Particle Methods

- An Overview -

DEM-Simulation of a cutting process (Source: Chair for Rock Mechanics, TU BAF)

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1 Introduction

What are particle methods?

Numerical calculation methods can be divided into:

- Explicit and implicit methods in terms of time discretization
- Continuum mechanical (mesh-based) and discontinuum mechanical (meshfree) methods in terms of spatial discretization

All methods can be executed independent of spatial discretization as implicit or explicit calculations. Typical representatives of mesh-based methods are FEM (Finite Element Method), REM (Rand Element Method) or VEM (Volume Element Method). Typical representatives of meshfree methods are DEM (Discrete Element Method), SPH (Smooth Particle Hydrodynamics) or MD (Molecular Dynamics).

While in classical continuum mechanics coherence of the body (continuum) is maintained (neighbor relations maintain resp. will be predefined), discontinuum mechanics allows the inspection of interactions of several single bodies (continua). Therefore these methods require an automated contact detection algorithm as well as corresponding contact laws, that get active during interaction (physical or as field force over far distances).

![Figure 1](source: Chair for Rock Mechanics, TU BAF)

Where are particle methods used?

- Simulation of granular media and hard rocks (DEM)
- Fluid simulations (SPH, LBM)
- Simulation of molecules and nanoparticles (MD)
- Calculations for astrophysics (SPH)
2 Discrete Elements

Discrete Element Method (DEM) is a particle method based on Newton’s laws of motion. Particles can move with six degrees of freedom (three for translational and three for rotational movement). Particles are rigid and hence not deformable. In principle particles can have every arbitrarily geometrical shape, in which sphere shape is most efficient from numerical point of view. When particles come in contact forces (depending on the chosen contact law) start acting on them. Furthermore external forces (e.g. gravity) can act on the particles. During a calculation contacts can be formed or break. Thus an efficient automated contact detection is one of the core components of a DEM-Software. Modelling with discrete elements take place in six phases:

1. Generation of particles + definition of boundary and initial conditions
2. Determination of contacts (between particles and between particles and boundary)
3. Calculation of forces $F$ and moments $M$ of all particles
4. Calculation of accelerations $\ddot{u}$ and $\dot{\omega}$, velocities $\dot{u}$ and $\omega$, and displacements $u$ and rotations of all particles
5. Calculation of new positions $x$ of all particles
6. Repeat step 2. to 5. with time step $\Delta t$ until stop criterion is achieved

Pioneer work to development of discrete resp. distinct elements was mainly provided by Peter A. Cundall, who improved and extended this method in numerous publications. Since 1995 DEM-Software PFC (Particle Flow Code) from company Itasca is available. With this code two-dimensional (PFC2D) as well as three-dimensional (PFC3D) models can be calculated. Following description mainly depends on manuals from this software.

2.1 Contact detection

Computing time for contact detection increases quadratically with the amount of particles in the model. Therefore it is necessary to use an optimized contact detection algorithm for DEM. Thereby all non-possible contacts will be sorted out in the first step. In the second step a more detailed procedure (computationally more intensive) analyzes, if remaining possible contacts are real contacts. When a real contact is found, contact law is applied to calculate the contact forces (see section 2.2).

Pre-sort processes can be divided in cellbased methods and methods with Verlet lists. In cellbased methods the model is splitted into smaller axis-aligned cells. The bigger the cells, the more possible contacts are pre-sorted. That leads to higher computing time. The smaller the cells, the more cells must be scanned, whereby also more computing time is needed. In the most DEM-Codes an optimal cell size is estimated by a heuristic method when the model is initialized. Applying method with Verlet lists a monitoring radius, including all possible contact neighbors, is assigned to every particle.
2.2 Contact laws

When two particles come in contact, acting forces were calculated by a contact law. Contact laws are built by different basic elements, that can be visualized with a circuit diagram. The basic elements are the spring $k_n$, the viscous dashpot $k_s$ and the frictional slider $k_f$ (also called frictional resistance or shear slider). In a pure elastic law contact force is described by two springs (see fig. 3) with stiffnesses $k_n$ for normal direction and $k_s$ for shear direction.

The normal contact force $F_n$ in the linear-elastic contact law is the product of constant normal stiffness $k_n$ and overlap $u_n$.

$$F_n = k_n u_n \quad (1)$$

The updated shear contact force $F_s^{new}$ arises from the sum of the shear contact force of the previous time step $F_s^{old}$ and the update value $\Delta F_s$.

$$F_s^{new} = F_s^{old} + \Delta F_s \quad \text{with} \quad \Delta F_s = -k_s \cdot \Delta u_s \quad (2)$$

At non-linear Hertz-Mindlin contact law stiffnesses $k_n$ and $k_s$ are calculated according to input parameters shear modulus $G$, Poisson ratio $\nu$, the radii of particles $R_1$ and $R_2$ and the amount of overlap $u_n$.

$$k_n = \left( \frac{G \sqrt{2R_d}}{(1-\nu)} \right) \sqrt{u_n} \quad \text{with} \quad R_d = \frac{2R_1 R_2}{R_1 + R_2} \quad (3)$$

$$k_s = \left( \frac{2(3G^2(1-\nu)R_d)^{\frac{1}{2}}}{2-\nu} \right) \cdot (F_n)^{\frac{1}{2}} \quad (4)$$

Every mechanical system “looses” energy (e.g. in terms of thermal energy with friction or plastic deformation). For taking account of this loss of energy during
particle contact in the model, a damping element is connected in parallel to the spring in the contact law. In the case of shearing energy disappears because of frictional sliding. Hence a frictional slider is connected in series in the contact law (see fig. 4).

![Figure 4: Contact law with viscous damping and frictional slider](image)

Damping acts proportional to velocity at normal contact force $F_n$, hence the amount of viscous damping $c_n \dot{u}_n$ must be added to $F_n$.

$$F_n = k_n u_n - c_n \dot{u}_n$$

(5)

Also the update of shear contact force $\Delta F_s$ must be added by the amount of viscous damping for shear direction $c_s \Delta \dot{u}_s$.

$$\Delta F_s = -k_s \cdot \Delta u_s - c_s \cdot \Delta \dot{u}_s$$

(6)

Here $c_n$ and $c_s$ are the normal and shear damping coefficients. $c_n$ (resp. $c_s$) is the product of damping ratio $\beta_n$ (resp. $\beta_s$) and critical damping constant $c_n^{crit}$ (resp. $c_s^{crit}$). $c_n$ is given by

$$c_n = \beta_n c_n^{crit} = 2 \beta_n \sqrt{mk_n},$$

(7)

where $m$ is the effective mass of the system. The calculation for $c_s$ happens analog.

The sliding behavior during shear movement is described by friction coefficient $\mu$. $\mu$ is defined as ratio of maximum shear contact force $F_s^{max}$ to normal contact force and limits shear contact force in case of sliding to

$$F_s^{max} = \mu |F_n|.$$  

(8)

2.3 Physics of DEM-Particles

In the following the description of physics of two DEM particles in contact is given. The used notations are shown in fig. 5.

Vectors $\vec{x}_1$ and $\vec{x}_2$ are position vectors from particle centres and $\vec{n}$ and $\vec{t}$ denote the normal resp. tangential unity vectors. The following relations arise (see fig. 5).

$$d_a = |\vec{x}_2 - \vec{x}_1|$$

$$\vec{n} = \frac{\vec{x}_2 - \vec{x}_1}{d_a}$$

$u_n = R_1 + R_2 - d_a$  

(9)
The position vector of contact point \( \vec{x}_c \) results as follows.

\[
\vec{x}_c = \vec{x}_1 + (R_1 - 1/2 \cdot u_n) \cdot \vec{n}
\]  
\( (10) \)

With stiffnesses \( k_n \) and \( k_s \) the normal and shear component of contact force \( F_c \) are determined. The normal component is calculated directly during creation of the contact. The shear component at this time is zero. It is increased (resp. decreased) by \( \Delta F_s \) at each time step.

\[
\Delta F_s = -k_s \cdot \Delta u_s \quad \text{and} \quad F_n = k_n u_n
\]  
\( (11) \)

The amount of shear displacement at each time step \( \Delta u_s \) is described at the end of this section in eq. (21). The contact force between two particles is then determined by

\[
\vec{F}_c = F_n \cdot \vec{n} + F_s \cdot \vec{t}.
\]  
\( (12) \)

Total force \( \vec{F} (\approx F_i) \), that acts on one particle, is composed of the sum of all contact forces from neighbor particles and gravity force \( \vec{F}_g = m \cdot \vec{g} \).

\[
\vec{F} = \sum_c \vec{F}_c + \vec{F}_g
\]  
\( (13) \)

Total force in \( i \)-direction \( (i \in \{1, 2, 3\}) \) \( F_i \), that acts on one particle, is determined by Newton’s second law (fundamental law of dynamics). \( F_i \) is calculated by multiplication of the particle mass \( m \) and the sum of its accelerations \( (\ddot{u}_i \text{ and } g_i) \).

\[
F_i = m \cdot (\ddot{u}_i + g_i)
\]  
\( (14) \)

Equation (15) arises from rearranging eq. (14).

\[
\ddot{u}_i = \frac{F_i}{m} - g_i
\]  
\( (15) \)

After double integration with respect to time \( t \) velocity \( \dot{u}_i \) and displacement \( u_i \) are determined.

\[
\dot{u}_i = \int \dddot{u}_i dt \quad \text{and} \quad u_i = \int \dot{u}_i dt
\]  
\( (16) \)
For description of rotational movement vector $\mathbf{r}_c$ is introduced. $\mathbf{r}_c$ connects the particle centre with the contact point. Therefore torsional moment $\mathbf{M}$ arises from the following relation.

$$
\mathbf{M} = \sum_c \left( \mathbf{r}_c \times \mathbf{F}_c \right) \quad \text{with} \quad \mathbf{r}_c = \mathbf{x}_c - \mathbf{x} \quad \text{and} \quad \mathbf{x} = \mathbf{x}_1 \quad \text{(for particle 1)} \quad (17)
$$

Torsional moment in $i$-direction $M_i$ is determined by multiplication of moment of inertia $J$ and angular acceleration $\dot{\omega}_i$.

$$
M_i = J \cdot \dot{\omega}_i \quad (18)
$$

Moment of inertia is given by $J = \frac{2}{5} \cdot mR^2$, because all particles are spheres. By rearranging and integration with respect to $t$ one gets angular acceleration $\omega_i$.

$$
\omega_i = \int \frac{5M_i}{2mR^2} dt \quad (19)
$$

With velocities $\mathbf{v}_1 = \mathbf{u}_i$ and $\mathbf{v}_2 = \mathbf{\dot{u}}_i$ and angular accelerations $\ddot{\omega}_1$ and $\ddot{\omega}_2$ relative velocity $\mathbf{v}_{rel}$ is computed.

$$
\mathbf{v}_{rel} = (\mathbf{v}_2 + \mathbf{\ddot{\omega}}_2 \times (\mathbf{x}_c - \mathbf{x}_2)) - (\mathbf{v}_1 + \mathbf{\ddot{\omega}}_1 \times (\mathbf{x}_c - \mathbf{x}_1)) \quad (20)
$$

Shear displacement at each time step $\Delta u_s$ from eq. (11) is calculated by multiplication of shear velocity $v_s$ with time step $\Delta t$. $v_s$ is calculated by relative velocity $v_{rel} = |\mathbf{v}_{rel}|$ by subtracting normal velocity.

$$
\Delta u_s = v_s \Delta t \quad \text{with} \quad v_s = v_{rel} - v_n \quad \text{and} \quad v_n = |\mathbf{v}_{rel} \cdot \mathbf{n}| \quad (21)
$$

### 2.4 Time integration

PFC calculates velocities $\dot{x}_i$ and $\omega_i$ at averaged time intervals $t \pm \Delta t/2$ and values for $x_i, \dot{x}_i, \omega_i, F_i$ and $M_i$ at primary intervals $t \pm \Delta t$. Numerical solution of integrations from eq. (16) and (19) is done with central difference quotient

$$
\dot{u}_i^{(t)} = \frac{1}{\Delta t} \left( u_i^{(t+\Delta t/2)} - u_i^{(t-\Delta t/2)} \right) \quad \text{and} \quad (22)
$$

$$
\dot{\omega}_i^{(t)} = \frac{1}{\Delta t} \left( \omega_i^{(t+\Delta t/2)} - \omega_i^{(t-\Delta t/2)} \right) .
$$

From the equations (22), (15) and (18) the “new” velocities at time $t + \Delta t/2$

$$
\dot{u}_i^{(t+\Delta t/2)} = \dot{u}_i^{(t-\Delta t/2)} + \left( \frac{F_i^{(t)}}{m} + g_i \right) \Delta t \quad \text{and} \quad (23)
$$

$$
\dot{\omega}_i^{(t+\Delta t/2)} = \omega_i^{(t-\Delta t/2)} + \left( \frac{5M_i^{(t)}}{2mR^2} \right) \Delta t .
$$

and the “new” particle positions at time $t + \Delta t$ are determined.

$$
x_i^{(t+\Delta t)} = x_i^{(t)} + \dot{u}_i^{(t+\Delta t/2)} \Delta t \quad (24)
$$

Again beginning at eq. (9) forces and displacements for next time step are computed with these new positions. For complete numerical description of motion and contact behavior see PFC manual (Itasca (2008)).
2.5 More complex approaches

The particle approaches based on rigid spheres can be extended in many respects:

**Clumps** can be formed out of two or more particles. With it more complex particle shapes can be created.

![Clumps](source: Chair for Rock Mechanics, TU BAF)

**Bonds** are linkages with that particles can be cohesively connected. In doing so several particles can be bonded to a grain (or cluster), that can break along cohesive bonds at according action. Hence it is possible to simulate solids like ceramics, concrete or hard rocks and also solid bridges between granular media.

![Bonds](source: Chair for Rock Mechanics, TU BAF)
Model boundaries are defined by planes and/or by particles itself. Planes can be rectangles, but they can also have more complex geometries (consisting of triangular partial planes). Particles can be used as boundary by fixing them (limitation of degrees of freedom) or by defining periodic boundaries. In the case of periodic boundaries particles, that touch the boundary box, are getting master particles. Every of these master particles generates one slave particle on the opposite side of the boundary. Thus the model is leaning on itself at the boundary.

**Figure 8:** DEM model with different boundaries, planes (left) and periodic boundary (right)
2.6 Examples

DEM permits to observe a solid body with respect of its deformation, stress and resistance behavior including crack propagation (see fig. 9).

Figure 9: Intrusion of a wedge into a solid body consisting of Voronoi-particles, with crack propagation (source: Chair for Rock Mechanics, TU BAF)

By using bonds cementations between sand grains, like they are present in sandstone, can be simulated. Following example shows a sample (left) and its broken bonds, that result from a shear fracture (visualized by grey cylinders).

Figure 10: Sandstone sample, setting (left) and cohesive bonds after shear fracture (right), (source: Chair for Rock Mechanics, TU BAF)
Further examples can be found in following list.

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**Table 1:** List of DEM applications with references
3 Smoothed Particles

The method of Smoothed Particle Hydrodynamics (SPH) was introduced by Monaghan (1988) for calculations of astrophysical phenomena. SPH was further developed and it is now also possible to simulate fluids with it. The continuum to be examined is approximated by discrete particles. Every SPH particle has a core, that is described by a kernel function $W(x, h)$ (also called smoothing kernel or interpolating kernel). Here $h$ is the effective area of a particle and $x$ is the distance to the centre. The kernel function in one-dimensional case can e.g. be a Gaussian distribution (see fig. 11).

\[ W(x, h) = \frac{1}{h\sqrt{\pi}} e^{-\frac{x^2}{h^2}} \]

Figure 11: Gauss kernel

The particles are attributed to physical quantities density $\rho$, position $\vec{x}$ and velocity $\vec{v}$. These must be recalculated in every time step.

3.1 Interpolation

The integral interpolant of any function $A(\vec{x})$ is defined as

\[ A(\vec{x}) = \int A(\vec{x}') W(\vec{x} - \vec{x}', h) d\vec{x}', \quad (25) \]

whereby it is integrated over the whole space and kernel function $W$ has following properties (Monaghan (1992)).

\[ \int W(\vec{x} - \vec{x}', h) d\vec{x}' = 1 \quad \text{and} \quad \lim_{h \to 0} W(\vec{x} - \vec{x}', h) = \delta(\vec{x} - \vec{x}') \quad (26) \]

The integral interpolant of field variable $A$ of one SPH particle at position $\vec{x}$ can be approximated by a summation interpolation of its neighbor particles ($b$).

\[ A(\vec{x}) = \sum_b A_b \frac{m_b}{\rho_b} W(\vec{x} - \vec{x}_b, h) \quad (27) \]

$m_b, \rho_b, A_b$ and $\vec{x}_b$ are mass, density, value of the field variable and position vector of neighbor particles. The gradient of $A$ is determined by

\[ \nabla A(\vec{x}) = \sum_b A_b \frac{m_b}{\rho_b} \nabla W(\vec{x} - \vec{x}_b, h). \quad (28) \]

Therefore, in SPH method no mesh is needed for calculation of partial derivatives (meshless method).
3.2 Physics of SPH-Particles

Required derivatives $\nabla \cdot \vec{v}$ and $\nabla p$ for mass conservation (continuity equation)

$$\frac{\partial \rho}{\partial t} + \rho \nabla \cdot \vec{v} = 0 \quad (29)$$

and momentum conservation

$$\frac{\partial \vec{v}}{\partial t} + \frac{1}{\rho} \nabla p = 0 \quad (30)$$

results with $\vec{v}_{ba} = \vec{v}_b - \vec{v}_a$ and $W_{ab} = W(\vec{x}_a - \vec{x}_b)$ from summation interpolation (see eq.(28)). It applies for all particles $a$

$$(\nabla \cdot \vec{v})_a = \sum_b \frac{m_b}{\rho_b} \vec{v}_{ba} \nabla W_{ab} \quad (31)$$

and

$$(\nabla p)_a = \sum_b \frac{m_b}{\rho_b} \vec{p}_b \nabla W_{ab}, \quad (32)$$

in which density is given by

$$\rho(\vec{x}) = \sum_b m_b \nabla W(\vec{x} - \vec{x}_b, h). \quad (33)$$

Consequently, discrete form of continuity equation is given by

$$\frac{\partial \rho_a}{\partial t} = \rho_a \sum_b \frac{m_b}{\rho_b} \vec{v}_{ab} \nabla W_{ab} \quad (34)$$

and discrete form of momentum conservation by

$$\frac{\partial \vec{v}_a}{\partial t} = -\frac{1}{\rho_a} \sum_b \frac{m_b}{\rho_b} \vec{p}_b \nabla W_{ab}. \quad (35)$$

The particle velocity $\vec{v}$ arises from derivative

$$\vec{v}_a = \frac{\partial \vec{x}_a}{\partial t} \quad (36)$$
3.3 Examples

SPH comes often into operation in computer animations with water, because with SPH very realistic results can be obtained.

![Figure 12: Water simulation with SPH (source: Bell et al. (2005))](image)

Another example shows the so-called “Millenium Simulation”, that simulates the formation of big scale mass distributions in space (galaxies and galaxy clusters). For this calculation more than 10 billion particles were used (see [http://www.mpa-garching.mpg.de/galform/millennium/](http://www.mpa-garching.mpg.de/galform/millennium/)).

![Figure 13: “Millenium Simulation” with $> 10^{10}$ particles (source: Springel et al. (2005))](image)
4 Lattice Boltzmann

The Lattice Boltzmann Method (LBM) is based on the theory of Ludwig Boltzmann (1844-1906). Boltzmann treats a gas as medium, that consists of interacting particles (molecules or atoms), that can be described with classical mechanics and a statistical treatment. This basic idea was adopted into LBM, where gases are simulated by streaming and collisions of particles (Sukop and Thorne (2006)). The approach was later extended on simulation of fluids in general (gases and liquids).

For LBM a lattice consisting of nodes is needed. In figure 14 the lattice array for a D2Q9 model (2 dimensions, 9 nodes) is illustrated. The vectors $\vec{e}_i$ denotes particle velocities in direction $i$. In the simplest approach particles have a uniform mass ($m_u$ - mass unit) and the lattice has uniform lattice spacing ($l_u$ - lattice unit).

4.1 Boltzmann equation

A system consisting of $N$ particles (molecules) can be described by a density function $F(\vec{x}, \vec{e}, t)$. When positions $\vec{x}$ and velocities $\vec{e}$ of all particles (molecules) are known at time $t$, so it is (at least hypothetical) possible to predict the mechanical behavior of the system. Assuming that an external force $f$ is acting on the particles, then they have positions $\vec{x} + \vec{e}dt$ and velocities $\vec{e} + f dt$ at time $t + dt$. When no collisions occurred, then it applies

$$F(\vec{x}, \vec{e}, t)d\vec{x}d\vec{e} = F(\vec{x} + \vec{e}dt, \vec{e} + f dt, t + dt)d\vec{x}d\vec{e}.$$  \hspace{1cm} (37)

If there are collisions within the time step, eq. (37) is added by a collision term $\Omega$, which describes changing rate between end and start condition of the system (Mohamad (2007)).

$$F(\vec{x}, \vec{e}, t)d\vec{x}d\vec{e} = F(\vec{x} + \vec{e}dt, \vec{e} + f dt, t + dt)d\vec{x}d\vec{e} + \Omega(F)d\vec{x}d\vec{e}dt$$  \hspace{1cm} (38)
The time derivative of eq. (38) gives
\[ \frac{dF}{dt} = \Omega(F), \] (39)
i.e. total changing rate of density function is equal to collision rate. When \( F \) is
a function of \( \vec{x}, \vec{e} \) and \( t \), then it applies
\[ dF = \frac{\partial F}{\partial \vec{x}} d\vec{x} + \frac{\partial F}{\partial \vec{e}} d\vec{e} + \frac{\partial F}{\partial t} dt \] (40)
Dividing eq. (40) with \( dt \), one gets
\[ \frac{dF}{dt} = \frac{\partial F}{\partial \vec{x}} \frac{d\vec{x}}{dt} + \frac{\partial F}{\partial \vec{e}} \frac{d\vec{e}}{dt} + \frac{\partial F}{\partial t}. \] (41)
With velocity \( \vec{e} = d\vec{x}/dt \), acceleration \( \vec{a} = d\vec{e}/dt \), second law of Newton \( \vec{a} = f/m \) (mass \( m \)) and eq. (39) it offers Boltzmanns law of motion (Mohamad (2007)).
\[ \frac{\partial F}{\partial t} + \frac{\partial F}{\partial \vec{x}} \vec{e} + \frac{f}{m} \frac{\partial F}{\partial \vec{e}} = \Omega \] (42)

4.2 Time relaxation according to Bhatnagar, Gross and Krook

The exact calculation of collision term \( \Omega \) in eq. (42) is very complicated because
of its complexity. Therefore \( \Omega \) is approximated by a simple operator, that does
not bring significant errors to the solution. Bhatnagar, Gross and Krook (BGK)
presented in 1954 a simplified model for the collision operator (Mohamad (2007)).
Thereby local equilibrium distribution function \( F^{eq} \) and a relaxation factor \( \tau \) were
introduced.
\[ \Omega = \frac{1}{\tau} (F^{eq} - F) \] (43)
With this approximation (see eq.(38)) and discretization (index \( i \)) one gets the
linearized BGK relaxation form of LB equation (Cook and Noble (2004)).
\[ F_i(\vec{x} + \vec{e}_i dt, t + dt) = F_i(\vec{x}, t) - \frac{dt}{\tau} (F_i(\vec{x}, t) - F^{eq}_i(\vec{x}, t)) \] (44)
Equation (44) consists of a streaming and a collision part. At every node there
are eight density distributions \( F_i \) and a residual distribution \( F_0 \). Equilibrium
conditions are, according to Sukop and Thorne (2006), given by
\[ F^{eq}_i(\vec{x}) = a_i \rho(\vec{x}) \left( 1 + 3 \frac{\vec{e}_i \cdot \vec{v}_f}{c^2} + \frac{9}{2} \frac{(\vec{e}_i \cdot \vec{v}_f)^2}{c^4} - \frac{3}{2} \frac{\vec{v}_f^2}{c^2} \right). \] (45)
Weightings \( a_i \) are 4/9 for the residual particles with \( i = 0 \), 1/9 for \( i \in \{1,2,3,4\} \)
and 1/36 for \( i \in \{5,6,7,8\} \). \( c \) is basic speed on nodes and macroscopic density \( \rho \)
is defined as sum over all directional densities \( (\rho = \sum_i F_i) \). Macroscopic velocity
\( \vec{v}_f \) is the average of microscopic velocities \( \vec{e}_i \) weighted with densities \( F_i \).
\[ \vec{v}_f = \frac{1}{\rho} \sum_{i=0}^{8} F_i \vec{e}_i \] (46)
4.3 Examples

A LBM application of Schenkengel and Vrettos (2011) simulates an induced soil liquefaction. The 2D model consists of $300 \times 60$ nodes and represents a slope with a slope angle of $18^\circ$. An explosion within the slope leads to stability failure, whereby the material liquefies. Following figure shows velocities of the material at times $t = 0, 01s$, $t = 0, 29s$ and $t = 1, 17s$ during liquefaction event.

![Image of 2D-LBM model of an explosion within a slope, velocities in the lattice illustrated by vectors and filled colored contours (source: Schenkengel and Vrettos (2011))](image)

**Figure 15:** 2D-LBM model of an explosion within a slope, velocities in the lattice illustrated by vectors and filled colored contours (source: Schenkengel and Vrettos (2011))

With the model of Schenkengel and Vrettos (2011) at first time rheological transition from rigid to liquid was solved and implemented in a model with a LBM approach.
Another example shows a phase separation of two fluids with different density.

Figure 16: Phase separation of two fluids (source: http://www.bgce.de/curriculum/projects/patilgmeiner/)
5 Molecular Dynamics

With Molecular Dynamics (MD) interactions between atomic particles (atoms, molecules, nanoparticles, etc.) can be simulated. The laws of classical mechanics find no more validity. For description of particle mechanics, laws of quantum mechanics must be used. Anyhow DEM and MD are very similar.

The first application of MD is dated to the year 1957 with a paper by Alder and Wainwright. Hence MD is the oldest applied particle method. Compared to DEM, where Newton’s equations serves as basic principle of description of particle motion, MD is based on Schrödinger equation. It is a very complex equation and can be solved analytically only in rarest cases. Even numerical approaches limit applications of the Schrödinger equation to very simple systems and few particles. So approximation procedures are used to simplify solving of the equation (Griebel et al. (2004)).

5.1 Schrödinger equation

In quantum-mechanical systems conclusions about the state of the system can be derived by a state function \( \Psi \) (also called wave function). A system consisting of \( N \) cores and \( K \) electrons with variables \( R_i \) resp. \( r_i \) is characterized by its state function as follows (Griebel et al. (2004)).

\[
\Psi = \Psi(R_1, ..., R_N, r_1, ..., r_K, t)
\]  

(47)

Variable \( t \) indicates time dependency of the state function. \( \Psi \) is due to solution of the Schrödinger equation (with \( R = R_1, ..., R_N \) and \( r = r_1, ..., r_K \)).

\[
\frac{\text{i}\hbar}{\partial t} \frac{\partial \Psi(R, r, t)}{\partial t} = \mathcal{H} \Psi(R, r, t)
\]  

(48)

Here, \( \text{i} \) is imaginary unit, \( \mathcal{H} \) is the Hamilton operator and \( \hbar = h/2\pi \) with \( h \), the Planck’s constant. The Hamilton operator describes temporal evolution of possible energy values in the system based on its potentials.

5.2 Potentials

Interactions between two particles, that depend only on particle distance, are described by pair-potentials. Such potentials are e.g. the gravity potential, the Coulomb potential (electrical point charge), the van-der-Waals potential (weak attraction at inert gases) and the Lennard-Jones potential (uncharged, unbound atoms).

The Lennard-Jones potential

\[
U(r_{ij}) = \alpha \varepsilon \left[ \left( \frac{\sigma}{r_{ij}} \right)^{n} - \left( \frac{\sigma}{r_{ij}} \right)^{m} \right], \quad m < n
\]  

(49)

with \( \alpha = \frac{1}{n-m} \left( \frac{n}{m} \right)^{\frac{1}{m-n}} \) is parameterized by \( \sigma \) and \( \varepsilon \). Here \( \varepsilon \) defines the magnitude of repulsion resp. attraction forces. Thus, materials of different stiffnesses can be
$rac{1}{2}(n-1)$.

\[\sigma \quad \varepsilon \]

Figure 17: Lennard-Jones potential with $\varepsilon = 1$ and $\sigma = 1$

simulated. $\sigma$ specifies zero-crossing of the potential. Figure 17 shows a Lennard-Jones potential for $n = 12$ and $m = 6$.

If a particle moves within a potential, the corresponding potential energy is due to

$$E_{\text{pot}}(R) = \sum_{i=1}^{N} \sum_{j=i+1}^{N} U_{ij}(r_{ij}),$$

(50)

where $r_{ij} = ||R_j - R_i||$ is the distance between particles. The potential function for the Lennard-Jones potential with $n = 12$ and $m = 6$ is

$$E_{\text{pot}}(R) = 4 \cdot \varepsilon \sum_{i=1}^{N} \sum_{j=1,j>i}^{N} \left[ \left( \frac{\sigma}{r_{ij}} \right)^{12} - \left( \frac{\sigma}{r_{ij}} \right)^{6} \right].$$

(51)

The corresponding force $\vec{F}_i$, which acts on particle $i$, results by creation of a gradient with respect to $R_i$.

$$\vec{F}_i = -\nabla_{R_i} E_{\text{pot}}(R)$$

(52)

For the Lennard-Jones potential this force is given by the equation

$$\vec{F}_i = 24 \cdot \varepsilon \sum_{j=1,j\neq i}^{N} \frac{1}{r_{ij}^2} \cdot \left( \frac{\sigma}{r_{ij}} \right)^6 \cdot \left( 1 - 2 \cdot \left( \frac{\sigma}{r_{ij}} \right)^6 \right) \vec{r}_{ij},$$

(53)

where $\vec{r}_{ij}$ is the directional vector between particles $i$ and $j$ (Griebel et al. (2004)).

5.3 Physics of MD-Particles

Physics of MD-Particles is in general the same as for DEM-Particles (see section 2.3). With Newton’s second law accelerations, velocities and displacements are determined by time integration. In doing so one gets new positions of the particles. Unlike in DEM simulations MD-Particles have a cutting radius, which results from potential range. As simplification no forces act on a particle, as long no further particles stay in its potential range.
5.4 Examples

In figure 18 a collision of two bodies is displayed. Particle velocities are coded with colors (red - high velocity, blue - low velocity).

![Figure 18: Collision of two bodies, temporal evolution of particle distribution](source: http://wissrech.ins.uni-bonn.de)

Lipid molecules typically form double layer membranes in the presence of water, because one end is hydrophilic and the other end is hydrophobic (e.g. oil film). These membranes spontaneously form bubbles or vesicles. Following figure shows a simulation of a fusion of such a vesicle with a lipid membrane.

![Figure 19: Fusion of one vesicle with a membrane of 2018 Diblock-Copolymers, vesicle diameter 40 nm](source: http://www.mpg.de)
References


