Research Article

Implementation of a Time-Domain Random-Walk Method into a Discrete Element Method to Simulate Nuclide Transport in Fractured Rock Masses

Yuexiu Wu, Quansheng Liu, Andrew H. C. Chan, and Hongyuan Liu

1 Key Laboratory of Rock Mechanics in Hydraulic Structural Engineering, Ministry of Education, Wuhan University, Wuhan, Hubei 430072, China
2 School of Engineering and ICT, University of Tasmania, Hobart, TAS 7005, Australia

Correspondence should be addressed to Yuexiu Wu; yuexiuwu@whu.edu.cn

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1. Introduction

1.1. Significance. With the development of nuclear waste disposal, nuclide transport with underground water in fractured rock masses has been an important issue in recent decades [1]. Especially after the nuclear leakage accident in Fukushima in Japan [2], this becomes more and more pressing. Therefore, it is essential to understand the principal transport mechanisms of radionuclides with underground water in fractured rock masses and evaluate the potential of radionuclide leakage from the repositories to the biosphere.

1.2. Literature Review about Nuclide Transport Simulation. Once the radionuclide is released from the container, it will flow with underground water in fractured rock masses to the biosphere. Therefore, water flow process in fractured rock masses should be studied firstly and foremost. Till now, five models have been found in literatures to simulate the water flow in fractured rock masses, that is, continuous model [3–5], dual-continuum model [6–8], equivalent continuum model [9–11], discrete fracture network (DFN) [12–14], and hybrid model [15–17]. In continuous model, rock masses are treated as continuum porous media with porosity and permeability and continuum-mechanics formulations can be used to describe flow and transport in media. It is a simple way but does not reflect the real flow path and velocity distribution in rock masses. In dual-continuum model, the rock matrix and fractures are represented as two overlapping and interactive continuum media with different mechanical and flow characteristics. However, this model cannot be applied to disconnected fractures in fractured media and the evaluation of fluid exchange between the matrix and the fractured domain is a complex task. In equivalent continuum model, fractured rock masses are treated as equivalent continua for large-scale analyses with equivalent mechanical and hydraulic properties that are measured or calculated in...
Representative Element Volume (REV). Nevertheless, REV may not exist for fractured rock masses in general. Moreover, flow rate in fractures is much larger than that in rock matrix. Thus, the DFN seems to be more suitable to simulate the water flow in fractured rock masses, irrespective of its high demand of computer resource. As a matter of fact, DFN has already been implemented into several numerical methods to simulate water flow in fractured rock masses, such as distinct deformation analysis (DDA) [18,18–22], numerical manifold method (NMM) [23,24], extended finite element method (XFEM) [25–27], and discrete element method (DEM) [28,29]. Compared with other numerical methods, DEM coupled with DFN shows the advantages of simulating water flow through fractured rock masses, since it is capable of simulating coupled hydromechanical effect and, especially, massive fractures.

Secondly, radionuclide transport process associated with water flow through fractured rock masses should be analyzed. According to former researches [30,31], the difficulty of discovering nuclide transport mechanism lies in solving the partial differential transport equation. As mentioned by Delay et al. [32], Lagrangian schemes (particle tracking methods) are more accurate and less computationally intensive than Eulerian schemes. The main Lagrangian schemes include random-walk particle tracking (RWPT) method [33,34], convolution-based particle tracking (CBPT) method [35,36], and continuous-time random-walk (CTRW) method [37,38]. For very heterogeneous media such as fractured rock mass, flow velocities may span several orders of magnitude. With the RWPT method, a large number of jumps are required in low-velocity areas to move the particles significantly, which may cause the calculations to become inefficient in terms of computational costs. The CBPT method is valid for mainly steady-state flow field and linear transport process. Some discrepancies are caused when the CTRW method is employed to the advection-dispersion equation. Aiming at removing these constraints, Delay and Bodin [39] firstly proposed a new method called time-domain random-walk (TDRW) method to simulate solute transport in heterogeneous media, which was further extended to simulate solute transport in fractured rock masses [40–42].

As can be seen from the literature review above, DEM and the TDRW method have the advantages of modelling water flow and solute transport in fractured rock masses. Thus, the TDRW method was firstly implemented into the DEM software UDEC in this paper.

1.3. Objectives and Organization of Paper. The primary objective of this paper is to propose a new numerical tool to simulate nuclide transport with underground water in fractured rock mass. In order to achieve this objective, the paper is organized as follows.

Theoretical introduction about hydromechanical coupling in UDEC is given in Section 2; Section 3 discusses nuclide transport equation and theory about time-domain random-walk method; the implementation of TDRW into UDEC is given in Section 4; verification of implementation program with analytical solution is given in Section 5; Section 6 discusses application of implementation program to DECOVALEX (DEmonstration of Coupled models and their VALidation against EXperiment) 2011 project to analyze the effect of matrix diffusion and stress on nuclide transport process; and conclusion is given in Section 7.

Throughout this study, it was concluded that, due to the effects of both matrix diffusion and stress, it took much more time for particles to outflow the fractured rock masses. Moreover, the proposed numerical method is a valuable numerical tool to study nuclide transport process through fractured rock masses, which has also wide applications in the field of underground water pollution, oil reservoir engineering, and so on.

2. Principle of Flow in UDEC

In UDEC [43], fractured rock masses are divided into many deformable blocks whose boundaries are fractures. Each deformable block is meshed into triangle elements, as shown in Figure 1. Motion is first calculated at the grid points of triangle elements within blocks and the stresses within the elements are then obtained according to the block material constitutive relations. Moreover, a fully coupled mechanical-hydraulic analysis can also be performed in UDEC, in which matrix is regarded as impermeable and fluid flows only through fractures.

2.1. Mechanical Behavior. For any grid point \( p \) in a triangle element, the total force is obtained as a sum of four terms:

\[
F_p = F^p + F^\gamma + F^\sigma + F^\alpha,
\]

where \( F^\sigma \) are the external applied loads, \( F^\sigma \) are the internal stresses contributed by zones adjacent to grid point \( p \), \( F^\gamma \) are the contact forces from all the adjacent blocks, and \( F^\alpha \) are gravity forces. The internal stress is calculated using the following equation:

\[
F^\sigma = \int_C \sigma_{ij} n_j ds,
\]

where \( \sigma_{ij} \) is element stress tensor and \( n_j \) is the unit vector outward normal to the integral path \( C \), which follows the closed polygonal line around the grid point \( p \). The gravity force is calculated using the following equation:

\[
F^\alpha = g_p m_p,
\]

where \( m_p \) is the lumped gravitational mass, defined as the sum of one-third of the masses of triangles connected to the grid point, and \( g_p \) is the gravitational acceleration.

The contact force \( F^\gamma \) exists only when grid points are on block boundaries (like grid point \( n \) in Figure 1) and is calculated according to the contact type and joint constitutive model. Taking the Coulomb slide model as an example, the contact force at the grid point from an adjacent block is divided into two components: \( F^\gamma_{\text{nc}} \) (normal contact force) and \( F^\gamma_{\text{sc}} \) (shear contact force):

\[
F^\gamma_{\text{nc}} := F^\gamma_{\text{sc}} - k_n \Delta u_n l,
\]
where $\Delta u_n$ is the normal displacement increment for each contact, $k_n$ is joint normal stiffness, and $l$ is the contact area, and

$$ F^c_p = F^c_p - k_n \Delta u_n l, $$  \hspace{0.5cm} (5)

where $\Delta u_s$ is the shear displacement increment and $k_s$ is joint shear stiffness. If the shear contact force exceeds the shear strength, it is modified according to the following equation:

$$ F^c_p = \text{sign} (\Delta u_s) (c + \tan \theta), $$  \hspace{0.5cm} (6)

where $c$ is joint cohesion and $\theta$ is joint frictional strength.

The motion of the grid point is obtained according to Newton's second law:

$$ \frac{d \dot{u}_p}{dt} = \frac{F_p}{m_p}, $$  \hspace{0.5cm} (7)

where $\dot{u}_p$ is the velocity of grid point $p$ and $t$ is time.

After the calculation of displacement of each grid point, the strain and stress of each triangle element are obtained according to the block constitutive model:

$$ \Delta e_{ij} = \frac{1}{2} (u_{i,j} + u_{j,i}) \Delta t, $$  \hspace{0.5cm} (8)

where $\Delta t$ is time step.

$$ \Delta \sigma_{ij} = \lambda \Delta e_{ij} \delta_{ij} + 2 \mu \Delta e_{ij}, $$  \hspace{0.5cm} (9)

where $\lambda$ and $\mu$ are Lamé's constant and $\Delta e_v$ is the bulk strain.

2.2. Fluid Flow in Fractures. Depending on the different contact type, the flow rate in contact domain is calculated differently. For a point contact (from a domain with pressure $p_1$ to a domain with pressure $p_2$),

$$ q = -k_c \Delta p, $$  \hspace{0.5cm} (10)

where $q$ is the flow rate, $k_c$ is a point contact permeability factor, and $\Delta p$ is the pressure difference calculated using the following equation:

$$ \Delta p = p_2 - p_1 + \rho_w g (y_2 - y_1), $$  \hspace{0.5cm} (11)

where $\rho_w$ is the fluid density; $g$ is the acceleration of gravity; and $y_2$ and $y_1$ are $y$-coordinates of the domain centers which are perpendicular to the joint surface. For edge-edge contact (an edge of one element contacts with an edge of another element), the flow rate in contact domain is calculated using the following equation:

$$ q = -k_j \frac{a}{l} \Delta p, $$  \hspace{0.5cm} (12)

where $k_j$ is a joint permeability factor (whose theoretical value is one-twelfth dynamic viscosity of fluid), $a$ is the contact hydraulic aperture, and $l$ is the length assigned to the contact between the domains. Fluid velocity can be obtained using (13) according to the flow rate calculated according to (10) and (12):

$$ u_f = \frac{q}{A}, $$  \hspace{0.5cm} (13)

where $A$ is the fluid crossing area.

2.3. Coupled Mechanical-Hydraulic Analysis. When the fluid flows in the contact domain, the fluid pressure is regarded as an external force applied on the contact domain, which
3. Nuclide Transport

3.1. Nuclide Transport Equation. It is known that nuclide transport is the consequence of several physical mechanisms such as advection, hydrodynamic dispersion, matrix diffusion, decay, and adsorption. As mentioned by Bodin et al. [42], the nuclide transport can be described in three media: (1) fracture, (2) matrix, and (3) stagnant zone. The stagnant zones in the fracture plane act as an additional “nonflowing” pore space available for solute diffusion. Thus, transport in the stagnant zone is ignored in this study. The nuclide transport equation in 2D fracture can be written as follows [42]:

\[
\frac{\partial c_f}{\partial t} + \lambda c_f = -\frac{u_f}{R_f} \frac{\partial c_f}{\partial x} + \frac{1}{R_f} \frac{\partial}{\partial x} \left( D_f \frac{\partial c_f}{\partial x} \right) + \frac{D_f}{bR_f} \frac{\partial c_m}{\partial y} \bigg|_{y=b},
\]

where \( c_f \) is the concentration of nuclide in fracture, \( u_f \) is the fluid velocity in fracture, \( b \) is half of the aperture, \( c_m \) is the concentration of nuclide in rock matrix, \( \lambda \) is the decay constant of nuclide, the term \( \lambda c_f \) represents the decay effect, \( D_f \) is the hydrodynamic dispersion coefficient in fracture, \( R_f \) is a retardation factor accounting for the sorption of nuclide onto fracture surface, the last term on the right of the equation represents the matrix diffusion effect, and \( x \) and \( y \) are the space coordinates along and perpendicular to the fracture, respectively, as shown in Figure 2. Not considering the effect of decay and matrix diffusion, the equation becomes the advection-dispersion-sorption (ADS) equation.

The retardation factor \( R_f \) is defined using the following equation [42]:

\[
R_f = 1 + \frac{K_f}{b},
\]

where \( K_f \) is the surface sorption coefficient of nuclide onto the fracture surface. The nuclide transport equation in rock matrix can be written as follows [42]:

\[
\frac{\partial c_m}{\partial t} + \lambda c_m = D_a \frac{\partial^2 c_m}{\partial y^2},
\]

where \( D_a \) is the apparent-diffusion coefficient of the nuclide in the matrix, which is calculated as \( D_a = D_e / (\theta_m + \rho_m K_m) \); \( D_e \) is effective diffusion coefficient; \( \theta_m \) is matrix porosity; \( \rho_m \) is matrix density; and \( K_m \) is volumetric-sorption coefficient of the nuclide in the matrix.

3.2. Time-Domain Random-Walk Method

3.2.1. Advection-Dispersion-Sorption Equation. In TDRW method, the solution of (16) can be divided into three steps: (1) advection-dispersion-sorption (ADS) transport equation, (2) decay effect, and (3) matrix diffusion effect. The ADS equation can be rewritten as (19) [42], considering the change of variable \( (\partial c_f / \partial t = (\partial c_f / \partial x)(\partial x / \partial t) = (\partial c_f / \partial x)(u_f / R_f) ) \):

\[
\frac{\partial c_f}{\partial x} = -\frac{R_f}{u_f^2} \frac{\partial}{\partial t} \left( u_f + \frac{\partial D_f}{\partial x} \right) c_f + \frac{R_f}{u_f^2} \frac{\partial^2}{\partial t^2} \left( D_f c_f \right). \tag{19}
\]

By applying an equivalent change of variable, (19) can be written with Fokker-Planck formalism:

\[
\frac{\partial \phi}{\partial x} = -\frac{1}{v^2} \frac{\partial}{\partial t} \left( \phi \right) + \frac{1}{2v^2} \frac{\partial^2}{\partial t^2} \left( \phi \right). \tag{20}
\]

According to the solution of random-walk approach, the mean and variance of nuclide transport time distribution for a displacement of \( dx \) are obtained using (21a) and (21b), respectively [42]:

\[
\mu_t (dx) = \frac{R_f}{u_f} \left( u_f + \frac{\partial D_f}{\partial x} \right) dx, \tag{21a}
\]

\[
\sigma_t^2 (dx) = \frac{2R_f}{u_f} d x. \tag{21b}
\]
For Peclet number $Pe = u_f dx / D_f$ larger than 10, it is shown that the travel time distribution is lognormal for a displacement of $dx$ [30]:

$$
\ln (\Delta t_f) = \mu_{ln} + Z_N \sigma_{ln},
$$

(22a)

$$
\mu_{ln} = \ln \left( \frac{\mu_4}{\sqrt{1 + \sigma_4^2 / \mu_4^2}} \right),
$$

(22b)

$$
\sigma_{ln}^2 = \ln \left( 1 + \frac{\sigma_4^2}{\mu_4^2} \right),
$$

(22c)

where $\Delta t_f$ is the travel time of the nuclide for a displacement $dx$ and $Z_N$ is a random number that follows a standard normal distribution.

For very low flow velocities in short fractures (i.e., the Peclet number $Pe$ is less than 10), the assumption of a lognormal distribution is not accurate. Bodin et al. [30] suggested that (22b) should be modified:

$$
\mu'_{ln} = \beta \mu_{ln} = \left(1 - \frac{1}{33 Pe}\right) \ln \left( \frac{\mu_4}{\sqrt{1 + \sigma_4^2 / \mu_4^2}} \right).
$$

(23)

The total travel time in a fracture of length $dx$ can be written as

$$
\Delta t_f = \exp (\mu_{ln} + Z_N \sigma_{ln}) \quad \text{(for } Pe > 10)\). \quad (24)
$$

When Peclet number $Pe$ is less than 10, the variable $\mu_{ln}$ is replaced by $\mu'_{ln}$.

3.2.2. Matrix Diffusion. To account for the interaction between ADS and matrix diffusion in a bond of length $dx$, the fluid velocity $u_f$ and dispersion coefficient $D_f$ in (22a) and (22b) were replaced by an apparent fluid velocity $u_f^*$ and an apparent dispersion coefficient $D_f^*$ [42]:

$$
u_f^* = R_{diff} u_f,
\quad D_f^* = f (u_f^*), \quad \text{like } D_f^* = \alpha u_f^*, \text{ or } D_f^* = D_f,
$$

(25)

where $\alpha$ is a dispersivity constant and $R_{diff}$ is a retardation factor accounting for matrix diffusion:

$$
R_{diff} = 1 + \frac{\Omega \sqrt{2 \sigma_4}}{R_f} \left( \frac{\exp (-\xi^2)}{\sqrt{\pi \text{erfc} (\xi)}} - \xi \right),
$$

(26)

where $\xi = (dx / u_f)(\sqrt{2 / \sigma_4})$, $\Omega = \sqrt{(\theta_m + \rho_m K_m) D_f / 2 b}$, $\theta_m$ is the matrix porosity, $\rho_m$ is the bulk density of matrix, $K_m$ is the volumetric-sorption coefficient of nuclide in matrix, and $D_n$ is the effective diffusion coefficient of nuclide in matrix.

The modified (22a) and (22b) with $u_f^*$ and $D_f^*$ in place of $u_f$ and $D_f$, respectively, were used to calculate the mean and variance of the log transform $\mu_{ln}$ (or $\mu'_{ln}$) and $\sigma_{ln}^2$ according to (22b) and (22c), respectively. The total travel time for the coupled ADS and matrix diffusion is calculated using (27) for $Pe > 10$ [42]:

$$
\Delta t_{fm} = \exp (\mu_{ln} + Z_N \sigma_{ln}) + \left( \frac{\Omega t_0}{R_f \text{erfc}^{-1} (U_{01})} \right)^2,
$$

(27)

where $t_0$ is the particle residence time in the fracture for pure advection delayed by sorption reactions onto the fracture walls, expressed as $t_0 = R_f (dx / u_f)$ and $U_{01}$ is a random number drawn from a uniform distribution between 0 and 1. For $Pe < 10$, the variable $\mu_{ln}$ in (27) is replaced by $\mu'_{ln}$.

3.2.3. Decay Effect. If the linear decay model is used, the nuclide mass decreases according to the following equation [42]:

$$
m_{n+1}^p = m_n^p \exp (-\lambda (t_{n+1} - t_n)),
$$

(28)

where $m_{n+1}^p$ and $m_n^p$ represent the masses of the nuclide at node $n + 1$ and $n$, respectively.

For simplicity, the decay effect is not considered here, which accords with the situation in DECOVALEX 2011 project.

3.2.4. Mass Partition Model at Fracture Intersection. Because the nuclide was tracked in fractures from the inflow boundary to the outflow boundary in TDRW method, the mass partition model at fracture intersection is vital in the calculation. There are three models: the perfect-mixing model [44], the stream-tube model [45], and the diffusional-mixing model [46]. The focus of this paper is not the mass partition law of solute at fracture intersection. For simplicity, the perfect-mixing model was adopted here:

$$
P_{ij} = \frac{Q_j}{\sum Q^-},
$$

(29)

where $P_{ij}$ is the probability of particle transition from an inlet bond $i$ to an outlet bond $j$; $Q_j$ is the flow rate in the outlet bond $j$; and $\sum Q^-$ is the sum of the flow rates over all the outlet bonds connected to the junction of interest. It can be seen from (29) that mass partition is only related to the outlet fluxes.

4. Implementation of TDRW into UDEC

In TDRW method, the fractures are regarded as the sum of many bonds and a continuous connection of the bonds from inlet to outlet boundaries is considered as a flow path (e.g., the dashed line shown in Figure 3). In a bond, the fluid flow is a 1D flow with a constant fluid velocity. Therefore, the first thing, before we do the particle tracking calculation, is to get the flow paths of particle transport.

In UDEC, jointed rock masses are divided into blocks by fractures. The geometric information of fracture and block, such as corner coordinates and their connection, is stored in list in files (e.g., BLOCK.FIN, DOMAIN.FIN, and CONTACT.FIN). According to the above files, the coordinates of
fracture intersections can be obtained easily, as well as their connective information. The TDRW method is implemented into UDEC using the FISH language, as shown in Figure 4.

After the hydromechanical calculation with UDEC, the final flow velocity at each fracture intersection can be obtained and this is the input data for TDRW simulation. At the beginning, nuclide particles are injected into the fractured rock masses through the inlet boundary. According to the inlet flow rate, the injection location of each particle is chosen. The particle transport time in each bond is calculated with (22a), (22b), (22c)–(28). At the fracture intersection, the perfect-mixing model is adopted to decide the next step of particles until they arrive at the outlet boundary. If the injected mass is $M_0$ and the total number of particles is $N$, mass of each particle is expressed as

$$m_p = \frac{M_0}{N}.$$  

(30)

If the collected particle number on the outlet boundary is $N_i$ in a period of time $dt$, the outlet concentration is written as

$$c_i = \frac{m_p N_i}{\sum Q_{out} dt}.$$  

(31)

where $\sum Q_{out}$ is the total outlet fluid flow rate.

The following steps are adopted to obtain the curve of the relative concentration versus time:

1. Calculate the maximum and minimum travel times of all particles.

2. Divide the travel time period (from minimum travel time to maximum travel time) into $N$ parts, in which the time interval is $\Delta t$.

3. Collect particles (number is $N_i$) in each time interval and calculate their concentration with (31).

5. Verification

Two numerical particle transport tests were performed to verify the implementation of the TDRW method into UDEC,
that is, particle transports in a single fracture and in a simple fracture network.

5.1. Analytical Solution of the Nuclide Transport Equation.
Analytical solution for nuclide transport in a single fracture has been deduced in literatures [47–49]. The analytical solution is to be used later to verify the numerical results obtained with combination of DEM and TDRW. For simplicity, the analytical solution for advection-dispersion transport equation with constant concentration injection is adopted, which can be written as follows:

\[
\frac{c_f}{c_0} = \frac{1}{2} \left[ \text{erfc}\left( \frac{x - u_f t}{2 \sqrt{D_f t}} \right) - \exp\left( \frac{u_f x}{D_f} \right) \text{erfc}\left( \frac{x + u_f t}{2 \sqrt{D_f t}} \right) \right],
\]  

(32)

where \(c_0\) is the constant concentration injection; and \(\text{erfc}(x)\) is a complementary error function.

5.2. Calibration Test 1: Particle Transport in a Single Fracture.
In the calibration test 1 of particle transport in a single fracture, the injection mass at the inlet boundary of the fracture was 0.001 g, and the total number of nuclide particles was 30000. The trace of the fracture was 10 m, fluid velocity was \(4.0 \times 10^{-4} \text{ m/s}\), hydrodynamic dispersion coefficient in the fracture was \(2.0 \times 10^{-5} \text{ m}^2/\text{s}\), and the aperture of the fracture was \(2.5 \times 10^{-4} \text{ m}\). Thus, the Peclet number \(\text{Pe} = u_f L/D_f\) was 200. The analytical solution can be obtained using (32). The corresponding numerical solution was obtained using the TDRW method implemented into UDEC, which is compared in Figure 5 with the analytical solution. From Figure 5, the maximum and minimum differences between numerical and analytical solutions are 11% and 0.03%, respectively.

The reason for this difference is that the numerical results are iterated. In general, the results from UDEC with the TDRW method implemented agreed well with the analytical solution.

5.3. Calibration Test 2: Particle Transport in a Simple Fracture Network.
The simple fracture network was shown in Figure 6. A hydromechanical calculation was performed with UDEC at first. The water heads at top and bottom boundaries were 100 m and 99 m, respectively, and two lateral sides were impermeable. The aperture of the fractures was \(2.5 \times 10^{-6} \text{ m}\). The other parameters were the same as those in test 1. The analytical solution was the summarizing solution of (32) for each fracture. Figure 7 was the comparison between the numerical solution and analytical solution for the particle transport in this simple fracture network. From Figure 7, the maximum and minimum differences between numerical and analytical solutions are 10.1% and 0, respectively.

From the comparisons shown in Figures 5 and 7, it is concluded that the numerical solution obtained using the
TDRW method implemented into UDEC was close to the analytical solution. Thus, it is proven that the TDRW method implemented into UDEC can well simulate nuclide particle transport in fracture networks.

### 6. Application to Particle Transport in Discrete Fracture Network

#### 6.1. Calculation Models and Material

The proposed method was used to evaluate the effect of matrix diffusion and stress ratio on particle transport process in the fracture network studied in the DECOVALEX 2011 project. The calculation model was shown in Figure 8, which contained 771 fractures. The hydromechanical boundary condition was depicted in Figure 9. The hydraulic heads at the top and bottom boundaries were 30 m and 10 m, respectively. On the two lateral sides, a linear distribution of hydraulic head was applied with a maximum value of 30 m and a minimum value of 10 m; in order to analyze the effect of matrix diffusion, the same hydromechanical boundaries were applied for the case with matrix diffusion and the case without matrix diffusion. Accounting for the effect of stress ratio, simulations without stress \((\sigma_v = \sigma_h = 0)\) and with different stress ratios were performed. For the cases with stress, a constant vertical stress of 5 MPa was specified on the top and bottom boundaries. And a horizontal stress, with ratios of vertical/horizontal stress varying from 1 to 2, 3, and 5, was applied, respectively \((\sigma_v/\sigma_h = 1, 2, 3, 5)\). For each calculation, 10,000 particles were injected at a time along the inflow boundary. At each injected point, which is the intersection of fracture and inflow boundary, the injecting particle number is proportional to the local flow rate. This is equivalent to injecting a constant concentration at the inflow boundary.

The block material property was \(K = 54.23\) GPa, and \(G = 34.11\) GPa.

The normal behavior of the fractures in the calculation model was nonlinear, as shown in Figure 10 [28]. Other parameters of the fractures were listed in Table 1 [28]. The aperture of the fractures followed a lognormal distribution with a mean value of 65 \(\mu\)m and a second moment of 1.0. The minimum and maximum apertures were 1 \(\mu\)m and 200 \(\mu\)m, respectively. The residual aperture was assumed to be 20% of the initial aperture.

The hydrodynamic dispersion coefficient in fractures was \(1.0 \times 10^{-7}\) m²/s. The hydromechanical calculation was done firstly and the fluid velocity could then be obtained. After that, the TDRW simulation was performed.

#### 6.2. Results and Discussion

##### 6.2.1. Fluid Flow

Flow rates in fractures which intersect the bottom, right, and left outlet boundaries with different applied stress were shown in Figures 11–13. The location in Figures 11–13 was the coordinates of intersections of fractures along the corresponding boundary. On the bottom outlet boundary, the maximum flow rate and average flow rate without stress applied were \(1.16 \times 10^{-8}\) m³/(s·m) and \(4.06 \times 10^{-8}\) m³/(s·m), respectively, which were larger than those with stress applied. From Figure 11, it was shown that the main outlet positions (intersections with flow rate larger than the average flowrate) changed comparing the results not considering stress with results considering stress effect. When the stress was applied, the main outlet positions were same for different stress ratio. However, the maximum and average values of flow rate decreased with the increasing of stress ratio. On the right boundary, the maximum and average values of flow rate in intersections without stress were \(8.59 \times 10^{-8}\) m³/(s·m) and \(3.13 \times 10^{-8}\) m³/(s·m), respectively, which were obviously larger than those considering stress ratio. On the left boundary, the maximum and average values of flow rate in intersections without stress were \(5.38 \times 10^{-8}\) m³/(s·m), \(3.12 \times 10^{-8}\) m³/(s·m), \(1.97 \times 10^{-8}\) m³/(s·m), respectively, which were obviously larger than those considering stress ratio. On the right boundary, the maximum and average values of flow rate in intersections without stress were \(2.58 \times 10^{-8}\) m³/(s·m), \(1.95 \times 10^{-8}\) m³/(s·m), \(1.0 \times 10^{-8}\) m³/(s·m), respectively, which were obviously larger than those considering stress ratio.

##### 6.2.2. Nuclide Transport

(1) Without Matrix Diffusion

After the nuclide transport calculations with different stress ratio, nuclides which outflow the outlet boundary at a certain time were collected. The ratio of number of leaking nuclides from the outlet boundary to the total number of nuclides was plotted versus traveling time (i.e., breakthrough curve) in Figure 14. From Figure 14, it was shown that the total traveling times with different stress boundaries were \(1.97 \times 10^4\) s, \(2.58 \times 10^4\) s, \(3.12 \times 10^4\) s, \(1.95 \times 10^4\) s, respectively.

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**Table 1: Material parameters of the fracture in the calculation model.**

<table>
<thead>
<tr>
<th>Mechanical properties</th>
<th>Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shear stiffness, (Ks) (GPa/m)</td>
<td>434</td>
</tr>
<tr>
<td>Friction angle (°)</td>
<td>24.9</td>
</tr>
<tr>
<td>Dilation angle (°)</td>
<td>5</td>
</tr>
<tr>
<td>Cohesion (MPa)</td>
<td>0</td>
</tr>
</tbody>
</table>

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**Figure 8:** Calculation model studied in the DECOVALEX 2011 project.
Inflow boundary

Right outlet boundary

0.3 MPa

0.1 MPa

Bottom outlet boundary

Figure 9: Hydromechanical boundary condition.

$\sigma_v$

$\sigma_h$

$h$

$0.1$ MPa

$0.3$ MPa

Figure 10: Relationship between normal displacement and normal stress.

$10^5$ s, and $2.4 \times 10^5$ s, respectively. The mean traveling time for each calculation was $3.29 \times 10^7$ s, $4.31 \times 10^7$ s, $5.21 \times 10^7$ s, $1.27 \times 10^8$ s, and $2.19 \times 10^8$ s, respectively. Through dividing the total traveling times without stress, the normalized total travel times with stress ratio 1, 2, 3, and 5 are 1.3, 1.58, 9.89, and 12.18.

The total and average travel times for all nuclides leaking out from the fractured rock masses increased with the increasing of stress ratio. The reason for this phenomenon was that fractures were compressed to be narrow and it was difficult for nuclides to leak.

(2) With Matrix Diffusion. In the case with matrix diffusion, the porosity of the matrix $\theta_m$ was 0.316%. For simplicity, advection, dispersion, and matrix diffusion were considered in this case, while sorption in the matrix was not allowed ($K_m = 0$ m$^3$/kg). The effective diffusion coefficient of the nuclide in the matrix is $1.0 \times 10^{-11}$ m$^2$/s. The breakthrough curves with different stress applied considering the effect of matrix diffusion were shown in Figure 15. From Figure 15, it was shown that the total traveling times with different stress boundaries were $3.03 \times 10^7$ s, $7.98 \times 10^7$ s, $1.086 \times 10^8$ s, $8.02 \times 10^8$ s, and $8.42 \times 10^8$ s, respectively. The mean traveling time for each calculation was $4.88 \times 10^9$ s, $1.77 \times 10^9$ s, $2.89 \times 10^9$ s, $5.2 \times 10^9$ s, and $7.7 \times 10^9$ s, respectively. Through dividing the total traveling times without stress, the normalized total travel times with stress ratio 1, 2, 3, and 5 are 2.63, 3.58, 26.5, and 27.8. The total and average travel times for all nuclides leaking out from the fractured rock masses increased with the increasing of stress ratio.

Comparing the breakthrough curves in Figure 14 with those in Figure 15, it could be seen that the total travel time for all particles outflowing from fractured rock masses without the matrix diffusion was much shorter than that with matrix
Figure 12: Flow rate distribution of fractures on the right outlet boundary.

Figure 13: Flow rate distribution of fractures on the left outlet boundary.

Figure 14: Breakthrough curve with different stress ratios without matrix diffusion.

7. Conclusion

Although the nuclide transport with underground water in fractured rock masses had been studied for decades by many researchers, the coupled hydromechanical effect and the effect of matrix diffusion and stress were still not discussed systematically. Aiming at solving these problems, the time-domain random-walk (TDRW) method was firstly implemented into the discrete element method (DEM) in this study. During each time step of hydromechanical calculation, hydraulic aperture of each fracture and fluid pressure on fracture surface were calculated according to (15) and (14) to consider the coupled hydromechanical effect. To present the effect of matrix diffusion on nuclide transport, (24) and (27)
were adopted to get the breakthrough curves. The implementation was then verified against the analytical solutions for nuclide transport in a single fracture and a single fracture network. After that, the implementation was applied to simulate the nuclide transport in a complex fracture network investigated in the DECOVALEX 2011 project. Finally, the effect of matrix diffusion and stress on nuclide transport was discussed in detail. Throughout this study, the following conclusions were drawn:

1. The implementation of the TDRW method into UDEC could well simulate the nuclide transport with underground water in fractured rock masses.

2. Through the hydromechanical calculation for DECOVALEX 2011, the closure of fractures increased with the increasing of applied stress and it caused decreasing of fluid flow in fractured rock masses. More time was needed for fluid or particle transport through a narrow path. This finally resulted in the fact that the total travel time of the nuclide particles with the external stress applied on the fracture network was longer than that without the applied stress.

3. According to the numerical results for the nuclide transport in the complex fracture network investigated in the DECOVALEX 2011 project, matrix diffusion has a significant effect on the nuclide transport. The travel time of the nuclide particle with the matrix diffusion modelled was much longer than that without the matrix diffusion.

Conflicts of Interest

The authors declare that the received funds do not lead to any conflicts of interest regarding the publication of this manuscript.

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