## UMENO LAB. [Modeling and Simulation of Nano-Micro Mechanophysics]

Center for Research on Innovative Simulation Software

Nano-Micro Mechanophysics

Department of Mechanical Engineering

http://www.cmsm.iis.u-tokyo.ac.jp

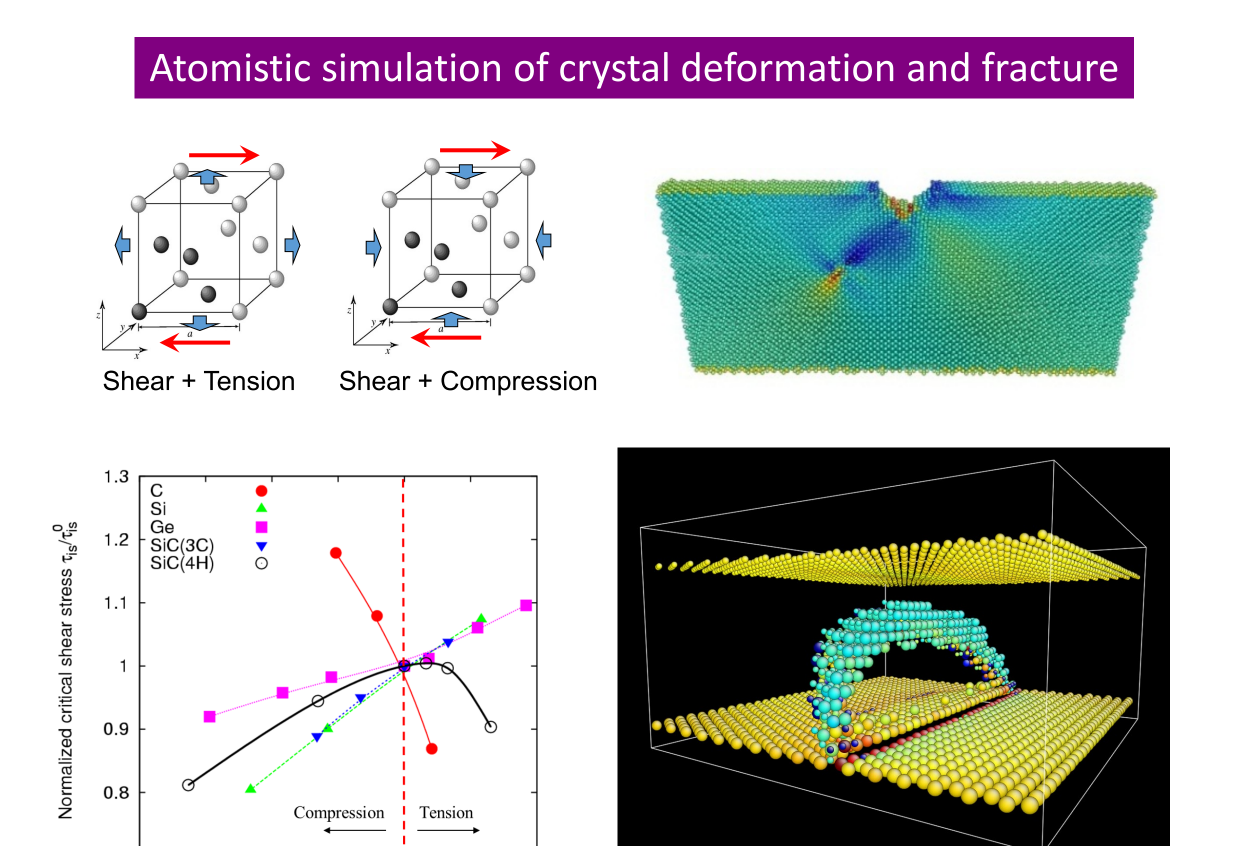
# **Challenges to Nano-Micro Mechanophysics and**

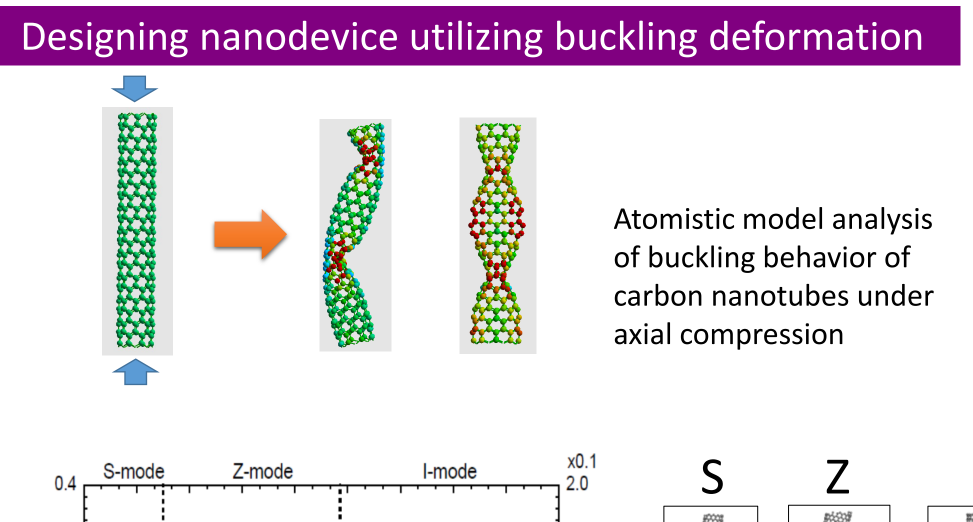


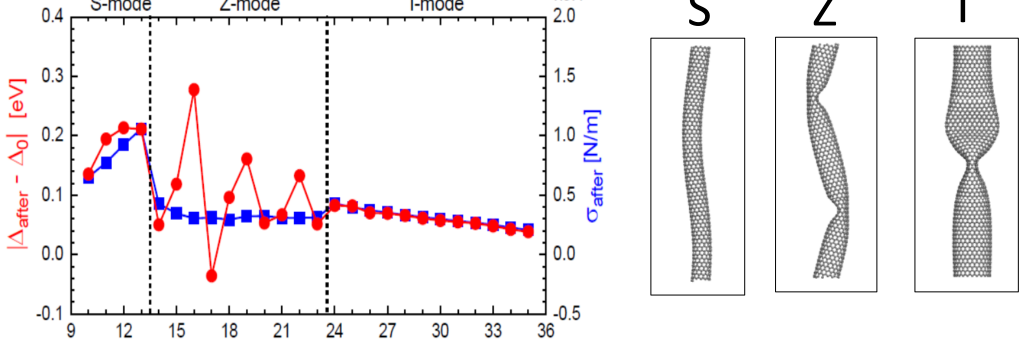
Cw-302

### **Multiscale Simulation**

We aim to reveal the mechanical and physical properties of nanomaterials by ab initio density functional theory calculations and molecular dynamics simulations. We also work on multiscale simulation based on knowledge of nano-microscale phenomena.





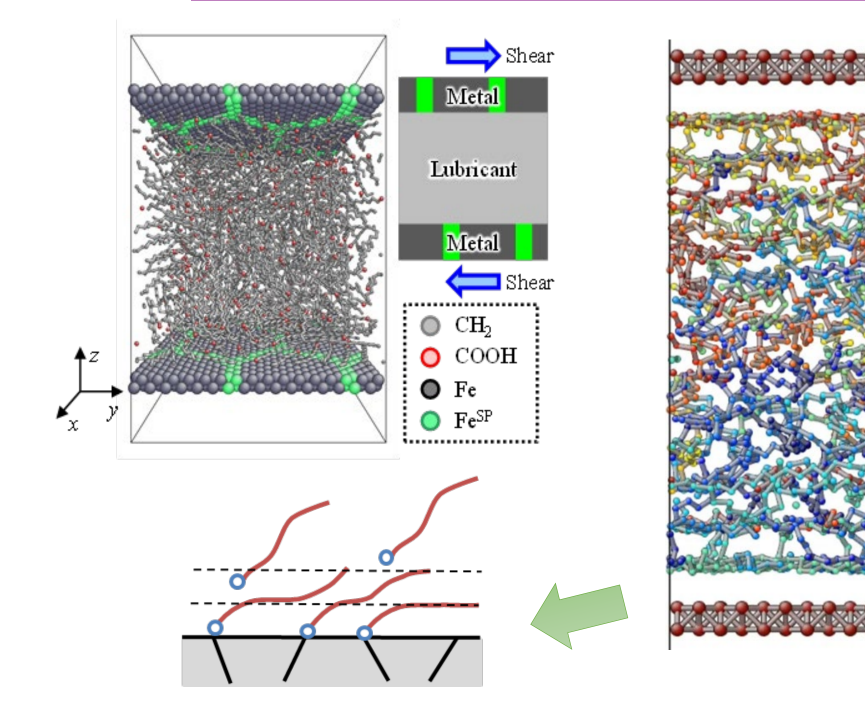


#### Normalized hydrostatic stress $\sigma_{\rm h}/\tau_{\rm is}^0$

Ideal shear stress (ISS) under compression/tension in covalent crystals

Molecular dynamics simulations of dislocation motion and crystal slip

#### Molecular simulation of boundary lubrication

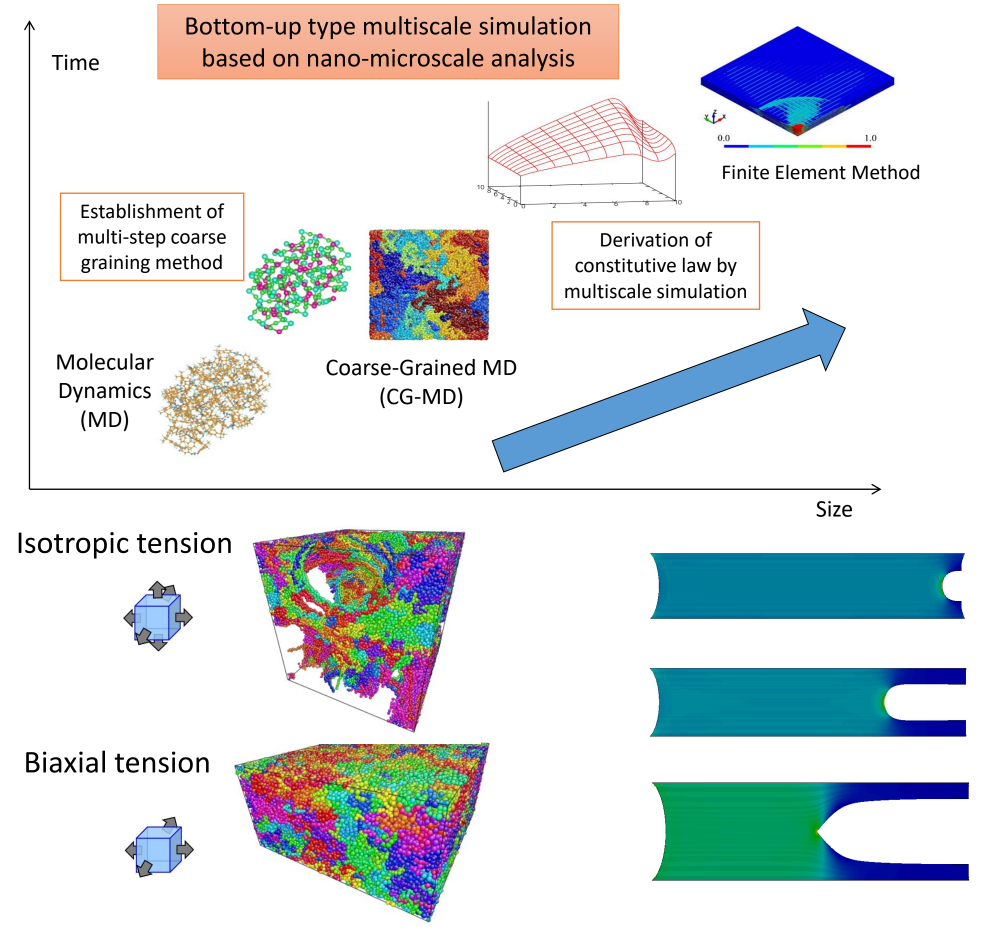


Coarse-grained molecular dynamics simulation of formation of lubricant molecule structure (polymer brush) on metal surface with nanograins

Nanotube diameter vs. buckling mode, buckling-induced band gap change, and post-buckling stress

Chiral index n

#### Multiscale simulation of polymer materials



FEM analysis of crack propagation

