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Assessment of the Energy Balance of Rock Masses through Discrete Element Modelling

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Abstract

With the increase of mining depth, rockbursts become one of the most serious hazards in mines. Studies of rockburst mechanisms can be done through the application of various numerical methods at the laboratory and site scales. Among the different methods available, the discrete element method (DEM) is now increasingly used thanks to its capability to explicitly model the initiation and propagation of fractures leading to failure. The paper deals with the evaluation and analysis of the energy components developing inside a rock mass by means of discrete element modeling. For this purpose, the code YADE Open DEM has been used. The rock mass is viewed as an assembly of bonded particles that interact through an elastic-brittle and frictional contact law. Interparticle breakage can occur by either tensile or shear mechanisms and interparticle friction is considered with respect to a Mohr-Coulomb criterion. Pre-existing fractures are explicitly modeled as frictional interfaces. To investigate the evolution of input and dissipated energy, various energy terms are evaluated for different loading paths. The energy terms include boundary works, elastic strain energy, friction and crack dissipation, kinetic energy and damping dissipation. The proposed energetic approach is verified by computing the energy balance of the system during simulations of compression, tension and shear tests.

Keywords: energy balance; rock mass; discrete elements method; Yade

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1. Introduction

Rockburst is a sudden and violent phenomenon resulting from high concentration of stresses. This phenomenon is dependent on the initial and induced states of stress, on geology as well as on the presence of discontinuities in the rock mass. In damage mechanics, failure is caused by the initiation and the propagation of one or a network of microcracks. These microcracks propagate and coalesce to form fractures, leading to the rock failure. To simulate the initiation and the propagation of cracks, two approaches are generally used: the continuum and the discontinuum approach. In the continuum approach, the numerical methods are based on the finite difference (FD), finite volume (FV) and finite element (FE) methods. In this case, the initiation and propagation of cracks are modeled through the remeshing techniques which depend on the initial mesh size used. To overcome this limitation, a new numerical method has been developed: the extended finite element method (XFEM). The XFEM is able to simulate the initiation and the propagation of cracks independently of the mesh [4, 1, 2, 7]. Discrete approaches such as the discrete element method (DEM) or the discontinuous deformation analysis (DDA) provide an alternative. Cundall and Strack [3] have introduced the DEM which models the medium as an assembly of interacting particles. Each particle is identified by its mass, radius and moment of inertia and interacts with its neighbors via predefined contact laws. Potyondy and Cundall [6] then have introduced the bonded particles model (BPM) that allows simulating the behavior of rock materials and fracturing processes. Lately, the FEM/DEM has been introduced as a hybrid numerical method which combines DEM and FEM techniques. This method was initiated by Munjiza et al. [5]. In order to simulate explicitly the initiation and the propagation of cracks, we have chosen to use the computer code Yade Open DEM [10] where we have implemented the computation of energy terms. Yade Open DEM is selected because it enables us to implement the undertaken development. To validate our implementation, laboratory scale tests are simulated. In this paper, various components of energy occurring in a rock system subjected to external solicitations have been proposed. These are external work, potential energy, kinetic energy, elastic strain energy, energy dissipated by cracks, energy dissipated by numerical damping and energy dissipated by friction between particles.

<table>
<thead>
<tr>
<th>Nomenclature</th>
</tr>
</thead>
<tbody>
<tr>
<td>$F_n, F_t$</td>
</tr>
<tr>
<td>$k_n, k_t$</td>
</tr>
<tr>
<td>$u_n, u_t$</td>
</tr>
<tr>
<td>$c$</td>
</tr>
<tr>
<td>$t$</td>
</tr>
<tr>
<td>$\phi_b$</td>
</tr>
<tr>
<td>$\phi_c$</td>
</tr>
<tr>
<td>$E_{el}$</td>
</tr>
<tr>
<td>$E_{ki}$</td>
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<tr>
<td>$E_{da}$</td>
</tr>
<tr>
<td>$E_{cr}$</td>
</tr>
<tr>
<td>$W_{ext}$</td>
</tr>
<tr>
<td>$W$</td>
</tr>
<tr>
<td>$\alpha$</td>
</tr>
<tr>
<td>$m$</td>
</tr>
<tr>
<td>$v$</td>
</tr>
<tr>
<td>$I$</td>
</tr>
<tr>
<td>$\dot{\theta}$</td>
</tr>
<tr>
<td>$\dot{F}$</td>
</tr>
<tr>
<td>$\dot{m}$</td>
</tr>
<tr>
<td>$E_{micro}$</td>
</tr>
<tr>
<td>$R_1, R_2$</td>
</tr>
</tbody>
</table>
The energy balance represents the difference between the total energy supplied to the system and that accumulated during the deformations and/or dissipated in friction and/or in cracking. Finally, this energy approach was successfully applied to numerical simulations of different laboratory tests. In this paper we limit ourselves to the calculations of the balance energy for the laboratory scale tests in order to check the robustness of the developments and to apply it in the simulation of large scale problems like underground excavation and to study rockburst phenomenon.

2. Model formulation

The simulations were performed with Yade Open DEM, a 3D code based on the DEM [10]. The rock is represented by an assembly of rigid spherical particles interacting one with each other according to an elastic-brittle contact law. The interaction force between particles is subdivided into a normal force $F_n$ and a tangential force $F_t$ (Fig.1). The normal force is computed such as:

$$F_n = k_n u_n$$

(1)

The normal stiffness is given by:

$$k_n = \frac{E_{\text{micro}} R_1 R_2}{R_1 + R_2}$$

(2)

And the tangential stiffness is computed via the ratio $k_n / k_t$ which is determined by the calibration of the material.

Under compressive loading (contact closure), the normal force can increase indefinitely without overlap threshold. Under tensile loading, the normal force is limited by the maximal admissible normal force $F_{n,\text{max}}$ defined as a function of the tensile strength ($t$) and the surface ($A_{\text{int}}$) between particles:

$$F_{n,\text{max}} = tA_{\text{int}}$$

(3)

If $F_n \geq F_{n,\text{max}}$, tensile failure occurs and a mode I crack is created.

The tangential force is calculated in an incremental manner as a function of the increment of shear displacement $\Delta u_t$ that developed at the contact, such as:

$$F_t^{(t+\Delta t)} = F_t^{(t)} + k_t \Delta u_t$$

(4)

with $F_t^{(t+\Delta t)}$ and $F_t^{(t)}$ tangential force at the instants $t + \Delta t$ and $t$, respectively.
The maximal admissible shear force $F_{t,\text{max}}$ is defined as a function of the normal compression force $F_n$, the cohesion $c$, the local friction angle $\phi_b$ and the interacting surface between two particles $A_{\text{int}}$ through a Mohr-Coulomb failure criterion:

$$F_{t,\text{max}} = F_n \tan(\phi_b) + cA_{\text{int}}$$  \hfill (5)

when $F_t \geq F_{t,\text{max}}$, shear failure occurs, a mode II crack is created and the interaction between particles become purely frictional for compressive regimes or disappears for tensile regimes. For purely frictional contacts, the maximum admissible shear force is calculated as a function of and the local residual frictional angle $\phi_c$ according to:

$$F_{t,\text{residual}} = F_n \tan \phi_c$$  \hfill (6)

The failure envelopes in tension and shear are illustrated in Fig. 2.

![Fig. 2. Failure envelope used for the contact in Yade.](image)

Because of the dynamic formulation of the method (explicit time domain integration), a global non-viscous damping is used to dissipate kinetic energy and facilitate convergence towards quasi-static equilibrium. This damping directly acts on the forces and torques considered in the equations of motion so that the displacements and rotations are calculated from the damped forces and damped torques such that:

$$\ddot{F}_d = -\alpha \ddot{F}$$  \hfill (7)

$$\ddot{M}_d = -\alpha \ddot{M}$$  \hfill (8)

with $\alpha$ the damping coefficient.

3. Energetical formulation

The DEM codes are explicit and based on the second Newton law. The different physical parameters are reachable throughout the simulation and the energetic calculation is possible. In order to calculate the balance energy with Yade, an energetic development is implemented in the code. In this development presented herein, various energy components are proposed: boundary works, elastic strain energy, friction and crack dissipation, kinetic energy and damping dissipation. These components are calculated for each time step. The increment of the elastic strain energy ($\Delta E_{el}$) is computed as:
\[ \Delta E_{el} = E_{el}^{(t+\Delta t)} - E_{el}^{(t)} = \frac{1}{2} \sum_{n_e^{(t+\Delta t)}} \left( \frac{F_n^2 + F_t^2}{k_n + k_t} \right)^{(t+\Delta t)} - \frac{1}{2} \sum_{n_e^{(t)}} \left( \frac{F_n^2 + F_t^2}{k_n + k_t} \right)^{(t)} \]  

with \( N_e(t) \) the number of elastic contacts at time \( t \).

The increment of the kinetic energy (\( \Delta E_k \)) is given by:

\[ \Delta E_k = \frac{1}{2} \sum_{n_p} \left( mv^2 + I\dot{\theta}^2 \right)^{(t+\Delta t)} - \frac{1}{2} \sum_{n_p} \left( mv^2 - I\dot{\theta}^2 \right)^{(t)} \]  

with \( N_p \) the total number of particles, \( m \) their mass, \( v \) their translational velocity, \( I \) their moment of inertia and \( \dot{\theta} \) their angular velocity.

The increment of energy dissipated by damping (\( \Delta E_{da} \)) is equal to:

\[ \Delta E_{da} = \sum_{n_p} \alpha \left( \langle \dot{F} | \dot{v} \rangle + \langle \dot{M} | \dot{\theta} \rangle \right) \Delta t \]  

where \( \alpha \) is the coefficient of numerical damping, \( \dot{F} \) and \( \dot{M} \) are respectively the resultant force and moment acting on the particle, \( \Delta t \) is the timestep, \( \langle x | y \rangle \) is the scalar product between two vectors \( x \) and \( y \).

The increment of energy dissipated during the process of cracking (\( \Delta E_{cr} \)) is calculated such that:

\[ \Delta E_{cr} = \frac{1}{2} \sum_{n_c} \left( \frac{F_n^2 + F_t^2}{k_n + k_t} \right) \]  

where \( N_c(t) \) is the number of contacts that broke during the timestep \( \Delta t \).

The increment of energy dissipated by friction between particles (\( \Delta E_{fr} \)) is obtained by:

\[ \Delta E_{fr} = \sum_{n_f} \left( \langle \dot{F}_f^{(t+\Delta t)} | \dot{F}_f^{(t)} \rangle \right) \]  

where \( N_f \) is the number of frictional contacts, \( F_f^{(t+\Delta t)} \) and \( F_f^{(t)} \) tangential force at the instants \( t + \Delta t \) and \( t \), respectively.

The increment of external work (\( \Delta W_{ext} \)) is calculated such that:

\[ \Delta W_{ext} = \sum_{N_w} \sum_{N_{pw}} \left( \dot{F}_f^{(t+\Delta t)} | \dot{u}^{(t+\Delta t)} - \dot{u}^{(t)} \right) + \sum_{N_b} \left( \dot{F}_b^{(t+\Delta t)} | \dot{u}^{(t+\Delta t)} - \dot{u}^{(t)} \right) \]  

where \( N_w \) is the number of walls, \( N_{pw} \) is the total number of particles for a given wall, \( N_b \) is the total number of particles on the other boundaries except the walls, \( \dot{F}_f^{(t+\Delta t)} \) is the force vector applied at instant \( t + \Delta t \) and \( \dot{u}^{(t+\Delta t)} \) and \( \dot{u}^{(t)} \) are the displacement vector at times \( t \) et \( t + \Delta t \), respectively.

The increment of the potential energy (\( \Delta E_p \)) is obtained by:

\[ \Delta E_p = \sum_{n_p} m \langle \ddot{g} | \ddot{v} \rangle \Delta t \]  

where \( m \) mass of particle, \( g \) gravitational acceleration, \( \ddot{v} \) translational velocity of particle and \( \Delta t \) time step.
The increment of energy balance ($\Delta W$) is the difference between on the one hand the input energy and secondly the energy accumulated and dissipated by the system. It is given by:

$$\Delta W = \Delta W_{\text{ext}} + \Delta E_p - (\Delta E_{\text{el}} + \Delta E_{\text{ki}} + \Delta E_d + \Delta E_{\text{fr}})$$  \hspace{1cm} (16)

Finally, equations (9) to (16) provide the set of energy components needed to assess the balance of a given system at each timestep $\Delta t$, and by integration the balance at each time $t$.

4. Applications

In order to validate the constituent equation and numerical implement, three laboratory tests are simulated: triaxial compression test, direct tension test and direct shear test. For these different tests, it is used a sample with dimensions of 1 x 2 x 1 m$^3$ constituted by 5000 particles calibrated on the Lac du Bonnet granite [8]. These simple configurations are chosen because analytical solutions exist or/and the results are understandable.

4.1. Triaxial compression test

A triaxial compression test with a confining pressure of 5 MPa was simulated. The test consists in two steps: hydrostatic loading with constant boundary velocity equal to 0.001 m/s until obtaining $\sigma_1 = \sigma_2 = \sigma_3 = 5$ MPa and deviatoric loading with axial velocity equal to 0.01 m/s. The value of the deviatoric loading velocity was chosen based on sensitivity analysis to ensure a quasi-static behavior. The different energy components were evaluated during the test as illustrated in Fig. 3 where the deviatoric stress versus axial strain has also been plotted. In the pre-peak stage, the external work and the elastic energy increase significantly with the increase of the deviatoric stress, some cracks appears in the sample and the energy is slightly dissipated by cracks and by friction between particles. In addition, the energy dissipated by damping is low. At the peak, the external work and the elastic energy reach also their maximal values. The number of cracks becomes important which results in a more and more increase of the energy dissipated by cracks, by friction between particles. In the post-peak stage, the external work slightly decreases and remains constant. The elastic energy becomes quasi-null. The number of cracks increases significantly, hence the energy dissipated by cracks increases and reaches its maximal value before remaining constant. The increase of the cracks number means the increase of the broken contacts. So the energy dissipated by friction between particles increases much because the friction between particles occurs when the contact is broken. It attains its maximum and remains constant. The energy dissipated by damping increases significantly (Fig. 3). In fact when the contact is broken the particles become free in the movement. Consequently, both damping force and moment become important in relation to the increase of translational and angular accelerations. The associated energy reaches its maximum value and remains constant. The kinetic energy $E_{\text{ki}}$ is low in the pre-peak region. At the peak, $E_{\text{ki}}$ attains its maximum. In the post-peak phase, the kinetic energy becomes low.

To verify the robustness of the proposed energy formulation, the balance energy ratio $W'$ defined as a ratio between the balance energy and the external work was computed $W' = W/W_{\text{ext}}$. The value of this ratio varies between 1% and 3%, which validates the implementation of energy balance for triaxial compression paths (Table 1).
Fig. 3. (a) Geometry and boundary conditions of the triaxial compression test; (b) Evolution of the deviatoric stress and the different energy components during a triaxial compression test.

Table 1. Value of different energetic components at the end of the triaxial compression test.

<table>
<thead>
<tr>
<th>Energetic components</th>
<th>External Work (W_{\text{ext}})</th>
<th>Elastic Energy (E_{\text{el}})</th>
<th>Kinetic Energy (E_{\text{ki}})</th>
<th>Energy Dissipated by Cracks (E_{\text{cr}})</th>
<th>Energy Dissipated by Friction (E_{\text{fr}})</th>
<th>Energy Dissipated by Damping (E_{\text{da}})</th>
<th>Energy Balance (W)</th>
<th>Error (W^*)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value (kJ)</td>
<td>1956</td>
<td>44</td>
<td>0.03</td>
<td>275</td>
<td>501</td>
<td>1105</td>
<td>29</td>
<td>1.5%</td>
</tr>
</tbody>
</table>

4.2. Direct tension test

In order to calculate the energy balance for a tensile loading path, a direct tension test was simulated. The test, illustrated in Fig. 4, was run with a loading velocity equal to 0.002 m/s. This velocity was chosen by performing a sensitivity analysis to ensure a quasi-static behavior. In the pre-peak region, the external work and the elastic energy increase as the axial stress with respect to axial strain. The cracks are not created in this stage so no energy is dissipated. The kinetic energy is low in this stage. At the peak, the external work and the elastic energy reaches their maximums. In the post-peak, failure occurs, the external work becomes constant, elastic energy decreases and then remains constant. The cracks are created when failure occurs, so the energy dissipated by cracks increase. The energy dissipated by friction between particles stills low. As in triaxial compression path, the kinetic energy reaches its maximum at the peak of loading and becomes again low in the post-peak stage. The different energy components and the balance energy ratio were evaluated. It has also been noted that the values of \(W^*\) are ranged between 2% and 3%, which remains acceptable (Table 2).

Fig. 4. (a) Illustration of direct tension test; (b) Evolution of the axial stress and the different energy components during the direct tension test.
Table 2. Value of different energetic components at the end of the direct tension test.

<table>
<thead>
<tr>
<th>Energetic components</th>
<th>External Work $W_{ext}$</th>
<th>Elastic Energy $E_{el}$</th>
<th>Kinetic Energy $E_{ki}$</th>
<th>Energy Dissipated by Cracks $E_{cr}$</th>
<th>Energy Dissipated by Friction $E_{fr}$</th>
<th>Energy Dissipated by Damping $E_{da}$</th>
<th>Energy Balance $W$</th>
<th>Error $W^*$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value (J)</td>
<td>1437</td>
<td>139</td>
<td>0.006</td>
<td>479</td>
<td>18</td>
<td>756</td>
<td>43</td>
<td>0.2%</td>
</tr>
</tbody>
</table>

4.3. Direct shear test

In order to evaluate the energy balance for a shear loading path, a direct shear test was simulated on a pre-fractured sample and subjected to a normal stress of 5 MPa. The test consists in two steps: applying first the normal stress until obtaining the equilibrium of the sample, and then applying the shear loading by applying the displacement velocity on the upper compartment of the shear box.

![Illustration of the direct shear test](image)

Fig. 5. (a) Illustration of the direct shear test [9]; (b) Evaluation of the shear stress and the different energy components for direct shear test.

For this test, the energy dissipated by damping and cracks are null since no cracks were created. The kinetic energy is so low during the simulation. External work, elastic energy and energy dissipated by friction between particles. In the first step of test (application of the normal stress), the external work and the elastic energy increase in the same way with the same values. Moreover the energy dissipated by friction between particles is null because the shear loading is not begun. In the second step of test (shearing process), the elastic energy becomes constant. Beyond the elastic limit, the external work and the energy dissipated by friction between particles increase with the same slope. In others words, during the sliding, the entire work of external forces is dissipated by friction. Finally, for shear loading the balance energy ratio $W^*$ remains inferior to 0.1% (Table 3).

![Energy components table](image)

Table 3. Value of different energetic components at the end of the direct shear test.

<table>
<thead>
<tr>
<th>Energetic components</th>
<th>External Work $W_{ext}$</th>
<th>Elastic Energy $E_{el}$</th>
<th>Kinetic Energy $E_{ki}$</th>
<th>Energy Dissipated by Cracks $E_{cr}$</th>
<th>Energy Dissipated by Friction $E_{fr}$</th>
<th>Energy Dissipated by Damping $E_{da}$</th>
<th>Energy Balance $W$</th>
<th>Error $W^*$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value (J)</td>
<td>14998</td>
<td>2271</td>
<td>3.12</td>
<td>0</td>
<td>12742</td>
<td>25</td>
<td>-43</td>
<td>0.02%</td>
</tr>
</tbody>
</table>
5. Conclusion

Through this paper, it is demonstrated that the balance of energy can be correctly evaluated for different loading paths with a 3D discrete element code. The expressions of the various components of energy involved in a given system are presented as well as the way they have been implemented in the code Yade Open DEM. The quasi-null value of the energy balance $W$ (or energy balance ration $W^*$ less than 3%) confirm the robustness of the code regarding the energetic problems. The different energy components were correctly computed. For example, the increment of energy dissipated by friction between particles is identical to the increment of the external work during the sliding stage of the direct shear test. After the validation of the energetic developments for different loading paths, we plan to apply this energy computation to underground excavations under quasi-static and dynamic conditions with possible correlations with geophysical measurements and later to simulate the rockburst phenomenon in the underground mines.

References