# WAVE PROPAGATION FROM CRACK EXTENSION BY MD AND FEM SIMULATIONS

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#### **COMPUTATIONAL MECHANICS 2012**

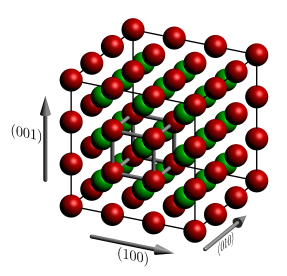
November 12 – 14, 2012 Špičák, Czech Republic

#### Introduction

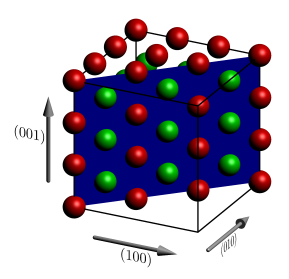
- ▶ We present results for molecular dynamic (MD) and finite element (FEM) simulations in 3D bcc iron crystals, with embedded central through crack (001)[110] of Griffith type, loaded in mode I.
- ► The sample geometry and border conditions in MD were chosen in such a way as to invoke a cleavage crack extension.
- Acoustic emission (AE) sources caused by the crack were analysed on both the atomistic and continuum level with FEM.

### Problem description

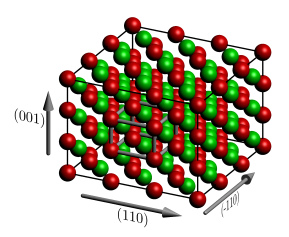
# Material (bcc iron)



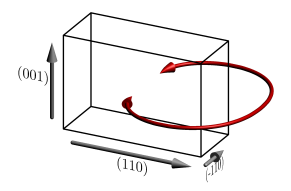
#### Orientation



#### Orientation



# Size and boundary conditions

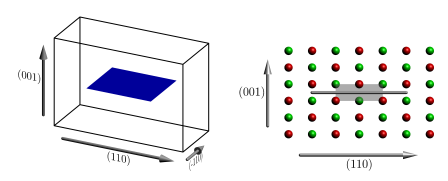


Plane strain

$$300\times150\times15$$
 atoms

$$299 a_0 \sqrt{2} \times 149.5 a_0 \sqrt{2} \times 14.5 a_0 \sqrt{2}$$

#### Crack description

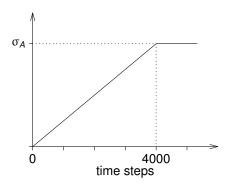


Size  $178a_0\sqrt{2} \times 14.5a_0\sqrt{2}$ 

Method bond restriction,

i.e. atomic interactions over crack plane are not allowed.

# Type of loading



Temperature: 0 K

Tension MODE: I

# Many-body interatomic potential

G.J.Ackland, D.J.Bacon, A.F.Calder, T.Harry:

Computer simulation of point defect properties in dilute Fe-Cu alloy using a many-body interatomic potential.

Philosophical Magazine A, 1997, Vol. 75, No. 3, 713–732

The energy of an assembly of *N* atoms is given by

$$E = \frac{1}{2} \sum_{i \neq j=1}^{N} V(r_{ij}) - \sum_{i=1}^{N} \left( \sum_{j \neq i=1}^{N} \phi(r_{ij}) \right)^{1/2}$$

 $V(r_{ij})$  - pair repulsive potential

 $\phi\left(r_{ij}\right)$  - many-body cohesive potential

### Integration of equations of motion

Newtonian equations of motion are solved by the central difference method.

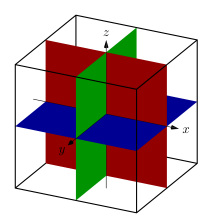
Time integration step:

$$1 \times 10^{-14} \text{ s}$$

# Simulation steps

- 1. Generation of the crystal containing the crack.
- 2. Fixation,

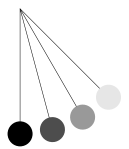
i.e. atoms in plane x = 0, y = 0, z = 0 can move only in a given plane.



Surface relaxation,

 i.e. set system to equilibrium state
 (minimum potential energy and kinetic energy nearly zero).

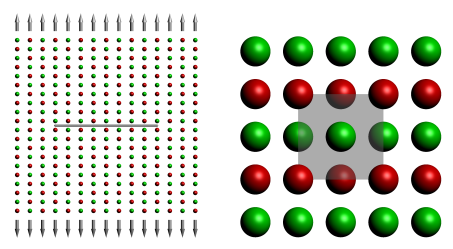
Pendulum method:



J.B.Gibson, A.N.Goland, M.Milgram, G.H.Vineyard: Phys. Rev., **120**, p.1229, 1960

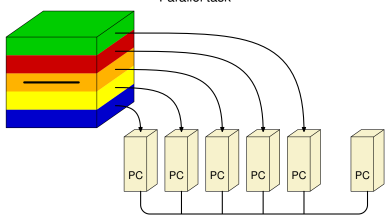
4. Remove fixation.

#### Loading symmetric loading, distributed in 2 surface layers



#### Simulation technique

# Simulation code has been written in *Fortran 90*. Parallel task



# Simulation code has been developed under system MPI (Message Passing Interface).

#### **Used MPI-functions:**

- MPI\_INIT, MPI\_FINALIZE,
- ▶ MPI\_COMM\_RANK, MPI\_COMM\_SIZE,
- MPI\_SEND, MPI\_RECV, MPI\_BCAST,
- MPI\_ISSEND, MPI\_IRECV,
- ► MPI\_WAIT.

#### Memory requirement:

$$6\times8 + 2\times8 = 64$$
 bytes/atom  $\rightarrow$  165 MiB

#### Disk requirement:

$$6\times8 = 48$$
 bytes/atom  $\rightarrow$  124 MiB

Hash: cell index method (link cell method)

M.P.Allen, D.J.Tildesley: Computer Simulation of Liquids. Oxford University Press, New York, 1987

D. Frenkel, B. Smit: Understanding Molecular Simulations. Academic Press, New York, 1996

#### Where was it computed?



	49 nodes	
0 0		_,

2x 6-cores Xeon E5645 2.4 GHz

MINOS

24 GiB

2x 600 GiB (15 k rpm, SAS)

1 Gb Ethernet, Infiniband

#### TARKIL

28 nodes

2x Quad Core Intel Xeon X5570

2.93 GHz 24 GiB

2x 300 GiB (15 k rpm, SAS)

1 Gb Ethernet, Infiniband 4x QDR

#### MANDOS

14 nodes

4x AMD Opteron 6274

2.5 GHz

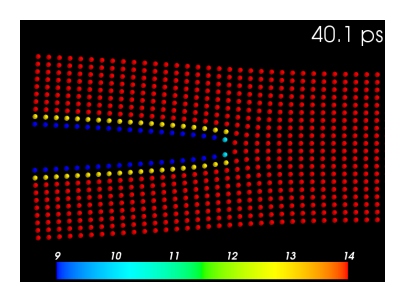
256 GiB

870 GiB, 27 TiB

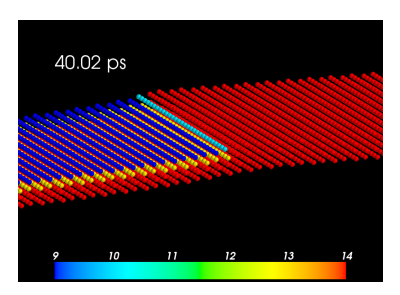
1 Gb Ethernet, Infiniband

#### Results - MD

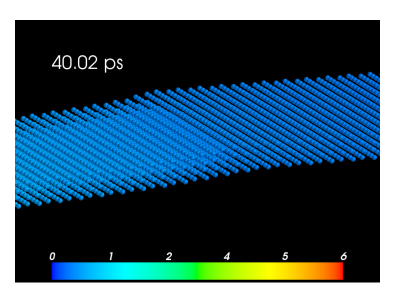
#### Coordination number



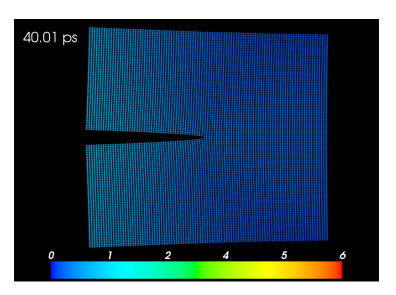
#### Coordination number



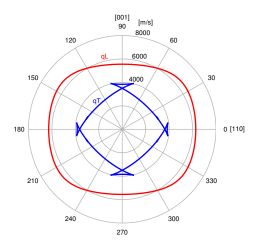
# Kinetic energy



# Kinetic energy



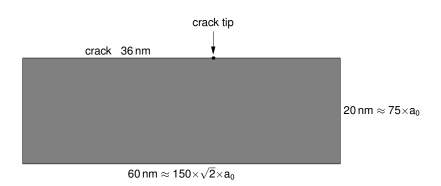
# Section of the ray (wave) surfaces in the (110) plane



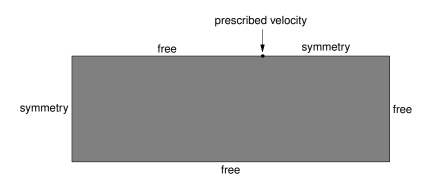
### FEM analysis

- COMSOL Multiphysics
- Structural Mechanics Module
- Plane strain
- Shape function: Lagrange (Quadratic)
- Time analysis
- Time: 0 5 ps, step 0.01 ps
- Time stepping method: Generalized alpha
- Relative tolerance: 10<sup>-7</sup>
- Absolute tolerance: 10<sup>-14</sup>
- Mesh: Quadrilateral elements (300 $\times$ 100), Size: 2 $\times$ 2 Å
- Number of degrees of freedom: ≈250000

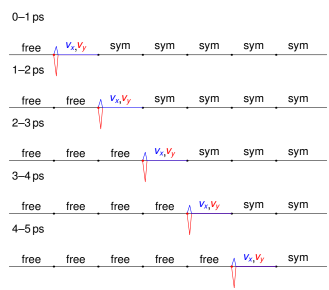
### Sample



### Boundary condition



#### Detail of crack tip



#### Results - FEM

### Modulus of velocity



#### Conclusion

- At higher applied loads, positive T-stress contributes to cleavage crack extension in MD. Under the ramp loading during 4000 time steps, the crack was initiated at the critical Griffith stress intensity.
- ► MD simulations show that cleavage crack initiation in the 3D bcc iron crystal forms an AE-source, where qL-waves dominate. However, qT-waves are also generated during a continuous bond breakage in the crystal, which is new knowledge from 3D modelling. The strongest pulse emission comes from stress relaxation at the crack front, after the crack initiation.
- ▶ Simplified modelling of the pulse emission by FEM shows that, besides the qL and qT-waves, Rayleigh waves can also be generated at the (001) free crack faces, in agreement with expectations according to continuum analysis.

#### Question time

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