## 22.53 Overview

- 1. To understand the motivation and the basic purpose of this course, one might ask:
  - Why "statistical processes"?
  - Why "atomistic simulations"?
- 2. The aim is to **analyze** and **understand** complex phenomena, e.g., polymorphism, melting, amorphization or glass transition, plastic deformation and fracture a host of microstructural processes in the case of materials applications. Other examples could be in fluid dynamics, biomolecular transport, etc.
- 3. Physical systems of interest tend to have many degrees of freedom (DOF), strongly coupled, and are usually **nonlinear**, **nonhomogeneous**, and **nonequilibrium**. They are difficult to study both theoretically and experimentally.
- 4. Partial answer to 1 above:
  - Systems which undergo strong fluctuations are best treated in statistical terms
  - Discrete-particle simulation provides the only tractable method of analysis.
- 5. An area of application of particular interest in this course is materials **modeling**, which is defined to be **theory and simulation**, combined with use of selected measurements for validation and parameterization.

Why the special focus?

- need for advance in fundamental understanding
- exploit high performance computing (HPC)
- emerging role of *computational* materials research, and increasing relevance of modeling across several length and time scales in a number of grand challenge-type initiatives launched by funding agencies

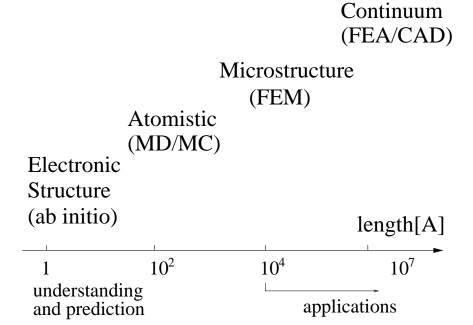


Figure 1: Four lengthscales in Multiscale Materials Modeling.

- 6. Multiscale Materials Modeling (MMM) hierarchy of simulation techniques Fig. 1 shows the four lengthscales in MMM.
  - Electronic Structure
    - Quantum Chemistry
      - \* Hartree-Fock (HF) and Configuration Interaction (CI)
      - \* Coupled-Cluster and Many-Body Perturbation Theory
    - Density Functional Theory (DFT)
      - \* Local Density Approximation (LDA)
      - \* Generalized Gradient Approximation (GGA)
    - Local Combination of Atomic Orbitals (LCAO), also called Tight-Binding
       (TB): orthogonal vs non-orthogonal, selfconsistent vs non-selfconsistent.
  - Atomistic
    - Lattice calculations: cohesive energy, phonon dispersion curve, phonon Boltzmann equation, etc.
    - Static Minimization

- \* Conjugate Gradient (CG)
- \* Simulated Annealing
- Molecular Dynamics (MD)
- Monte Carlo (MC)
- Quantum mechanical extensions such as Path Integral Monte Carlo (PIMC)
- Microstructural or Mesoscale
  - Kinetic Monte Carlo (KMC), a.k.a. Nodal Dynamics in dislocation context
  - Finite Element (FEM)
- Continuum
  - Finite Element Analysis / Computer Aided Design (FEA/CAD)
  - Computational Fluid Dynamics (CFD)

The final step is Product and Process Design, and of course, Marketing.

Note: applications lie mostly on the micron scale and above, whereas understanding and predictions normally occur at the molecular level and below. Effective linking of adjacent techniques is a major challenge.

A central theme across several lengthscales is **microstructure**, which can be as small as a point defect and as large as a visible crack on an airplane (see Fig. 2). Different microstructures interact in complicated manner depending on temperature, stress and chemical conditions. More often than not, *microstructures control the response of the material*.

7. In this course we will discuss primarily MD and MC. A simple definition is:

each is a method of sampling the configuration of an N-particle system using specific interatomic interaction models

What are the frontiers? Materials modeling ...

Current developments: Workshop series ITP-UCSB '96 ... ASCI - Accelerated Strategic Computing Initiative ... SSI - Strategic Simulation Initiative ...

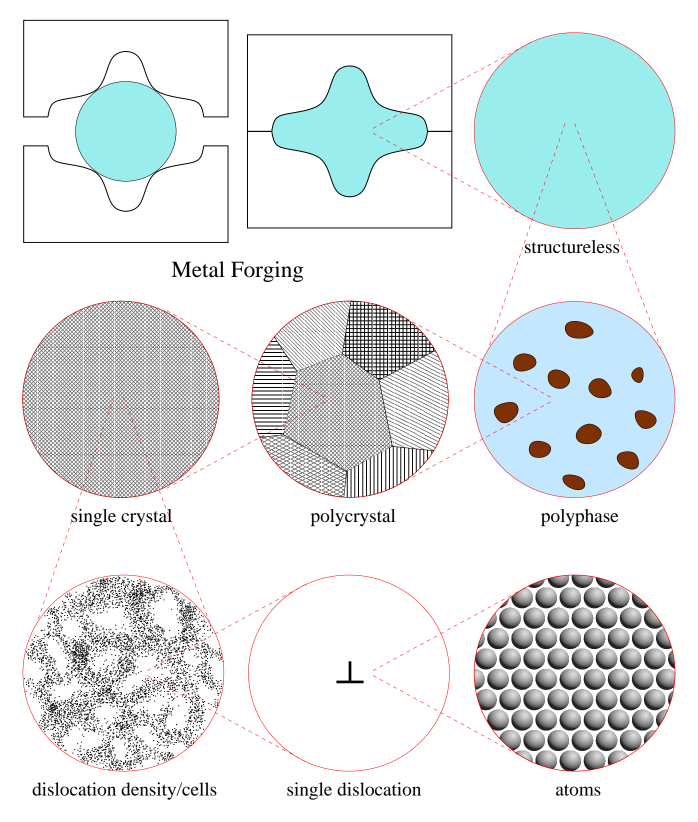


Figure 2: Microstructure across several lengthscales.

|                               | MC                                    | MD   |
|-------------------------------|---------------------------------------|--|
| driving factor for            | potential energy                      | interatomic forces   |
| particle configuration change |                                       |  |
| type of dynamics              | stochastic                            | Hamiltonian  |
| type of sampling              | probabilistic                         | deterministic  |
| space sampled                 | configuration space                   | phase space  |
|                               | $\mid \{\mathbf{r}_1,,\mathbf{r}_N\}$ | $\mid \{(\mathbf{r}_1,\dot{\mathbf{r}}_1),,(\mathbf{r}_N,\dot{\mathbf{r}}_N\}\mid$ |

Table 1: Comparison of Molecular Dynamics (MD) and Monte Carlo (MC) methods.

Codes: - Haile [1], Li Ju, term project

## References

[1] J.M. Haile, Molecular Dynamics Simulation: Elementary Methods (Wiley, New York, 1997).