The Stable Explicit Time Stepping Analysis with a New Enrichment Scheme by XFEM

Xue-cong Liu¹, Qing Zhang¹,*, Xiao-zhou Xia¹

Abstract: This paper focuses on the study of the stability of explicit time integration algorithm for dynamic problem by the Extended Finite Element Method (XFEM). A new enrichment scheme of crack tip is proposed within the framework of XFEM. Then the governing equations are derived and evolved into the discretized form. For dynamic problem, the lumped mass and the explicit time algorithm are applied. With different grid densities and different forms of Newmark scheme, the Dynamic Stress Intensity Factor (DSIF) is computed by using interaction integral approach to reflect the dynamic response. The effectiveness of the proposed scheme is demonstrated through the numerical examples, and the critical time stepping in different situations are listed and analyzed to illustrate the factors that affect the numerical stability.

Keywords: XFEM, DSIF, Newmark scheme, time stepping, stability.

1 Introduction

The fracture analysis of structures and components has been widely applied and highly valued in recent years, and modeling discontinuities like crack is one of the important parts. In order to model the behavior of discontinuities, the way of re-meshing is used which is to align the element edges with the discontinuities by classic finite element method (FEM). However, for the case of crack arbitrary growth, the mesh changes at every step and incurs high computation cost. Besides, other solutions such as meshfree method, boundary element method and extended finite element method (XFEM) are available, and XFEM is considered to be the most applicable.

As proposed by Belytschko and Black (1999); Moes, Dolbow and Belytschko (1999); Belytschko and Moe (2001), XFEM becomes a dominant numerical scheme nowadays. XFEM is based on the concept of partition unity, and the crack can be modeled independent of finite element mesh. All the elements are divided into the normal parts

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and the enriched parts. For the enriched parts, the elements can be influenced directly by crack, so the enrich functions are introduced. As the most used enrich functions, Heaviside function and the Westergaard stress function are used frequently for the cracks and cracks’ tips, respectively.

Dealing with the dynamic fracture, Belytschko, Chen, Xu et al (2003) proposed a kind of tip element in which the crack opens linearly and developed a propagation criterion with loss of hyperbolic. Later on, a singular enrichment function for tips is proposed for the elastodynamic cracks with explicit time integration scheme [Belytschko and Chen (2004)]. In order to deeply study the stability and energy conservation and to get a more accurate result, Réthoré, Gravouil and Combescure (2005a, 2005b) combined Space and Time XFEM (STX-FEM) to obtain a unified space-time discretization, and concluded that the STX-FEM is a suitable technique for dynamic fracture problems. On the other hand, a new lumping technique for mass matrix was proposed in order to be more suitable for dynamic problems by Menouillard, Réthoré, Combescure et al (2006); Menouillard, Réthoré, Noes et al (2008) and the robustness and the stability of this approach has been proved.

As we noticed, the previous studies are all based on the classical enrichment scheme, and a large number of additional degrees of freedom (DOFs) are required. In the meantime, various improved enrichment methods have been studied. Song, Areias and Belytschko (2006) has reinterpreted the conventional displacement field, described discontinuities by using phantom nodes and superimposed extra elements onto the intrinsic grid for dynamic fracture problems. The method doesn’t require subdomain integration for the discontinuous integrand and has a highly efficient but nevertheless quite accurate formulation. Further, Duan, Song, Menouillard et al (2009) has shown its practicability on the shell problem as well as three-dimension problem [Song and Belytschko (2009)]. Besides, changing the basic enrichment function is another solution. Menouillard, Song, Duan et al (2010) proposed a new enrichment method with only a singular enrichment, which shows great accuracy for stationary cracks. The similar research has been done by Rabczuk, Zi, Geretenberger et al (2008). Without crack tip enrichment, the Heaviside function has also been improved. Nistor, Pantale and Caperaa (2008) used only Heaviside function to model the dynamic crack growth. Kumar, Singh, Mishra et al (2015) presented a new approach based Heaviside function along with a ramp function which contains information like crack length and angle. A similar method was proposed by Wen and Tian (2016); Tian and Wen (2016), which is based on an extra-dof-free partition of unity enrichment technique, and no more extra DOFs are added in the dynamic crack growth simulation.

For all the study discussed above, the stability of the method is always concerned. Generally, using explicit scheme for dynamic problem, one goes through a very small time stepping that leads to high computation cost, while with a larger one the numerical result may be divergent. So in the present paper, we will focus on the stability of the numerical scheme. Based on the analytical solution of the displacement fields near crack tip, a new enrichment scheme is used for the elements influenced by crack tip. The Newmark scheme is adopted for time integration, and different parameters are tested to investigate their influence on the stability. In addition, DSIF is calculated as an important parameter which represents the variation of the stress field around the crack tip, and also
can determine the stability of the simulation.

This paper is organized as following: Section 2 illustrates the governing equations and the XFEM formulation; The explicit time algorithm and the lumping technique are introduced in Section 3; The DSIF is shown in Section 4; In Section 5, several numerical examples are provided, the feasibility and stability of the simulation are discussed.

2 Governing equations and XFEM formulation

2.1 Governing equations

Let $\Omega$ be a homogeneous two-dimensional domain with cracks. $\Omega$ is bounded by $\Gamma$, which is composed of $\Gamma_u, \Gamma_t$, and $\Gamma_c$ as described in Fig.1. $\Gamma_u$ denotes the prescribed displacements and $\Gamma_t$ denotes the traction in the domain. $\Gamma_c$ is referred to as the crack surface and assumed to be traction-free. The strong form of the linear momentum equation and the boundary conditions give

$$\nabla \sigma + b = \rho \ddot{u} \quad \text{in} \quad \Omega$$  \hspace{1cm} (1a)

$$u = \bar{u} \quad \text{on} \quad \Gamma_u$$ \hspace{1cm} (1b)

$$\sigma \cdot n = \bar{t} \quad \text{on} \quad \Gamma_t$$ \hspace{1cm} (1c)

$$\sigma \cdot n = 0 \quad \text{on} \quad \Gamma_c$$ \hspace{1cm} (1d)

$$\varepsilon = \nabla_u$$ \hspace{1cm} (1e)

$$\sigma = C:\varepsilon$$ \hspace{1cm} (1f)

where $\sigma$ is prescribed the Cauchy stress tensor, $u$ and $\ddot{u}$ are the vector of displacement field and acceleration field, respectively. $b$ is the body force vector, $\rho$ is the mass density, $\bar{u}$ is the prescribed displacement, $n_r$ is the unit outward normal, $\nabla_u$ is the symmetric operator, $C$ is the elastic module tensor and $\varepsilon$ is the strain tensor. The small strains and displacements are considered here as shown in Eq.(1e). Combine the equilibrium momentum equation and the constitutive relation into the weak form, we have

$$\int_{\Omega} \delta \nabla_u : C \nabla_u \Omega + \int_{\Omega} \rho \ddot{u} \delta u \Omega = \int_{\Omega} \delta \ddot{u} b \delta u \Omega + \int_{\Gamma_t} \delta \ddot{u} \bar{t} \delta \Gamma$$ \hspace{1cm} (2)

2.2 The XFEM formulation

Consider a typical finite element mesh with four-node elements as shown in Fig.2, the geometry of crack is independent of the mesh. As in the classical XFEM, each node enriched by Heaviside function $H(x)$ have two additional DOFs. $H(x)$ is defined as a unit magnitude for the elements cut completely by crack, and takes $\pm 1$ on the two sides of the crack. For the case of the nodes with crack tip enrichment, eight additional DOFs are needed for each. The basic enrichment functions are inspired from the analytic solution of near tip displacement fields of mode I and mode II cracks, and can be written as the
functions \( \psi_k (r, \theta) \) [Belytschko and Black (1999)]

\[
\psi_k (r, \theta) = \sqrt{r} \left\{ \cos \left( \frac{\theta}{2} \right), \sin \left( \frac{\theta}{2} \right) \cos(\theta), \cos \left( \frac{\theta}{2} \right) \sin(\theta) \right\}, \quad k = 1, 2, 3, 4
\]

(3)

where \((r, \theta)\) are the local polar coordinates at the crack tip. We note that the second function in Eq. (3) is commonly referred to branch function, while the others are continuous and added to improve the accuracy. So \( \psi_k (r, \theta) \) can be presented with other function set or just with the branch function, and it is well-documented and verified by Dolbow, Noes and Belytschko (2000).

**Figure 1:** Domain with cracks and prescribed boundary conditions

**Figure 2:** Typical discretization of a domain with crack and enriched nodes by XFEM

As mentioned above, the enrichment functions are developed based the displacement fields near the crack tip, and can have different forms. In the present paper, a new form of enrichment functions is used, which derives from the displacement fields directly. By shifting the enrichment functions, we are able to correspond the enriched nodes’ displacement to the true displacement with XFEM. The displacement can be written approximately as
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\[ \mathbf{u}(x) = \sum_{i=1}^{I} N_i(x) \mathbf{u}_i + \sum_{j=1}^{J} N_j(x) \left[ \mathbf{H}(x) - \mathbf{H}(x_j) \right] \mathbf{a}_j + \sum_{k=1}^{K} N_k(x) \mathbf{b}_k \left[ \mathbf{F}(x) - \mathbf{F}(x_k) \right] \mathbf{b}_k \] (4)

where \( I \) is the set of all nodes in the mesh, \( J \) is the set of nodes enriched by Heaviside function and \( K \) is enriched by \( \mathbf{F}(x) \). \( N_i(x) \) is the standard FEM shape functions. \( \mathbf{a}_j \) and \( \mathbf{b}_k \) are the addition DOFs associated with \( \mathbf{F}(x) \) and \( \mathbf{F}(x_k) \), respectively. \( \mathbf{b}_k \) is the transformation enrichment function in matrix form

\[ \mathbf{F}(x) = \frac{1}{2G} \sqrt{\frac{r}{2\pi}} \begin{bmatrix} \cos \frac{\theta}{2} (\kappa - \cos \theta) & \sin \frac{\theta}{2} (\kappa - 2 + \cos \theta) \\ \sin \frac{\theta}{2} (\kappa - \cos \theta) & \cos \frac{\theta}{2} (\kappa - 2 + \cos \theta) \end{bmatrix} \] (5)

where \( G \) is the shear modulus of material, \( G = \frac{E}{2(1+\nu)} \). \( E \) is the modulus of elasticity, \( \nu \) is the Poisson’s ratio. \( \kappa \) is the Kolosov constant, \( \kappa = \begin{cases} 3-4\nu, & \text{planestrain} \\ 3-\nu, & \text{planestress} \end{cases} \).

It is important to point out that although \( \mathbf{F}(x) \) still has 4 base functions, the number of additional DOFs is reduced from 8 to 2 for the nodes enriched by crack tip. The two base functions in first row of matrix \( \mathbf{F}(x) \) are associated with the horizontal axis in the local coordinate at crack tip, and the second row corresponds to the vertical axis.

Without concerning the damping effect, we substitute the displacement field Eq.(4) into the weak form Eq.(2). It then yields a system of linear algebraic matrix equations, which can be expressed as

\[ \mathbf{M} \dot{\mathbf{U}} + \mathbf{KU} = \mathbf{f} \] (6)

where \( \mathbf{M} \) is the mass matrix, \( \mathbf{K} \) is the stiffness matrix and \( \mathbf{f} \) is the force matrix:

\[ \mathbf{M} = \begin{bmatrix} \mathbf{M}^{uu} & \mathbf{M}^{ua} & \mathbf{M}^{ut} \\ \mathbf{M}^{au} & \mathbf{M}^{aa} & \mathbf{M}^{at} \\ \mathbf{M}^{tu} & \mathbf{M}^{ta} & \mathbf{M}^{tt} \end{bmatrix} \] (7.a)

\[ \mathbf{K} = \begin{bmatrix} \mathbf{K}^{uu} & \mathbf{K}^{ua} & \mathbf{K}^{ut} \\ \mathbf{K}^{au} & \mathbf{K}^{aa} & \mathbf{K}^{at} \\ \mathbf{K}^{tu} & \mathbf{K}^{ta} & \mathbf{K}^{tt} \end{bmatrix} \] (7.b)

\[ \dot{\mathbf{U}} = \begin{bmatrix} \dot{\mathbf{u}} \\ \dot{\mathbf{a}} \\ \dot{\mathbf{t}} \end{bmatrix}, \quad \mathbf{U} = \begin{bmatrix} \mathbf{u} \\ \mathbf{a} \\ \mathbf{t} \end{bmatrix}, \quad \mathbf{f} = \begin{bmatrix} \mathbf{f}^{uu} \\ \mathbf{f}^{ua} \\ \mathbf{f}^{at} \end{bmatrix} \] (7.c)

The sub-matrices and vectors that come in the foregoing equations are defined as below for four-node element (\( i, j = 1, 2, 3, 4 \)):
\[
M_{ij}^{rs} = \int_{\Omega} \rho \left( N_i \right)^T N_j \Omega, \quad r, s = u, a, t \\
K_{ij}^{rs} = \int_{\Omega} \left( B_i \right)^T B_j \Omega, \quad r, s = u, a, t \\
f_i^{rs} = \int_{\Omega} N_i b d\Omega + \int_{\Gamma_r} N_i \tilde{t} d\Gamma \\
f_i^{rs} = \int_{\Omega} N_i [H(x) - H(x_i)] b d\Omega + \int_{\Gamma_r} N_i [H(x) - H(x_i)] \tilde{t} d\Gamma \\
f_i^{rs} = \int_{\Omega} N_i \beta_T [F(x) - F(x_i)] b d\Omega + \int_{\Gamma_r} N_i \beta_T [F(x) - F(x_i)] \tilde{t} d\Gamma
\]

where \( N_i^a \) and \( N_i^b \) are the matrices of additional shape functions in XFEM, \( B_i^a \) and \( B_i^b \) are the matrices of shape function derivatives and can be expressed as:

\[
N_i^a = \left[ \begin{array}{cc} H(x) - H(x_i) & 0 \\ 0 & H(x) - H(x_i) \end{array} \right]
\]

\[
N_i^b = N_i \beta_T [F(x) - F(x_i)]
\]

\[
B_i^a = \left[ \begin{array}{cc} \left( N_i (H(x) - H(x_i)) \right)_x & 0 \\ 0 & \left( N_i (H(x) - H(x_i)) \right)_y \end{array} \right]
\]

\[
B_i^b = \left[ \begin{array}{cc} \frac{\partial}{\partial x} & 0 \\ 0 & \frac{\partial}{\partial y} \end{array} \right] \left( N_i \beta_T [F(x) - F(x_i)] \right)
\]
3 The explicit time integration

3.1 Time integration

As the most commonly used for dynamic problems, the Newmark scheme is chosen as the time integration algorithm. As we know, the time integration algorithm can be divided into two types: the explicit and the implicit. With the implicit method, there is no intrinsic limit to the time step, but it needs to solve the global equations by iterating in each step. For dynamic problems, lots of iterative calculations are needed, using the implicit scheme can bring many disadvantages such as vast computation and low efficiency. Compared with the implicit scheme, the explicit scheme which is chosen in the present paper can solve the equations independently with no iterative.

For explicit scheme, two parameters $\beta$ and $\gamma$ are considered. Combine the Newmark scheme and Eq.(6), the derived equation can be given as

$$U_{t+\Delta t} = U_t + \Delta t \dot{U}_t + \Delta t^2 \left( \frac{1}{2} - \beta \right) \ddot{U}_t + \Delta t^2 \beta \dddot{U}_{t+\Delta t} \tag{13.a}$$

$$M \ddot{U}_{t+\Delta t} = F_{t+\Delta t} - KU_{t+\Delta t} \tag{13.b}$$

$$\dot{U}_{t+\Delta t} = U_t + \Delta t (1 - \gamma) \dot{U}_t + \Delta t \gamma \dddot{U}_{t+\Delta t} \tag{13.c}$$

where $\Delta t$ is the time step, $U_t$ is the vector of displacement, $\dot{U}_t$ and $\dddot{U}_t$ are the vector of velocity and acceleration at time $t$, respectively.

For a numeric scheme, the stability, consistency and convergence are the main reference standards. From now on, we will focus on the stability of Newmark scheme because of the instability is a sufficient condition for non-convergence. The stability conditions of Newmark scheme are deduced in detail by Réthoré, Gravouil and Combescure (2004) with their custom notations:

1. if $\frac{1}{2} \leq \gamma \leq 2\beta$, it is an unconditionally stable scheme.

2. If $\frac{1}{2} \leq \gamma$ and $2\beta \leq \gamma$, the stable condition is $\Delta t \leq \frac{1}{\phi_{max} \sqrt{\frac{\gamma}{2} - \beta}}$.

where $\phi_{max}$ is the maximum frequency of the structure.
As mentioned by Réthoré, Gravouil and Combescure (2004), if the parameters are chosen to be $\beta = \frac{1}{4}$ and $\gamma = \frac{1}{2}$, the updating equations are unconditionally stable. However, according to Eq.(13), if the acceleration item $\dot{U}_{t+\Delta t}$ exists in Eq.(13.a), the equations still need to be solved iteratively. Thus, for an explicit time integrator, $\beta = 0$ is used in the present paper. We noticed that if $\beta = 0$ and $\gamma = \frac{1}{2}$, the Newmark scheme evolves into the central difference method.

Furthermore, due to such a restriction of stability condition, there must be a critical time step $\Delta t_c$. With the time step $\Delta t$ beyond the critical value, the numerical instability and convergence problem will happen at some point. In contrast, the numerical results is very stable within the critical time. We will focus on figuring out the critical time step, and finding out the factors that can affect it in this paper. Tests with different grid densities and different parameters in the Newmark scheme will be conducted.

### 3.2 The lumped mass

The matrix above in Eq.(8) is known as the consistent mass matrix, which includes standard terms, block-diagonal enriched terms, and coupling terms [Menouillard, Réthoré, Combescure et al (2006)]. However, for the problem of dynamic, the lumped mass matrix is used more frequently in order to simplify the numerical calculations. Due to the existence of additional DOFs which have no clear physical significance, the distribution of mass is not just as a simple average as in traditional FEM. Menouillard, Réthoré, Noes et al (2008) had in-depth study of the lumping technique for the mass matrix based on the conservation of mass and momentum, and proved its effectiveness with explicit scheme for dynamics by XFEM. Besides, the lumping technique was also researched by Zi, Chen, Xu et al (2005); Elguedj, Gravouil and Maigre (2009); Song and Belytschko (2009); Jim, Zhang, Fang et al (2016). In this paper, the lumped mass proposed by Menouillard, Réthoré, Combescure et al (2006) is used

$$m_{\text{diag}} = \frac{m}{n_{\text{node}}} \frac{1}{\text{mes}(\Omega)} \int_{\Omega} H^2 \text{d}\Omega$$

(14)

where $\Omega$ is the element being considered, $m$ is the element’s mass, $\text{mes}(\Omega)$ is the area of element in 2D, $n_{\text{node}}$ is the number of nodes in element, and $H$ is the Heaviside function.
4 DSIF

In this section, the DSIF is illustrated. Based on energetic consideration, the SIF is used as a parameter of the strength of singularity, and some questionable relevant quantities of crack tip such as stress fields are avoided. There are a few schemes to calculate the SIF, such as the displacement extrapolation method, the virtual crack extension method, the virtual crack closure method and the interaction integral method. Due to the research of Nagashima, Omoto and Tani (2003), the interaction integral method has the highest accuracy and is used here. In the interaction integral method, the auxiliary fields are introduced and superimposed onto the actual fields.

For dynamic loading case, an item related to inertia is added, and the interaction integral with force-free on crack surface can be given as

$$I = \int_A \left( \sigma_{ij} u_{i,1}^{\text{aux}} + \sigma_{ij}^{\text{aux}} u_{i,1} - \sigma_{ij}^{\text{aux}} \delta_{ij} \right) q_{ij} \, dA + \int_A \rho u_i u_{i,1}^{\text{aux}} q \, dA \quad (15)$$

where \(\sigma_{ij}, \varepsilon_{ij}, u_i\) and \(\sigma_{ij}^{\text{aux}}, \varepsilon_{ij}^{\text{aux}}, u_i^{\text{aux}}\) are the actual state and the auxiliary state, respectively. \(I\) is the interaction integral between the actual state and the auxiliary state, \(A\) is the integral domain. \(q\) is the weight function which is going to be 0 outside the contour boundary and is one inside in the present paper. The DSIF can be written as

$$\begin{cases} K_I = I^{\text{mode I}} \cdot E^*/2 \\ K_{II} = I^{\text{mode II}} \cdot E^*/2 \end{cases} \quad (16)$$

where \(E^*\) is equal to \(E\) for the plane stress and for the plane strain \(E^* = E/(1 - \nu^2)\).

The basic algorithm used here for the DSIF is concluded as following:

1. Give an integral rang \(R\), then search for all the integral elements;
2. Loop through all the integral elements;
3. Loop through all the Gauss points in each integral element;
4. Calculate the actual state and the auxiliary state of each Gauss point;
5. Get the value of DSIF through Eq.(15) and Eq.(16).

where \(R\) is the ratio between the actual integral radius \(r\) and the minimum size \(L_{\text{min}}\) of all elements as shown in Fig.3.
Figure 3: The integral elements for DSIF

5 Numerical examples

5.1 Stationary mode I crack

First, in order to illustrate the effectiveness of the new enrichment scheme presented above, we study the problem of an infinite plate contains a semi-infinite crack as shown in Fig.4. A prescribed vertical loading of $\sigma_0 = 500\text{MPa}$ is applied to the upper surface. The evolution of the loading is a type of Heaviside step function. For the geometry of configuration, a rectangular plate of size $L \times 2H = 10\text{m} \times 4\text{m}$ with an initial edge crack of length $a = 5\text{m}$ is used. Some material parameters are: Young’s modulus $E = 210\text{GPa}$, Poisson’s ratio $\nu = 0.3$, the density $\rho = 8000\ \text{kg/m}^3$. A mesh of $39 \times 99$ uniform square is used for tests. A theoretical solution of this problem with a stationary crack was obtained by [Freund (1990)] and it is given by

$$K_i(t) = 2\sigma_0 \frac{\sqrt{c_d t(1-2\nu)}}{1-\nu} \quad (17)$$

where $c_d$ is the dilatational wave speed, $t = 0$ is the time that the stress wave reach the crack tip from the edge. The theoretical solution is used to compare our present results, however, it has some limitation, such as being effective only in $t < 3t_c$ when the
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reflected stress wave reaches the crack tip, where \( t_c = \frac{H}{c_d} \).

Fig. 5 presents the values of DSIF with different integral domains. The time step is chosen as \( \Delta t = 0.1 \mu s \). The DSIF is normalized by \( K_0 = \sigma \sqrt{a} \) and the numerical time is normalized by \( t_c \). At the beginning, time \( t < t_c \), due to the stress wave has not reach the integral domain, the value of DSIF is 0. Then, after stress wave reaches the crack tip, the results with different integral domains are in good agreement with each other as shown in Fig. 5. The theoretical solution for comparison has also been plotted, and shows good accuracy to the present result.

Fig. 6 presents the results of DSIF with different time stepping \( \Delta t \) while \( R = 5 \). It shows good consistency and the results are not sensitive to the time step. The results inspire us to improve the computational efficient with a larger step time which is less than the critical time. However, with a much larger time stepping for test, \( \Delta t = 20 \mu s \), the numerical result is rapidly divergent. As a consequence, it must have a critical time stepping, which we will discuss it later.

In addition, the result of DSIF with four crack tip enrichment functions of standard XFEM is also plotted for comparison in Fig. 6. It can be seen that the new enrichment scheme offers the almost same accuracy as XFEM with standard enrichment functions.

![Figure 4: The geometry and loading of a homogeneous material plate with crack](image-url)
5.2 Finite size edge plate with an arbitrarily oriented central crack

In this example, a plate with a central crack under uniaxial tension is considered. As shown in Fig.7, the dimensions of the plate are \(2h=0.04\) m and \(2b=0.02\) m, and the length of the central crack is \(2a=0.0048\) m. The material’s properties are: \(E=199.99\) GPa, \(\nu=0.3\), \(\rho=5000\) kg/m\(^3\). A prescribed vertical loading of \(\sigma_0=100\) MPa is applied to the upper surface and the lower surface. A mesh of \(49 \times 99\) uniform elements is used.

First of all, a horizon central crack \(\theta=0^\circ\) is considered. For the cases of different integral domains, Fig.8 shows that the range \(R\) has little effect on the crack tip’s DSIF. The results agree very well with the conclusion that the SIF under different integral path are the same. The results are also compared with the standard XFEM with four crack tip enrichment functions, and shows good accuracy to the present result.

Fig.9 presents the numerical results with different time step, the same conclusion can be drawn as in Fig.6. In additional, the numerical result given by Song, Areias and Belytschko (2006) is compared, and shows the similar accuracy.

Secondly, the central cracks of different inclined angles are considered. The length of crack is the same \(2a=0.0048\) m, and the angles, \(\theta=15^\circ, 30^\circ, 45^\circ, 60^\circ, 75^\circ\) are tested. The problem has been discussed by Phan, Gray and Salvadori (2010) with Symmetric-Galerkin Boundary Element Method and by Liu, Bui, Zhang, et al (2012) with Smoothed Finite Element Method. For the case of mode I, as depicted in Fig.10a, the values in the
peak of DSIF curves decrease by the increase of $\theta$ for a small period of time after the stress wave arrive in the tip. Fig.10b reveals that DSIF in mode II are practically the same for the pair of $\theta = 15^\circ$ and $\theta = 75^\circ$, and the pair of $\theta = 30^\circ$ and $\theta = 60^\circ$. At the meanwhile, the curve of $\theta = 45^\circ$ has the highest peak value. The results are compared with Phan, Gray and Salvadori (2010) and shown the good accuracy.

Figure 7: The rectangular plate with crack of different angle

Figure 8: The DSIF with different integral path of the left crack tip
Figure 9: The DSIF of the left crack tip with different time increments

Figure 10: The DSIF of crack tip with different rotation angle: (a) Mode I; (b) Mode II

5.3 The stable explicit time stepping analysis

This part focuses on the main factors that influence the critical time step. The grid density and iteration form are the two main subjects considered here. The experiment configuration model is presented in Fig.7 with θ=0°. The material properties and the other parameters are the same as those used in last example. The method of numerical approximation is used to get the critical time Δt_c.

Firstly, the results with different grid densities were obtained. Three uniform meshes are considered, which are of CCT: 49×99, CCT1: 24×49, CCT2: 13×24 elements. With parameters \( \beta = 0, \gamma = 1/2 \) of Newmark scheme, the critical time step of different meshes can be turned out. As shown in Fig.11(a), the critical time we got is about \( \Delta t_c = 4.825 \times 10^{-8} \) s with 49×99 elements. When the time step Δt is less than \( \Delta t_c \), the numerical calculation results are completely consistent and do not produce divergence. Conversely,
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divergence is presented in the calculation when $\Delta t > \Delta c$. The divergence occurs at about 4.750 $\mu$s, 7.154 $\mu$s, 11.495 $\mu$s, 15.141 $\mu$s when $\Delta t$ is 5.000×10$^{-8}$ s, 4.900×10$^{-8}$ s, 4.850×10$^{-8}$ s, 4.838×10$^{-8}$ s, respectively. As a comparison, Fig.11(b) is presented with the mesh of 24×49. It is seen that the critical time is 10.025×10$^{-8}$ s which is improved than the one in Fig.11(a). The divergence occurs at about 4.095 $\mu$s, 9.494 $\mu$s, 12.090 $\mu$s, 17.085 $\mu$s when $\Delta t$ is 10.500×10$^{-8}$ s, 10.100×10$^{-8}$ s, 10.075×10$^{-8}$ s, 10.050×10$^{-8}$ s, respectively.

Figure 11: Numerical stability with different time stepping (R=5, $\beta$ =0, $\gamma$ =1/2): (a) CCT: 49×99, (b) CCT1: 24×49.

Table 1: The critical time stepping for different densities of grid (R=5, $\beta$ =0, $\gamma$ =1/2)

<table>
<thead>
<tr>
<th></th>
<th>CCT: 49×99</th>
<th>CCT1: 24×49</th>
<th>CCT2: 13×24</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta c$ ($\mu$s)</td>
<td>4.825×10$^{-2}$</td>
<td>10.025×10$^{-2}$</td>
<td>16.568×10$^{-2}$</td>
</tr>
<tr>
<td>$\Delta c_{\text{lump}}$ ($\mu$s)</td>
<td>5.479×10$^{-2}$</td>
<td>11.012×10$^{-2}$</td>
<td>18.138×10$^{-2}$</td>
</tr>
<tr>
<td>$\Delta c / \Delta c_{\text{lump}}$</td>
<td>88.064%</td>
<td>91.037%</td>
<td>91.344%</td>
</tr>
</tbody>
</table>

To clarify this case further, we repeated the above steps with CCT2: 13×24 elements, and the comparison results are shown in the Table 1. The critical time step is about 16.568×10$^{-8}$ s in the case of CCT2, which is larger than the case of CCT1. It is hence concluded that the critical time step decreases with the increase of grid density. Besides, the critical time step of the standard FEM for the lumped mass is also listed. With more elements, the critical time step $\Delta c_{\text{lump}}$ is decreased, and this is consistent with the case of $\Delta c$. The values of $\Delta c / \Delta c_{\text{lump}}$ are similar, which range from 88.064% to 91.344%. As Menouillard, Réthoré, Combescure et al (2006); Elguejd, Gravouil and Maigre (2009) suggested, $\Delta t = \frac{2}{3} \Delta t_{\text{lump}}$ for the stationary crack, the value 2/3 is within the numerical range listed in this paper. So the numerical stability can be guaranteed.
In addition, we took into account the effect of Newmark scheme for the critical time step. Four cases are concerned. Before studying the impact of iterative format on critical time step, all the cases are listed under the same conditions: CCT1, a mesh of 24×49 elements, R=5, Δt=5×10⁻⁸s. We listed the first 30 microseconds with different parameter values γ in Fig.12. An approximately identical result can be obtained. The stability conditions of the Newmark scheme are also verified directly.

In Fig.11(b), we presented the test result with γ=1/2. As a comparison, the result with γ=2/3 is shown in Fig.13. The divergence occur at about 6.030µs, 12.848µs, 75.168µs when Δt is 9.000×10⁻⁸s, 8.800×10⁻⁸s, 8.700×10⁻⁸s, respectively. The critical time we obtained is about Δt_c=8.685×10⁻⁸s, which is smaller than the case of γ=1/2. For further investigation, the cases of γ=3/4, γ=1 are tested. The results are listed in Table.2. The critical time step of the standard FEM for the lumped mass are also listed. In Table.2, it is seen that the critical time step decreases with the increase of γ. So does Δt_c/lump. Furthermore, we observed that the values of Δt_c/lump are nearly the same (about 91%), and the parameter γ have nearly no influence on the values of Δt_c/lump.

**Figure 12:** Numerical results with different parameters γ (CCT1: 24×49, R=5, β=0, Δt=5×10⁻⁸s)
Figure 13: Numerical stability with different time stepping (CCT1: 24 × 49, R=5, β=0, γ =2/3)

Table 2: The critical time stepping for different parameters γ (CCT1: 24 × 49, R=5, β=0)

<table>
<thead>
<tr>
<th></th>
<th>γ = 1/2</th>
<th>γ = 2/3</th>
<th>γ = 3/4</th>
<th>γ = 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Δt</td>
<td>10.025×10^{-2}</td>
<td>8.685×10^{-2}</td>
<td>8.190×10^{-2}</td>
<td>7.100×10^{-2}</td>
</tr>
<tr>
<td>Δt_{fem}</td>
<td>11.012×10^{-2}</td>
<td>9.537×10^{-2}</td>
<td>8.992×10^{-2}</td>
<td>7.787×10^{-2}</td>
</tr>
<tr>
<td>Δt_{lump}/Δt_{fem}</td>
<td>91.037%</td>
<td>91.066%</td>
<td>91.081%</td>
<td>91.178%</td>
</tr>
</tbody>
</table>

6 Conclusions

In the present paper, we carried out some numerical experiments of the stable explicit time stepping within the XFEM framework. A new enrichment scheme for crack tip is proposed and its applicability and availability has been sufficiently verified. The DSIF is used as an important parameter of the dynamic response and is also a parameter of judging the stability of numerical method. Objective to studying the factors that can affect the stability, different densities of grid and different parameters of Newmark scheme have been tested. The conclusions are shown as:

- The grid density and the form of iterative method have obvious effects on stability;
- The critical time stepping Δt_c decreases with the increase of grid density;
- The critical time stepping Δt_c decreases with the increase of the parameter γ between 0.5 and 1 of Newmark scheme;
A similar conclusion can be obtained by the standard FEM with the lumped mass, and the values of $\Delta t_c / \Delta t_{\text{lump}}$ are relatively stable. Furthermore, the simulation results are found in good agreement with each other when they are stable. Therefore, increasing time stepping appropriately in the range of critical value can improve the computational efficiency.

Acknowledgment: The authors are grateful to the National Natural Science Foundation of China (No.11672101, No.11372099), the 12th Five-Year Supporting Plan Issue (No. 2015 BAB07B10), Jiangsu Province Natural Science Fund Project (No. BK 20151493) and the Postgraduate Research and Innovation Projects in Jiangsu Province (No.2014B 31614) for the financial support.

References


