)} d\eta + p_0(t). \tag{71}
\]
where the constant $p_0$ is defined from (35) and (37)
\[ p_0(t) = -\frac{1}{L(t)} \int_0^1 \frac{\partial w(t,\eta)}{\partial \eta} d\eta \tag{72} \]
Note that the integral converges due to condition (46). The physical interpretation of this is that it refers to a bounded value of particle velocity at the fracture inlet.

5 Self-similar solution
Let us search for a solution of the problem described in the previous section in the following manner:
\[ w(t,x) = \psi(t) \hat{w}(x), \quad p(t,x) = \frac{\psi(t)}{L^{m-1}(t)} \hat{p}(x), \tag{73} \]
\[ q(t,x) = \frac{\psi^4(t)}{L^m(t)} \hat{q}(x), \quad v(t,x) = \frac{\psi^3(t)}{L^m(t)} \hat{v}(x), \quad \phi(t,x) = \frac{\psi^3(t)}{L^m(t)} \hat{\phi}(x). \tag{74} \]
As shown in Appendix A, such a separation of variables enables one to reduce the problem to the time-independent form in the case when $\psi(t)$ is a power law or exponential function of time. Such a formulation will be called henceforth the self-similar formulation. Respective spatial components of the solution (depending exclusively on $x$) are marked by 'hat'-symbol.

Below we demonstrate the basic assumptions and features of the universal algorithm, considering first the self-similar formulation of the problem.

5.1 Problem formulation
As follows from the self-similar formulation given in Appendix A, a set of governing equations can be written in general form as:

- equation determining the reduced velocity following from (62) (self-similar equivalents: (103) or (110))
\[ \hat{\phi}(x) = \frac{L(\hat{w})}{\hat{w}(x)} \int_x^1 \left( \chi \hat{\phi}(\xi) + \kappa \hat{q}_l(\xi) \right) d\xi, \tag{74} \]

- equation allowing the computation of the crack opening, which follows from (65) and the respective inverse elasticity operator (self-similar versions: (95) – (93))
\[ \hat{w} = B_{w} \left( \frac{1}{\hat{w}^2} \left( \hat{\phi} + xL(\hat{w})C_A \right) \right), \tag{75} \]

- boundary conditions:
\[ \hat{w}(0)\hat{\phi}(0) = \hat{q}_0, \quad \hat{w}(1) = 0, \quad \hat{\phi}(1) = 0, \tag{76} \]

- solvability condition allowing one to determine the value of the parameter $\hat{v}_0 = L(\hat{w})$, which follows immediately from (74) and (76). It is equivalent to the global fluid balance equation (45) (self-similar counterparts: (106) or (111))
\[ \frac{1}{L(\hat{w})} \hat{q}_0 - \chi \int_0^1 \hat{w}(x)dx - \kappa \int_0^1 \hat{q}_l(x)dx = 0. \tag{77} \]
For the PKN model operator $B_w$ from (75) has one of the following alternative forms:

$$B_w \hat{w}(x) = \int_x^1 \frac{\hat{\phi}(\eta) + \eta \mathcal{L}(\hat{w}) C_A}{\hat{w}^2(\eta)} d\eta,$$  \hspace{1cm} (78)

or

$$B_w \hat{w}(x) = \left[ 3 \int_x^1 (\hat{\phi}(\eta) + \eta \mathcal{L}(\hat{w}) C_A) d\eta \right]^{1/3}. \hspace{1cm} (79)$$

For the KGD variant of the problem it is directly defined by relation (93) which is a self-similar equivalent of (39):

$$B_w \hat{w}(x) = \frac{2}{\pi} \int_0^1 \frac{\hat{\phi}(\eta) + \eta \mathcal{L}(\hat{w}) C_A}{\hat{w}^2(\eta)} K(\eta, x) d\eta + \frac{4}{\sqrt{\pi}} \hat{K}_I \sqrt{1 - x^2}, \hspace{1cm} (80)$$

where the kernel $K(\eta, x)$ is defined in (40).

Note that the general form of the speed equation (49) and the additional boundary condition (46) for the KGD model ((104) in self-similar formulation) are satisfied by the system (74) – (77) automatically. Positive constants $\chi$ and $\kappa$ from equation (74) depend on the type of the function $\psi$ defining the self-similar solution and given by (110) and (103), respectively. All the values of different parameters used in the above equations are collected in Tables 1, 2.

| Type of the self-similar law | $\chi$ | $\kappa$ |
|-----------------------------|--------|--------|
| $\psi(t) = e^{\gamma t}$    | $C_A \frac{m+2}{3\gamma}$   | $C_A \frac{m+1}{3\gamma}$   |
| $\psi(t) = (1 + t)^\gamma$  | $C_A \frac{(m+1)\gamma}{3\gamma+1} + 1$   | $C_A \frac{m+1}{3\gamma+1}$   |

Table 2: Values of the auxiliary parameters in the self-similar formulations reflecting different time dependent behaviours.

### 5.2 Numerical algorithm for the self-similar solution

The solution of the self-similar problem formulated above is sought in the framework of the universal algorithm. The universality refers here to the fact, that only some parameters in respective blocks should be changed to adjust the solver to work with different variants of the problem (PKB, KGD in both considered regimes). The algorithms consists of the following iterative steps:

- In the first stage we accept some initial approximation of the crack opening, $\hat{w} = \hat{w}\text{^{(i-1)}}$. Equations (77) and (74) are utilized to compute $\mathcal{L}\text{^{(i)}}(\hat{w})$ and the reduced velocity $\hat{\phi}\text{^{(i)}}$. In general (77) yields $\mathcal{L}(\hat{w})$ which, when substituted into (74), enables the integration of the latter to obtain $\hat{\phi}$. In the manner of the so-called $\varepsilon$-regularization technique [44, 38] the integration is carried out over truncated spatial interval $x \in [0, 1 - \varepsilon]$, where $\varepsilon$ is a small parameter. The boundary condition (76) is replaced by the condition resulting from the asymptotics (60), specified at $x = 1 - \varepsilon$. The regularized boundary condition is introduced in the form:

$$\hat{\phi}_N = s_1(\zeta_0, \zeta_1) \hat{\phi}_{N-1} + s_2(\zeta_0, \zeta_1) \hat{\phi}_{N-2},$$  \hspace{1cm} (81)

where the subscripts of $\hat{\phi}$ refer to the indices of nodal points of the spatial mesh (containing $N$ nodes). The values of multipliers $s_1(2)(\zeta_0, \zeta_1)$ depend on the particular asymptotic behaviour of

\footnote{A comprehensive description of this form of the $\varepsilon$ - regularization technique, together with methods for its implementation, can be found in [38].}
the function $\hat{\phi}$ (see (60) and Table 1). As a result, function $\hat{\phi}^{(i)}(x)$ and the constant $\mathcal{L}^{(i,1)}(\hat{w})$ computed at this stage satisfy, together with predefined $\hat{w}^{(i-1)}$: i) the fluid balance equation (77), ii) the continuity equation (74), iii) the regularized boundary condition for $\phi$ (81) (equivalent to (76)3), iv) the influx boundary condition (76)1 - indirectly, through the fluid balance equation.

• At the second stage of each iterative loop, the values of $\hat{\phi}^{(i)}$ obtained previously are utilized to compute the next iteration $\hat{w}^{(i)}$ from (75). Note, that by the properties of the operator $\mathcal{B}_w$ corresponding to the KGD model, condition (105) is satisfied automatically.

While computing respective integral operators $\mathcal{B}_w$, it is crucial to preserve appropriate asymptotic behaviour of the integrands, resulting from (47), (60). Moreover, at this stage $\mathcal{L}^{(i,2)}(\hat{w})$ is considered a natural regularization parameter, chosen to satisfy the influx boundary condition (76)1. Hence, $\hat{w}^{(i)}$ computed at this stage satisfies respective elasticity relation and boundary conditions: (76)2 through the imposed asymptotics

$$\hat{w}_N = s_1(\alpha_0, \alpha_1)\hat{w}_{N-1} + s_2(\alpha_0, \alpha_1)\hat{w}_{N-2},$$

and (76)3.

• The aforementioned two stages of the iterative loop are repeated until all components of the solutions $\hat{\phi}$, $\hat{w}$ and $\mathcal{L}(\hat{w})$ have converged with the prescribed tolerances.

Remark 4. As shown in [38], the relationships (81) and (82) allow one to determine numerically the multipliers of the first two leading terms of the asymptotics of $\hat{\phi}$ and $\hat{w}$.

Remark 5. Similarly, as shown in [69], the performance of the algorithm improves significantly when instead of the dependent variables $\hat{w}$ and $\hat{\phi}$ one uses the difference between them and their leading asymptotic terms:

$$\Delta \hat{w} = \hat{w} - w_0(1 - x^m)^{\alpha_0}, \quad \Delta \hat{\phi} = \hat{\phi} - \phi_0(1 - x^m)^{\zeta_0}.$$ 

Then the leading asymptotic terms in the left and right-hand sides of the equations (74) and (75) are canceled analytically and the functions $\Delta \hat{w}$ and $\Delta \hat{\phi}$ are computed in the iterative process. Moreover, while searching for the regularization parameter $\mathcal{L}(\hat{w})$ we take into account its relation to the respective coefficients in the asymptotic expansion of the solution (compare (52) – (54)). This, in turn, leads to nonlinear equations solved with the Newton-Raphson method. Finally, the qualitative asymptotic behaviour of the new dependent variables $\Delta \hat{w}$ and $\Delta \hat{\phi}$ is also known in advance and the respective exponents should be appropriately adopted in the $\varepsilon$-regularization technique.

Remark 6. When computing the integral operators (74) – (75), we use the test (weight) functions coinciding with the leading asymptotic terms of the integrands. This provides better accuracy and efficiency when integrating.

Remark 7. Note, that, due to the modular algorithm architecture, the subroutine for computing the crack opening can be easily replaced in accordance with the chosen elasticity model.

5.3 Analysis of the algorithm performance for the self-similar formulation

Let us now analyze the performance of the algorithm presented above. Since the self-similar variant of the algorithm constitutes the integral part of the algorithm for the transient problem, it is the first major step towards constructing the general numerical scheme. Its analysis enabled us to overcome the basic difficulties, and to optimize the computations. In the following four subsections we investigate the properties of the universal algorithm against several analytical benchmarks and compare the results to the data available in literature.
5.3.1 Analysis of the algorithm - PKN model

For the PKN model, we use two different benchmark solutions. The first of them is built on the formula (113) for three base functions of the type (119). All the resulting quantities can be obtained by the way described in Appendix B. Note that this benchmark solution, called hereafter benchmark I, assumes a predefined non-zero leak-off function. The second benchmark example is taken from [51] (p.7-8, eqs.(38) – (39)). In this case there is no leak-off in the formulation. This solution is called from now on benchmark II.

In the following we analyze the accuracy of computations described by two parameters: \( \delta \hat{w} \) and \( \delta \hat{\phi} \) - the maximal relative errors of the crack opening and the reduced velocity, respectively. The corresponding results are presented in Fig.1 a) and Fig.1 b). These errors were estimated for a number of nodal points varying from 10 to 300, where the spatial mesh density was increased at both sides of the interval (in a manner described in [69]). For comparison, we additionally show in Fig.1 a) the accuracy of computations performed by the extremely efficient integral solver proposed in [69] for the PKN model. The latter algorithm does not utilize the reduced velocity. Its computation needs additional post-processing, and thus \( \delta \hat{\phi} \) is not shown in this case.

![Figure 1](image)

Figure 1: Relative errors of the self-similar solution for: a) the crack opening \( \hat{w} \), b) the reduced particle velocity \( \hat{\phi} \). Dashed lines in a) correspond to the results obtained by integral solver described in [69].

The results presented in Fig.1 demonstrate that the solution error depends on the type of benchmark. In this particular case there is a clear explanation for this fact, as benchmark II excludes an error introduced by the numerical integration of the leak-off function. In the analyzed range of \( N \), a clear trend of accuracy improvement with growing mesh density can be observed. However, the error reduction becomes slower with \( N \) growth.

The comparison of the results with those for the integral solver shows that the latter can provide better accuracy for smaller \( N \). It also gives greater potential for the solution improvement for large values of \( N \). On the other hand, for both benchmarks, there exists an intermediate interval where the solver based on the universal algorithm gives better accuracy. Summarizing, for both benchmarks the overall accuracy is extremely high, comparable with that provided by the integral solver and much better that the level of accuracy reported in [35].

Remark 8. Computations shown in this subsection were done with the operator \( B_w \) defined in (79) as the respective algorithm provided better accuracy. When using the alternative operator (78), the accuracy was up to one order of magnitude worse for both the crack opening, \( w \), and the reduced partial velocity, \( \phi \). However, when moving forward to the transient state scheme, we observed no difference
in accuracy between these approaches. The reason for this is rather clear: the major error in the computations (and thus the accuracy limiting factor) in this case comes from the FD representation of the temporal derivative, even though we take its more accurate approximation than that used in the standard hydrofracturing algorithms (compare [3]).

5.3.2 Analysis of the algorithm - accuracy of computations for the KGD model

In this subsection we estimate the solution accuracy for two analytical benchmarks (for the fluid and toughness driven regimes respectively) given in Appendix B.

The benchmark solution for the fluid driven regime is based on the representation (122) composed of three terms. For the toughness driven mode we apply the basis (125) for four terms. The computations for different numbers of nodal points $N$ ranging from 20 to 300 were performed. The lower limit of $N$ was set to 20 instead of 10, as the proper numerical computation of the inverse elasticity operator (93) necessitates finer meshing than its equivalent in the PKN model (identity operator). Again, the mesh density was increased at both ends of the spatial interval.

![Figure 2: Relative errors of the self-similar solution for: a) the crack opening $\hat{w}$, b) the reduced particle velocity $\hat{\phi}$.](image)

Similarly as for the PKN model, the results show that the accuracy depends on the type of the benchmark. In the analyzed range of $N$ the increase of mesh density gives monotonous reduction of the error of computations. This trend attenuates with growing $N$, however for $N < 300$ it is still far away from the stabilization level. In the benchmarks under consideration it is sufficient to take merely 20 nodal points to have the accuracy of the level $10^{-5}$ for both regimes. Thus, the universal algorithm shows the same quality of performance in terms of the accuracy and efficiency as both the KGD and PKN models (the latter being slightly less challenging for computations). The toughness driven benchmark considered here does not refer to any of the boundary cases, namely neither to the small or to the large toughness regimes. These will be discussed later on.

5.3.3 Fluid driven KGD model for impermeable rock - comparison with other results.

Having identified the level of solution accuracy provided by the implemented algorithm, we will now perform similar analysis against various classical results (both numerical and semi-analytical) available in the literature. We start with the numerical solution given in terms of series approximation in [2].
The authors analyzed the fluid driven regime (KGD model) for a number of shear-thinning fluids. The constant influx and impermeability of the rock formation ($\dot{q}_I = 0$) were also assumed. In our case only the data for the newtonian fluid ($n=1$) will be used. For the same problem, a semi-analytical approximate solution has been recently proposed in [46]. Finally, one can find in [21] a simple approximations for the crack opening and the net fluid pressure, originally introduced in [1].

In the following we compare our numerical results with those given in the recalled papers. In order to make sure that the accuracy of our computations is at least of the order $10^{-6}$, we take a mesh composed of 300 nodal points (compare Fig. 2) whose density was increased at both ends of the interval. We compare the results in terms of: i) self-similar crack opening, ii) self-similar fluid pressure, iii) self-similar particle velocity and (iv) self-similar fluid flow rate. The explicit formulae for the first three dependent variables are given in [46]. Although in [2] there is no data for the particle velocity, it can be easily obtained through the flux and the fracture opening ($\dot{v} = \dot{q}/\dot{w}$) or the fracture opening and pressure derivative ($\dot{v} = -\dot{w}^2\dot{p}'$). Similarly, the lacking data for the fluid flow rate in [46] can be recreated as $\dot{q} = \dot{w}\dot{v}$. Unfortunately, there are no direct formulae in [21] for either the fluid flow rate or the fluid pressure derivative. Thus, aiming at a fair comparison, we do not show here results for $\dot{v}$ and $\dot{q}$ (which could theoretically be obtained by pressure differentiation). As for the results by the universal algorithm, the particle velocity is retrieved directly from the reduced particle velocity, $\dot{\phi}$, while the computation of the fluid pressure necessitates additional post-processing (integration).

![Figure 3: Results for the self-similar problem: a) the crack opening $\dot{w}$, b) the fluid pressure $\dot{p}$.](image)

The respective results are shown in Fig. 3 – Fig. 4. To make the graphs more legible, we do not depict here the approximation from [11], which is discussed later on.

Considering the crack opening and fluid pressure, one can see that the curves corresponding to the universal algorithm and the solution presented in [2] are indistinguishable from each other in the used scale (the $x$ interval in Fig. 4 is truncated, since the pressure tends to infinity for $x \to 1$). The solution found in [46] provides a very good approximation for the fracture aperture, however for the fluid pressure it deviates from other results. When analyzing the particle velocity, we decided to use two methods of $\dot{v}$ computation for the data from [2], as each of them produces a slightly different result. The method refereed to as Adachi 2002(1) utilizes $\dot{w}$ and $\dot{p}'$ ($\dot{v} = -\dot{w}^2\dot{p}'$), while for Adachi 2002(2) we employed $\dot{q}$ and $\dot{w}$ ($\dot{v} = \dot{q}/\dot{w}$). Both methods are equivalent in the case of an exact solution. It shows that the values given by the universal algorithm are in a good agreement with those by Adachi 2002(1) except for the fracture tip, where the apparent deterioration of the latter solution takes place. However, in this region Adachi 2002(2) turns out to be perfectly consistent with our results. The solution from [46] is hardly
Figure 4: Results for the self-similar problem: a) particle velocity $\hat{v}$, b) fluid flow rate $\hat{q}$.

distinguishable from that by the universal algorithm for $x > 0.8$, but deviates when $x$ decreases.

Finally, for the fluid flow rate, $\hat{q}$, we observe a good agreement between our data and that by Adachi 2002(1) (series approximation of $\hat{q}$ given in the paper) over the whole interval. Results by Adachi 2002(2) (flux recreated as and $\hat{q} = -\hat{w}^3\hat{p}'$) and [46] diverge from ours for decreasing $x$.

The above analysis confirms the credibility of our solution, which together with the previous accuracy estimation allows us to treat it now as a numerical benchmark in and of itself. Following the idea from [1] and [46], we propose a new improved approximation of the dependent variables analyzed above, which provides higher accuracy than other known semi-analytical formulae and can be treated as the reference data when testing other numerical algorithms.

Namely, we express the fracture opening, $\hat{w}$, the fluid pressure, $\hat{p}$, and the particle velocity, $\hat{v}$, in the following manner:

$$\hat{w}(x) = \sqrt{3}(1 - x^2)^{2/3} + 0.3 \left[ \sqrt{1 - x^2} - \frac{2}{3}(1 - x^2)^{3/2} - x^2 \ln \frac{1 + \sqrt{1 - x^2}}{x} \right],$$  \hspace{0.5cm} (83)

$$\hat{p}(x) = \frac{B(1/2, 2/3)}{\sqrt{3}\pi} 2F_1(1/6, 1; 1/2; x^2) + \sum_{i=1}^{6} p_i(1-x)^{(6-i)/2},$$ \hspace{0.5cm} (84)

$$\hat{v}(x) = \frac{2}{3} \left[ 1 + \sum_{i=1}^{5} v_i(1-x)^i \right],$$ \hspace{0.5cm} (85)

where $B$ is the beta function, $2F_1$ denotes the Gauss hypergeometric function, and the respective multipliers from [84] and [85] assume values: $p_1 = 0.274395$, $p_2 = -0.56408$, $p_3 = 0.547395$, $p_4 = -0.15621$, $p_5 = -0.02495$, $p_6 = -0.007285$, $v_1 = -0.1$, $v_2 = 0.10542$, $v_3 = 0.02875$, $v_4 = -0.02739$, $v_5 = 0.0752$.

Respective approximation for the fluid flow rate, $\hat{q}$, can be easily obtained from the product of the fracture opening [83] and particle velocity [85]. Note that the term multiplied by 0.3 in [83] is exactly the special term, $w_N$, used in representation [122] for the pressure derivative correction.

In Fig. 5 – Fig. 6 we show comparisons between the improved approximation and the results known from [2], [46], [1], referring them all to our numerical reference solution. For the crack opening, $w$, the particle velocity, $v$, and the fluid flow rate, $\hat{q}$, their relative deviations $\delta w$, $\delta v$ and $\delta \hat{q}$ from the numerical solution are given. Again, $\hat{v}$ and $\hat{q}$ from [2] are computed in two alternative ways described above. For the fluid pressure we show the absolute difference, $\Delta p$, as the pressure curve intersects the $x$-axis.
As can be seen, the new representation (83) imitates the crack opening with an accuracy of the order $10^{-3}$ (the maximal value of deviation is less than 0.2%). The same level of error of approximation is obtained for the fluid flow rate. For the fluid pressure, the improved approximation gives a maximal deviation from the numerical benchmark of the order $10^{-5}$. Finally, for the particle velocity, the error of the approximation is of order $10^{-4}$.

For all the considered parameters, the improved approximation gives a better agreement with the accurate numerical solution than any of the known results. Note, that this analysis also reveals the level of accuracy of all previously reported approximations. It shows that the solution from [2] is better than the one proposed in [46] in an average sense. However, its quality deteriorates near the fracture tip, where the latter exhibits the correct tip asymptotics. For the particle velocity and fluid flow rate, the solution from [2] could be improved by merging two representations obtained by two different ways of defining $\hat{q}$: series approximation near the fracture tip and $\hat{q} = -\hat{w}^3\hat{p}'$ on the rest of the interval. On the
other hand, the approximation from [1] preserves the tip asymptotic behaviour much better than the series approximation [2]. However, it gets worse when moving away from the fracture tip.

Remark 9. Respective formulae defining the improved approximation should be treated independently, i.e. when trying to recreate the particle velocity by using (83) and differentiating (84), one cannot expect the same accuracy as for (85).

5.3.4 Toughness driven KGD model for impermeable rock – comparison with other results.

For the toughness driven regime of the KGD model, it is difficult to find formulae in the literature which could be directly compared with our solution in the same way as done above. Such solutions are either not complete, in a sense they do not describe all the components analyzed above (rather only the crack opening and the fluid pressure), or one can find only the values of a few first multipliers of the respective base functions approximating the solution. As a result, it is practically impossible to provide a fair and credible comparison. In some cases, in order to rebuild the solution given by the author, one needs also to repeat the respective numerical algorithm. Then, the quality of the comparison would essentially depend on the algorithm implementation.

We compare our numerical results with those few sources available. First, we start with the classical example from [64], where the authors provide a number of numerical results obtained by the method of series approximation for different variants of the problem (p.300, Table 1). Unfortunately, the format of the data records (two or three decimal digits), and the fact that multipliers for merely three leading terms are given, allow us to consider the reconstructed solution rather as a rough approximation. We recreated that solution for the data corresponding to the first case ($\alpha = 1, \lambda = 2/3$) from the aforementioned Table 1. The self-similar stress intensity factor $\hat{K}$ was computed in accordance with the given coefficients. The utilized spatial mesh was composed of 300 nodal points with the density refined near the end points. The comparison of our results with those from [64] is given in Fig. 7.

![Figure 7: Spence & Sharp solution - comparison of the numerical solution with data provided in [64]: a) the crack opening, $\hat{w}$, b) relative deviation between crack openings.](image)

Figure 7: Spence & Sharp solution - comparison of the numerical solution with data provided in [64]: a) the crack opening, $\hat{w}$, b) relative deviation between crack openings.

Here only the crack opening, $w$, and its relative deviation are shown. The representation of the net fluid pressure based on only three terms is far from completeness (and does not reflect the proper asymptotic behaviour). Thus the comparison would be unfair in this case. One can see that the maximal relative error of the solution from [64] amounts to 3.5% (compare Fig. 7b)) and is located at some distance away from the fracture tip.
away from the crack tip. The asymptotic behaviours of both solutions in the near-tip region coincide with each other, being directly embedded into the respective numerical schemes.

Another reference solution used in this subsection is the one given in [22] for the large toughness (small viscosity) regime. The author provides the data sufficient to recreate the crack opening and the net fluid pressure (Table II - pp. 1458, with respective formulae for the opening and pressure approximations), for three different values of the fluid behaviour index. Since in this paper we consider Newtonian fluids, only the data corresponding to the case \( n = 1 \) was taken (the correct value of the first coefficient should be two times smaller than that reported: \( 6.05 \cdot 10^{-3} \)). As previously, we carried out the computations on a mesh composed of 300 nodal points. This, according to the characteristic given in Fig. 2, should provide an error of the level \( 10^{-7} \).

The relative difference between our numerical solution and the one reconstructed from the data given in [22] is shown in Fig. 8. The latter is denoted in the pictures as the '10-terms' solution. For comparison we present also the less accurate two-terms approximate solution available in [23] (described in the picture as the '2-terms' solution).

![Graphs showing relative deviation of crack opening and net fluid pressure](image)

Figure 8: Comparison of the numerical results with data provided in [22] and [23]: a) relative deviation of the crack opening, \( \hat{w} \), b) relative deviation of the net fluid pressure, \( \hat{p} \).

It shows that the solution based on 10 terms yields a very good accuracy. For the fracture aperture, \( \hat{w} \), its relative deviation from our solution is of the order \( 10^{-5} \). The lowest discrepancy between respective solutions can be observed in the near-tip region, which demonstrates high quality of our numerical computations. Indeed the solution from [22] represents the accurate asymptotics there. For the net fluid pressure, \( \hat{p} \), the relative error is of one order greater. Similar analysis for the two-terms approximation from [23] gives the level of \( 10^{-3} \) for the crack opening, and \( 10^{-2} \) of the net fluid pressure. We believe that Fig. 8 reveals the accuracy of semi-analytical solutions solutions proposed in [22] and [23]. It is worth mentioning that the error measure introduced in [22], the quadratic global error \( e^{(N)} \) (equation (E1), pp.1471 therein), is of the level \( 10^{-6} \) for \( N = 10 \) (see figure E2, pp.1472 therein). This establishes, in this particular case, a relationship between the recalled measure and the relative error of the solution.

In the foregoing analysis, we have compared our numerical solution with known results confirming its high accuracy. Now, we analyse the performance of our algorithm for different values of the normalised self-similar stress intensity factor. Special attention is paid to the problem of small toughness as it constitutes the most challenging case for the computations, which was underlined in [41]. In such a

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limiting variant of the problem, it is convenient to have a reference to the fluid-driven solution discussed in Fig. 3 – Fig. 4.

For this reason let us impose the same case as recalled for the influx magnitude. Then, by changing (decreasing) the values of $K_I$, we investigate different variants of the problem. It is not a surprise that for sufficiently small $K_I$ the numerical results converge to the fluid driven reference solution. The relative deviations between the latter and the solutions for three variants of $K_I$: 0.1, 0.05, 0.01, are shown in Fig. 9. As can be seen, the greatest discrepancies can be observed in the vicinity of the crack tip, where the respective asymptotics do not correspond with each other. It is enough to take $K_I = 0.01$ to have the relative deviation from the fluid driven solution of the order $10^{-6}$ along almost the entire spatial interval. This fact calls into question the sensibility of conducting computations for the toughness driven regime with such small values of $K_I$, especially as the efficiency of computations deteriorates as $K_I$ decreases.

Remark 10. Note that in case of very small toughness, the parameter $\varepsilon$ in the $\varepsilon$-regularization technique should be extremely small to capture the tip asymptotics. Indeed, in our computations it was $\varepsilon = 10^{-9}$. Moreover, when decreasing $K_I$ to very low values, we arrive at a situation where the relative difference between the small toughness solution and the fluid driven one is of the same order as the error of computation.

In Fig. 10 we present the evolution of the crack opening at inlet, $\hat{w}(0)$, and the crack propagation speed, $\hat{v}(1)$, for continuously growing $K_I$, ranging from 0 (fluid driven solution) to 10 (large toughness solution). It shows that, for greater values of the self-similar stress intensity factor, $\hat{w}(0)$ increases as a linear function of $K_I$, while the crack propagation speed is inversely proportional to this parameter:

$$\hat{w}(0) = O(K_I), \quad \hat{v}(1) = O(K_I^{-1}), \quad K_I \to \infty.$$  

To complete the analysis, in Fig. 11 – Fig. 12 we depict the data for $K_I = 0.1, 1, 2, 3$. As one can expect, an increase in $K_I$ entails growth in the fracture width with simultaneous deceleration of the crack. When analyzing the pressure graphs, one can see that the average value of $\hat{p}$ is growing, which is a result of the increasing contribution of the square root term in the crack opening (this single term gives a constant over $x$ component of pressure). The values of pressure derivatives decrease with $K_I$ growth. This, together with the counteracting trend for $\hat{w}$, enables us to define two extreme regimes: i) fluid driven regime (zero toughness), ii) storage regime (infinite toughness). The latter is illustrated for
6 Solution of the problem in transient regime

In this section we discuss an extension of the algorithm presented above to the time-dependent variant of the problem. The main assumptions and blocks of the algorithm remain the same. The new features introduced here are subroutines for approximating the temporal derivative and the crack length computation. Presented numerical examples demonstrate the performance of the general variant of the algorithm.

Figure 10: Results for different values of $\hat{K}_I$ ($\hat{K}_I = 0$ corresponds to the fluid driven regime): a) crack opening at the inlet, $\hat{w}(0)$, b) crack propagation speed, $\hat{v}(1)$.

Figure 11: Solutions for different values of the self-similar stress intensity factor, $\hat{K}_I$: a) the crack opening, $\hat{w}$, b) the net fluid pressure, $\hat{p}$.

the fluid flux in Fig. 12 b) by the data for $\hat{K}_I = 10^3$. One can see that the transient state between two boundary cases is relatively small.
fluid driven
$\hat{K}_I = 0$
$\hat{K}_I = 1$
$\hat{K}_I = 2$
$\hat{K}_I = 3$

Figure 12: Solutions for different values of the self-similar stress intensity factor, $\hat{K}_I$: a) the particle velocity, $\hat{v}$, b) the fluid flow rate, $\hat{q}$.

6.1 Problem formulation

Analogously as it was for the self-similar variant of the problem, let us write the basic system of equations, collecting them in the order of employment in the algorithm. The fundamental difference between the scheme presented in the previous section and the one for the transient problem is the introduction of a mechanism for approximating the temporal derivative, which shall be accounted for while computing the reduced velocity. Here the initial condition is also implemented. Finally, the crack length should be estimated for both main stages of the algorithm (for $\phi$ and $w$).

We consistently use the following representation of the temporal derivative of the crack opening:

$$\frac{\partial w}{\partial t} \bigg|_{t_i} = \frac{2}{t_{i+1} - t_i} \left( w(x, t_{i+1}) - w(x, t_i) \right) - \frac{\partial w}{\partial t} \bigg|_{t_i}.$$  \hspace{1cm} (86)

This representation has already been utilized in [69], where its advantages were discussed. The representation (86) yields an accuracy of $O((\Delta t)^2)$, while the standard finite difference provides the order of $O(\Delta t)$ only. Note that in every time step the value of the derivative for the previous time instant is known. At the initial time it is taken from the initial conditions and the continuity equation. Then, by formula (86) one can update $w^t_i$ for $t_{i+1}$.

Below we itemize the basic set of equations employed in the universal algorithm for the transient regime:

- equation (86) to compute the temporal derivative,
- equation defining the reduced velocity in the following form (compare (62))

$$\phi = \frac{L}{w} \int_x^1 \left( \frac{\partial w}{\partial t} + \frac{C_A L(w)}{L^{m+1}} w + q_l \right) d\eta,$$  \hspace{1cm} (87)

- solvability condition resulting from (87) and (63), equivalent to the global fluid balance condition (45) utilized to compute the crucial parameter $L(w)$

$$\int_0^1 \frac{\partial w(t, x)}{\partial t} dx - \frac{q_0(t)}{L(t)} + C_A \frac{L(w)}{L^{m+1}(t)} \int_0^1 w(t, x) dx + \int_0^1 q_l(t, x) dx = 0,$$  \hspace{1cm} (88)

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• equation to compute the crack opening obtained by merging with the respective form of the inverse elasticity operator \( B_w \) (compare (67), (68) or (39))

\[
w = B_w \left( \frac{L(t)}{u^2} \left( \phi + x(L(w) \frac{C_A}{Lm(t)}) \right) \right),
\]  

(89)

• boundary conditions: (43), (61) and (63),

• initial conditions (44),

• relation to compute the crack length: (55), (57) or (58), respectively.

### 6.2 Numerical algorithm for the transient regime

The solution to the transient variant of the problem, described by the system of equations collected above, is sought in the framework of an iterative algorithm. The main idea and assumptions of the numerical scheme are the same as for the self-similar formulation. By analogy to the description given in subsection 2.2, we can define the following stages of computations when looking for an unknown solution in the time instant \( t_{i+1} \):

- **Preliminary step.** The process is initiated by specifying the first approximation of the crack opening \( w^{(j-1)}(t_{i+1}, x) \). One can use here the preconditioning based on the temporal derivative \( w'_t(t_i, x) \) and the initial condition \( w(t_i, x) \). The first approximation of the crack length \( L^{(j-1)}(t_{i+1}) \) can be also easily computed by preconditioning based on the value of the parameter \( L(w_i) \) from the previous time step (or initial conditions).

- **First step.** According to (86) the temporal derivative of the crack opening is computed. Note, that when obtaining the final solution \( w(t_{i+1}, x) \), one automatically has its temporal derivative too. Next, equation (88) yields \( L^{(j)}(w_{i+1}) \), which substituted into (87) gives the reduced particle velocity \( \phi_{i+1}^{(j)} \). The integration in (87) is carried out with application of the \( \varepsilon \)-regularization technique, where the regularized tip condition has the form of (81). As a result, functions \( L^{(j)}(w_{i+1}) \) and \( \phi_{i+1}^{(j)} \) computed at this stage satisfy: i) fluid balance equation (88), ii) continuity equation (87), iii) regularized boundary condition for \( \phi \) (see (81)) which is an equivalent of (61), iv) the influx boundary condition (63) indirectly through the fluid balance equation.

- **Second step.** The crack length is updated by substituting \( L^{(j)}(w_{i+1}) \) into one of (55) – (58). Then, the next approximation of the crack opening \( w^{(j)}_{i+1} \) is obtained from (89). The technique of numerical computation of the operator \( B_w \) is exactly the same as it was for the self-similar variant of the problem. Also here, \( L^{(j)}(w_{i+1}) \) is considered a natural regularization parameter, used to satisfy the influx boundary condition (63).

The aforementioned two steps of the algorithm are repeated until respective components of the solution have converged to within a prescribed tolerance.

**Remark 11.** The modular algorithm architecture enables us to easily introduce the subroutine for the crack length computation as an additional block. Naturally, this block was not present in the self-similar variant of the algorithm.

### 6.3 Analysis of the algorithm performance for the transient regime

In this part of the paper we present a brief investigation into the performance of the universal algorithm for various elasticity models. The aim of this analysis is just to highlight its main peculiarities.
6.3.1 Algorithm performance for the PKN model

We utilize for the computations the benchmark solution described in subsection 2.3.1 as benchmark I for the time dependent term of the power law type (see (107)), where $\gamma = 1/3$. For the transient regime, the accuracy and performance of the algorithm depends on discretization of both independent variables. For this reason, four different variants of spatial and temporal meshing are considered. The number of spatial mesh points is denoted by $N$, while $M$ stands for the number of predefined time steps. We analyze respective combinations for $N = 20$, $N = 100$ and $M = 50$, $M = 100$. The time stepping strategy was taken from [69] (p.162 formula (60)). As previously for the self-similar problem, the spatial mesh density was increased at both ends of the interval.

In the analysis we use the following measures of the solution accuracy: i) the relative error of the crack opening, $\delta w$, ii) the relative error of the reduced particle velocity, $\delta \phi$, iii) the relative error of the crack length, $\delta L$, iv) and finally the relative error of the temporal derivative of the crack opening, $\delta w_t$.

![Figure 13: PKN model. The relative errors of solution for $N = 20$ (spatial mesh), $M = 50$ (temporal mesh): a) the error of crack opening, $\delta w$, b) the error of reduced particle velocity, $\delta \phi$.](image)

![Figure 14: PKN model. The relative errors of solution for $N = 100$ (spatial mesh), $M = 50$ (temporal mesh): a) the error of crack opening, $\delta w$, b) the error of reduced particle velocity, $\delta \phi$.](image)
The computational errors for the crack opening, reduced velocity and the temporal derivative are shown in Fig. 13 – Fig. 18.

When analyzing the solution errors for \( w \) and \( \phi \), one can see that the relation between \( N \) and \( M \) is of crucial importance, while in general the finer meshing produces better results. For example, for \( N = 20 \) there is almost no difference in accuracy between \( M = 50 \) and \( M = 100 \). In this case, the overall solution error is limited by the accuracy resulting from a coarse spatial meshing. However, better stabilization of the error in time can be seen for \( M = 100 \). On the other hand, when the spatial meshing is appreciably improved (\( N = 100 \)), the same change in the time step (from \( M = 50 \) to \( M = 100 \)) yields results up to one order of magnitude better. In all the investigated cases one can see a very low error of \( w \) and \( \phi \) when \( x = 0 \). It shows very good fulfillment of the influx boundary condition (63).

\[
\begin{align*}
\text{Figure 15: } & \text{PKN model. The relative errors of solution for } N = 20 \text{ (spatial mesh), } M = 100 \text{ (temporal mesh): a) the error of crack opening, } \delta w, \text{ b) the error of reduced particle velocity, } \delta \phi. \\
\end{align*}
\]

The graphs for evolution of the crack length error, \( \delta L \), over time are collected in Fig. 17 for all the considered discretization variants. It shows that the accuracy of the crack length is primarily determined by the quality of spatial meshing. It is directly related to the quality of computation of the parameter \( \mathcal{L}(w) \) defining the crack propagation speed (compare (52) – (54)), which in fact is based on the leading asymptotic term(s) of the crack opening. Indeed, better accuracy for the crack opening \( w \) near the crack tip corresponds to better accuracy for the crack length \( L \). Note that a low sensitivity of the results to the time step density for a fixed \( N \) is a direct consequence of using the general relation (51) following from the speed equation.

In the end of this subsection let us discuss the issue of approximating the temporal derivative of the crack opening by formula (86). In Fig. 18 we show the relative errors of \( w_t' \) for two ways of computing it: a) by formula (86), b) by the two-point finite difference (FD). The presented example involves \( N = M = 100 \). As can be seen, although the character of the \( \delta w_t' \) distribution is similar for both variants, the first one gives an error two orders of magnitude lower than the second. For the coarser temporal meshing (\( M = 50 \)) we obtained approximately two times more errors (we do not show this example in separate figures), however the mutual relation between both cases remained the same.

The advantages of using formula (86) in computations instead of the simplest FD scheme becomes less pronounced when one uses a rough spatial mesh. For example, when \( N = 20 \) (\( M = 50, 100 \)) the errors for both approximations are of a similar order (2-4%) to that shown in Fig. 15b).

In general, approximation (86) is not worse than that the two-points FD, giving increasingly greater superiority when refining the computational mesh. Note that in our analysis the FD approximation \( w_t' \) is in fact the post-processing. It is difficult to speculate to what degree it would deteriorate the overall
solution accuracy when implemented in the algorithm instead of \( \delta w \). Such a replacement, however, would neither introduce any simplification to the numerical scheme nor decrease the computational cost. The only imaginable benefit of this would be a slight saving in the memory (as there is no need to store the values of \( w'_t \) from the previous time step).

6.3.2 The algorithm performance for the fluid driven KGD model

In the following we present an analysis of the fluid driven KGD model, in a way analogous to that implemented in the previous chapter. To this end, we utilize the benchmark example already employed in subsection 2.3.2 for the self-similar variant of the problem. The time dependent term taken here, to construct the transient solution, is the power law type (see (107)), with the parameter \( \gamma = 1/3 \). The analyzed variants of mesh densities are defined by \( N = 30, N = 100 \) and \( M = 50, M = 100 \). This time we do not use the the lower value of \( N \) (\( N = 20 \)), as the numerical computation of the inverse elasticity operator \( \delta L \) in such a case becomes more sensitive to the mesh density near the crack tip. As we do not want to include in this paper an additional analysis of the influence of this parameter (mesh density
Figure 18: *PKN model*. The relative error of the solutions temporal derivative for \( N = 100 \) (spatial mesh), \( M = 100 \) (temporal mesh): a) improved temporal approximation b) ordinary finite difference.

in the near-tip region) on the computations we decided to take \( N = 30 \), providing 'fair' comparison for both spatial meshes. However even for \( N = 20 \) the results are still very good, and can be of the same order as for \( N = 30 \) when appropriately adjusting the mesh density near the tip. To access the accuracy, we analyze the same parameters as previously for the PKN model. The results are displayed in Fig. 19 – Fig. 24.

Figure 19: *KGD model fluid driven regime*. The relative error of the solution for \( N = 30 \) (spatial mesh), \( M = 50 \) (temporal mesh): a) the error of crack opening, \( \delta w \), b) the error of reduced particle velocity, \( \delta \phi \).

When analyzing the error of the crack aperture, one can see that a mere 30 points of spatial meshing produces sufficient potential to improve the accuracy by taking more time steps. Indeed, for a fixed number of time steps, \( M = 50 \), the level of the relative error for the crack opening, \( \delta w \), is the same for both \( N = 30 \) and \( N = 100 \) (only some improvement in smoothness of the error distribution can be observed). On the other hand, by using more time steps (\( M = 100 \) instead of \( M = 50 \)), the error of the crack opening, \( w \), can be reduced by an order of magnitude.
Figure 20: KGD model fluid driven regime. The relative error of the solution for $N = 30$ (spatial mesh), $M = 100$ (temporal mesh): a) the error of crack opening, $\delta w$, b) the error of reduced particle velocity, $\delta \phi$.

The situation is quite different, however, for the reduced velocity, $\phi$. In this case, $N = 30$ points of spatial meshing is not enough to provide optimal results especially near the crack tip. By taking $N = 100$, one can appreciably improve the accuracy of the reduced velocity (up to one order of magnitude) and prevent the escalation of the error near the fracture tip. For any particular number of the spatial mesh points, $N$, there exists an optimal number (within assumed time stepping strategy) of the time steps, $M$, at which the maximal achievable accuracy is obtained (saturation level). Further increases in $M$ do not produce better results unless the spatial mesh is refined.

Finally, the error in the crack length, $L$, is almost the same for a fixed number of the time steps $M$ regardless of the spatial mesh under consideration (respective curves in Fig. 20 are hardly distinguishable). The explanation of this fact lies in the quality of computation for the parameter $L(w)$ defining the crack speed: the better accuracy the of $w$ (especially near the crack tip), the better the accuracy of $L$. It is notable that, although this trend is the same as for the PKN model, its realization is obtained by different means: by increasing the number of the time steps $M$ in the PKN case, and by decreasing the step size for the spatial discretization (increasing $N$) for the KGD formulation.

Similarly to that done for the PKN model, we provide here a brief discussion on the application of the time derivative formula (86), comparing it with the standard two-points FD approach. In Fig. 24, we display the distributions of the relative error of $w'$ for $N = M = 100$. Again, the superiority of approximation (86) is clear. The values of $\delta w'$ are of the same level as for the PKN model, however the error distribution for the improved temporal approximation becomes non-uniform. When changing the mesh densities the same tendencies were observed as in the PKN case. Thus, the conclusions drawn beforehand also hold true here.

6.3.3 The algorithm performance for the toughness driven KGD model

In the last part of this subsection we investigate the algorithm performance for a transient regime of the toughness driven KGD model. The benchmark used in this case is constructed in the same way as that for the fluid driven regime. The time dependent term is the same as previously taken for the fluid driven variant. In the analysis of the accuracy of computations, the combinations of spatial and temporal meshing remain the same as in the case of the fluid drive regime. The results are displayed in Fig. 25 – Fig. 29.
The error distributions for $N = 30$ show that, for this number of nodal points, the solution cannot be further improved by taking more time steps (the limiting factor here is the spatial meshing). When analyzing $N = 100$ it turns out that, not only does the error of the solution decrease, but there also exists the potential for increasing the accuracy by taking more densely packed time steps. The graph for $\delta L$, Fig. 29, exhibits rather a surprising result. First, we can see the fluctuations in the crack length error with time for $N = 30$. Respective curves for $N = 100$ are already smooth, however, the level of the error, $\delta L$, does not decrease. As mentioned previously, the quality of the fracture length computation depends on the accuracy of the parameter $\mathcal{L}(w)$ defining the crack speed. In the toughness driven regime, this value utilizes the multipliers of the first two leading terms of the asymptotic expansion of $w$ (instead of the only one leading term in the PKN or the fluid driven KGD models) – see \cite{54}. In this way the
second term, which by its nature is approximated with lower accuracy than the leading one, appreciably affects the error of $L(t)$, limiting its potential for improvement with increasing $N$.

Finally, let us complete the discussion by presenting the results for the temporal derivative of the crack opening. The relative error of $w'_t$ for both methods of approximation (formula (86) and the two-point FD scheme, respectively) are shown in Fig. 30 ($N = M = 100$). As can be seen, the relative error for the improved temporal approximation is of the order $10^{-3}$ – one order of magnitude worse than that previously revealed for the PKN and fluid driven KGD models. However it is still much lower than that of the classic FD approach (see Fig. 30b)). One can also observe in Fig. 30b) a pronounced growth of $\delta w'_t$ at both ends of the spatial interval, which magnifies with time. This trend is caused, to a large extent, by the behaviour of $w'_t$ itself, which yields a time asymptote proportional to $(a + t)^{\gamma - 1}$ (compare (107)). Indeed, in Fig. 31 we show the absolute values of the error of the temporal derivative $\Delta w'_t$, and for formula (86) one obtains its stable (or even decreasing with time) levels at both ends of the spatial interval.
Figure 25: *KGD model toughness driven regime.* The relative error of the solution for $N = 30$ (spatial mesh), $M = 50$ (temporal mesh): a) the error for the crack opening, $\delta w$, b) the error for the reduced particle velocity, $\delta \phi$.

Figure 26: *KGD model toughness driven regime.* The relative error of the solution for $N = 30$ (spatial mesh), $M = 50$ (temporal mesh): a) the error for the crack opening, $\delta w$, b) the error for the reduced particle velocity, $\delta \phi$.

The general trends in the quality of $w_t'$ approximation for the coarser meshes were the same as previously obtained for the PKN and fluid driven KGD models.
7 Discussion and Conclusions

In this paper the classic problem of hydraulic fracture, considered in its general form, has been revisited and reformulated in terms of a new pair of the dependent variables, with both having a clear physical sense: the crack opening, $w$, and the reduced particle velocity, $\phi$. The latter is directly related to the average (over the fracture cross section) speed of fluid flow. It was shown that the new formulation is mathematically complete and well defined. In particular, new equations relating the crack propagation speed to the terms of the crack opening tip asymptotics were evaluated, using the speed equation, for various elasticity operators and fracture propagation regimes. Self-similar formulations of the problem for different hydraulic fracture models have been given. In this setting, a universal algorithm for the
The crack length error, $\delta L$.

The relative error of the solutions temporal derivative for $N = 100$ (spatial mesh), $M = 100$ (temporal mesh): a) improved temporal approximation b) two-point finite difference.

computational simulation of hydraulic fracture has been developed. It enables one to account for various elasticity models, fluid flow and fracture propagation regimes within the framework of a unified scheme.

The proposed algorithm has a modular architecture and consists of two basic modules:

- the first one, a *universal* one, computes the reduced particle velocity, and is the same regardless of the variant of the problem under consideration,

- the second one, evaluating the crack opening, should be adjusted depending on which elasticity model is used. In order to account for a specific crack propagation (fluid/toughness driven) regime, one only needs to adjust the values of the respective parameters which describe the asymptotics of the crack opening and the reduced velocity near the crack tip. Thus, preliminary knowledge of asymptotic behaviour of the solution is needed.
The crack propagation speed is computed using respective explicit formula, (51), derived from the speed equation with utilization of the solvability condition (the fluid balance equation).

Various numerical techniques are utilized in the proposed method with the reduced particle velocity as the main component. The key points of the algorithm are: (i) proper handling of the independent variables (appropriate spatial and temporal meshing), (ii) relevant regularization techniques, in particular the so-called $\varepsilon$-regularization and operator regularization of the governing equations, taking the solvability condition into account when necessary, (iii) explicit formula for the crack propagation speed for the fracture front tracing, (iv) rigorous utilization of the solution tip asymptotics, (v) improved approximation of the temporal derivative of the crack opening.

Extensive analysis of performance of the algorithm, for both its self-similar and the general time-dependent formulations, has been done and comparison made with analytical benchmark solutions developed for various models of hydraulic fracture. Some of them were adopted from the authors previous papers, while others (KGD models) are discussed in Appendix B. Also, reference solutions available in the literature have been used. The following conclusions can be drawn:

(i) the algorithm is numerically stable regardless of the hydraulic fracture model used;

(ii) the accuracy of produced results is appreciably better than that of other solutions available in the literature;

(iii) in most cases the computational cost is very low. A properly distributed spatial mesh composed of several dozen of points provides accuracy better than 0.1%. Only the case of very small toughness may necessitate finer meshing.

To summarize, the universal algorithm developed in this paper is capable of tackling various hydraulic fracturing models under different crack propagation regimes. Its flexibility is a result of its adaptive character and modular code architecture. The method of tracing the fracture front, based on the speed equation, is stable and accurate. The key issue related to its realization is application of the explicit formulae for the crack propagation speed and the solution tip asymptotics. Additionally, taking advantage of the algorithms accuracy, semi-analytic formulae for solving the KGD model in the fluid driven regime have been evaluated, with a solution which yields all necessary components: the crack

\[
\Delta w'_t = \frac{x \times 10^{-4}}{t} 
\]

Figure 31: KGD model - toughness driven regime. The absolute error of the solutions temporal derivative for $N = 100$ (spatial mesh), $M = 100$ (temporal mesh): a) improved temporal approximation b) two-point finite difference.
opening, \( w \), the particle velocity, \( v \), the fluid pressure, \( p \) (the fluid flux should be computed as \( q = wv \)). It was shown that the new algorithm is more accurate than any other available in the literature.

Although we restrict ourselves in this paper only to the Newtonian fluids, the algorithm may be easily adapted to other rheological models. This approach can also be extended to 2D fractures.

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Appendix A: Self-similar solutions

A1: General representation

Let us assume the following separation of variables for the crack opening and the net pressure:

\[ w(t, x) = \psi(t) \hat{w}(x), \quad p(t, x) = \frac{\psi(t)}{L^{m-1}(t)} \hat{p}(x). \tag{90} \]

where \( \psi(t) \) is a smooth continuous function of time and will be specified later. As a consequence, the qualitative asymptotic behaviour of the respective spatial functions in (90) remain the same as their time dependent counterparts (e.g \( \hat{w}(x) \) complies with (47)). The elasticity equations (35), (36) and (39) are transformed to:

\[ \hat{w}(x) = A^{-1} \hat{p}(x), \tag{91} \]

\[ \hat{\bar{w}}(x) = \hat{\bar{p}}(x), \tag{92} \]

\[ \hat{\bar{w}} = -\frac{2}{\pi^2} \int_0^1 \frac{d\hat{\bar{p}}}{ds} K(s, x) ds + \frac{4}{\sqrt{\pi}} \hat{K}_I \sqrt{1 - x^2}, \tag{93} \]

for the PKN and KGD models, respectively. \( \hat{K}_I \) is a self-similar stress intensity factor:

\[ \hat{K}_I = \frac{1}{\pi^{3/2}} \int_0^1 \frac{\hat{p}(\eta)}{\sqrt{1 - \eta^2}} d\eta. \tag{94} \]

Equation (93) is the inversion of the self-similar form of the operator (37):

\[ \hat{p}(x) = -\int_0^1 \frac{d\hat{w}}{d\eta} \frac{\eta d\eta}{\eta^2 - x^2}. \tag{95} \]
The fluid flow rate, particle velocity and reduced particle velocity functions can be expressed as:

\[ q(t, x) = \frac{\psi^4(t)}{L^m} \hat{q}(x), \quad v(t, x) = \frac{\psi^3(t)}{L^m(t)} \hat{v}(x), \quad \phi(t, x) = \frac{\psi^3(t)}{L^m(t)} \hat{\phi}(x), \quad (96) \]

\[ \hat{q}(x) = -\hat{w}^2 \frac{d\hat{p}}{dx}, \quad \hat{v}(x) = \hat{\phi}(x) + x\hat{v}_0, \quad (97) \]

where

\[ \hat{v} = -\hat{w}^2 \frac{d\hat{p}}{dx}, \quad \text{or} \quad \frac{d\hat{p}}{dx} = -\frac{1}{\hat{w}^2} \left( \hat{\phi}(x) + x\hat{v}_0 \right). \quad (98) \]

Based on these assumptions, equation (62) can be transformed into:

\[ \frac{d\psi}{dt} \hat{w} + \frac{\psi^4}{L^{m+1}} \frac{d}{dx} \left( \hat{w} \hat{\phi} \right) + \frac{L'}{L} \hat{\psi} \hat{w} + q_l = 0. \quad (99) \]

Functions \( \psi(t), L(t) \) and \( q_l(t, x) \) should be properly specified in order to eliminate the time variable from the above equation. Then, the problem reduces to the time-independent form.

**A2: Self-similar solution of exponential type**

Let the function \( \psi(t) \) have the form:

\[ \psi(t) = e^{\gamma t}, \quad (100) \]

where \( \gamma > 0 \) is an arbitrary constant. Then, substituting (96) into (55), and taking the initial crack length as:

\[ L(0) = \left[ \frac{(m+1)CA\hat{w}}{3\gamma} \right]^{\frac{1}{m+1}}, \]

one can derive the following relation for \( L(t) \):

\[ L(t) = \left[ \frac{(m+1)CA\hat{w}}{3\gamma} \right]^{\frac{1}{m+1}} e^{\frac{3\gamma}{m+1} t}. \quad (101) \]

Then, if one assumes that the leak-off function complies with the representation:

\[ q_l(t, x) = e^{\gamma t} \hat{q}_l(x), \quad (102) \]

the governing equation (99) after simple transformations can be reduced to:

\[ \frac{1}{CA\hat{w}} \frac{d}{dx} \left( \hat{w} \hat{\phi} \right) = -\frac{m+4}{3} \hat{w} - \frac{m+1}{3\gamma} \hat{q}_l. \quad (103) \]

In this way we obtain an ordinary differential equation equipped with the following boundary conditions:

\[ \hat{w}(1) = 0, \quad \hat{\phi}(1) = 0, \quad \hat{w}(0)\hat{\phi}(0) = \hat{q}_0. \quad (104) \]

Additionally, for the KGD model the following symmetry condition holds:

\[ \frac{d\hat{w}}{dx} \bigg|_{x=0} = 0. \quad (105) \]

The equivalent of the fluid balance equation (45) is:

\[ \frac{1}{CA\hat{w}} \hat{q}_0 - \frac{m+4}{3} \int_0^1 \hat{w} dx - \frac{m+1}{3\gamma} \int_0^1 \hat{q}_l dx = 0. \quad (106) \]
A3: Self-similar solution of the power law type

In this variant of the self-similar solution let us take:

\[ \psi(t) = (a + t)\gamma, \quad (107) \]

where \( a \geq 0 \) and \( \gamma > 0 \) are some constants. In the same way as previously, one can obtain a relation for the crack length in the form:

\[ L(t) = \left[ \frac{(m + 1)C_A L(\hat{w})}{3\gamma + 1} \right]^{\frac{1}{m+1}} (a + t)^{\frac{3\gamma + 1}{m+1}}, \quad (108) \]

provided that:

\[ L(0) = \left[ \frac{(m + 1)C_A L(\hat{w})}{3\gamma + 1} \right]^{\frac{1}{m+1}} a^{\frac{3\gamma + 1}{m+1}}. \]

Additionally if the leak-off function can be expressed as:

\[ q_l(t, x) = (a + t)^{\gamma - 1} \hat{q}_l(x), \quad (109) \]

equation (99) converts to:

\[ \frac{1}{C_A L(\hat{w})} \frac{d}{dx}(\hat{w} \hat{\phi}) = -\left[ \frac{(m + 1)\gamma}{3\gamma + 1} + 1 \right] \hat{w} - \frac{m + 1}{3\gamma + 1} \hat{q}_l. \quad (110) \]

Boundary conditions for the above differential equation remain the same as (104) - (105). Finally, the global fluid balance equation can be transformed to:

\[ \frac{1}{C_A L(\hat{w})} \hat{q}_0 - \left[ \frac{(m + 1)\gamma}{3\gamma + 1} + 1 \right] \int_0^1 \hat{w}dx - \frac{m + 1}{3\gamma + 1} \int_0^1 \hat{q}_l dx = 0. \quad (111) \]

Appendix B: Construction of benchmarks solutions

In the following we introduce a set of analytical benchmark solutions for the considered problem. The ideas behind their construction are the same regardless of the elasticity model in use, and have already been employed in [51, 69, 38] for PKN model. The basic idea is to use an analytical solution to the self-similar problem defined in Appendix A, and extended it into the time-dependent form using relations (90), (96) – (97) and (101) or (108).

Let us concentrate now on finding some examples of analytical solutions to the self-similar equation:

\[ \frac{1}{L(\hat{w})} \frac{d}{dx}(\hat{w} \hat{\phi}) = -\chi \hat{w} - \kappa \hat{q}_l, \quad (112) \]

which is a generalization of (103) and (110). The corresponding constants \( \chi \) and \( \kappa \) in (103) can be determined by direct comparison with equations (103) and (110), and are given in Table 2.

The boundary conditions (104) are to be satisfied, together with their respective form of the balance equation (106) or (111). Depending on the elasticity model one of the operators (91) shall be in use.

The general idea behind constructing a benchmark solution is quite straightforward. At first, assume that the crack opening function can be expressed as a weighted sum of properly chosen base functions:

\[ \hat{w}(x) = \sum_{i=0}^{N} \lambda_i \hat{w}_i(x). \quad (113) \]
The functions \( \hat{w}_i(x) \) should be selected in a way which enables one to: i) comply with the respective asymptotic representation (47), ii) analytically compute the pressure operators (91), iii) satisfy the boundary conditions (104) (and (105) for KGD). Provided that ii) is fulfilled, the pressure function can be calculated in a closed form from (91) to give:

\[
\hat{p}(x) = \sum_{i=0}^{N} \lambda_i \hat{p}_i(x),
\]

where each of the functions \( \hat{p}_i(x) \) corresponds to the respective function \( \hat{w}_i(x) \). Then, according to (98)\textsuperscript{1} the particle velocity is defined:

\[
\hat{v}(x) = -\left[ \sum_{i=0}^{N} \lambda_i \hat{w}_i(x) \right]^2 \sum_{i=0}^{N} \lambda_i \frac{d}{dx} \hat{p}_i(x),
\]

and consequently we can define its leading term as:

\[
\hat{v}_0 = \hat{v}(1) = C_A \mathcal{L}(\hat{w}). \tag{116}
\]

The reduced velocity is determined using (115) - (116) according to (97)\textsuperscript{2}:

\[
\hat{\phi}(x) = -\left[ \sum_{i=0}^{N} \lambda_i \hat{w}_i(x) \right]^2 \sum_{i=0}^{N} \lambda_i \frac{d}{dx} \hat{p}_i(x) - xC_A \mathcal{L}(\hat{w}). \tag{117}
\]

Next, by substituting (113) and (117) into equation (112), we can define the benchmark leak-off function \( \hat{q}_l(x) \). Finally the value of \( \hat{q}_0 \) is determined by substituting (113) and (117) into the boundary condition (104)\textsuperscript{3}:

\[
\hat{q}_0 = -\left[ \sum_{i=0}^{N} \lambda_i \hat{w}_i(0) \right]^3 \sum_{i=0}^{N} \lambda_i \frac{d\hat{p}_i}{dx}\bigg|_{x=0}. \tag{118}
\]

In this way, the analytic benchmark solution is fully defined by (113), (117), corresponding leak-off function and corresponding influx value (118). Clearly, the fluid balance equation ((106) or (111)) is satisfied automatically.

**B1: PKN model**

For the PKN model let us adopt the following \( N + 1 \) test functions \((i = 0, 1, ..., N)\):

\[
\hat{w}_i(x) = (1 - x)^{i+1/3}, \quad i \leq N - 1, \quad \hat{w}_N(x) = e^x(1 - x)^{N+1/3}. \tag{119}
\]

The base function \( \hat{w}_N(x) \) was taken in the specified form in order to introduce an additional non-linear effect to the benchmark, without violating the asymptotic behaviour.

By applying representation (119) in (113) and elasticity operator (91)\textsuperscript{1} one obtains the formula for the pressure function, \( \hat{p}(x) \), which after differentiation yields:

\[
\frac{d}{dx} \hat{p}(x) = -\left[ \sum_{i=0}^{N-1} \lambda_i (i + 1/3)(1 - x)^{i-2/3} + \lambda_N (N + x - 2/3)e^x(1 - x)^{N-2/3} \right]. \tag{120}
\]

Then by formulae (115)-(118) one can construct the benchmark solution, taking \( \mathcal{L}(\hat{w}) = \lambda_0^3 \).
In general, the leak-off behaviour near the crack tip can be controlled by the powers of the base functions \( \hat{w}_i \), for \( i > 1 \) (e.g. if one wants to mimic the Carter law, each power starting from the second one should be 1/6 greater than the previous - compare [38]). The representation used in this paper gives:

\[
\hat{q}_t(x) = O((1 - x)^{\alpha_0}), \quad x \to 1,
\]

which is the same as the asymptotics of the crack opening itself. However, even in such a case, proper manipulation of the multipliers can provide better behaviour of \( \hat{q}_t \).

**B2: KGD model: fluid driven regime**

For this model we assume that the self-similar crack aperture is defined by the following base functions:

\[
\hat{w}_i(x) = (1 - x^2)^{\alpha_i} C_{2(i+1)-2}^{\alpha_i-1/2}(x), \quad i \leq N - 1, \quad \hat{w}_N(x) = \sqrt{1 - x^2} - \frac{2}{3} (1 - x^2)^{3/2} - x^2 \ln \frac{1 + \sqrt{1 - x^2}}{x},
\]

where \( C_{2(i+1)-2}^{\alpha_i-1/2}(x) \) is the ultraspherical or Gegenbauer polynomial. The term \( \hat{w}_N(x) \) was introduced to obtain a non-zero pressure gradient for \( x = 0 \). In our case we take in computations \( N = 2 \), where \( \alpha_0 = 2/3, \alpha_1 = 5/3 \). Note that:

\[
\hat{w}_N(x) = O\left((1 - x^2)^{5/2}\right), \quad x \to 1.
\]

A representation similar to (122) (except for the last term) was used in [2] to define the base functions for the series approximation of solution. This general representation was utilized in order to solve the problem for a variety of shear-thinning fluids. In a similar way (122) with the proper values of \( \alpha_i \) can be employed to construct benchmark solutions for non-Newtonian fluids.

By applying (122) in (95) and subsequent differentiation one obtains an analytical formula for the pressure gradient:

\[
\frac{d}{dx} \hat{p}(x) = -\sum_{i=0}^{N-1} \lambda_i \alpha_i (2\alpha_i - 1) B(1/2, \alpha_i) x \cdot _2F_1(3/2 - \alpha_i, 2; 3/2; x^2) + \lambda_N (x - \pi/4),
\]

where \( B(1/2, \alpha_i) \) is the beta function, and \( _2F_1 \) is the Gauss hypergeometric function.

Then from (115) – (118) we have the complete benchmark solution, where \( L(\hat{w}) = \lambda_0^2 \).

**B3: KGD model: toughness driven regime**

In the case of the toughness driven KGD model we consider the benchmark where the crack opening is represented by a sum of four functions:

\[
\hat{w}_i(x) = (1 - x^2)^{\frac{i}{2}}, \quad i = 1, 2, \quad \hat{w}_3(x) = (1 - x^2)^{\frac{3}{2}} \ln (1 - x^2), \quad \hat{w}_4(x) = 2\sqrt{1 - x^2} + x^2 \ln \frac{1 - \sqrt{1 - x^2}}{1 + \sqrt{1 - x^2}}.
\]

The above representation is consistent with (117) and Table I. The special term \( \hat{w}_N \) gives a non-zero pressure gradient at \( x = 0 \) and:

\[
\hat{w}_N(x) = O\left((1 - x^2)^{3/2}\right), \quad x \to 1.
\]

Obviously it can be replaced by \( \hat{w}_n \) from (122) if convenient. Also further terms of type \( \hat{w}_i \) from (122) can be used here, provided that the powers \( \alpha_i \) are greater than those from the leading terms of the asymptotics.
Then by operator $\mathcal{H}_2$ one obtains respective pressure components $\hat{p}_i$ from (114), where $\hat{p}_0(x) = \pi/2$. Thus, the non-zero components of the pressure derivative are:

$$
\hat{p}'_1(x) = 0, \quad \hat{p}'_2(x) = -\frac{2}{x} \left[ -\frac{1}{1-x^2} + 2 F_1(-1/2, 1; 1/2; x^2) \right],
$$

(127)

$$
\hat{p}'_3(x) = \pi x (5 - 6 \ln 2) + 3 \pi \arcsin(x) \left( \sqrt{1-x^2} - \frac{x^2}{\sqrt{1-x^2}} \right), \quad \hat{p}'_4(x) = -\pi^2
$$

Finally, by applying (125) and (127) in (115) – (118) one constructs the benchmark solution, where $\mathcal{L}(\hat{\omega}) = \lambda_0^2 \lambda_1$. Moreover, decreasing/increasing this value, one tests the algorithms on the small/large toughness regimes. Finally, in the case of KGD problems (both regimes), the leak-off function, $q_l$, constructed above for the benchmarks, generally speaking, will behave in the same manner as in (121).