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Miguel Molinos^a, and Pedro Navas^{a1}

a ETSI Caminos, Canales y Puertos, Universidad Politéctnica de Madrid.
 c. Prof. Aranguren 3, 28040 Madrid, Spain

Abstract

The objective of this work is to introduce an alternative technique to address the fracture process of brittle and quasi-brittle materials under the Material Point Method (MPM) framework. With this purpose the eigensoftening algorithm, developed originally for the Optimal Transportation Meshfree (OTM) approximation scheme, is extended to the MPM with the aim of present a suitable alternative to the existing fracture algorithms developed for the MPM. The good fitting in the predictions made by the eigensoftening algorithm against both analytical and experimental results proofs the well performance of the method under challenging loads.

Keywords: Quasi brittle fracture, Local-max-ent approximation, Material Point Method, Solid Dynamics

1. Introduction

Presence of cracks are a violation of the continuity requirement in Finite Element Method (FEM). To overcome it, numerous of numerical artifacts has been proposed with the aim of reproducing such a complex behaviour. This techniques vary from employing cohesive approaches [1, 2], by adaptively inserting cohesive elements [3, 4, 5] at solid elements boundaries, or handling arbitrary cracks paths by level set representation of the fracture surface [6].

The simulation of fracture propagation in a more accurate and effective way can be considered as one of the original drivers for developing novel spatial discretization methods such as meshfree techniques. Some examples of it are the Material Point Method (MPM) [24, 25, 20, 23], the Element-Free Galerkin Method (EFGM) [7, 8, 9, 10], the Smoothed Particle Hydrodynamics (SPH) [11, 12], the Optimal Transportation Meshfree (OTM) [13, 14, 15, 16] or Peridynamics [17, 18] among others.

¹Corresponding author: p.navas@upm.es

Fracture can be described in two ways. One is to remove the restriction 15 of the single-valued velocity field close to the crack by using two or more sets of nodes [19]. This method is based in to assign different labels to distinguish if the material points and nodes are in the same side of the crack or not. Under this approach the crack surface is described with line segments in 2D and triangle patches in 3D cases. This method was named as "CRAcks with Material Points (CRAMP)" [20]. In this method, the criteria for crack propagation is based on such parameters as energy release rate analyzed by Tan & Nairn (2002)[21], and the stress intensity factor or the J-integral discussed by Guo & Nairn (2004)[22]. The other approach is to use failed material points to describe the crack evolution. In this method, the formation of failed points describes the nucleation of cracks, and thereafter its propagation and branching. Consequently, the position of the crack does not need to be explicitly stated. This represents a significant advantage over the "CRAMP". Under this approach, the prediction the failure evolution employing a decohesion model has been discussed by Chen et al. [23] or Schreyer et al. [24]. And has been successfully employed to simulate the fracture of brittle materials by Chen et al. [25, 26] and Sulsky & Schreyer [27]. Similar to this approach, Schmidt et al. [29] introduced the concept of eigenfracture, where they approximate the crack set by means of eigen-deformations, wich enable the material to develop displacement jumps at no cost of local elastic energy. Later, the eigenerosion approach to brittle facture was developed by Pandolfi et al. [30, 15]. In this technique the erosion of the material point means that each material point can be either intact or be completely failed or eroded and has no load bearing capacity. This method has been successfully applied to simulate high complex phenomena such dynamic fragmentation of metals [16], nonetheless in quasi-brittle materials, simulations performed with this approach exhibit an overestimation of tensile stress and the strain peaks. To overcome this limitation, Navas et al. [35?] introduced the concept of eigensoftening. Engineered for quasi-brittle materials, this approach is devoted to capture the gradual rather than abrupt dissipation of the fracture energy. It is inspired in the concept of the crack band model [37], since energy dissipation is thorough the softened (or failed) volume. Due to the similarities between the OTM, where this procedure was developed, and the MPM, we propose

The eigensoftening algorithm together with a explicit predictor corrector and the local *max-ent* approximation technique offers a new perspective in the MPM fracture simulation.

To overcome the difficulties of the e

In the present research we will introduce the

In the current work we attempt to model the fracture propagation employ-

ing the MPM and introducing both eigenerosion and eigensoftening [35, 36] and the concept of the eigenfracture under this framework.

Even thought the method has been successfully applied to dynamic fragmentation of metals, quantitative validations, such as stress or strain levels near or at the crack set compared to experimental measures are still lacking. Having in mind that the eigenerosion approach was intended for perfectly brittle fracture, an eigensoftening concept is developed for quasi-brittle materials.

This document is structured as follows

55 2. The meshfree methodology

The aim of this section is to provide an overview of the special techniques employed to face the fracture problem under the MPM framework. In consequence it is structured as follows: first in 2.1 the explicit-predictor algorithm for the MPM will be exposed, next the local *max-ent* approximants are introduced in 2.2 as an accurate alternative technique to interpolate data between particles and nodes, and finally the fracture algorithms based in the eigendeformations are presented in 2.3.

2.1. The Material Point Method

The MPM [31] is a meshfree Lagrangian-Eulerian method where particles carries on all the physical information and a set of background nodes is employed to compute the equilibrium equation. Since the MPM possesses the advantages of both Lagrangian and Eulerian descriptions, no element distortion exists in the MPM, therefore it is an appropriate and efficient method in solving problems involving extremely large deformation and moving discontinuities such fracture evolution. For the spatial discretization, two sets of points are introduced in the MPM. First, the nodes. This points are considered fixed in the space and are in charge of computing all the kinematic fields such forces f_I , accelerations a_I and velocities v_I . And second the material points or particles. They are in charge of the discretization of the continuum, and store the local state $(\sigma_p, \varepsilon_p)$. Without loosing generality, the MPM algorithm can be described with three main steps: (i) a variational recovery process, where particle data is projected to the grid nodes, (ii) an Eulerian step, where balance of momentum equation is expressed as a nodal equilibrium equation thorough a FEM-like procedure, and finally (iii) a Lagrangian advection of the particles. In the present research a explicit predictor-corrector time integration scheme is adopted. The purpose of this choice is motivated due its proved robustness and stability in numerical calculations. In the first stage (i), the nodal velocity predictor is computed

following (1),
$$\vec{v}_{I}^{k+1} = \frac{N_{Ip}^{k} m_{p} (\vec{v}_{p}^{k} + (1 - \gamma) \Delta t \vec{a}_{p}^{k})}{m_{I}}$$
 (1)

This way of computing the nodal predictor is both numerically stable and minimize the computational effort. Once nodal velocity are obtained, the essential boundary conditions are imposed. And in the following, a Eulerian phase (ii) is computed in the set of nodes in a FEM-like way, where nodal forces \vec{f}_I^{k+1} are computed thorough the equilibrium equation. Next the nodal velocities are corrected in a *corrector* stage,

$$\vec{v}_I^{k+1} = \vec{v}_I^{pred} + \gamma \Delta t \frac{\vec{f}_I^{k+1}}{\mathbf{m}_I^{k+1}}$$
 (2)

Finally updated the particles are advected in the Lagrangian stage (iii) using nodal values as,

$$\vec{a}_p^{k+1} = \frac{N_{Ip}^k \vec{f}_I^k}{\mathbf{m}_I^k} \tag{3}$$

$$\vec{v}_p^{k+1} = \vec{v}_p^n + \Delta t \, \frac{N_{Ip}^k \, \vec{f}_I^k}{\mathbf{m}_I^k} \tag{4}$$

$$\vec{x}_p^{k+1} = \vec{x}_p^n + \Delta t \ N_{Ip}^k \ \vec{v}_I^k + \frac{1}{2} \Delta t^2 \ \frac{N_{Ip}^k \ \vec{f}_I^k}{\mathbf{m}_I^k}$$
 (5)

The complete pseudo-algorithm it is summarized in Appendix A.

5 2.2. Spatial discretization: Local-max-ent approximants

Local maximum-entropy (or local max-ent) approximation scheme was introduced by Arroyo & Ortiz (2006)[32] as a bridge between finite elements and meshfree methods. The key idea of the shape functions is to interpret the nodal of a shape function N_I as a probability. This allow us to introduce two important limits: maximum-entropy (max-ent) limit, and the Delaunay triangulation which ensures the minimal width of the shape function. The first limit ensures a unbiased statistical inference based on the nodal data as ensures the Jayne's[33] principle of maximum entropy, the second limit warrant the least width shape function support. To reach to a compromise

between two competing objectives, a Pareto set is defined as,

(LME)_{\beta} For fixed
$$\vec{x}$$
 minimise $f_{\beta}(\vec{x}_p, N_I) = \beta U(\vec{x}_p, N_I) - H(N_I)$

subject to
$$\begin{cases} N_I \geq 0, & \text{I=1, ..., n} \\ \sum_{I=1}^{N_n} N_I = 1 \\ \sum_{I=1}^{N_n} N_I \vec{x}_I = \vec{x} \end{cases}$$

where $H(N_I)$ is the entropy of the system of nodes following the definition given by Shannon (1948) [34], and $U(\vec{x}_p, N_I) = \sum_I N_I |\vec{x}_p - \vec{x}_I|^2$ a magnitude of the shape function width. The regularization o thermalization parameter between the two criterion β has Pareto optimal values in the range $(0, \infty)$. The unique solution of the local max-ent problem LME $_\beta$ is,

$$N_I^*(\vec{x}) = \frac{\exp\left[-\beta |\vec{x} - \vec{x}_I|^2 + \vec{\lambda}^* \cdot (\vec{x} - \vec{x}_I)\right]}{Z(\vec{x}, \vec{\lambda}^*)}$$
(6)

where $Z(\vec{x}, \vec{\lambda}^*)$ is the partition function defined as,

$$Z(\vec{x}, \vec{\lambda}) = \sum_{I=1}^{N_n} \exp\left[-\beta |\vec{x} - \vec{x}_I|^2 + \vec{\lambda} \cdot (\vec{x} - \vec{x}_I)\right]$$
 (7)

and evaluated in the unique minimiser $\vec{\lambda}^*$ for the function $\log Z(\vec{x}, \vec{\lambda})$. The traditional way to obtain such a minimiser is using (8) to calculate small increments of $\partial \vec{\lambda}$ in a Newton-Raphson approach. Where **J** is the Hessian matrix, defined by:

$$\mathbf{J}(\vec{x}, \vec{\lambda}, \beta) \equiv \frac{\partial \vec{r}}{\partial \vec{\lambda}} \tag{8}$$

$$\vec{r}(\vec{x}, \vec{\lambda}, \beta) \equiv \frac{\partial \log Z(\vec{x}, \vec{\lambda})}{\partial \vec{\lambda}} = \sum_{I}^{N_n} p_I(\vec{x}, \vec{\lambda}, \beta) (\vec{x} - \vec{x}_I)$$
 (9)

The value of **J** can be employed to get the first derivatives of the shape function ∇N_I^* ,

$$\nabla N_I^* = N_I^* \left(\nabla f_I^* - \sum_{J}^{N_n} N_J^* \, \nabla f_J^* \right) \tag{10}$$

where

$$f_I^*(\vec{x}, \vec{\lambda}, \beta) = -\beta |\vec{x} - \vec{x}_I|^2 + \vec{\lambda}^* (\vec{x} - \vec{x}_I)$$
(11)

Employing the chain rule over (10), rearranging and considering β as a constant, Arroyo and Ortiz [32] obtained the following expression for the gradient of the shape function.

$$\nabla N_I^* = -N_I^* (\mathbf{J}^*)^{-1} (\vec{x} - \vec{x}_I)$$
 (12)

The regularization parameter β of LME shape functions may be controlled by adjusting a dimensionless parameter, $\gamma = \beta h^2$ [32], where h is defined as a measure of the nodal spacing. Since N_I is defined in the entire domain, in practice, the shape function decay $\exp(-\beta \vec{r})$ is truncated by a given tolerance, 10^{-6} , for example, would ensure a reasonable range of neighbours, see [32] for details. This tolerance defines a limit values of the influence radius and is used thereafter to find the neighbour nodes of a given integration point.

2.3. Fracture modelling approach

Within the context of MPM formulation, fracture can be modelled by failing particles according to a suitable criterion. When material points are failed, they are assumed to have a null stress tensor. Navas et al. (2017)[35] developed a eigensoftening algorithm as an extension for quasi-brittle materials of the eigenerosion proposed by Pandolfi & Ortiz (2012)[30] for fracture of brittle materials. A comparison between both in [35] shows that the eigenerosion algorithm significantly overestimates the tensile stress and the strain peaks, while it captures the forces and crack patterns accurately. On the other hand eigensoftening algorithm agree very well with experimental results in all the aspects. Furthermore, this algorithm has also proof its accuracy for complex fracture patters such the present in fiber reinforces concrete (FRC), [36].

$$G_p^{k+1} = \frac{C_{\epsilon}}{m_p^{k+1}} \sum_{x_q^{k+1} \in B_{\epsilon}(x_p^{k+1})} m_q W_q^{k+1}$$

$$m_p^{k+1} = \sum_{x_q^{k+1} \in B_{\epsilon}(x_p^{k+1})} m_q$$

$$\tag{13}$$

$$m_p^{k+1} = \sum_{x_q^{k+1} \in B_{\epsilon}(x_p^{k+1})} m_q \tag{14}$$

where $B_{\epsilon}(x_p^{k+1})$ is the sphere of radius ϵ centered at x_p^{k+1} known as the ϵ -neighborhood of the material point, m_p^{k+1} is the mass of the neighborhood at step k+1, W_q^{k+1} is the current free-energy density per unit mass as the material point x_q^{k+1} of the neighborhood, finally C_{ϵ} is a normalizing constant. This configuration conveniently is sketched in Figure 1. The material

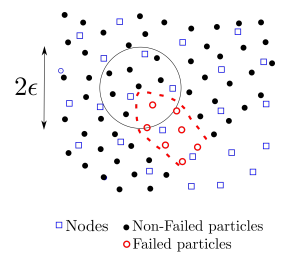


Figure 1: Scheme of a linear cohesive law, where the shaded area is G_f , f_t is the tensile strength, and w_c is the critical opening displacement.

point fails when G_p^{k+1} surpasses a critical energy release rate that measures the material-specific energy, G_F . The convergence of this approach has been analyzed by Schmidt *et al.* (2009)[29], who proof that it converges to the Griffith fracture when discretization size tends to zero. It is necessary to point out that when a material point overpass the critical energy, its contribution to the internal forces vector is set to zero, but its contribution to the mass matrix is maintained. The mass of a material point is discarded only when an eroded material point is not connected to any nodes.

As can be noticed, in the eigenerosion algorithm an energetic criterion is adopted. Due to that fact, unrealistic stress concentration (higher than tensile strength) in quasi-brittle materials, see [35]. To overcome this limitation, the aforementioned authors proposed the concept of eigensoftening to take in to account the gradual failure in quasi-brittle materials. The concept is inspired in the cohesive fracture widely employed in the context of FEM [3]. This gradual failure criterion is plotted in figure, where a linear decreasing cohesive law is presented to illustrate the concept here described. In the picture, the shaded area represents the static fracture energy per unit of area, G_F . Notice how a cohesive crack appears when the maximum tensile strength, f_t is reached. Once the opening displacement w takes the value of the critical crack displacement w_c , a stress-free crack is attained. For intermediate values, w_n , a damage value between zero and one represents the extension to which the material has failed. For the eigensoftening algorithm, a strength criterion for crack initialization was adopted. Particularly the

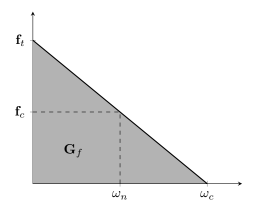


Figure 2: Scheme of a linear cohesive law, where the shaded area is G_f , f_t is the tensile strength, and w_c is the critical opening displacement.

maximum principal stress theory for brittle fracture was considered by authors [35]. With that purpose, the variation of the averaged strain energy density in the ϵ -neighborhood of the material point \vec{x}_p^{k+1} can be obtained as,

$$\delta W_{\epsilon,p} = \frac{\partial G_p}{C_{\epsilon}} = \frac{1}{m_p} \sum_{x_q^{k+1} \in B_{\epsilon}(x_p^{k+1})} m_q \sigma_{q,I} \delta \epsilon_q$$
 (15)

where $\sigma_{q,I}$ is the maximum principal stress of each material point in the ϵ -neighborhood. Here [35] introduces an effective strain ε_q , such the variation of the local strain energy can be obtained as $\delta W_q = \sigma_{q,1} \delta \varepsilon_q$. Now with the assumption that the effective strain of each material point at every time step is constant in the neighborhood of \vec{x}_p^{k+1} , the equation (15) can be simplified as follows

$$\delta W_{\epsilon,p} = \frac{\delta \epsilon_p}{m_p} \sum_{x_q^{k+1} \in B_{\epsilon}(x_p^{k+1})} m_q \sigma_{q,I}. \tag{16}$$

Consequently it is possible to define an equivalent critical stress at the material point \vec{x}_p^{k+1} :

$$\sigma_{\epsilon,p} = \frac{1}{m_p} \sum_{x_q^{k+1} \in B_{\epsilon}(x_p^{k+1})} m_q \sigma_{q,I}, \tag{17}$$

where m_p can be computed as:

$$m_p = \sum_{x_q^{k+1} \in B_{\epsilon}(x_p^{k+1})} m_q.$$
 (18)

This definition of the equivalent critical stress allow us to link it with the averaged strain energy as $\delta W_{\epsilon,p} = \sigma_{\epsilon,p} \ \delta \varepsilon_p$. The fracture criterion is defined

as once $\sigma_{\epsilon,p}^{k+1}$ surpasses the tensile strength, f_t , the softening behaviour is activated, which in turn causes a reduction of the initial forces as,

$$f_I^{int} = \sum_p (1 - \chi_p) \ \sigma_p^{k+1} \cdot grad(N_{Ip})$$
 (19)

where χ_p is the damage variable of each material point p, ranges between zero (an intact material) and one (completely failed material points). For the case of a linear softening such the sketched in the Figure 2, it is calculated as,

$$1 - \chi = \frac{f_n}{f_t} = 1 - \frac{w_n}{w_c} \to \chi = \frac{w_n}{w_c}.$$
 (20)

Following Bazant [37] crack-model, [35] introduced a band width h_{ϵ} , which has values between two and four times the maximum size of the aggregates for the concrete. The effective fracture strain $\varepsilon_{\epsilon,f}$ is defined as the difference between the strain at crack initialization, $\varepsilon_1(\vec{x}_p^0)$, and the current strain, $\varepsilon_1(\vec{x}_p^{k+1})$, for material point p. Also, $\varepsilon_{\epsilon,f}$ can be represented as the current crack opening w_n within the band width, h_{ϵ} . Therefore,

$$\varepsilon_{\epsilon,f} = \varepsilon_1(\vec{x}_p^{k+1}) - \varepsilon_1(\vec{x}_p^0) = \frac{w_n}{h_{\epsilon}}$$
(21)

Introducing (21) in (20), the damage variable can be computed as,

$$\chi = \frac{\varepsilon_{\epsilon,f} \ h^{\epsilon}}{w_c} \tag{22}$$

for the case of a linear softening behaviour. In a more general case, the damage variable can be expressed in terms of the following variables,

$$\chi = \chi(\varepsilon_{\epsilon,f}, h^{\epsilon}, f_t, w_c, G_f) \tag{23}$$

3. Cases of study and discussion

3.1. Edge-cracked square panel in mode I

This problem is devoted to compare the develop of both linear shape functions and the LME

3.2. Drop-weight impact test

4. Conclusions

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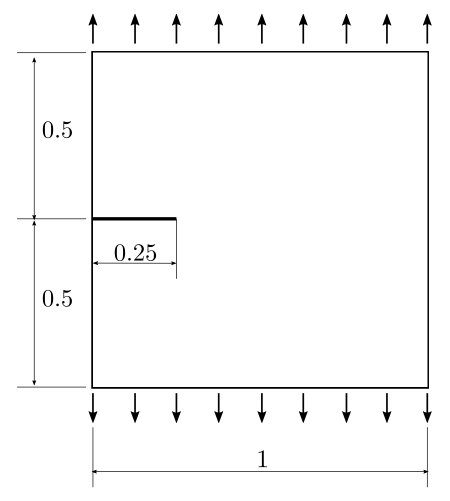


Figure 3: Geometry and boundary condition of the drop-weight impact test.

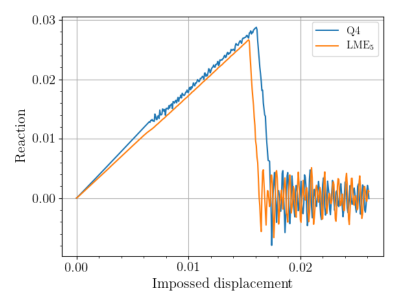


Figure 4: Geometry and boundary condition of the drop-weight impact test.

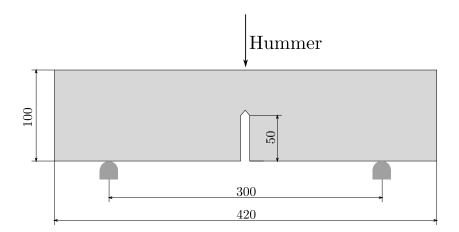


Figure 5: Geometry and boundary condition of the drop-weight impact test.

Appendix A. Explicit Predictor-Corrector algorithm

Appendix B. Eigensoftening Algorithm

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Algorithm Explicit Predictor-Corrector scheme

Update mass matrix:

$$\mathbf{m}_I = N_{Ip}^k \ m_p,$$

Explicit Newmark Predictor:

$$\vec{v}_I^{pred} = \frac{N_{Ip}^k m_p(\vec{v}_p^k + (1 - \gamma) \Delta t \vec{a}_p^k)}{m_I}$$

Impose essential boundary conditions: At the fixed boundary, set $\vec{v}_I^{pred} = 0$. Deformation tensor increment calculation.

$$\Delta \varepsilon_p^{k+1} = \Delta t \ \dot{\varepsilon_p}^{k+1} = \Delta \left[\vec{v}_I^{pred} \otimes grad(N_{Ip}^{k+1}) \right]^s$$

Update the density field:

$$\rho_p^{k+1} = \frac{\rho_p^k}{1 + tra\left[\Delta \varepsilon_p^{k+1}\right]}.$$

Compute stress field and update damage parameter: Balance of forces calculation: Calculate the total grid nodal force $\vec{f}_I^{k+1} = \vec{f}_I^{int,k+1} + \vec{f}_I^{ext,k+1}$. Explicit Newmark Corrector:

$$\vec{v}_I^{k+1} = \vec{v}_I^{pred} + \gamma \ \Delta t \ \frac{\vec{f}_I^{k+1}}{\mathbf{m}_I^{k+1}}$$

Update particles lagrangian quantities:

$$\begin{split} \vec{a}_{p}^{k+1} &= \frac{N_{Ip}^{k} \vec{f}_{I}^{k}}{\mathbf{m}_{I}^{k}} \\ \vec{v}_{p}^{k+1} &= \vec{v}_{p}^{n} + \Delta t \ \frac{N_{Ip}^{k} \ \vec{f}_{I}^{k}}{\mathbf{m}_{I}^{k}} \\ \vec{x}_{p}^{k+1} &= \vec{x}_{p}^{n} + \Delta t \ N_{Ip}^{k} \ \vec{v}_{I}^{k} + \frac{1}{2} \Delta t^{2} \ \frac{N_{Ip}^{k} \ \vec{f}_{I}^{k}}{\mathbf{m}_{I}^{k}} \end{split}$$

Reset nodal values

```
Algorithm 2 Compute damage parameter \chi_p^{k+1}
Require: Particle status
    Number of particles: N_p
    \epsilon-neighbourhood of each particle p: B_{\epsilon,p}
Require: Material data
    Tensile strength: f_{t,p}
    Bandwidth of the cohesive fracture: h_{\epsilon,p}
    Critical opening displacement: w_c
Ensure: Return damage parameter \chi := \{\chi_p\}
    \chi_p \leftarrow \chi_p^k
    for p to N_p do
         if \chi_p = 0 \ \epsilon_{f,p} = 0 then
               for q \in B_{\epsilon,p} do
                     if \chi_q < 1 then
\sum m_p \sigma_{p,I} \leftarrow \sum m_p \sigma_{p,I} + m_q \sigma_{q,I}
                     end if
                     m_p \leftarrow m_p + m_q
               end for
               \sigma_{p,\epsilon} \leftarrow \frac{1}{m_p} \sum_{p} m_p \sigma_{p,I}
if \sigma_{p,\epsilon} > f_{t,p} then
                    \epsilon_{f,p} = \epsilon_{I,p}
               end if
         else[\chi_p \neq 1 \ \epsilon_{f,p} > 0]
              \chi_p^{k+1} \leftarrow \min\left\{1, \max\{\chi_p^k, \frac{(\epsilon_{I,p} - \epsilon_{f,p}) \ h_{\epsilon,p}}{w_c}\}\right\}
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end if

end for

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