

機器學習於材料資訊的應用

Machine Learning on Material Informatics

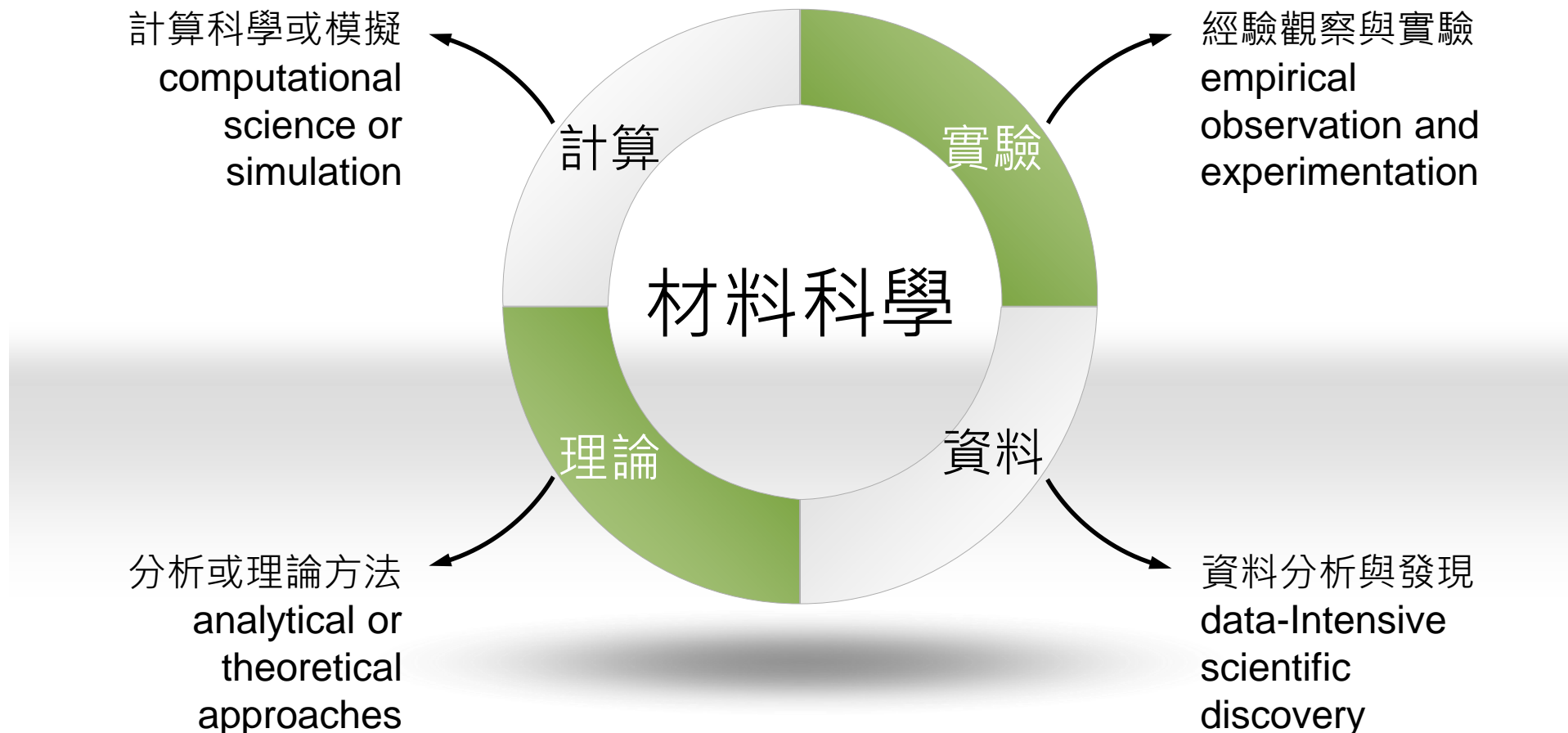
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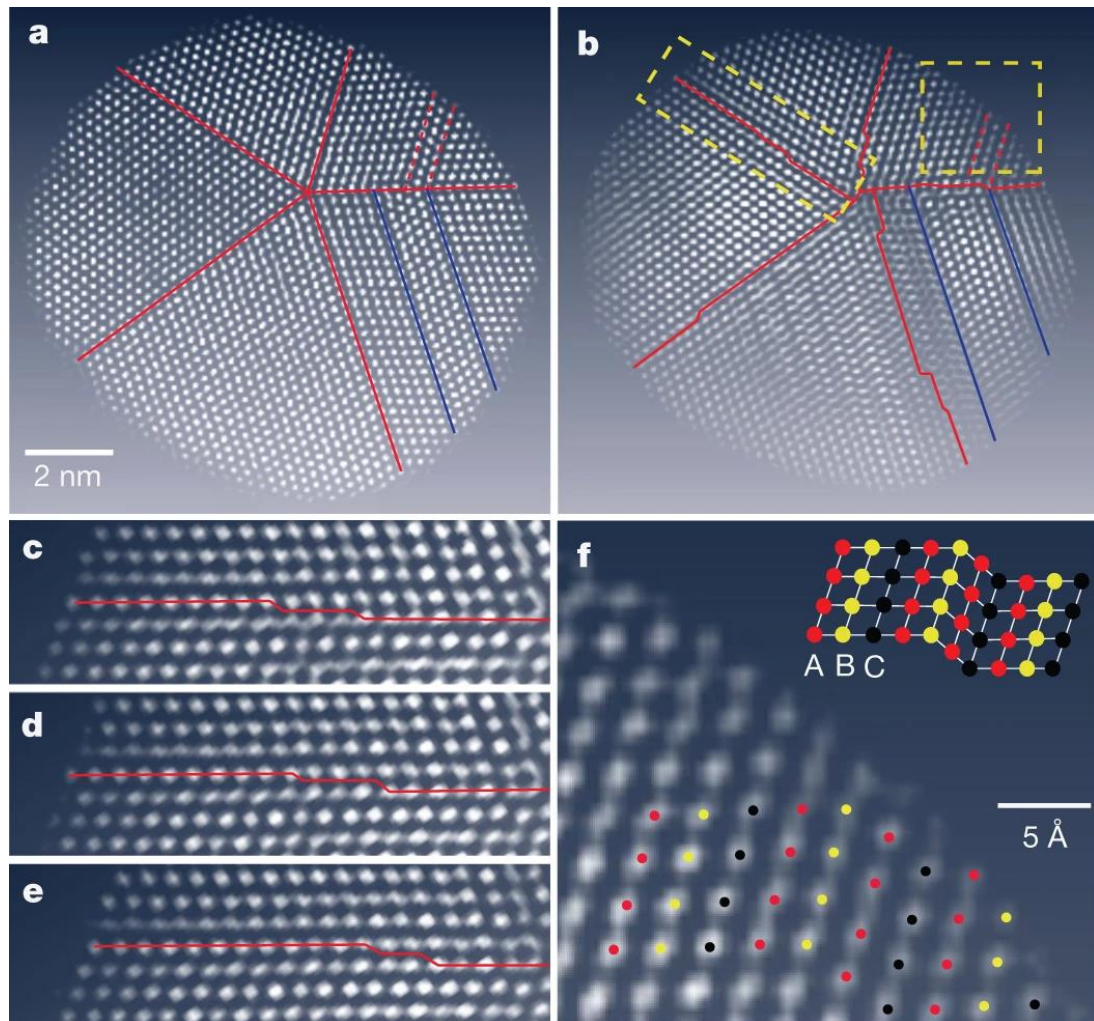
楊安正(AN-CHENG YANG)

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材料科學的分支

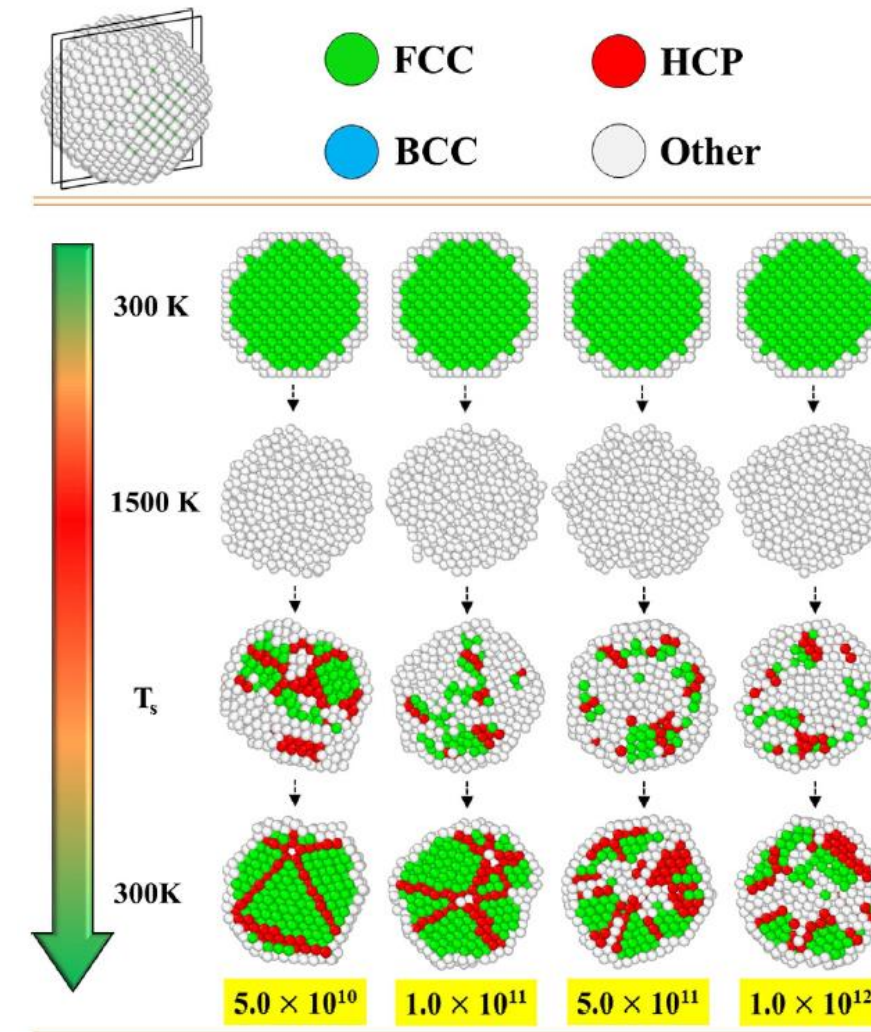


實驗與測量



Chen, C., Zhu, C., White, E. et al. Three-dimensional imaging of dislocations in a nanoparticle at atomic resolution. *Nature* 496, 74–77 (2013).

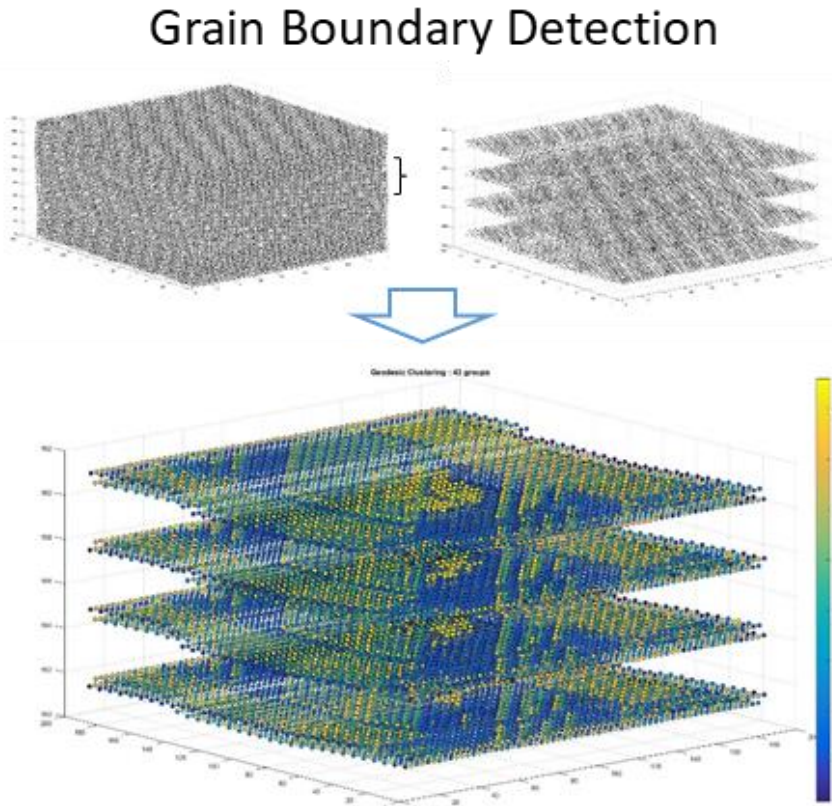
計算或模擬



Vo, T.Q., Kim, B. A molecular dynamics study on cooling rate effect on atomic structure of solidified silver nanoparticles. *Eur. Phys. J. D* 73, 183 (2019).

Case1-晶粒邊界辨識

晶粒邊界分群機器

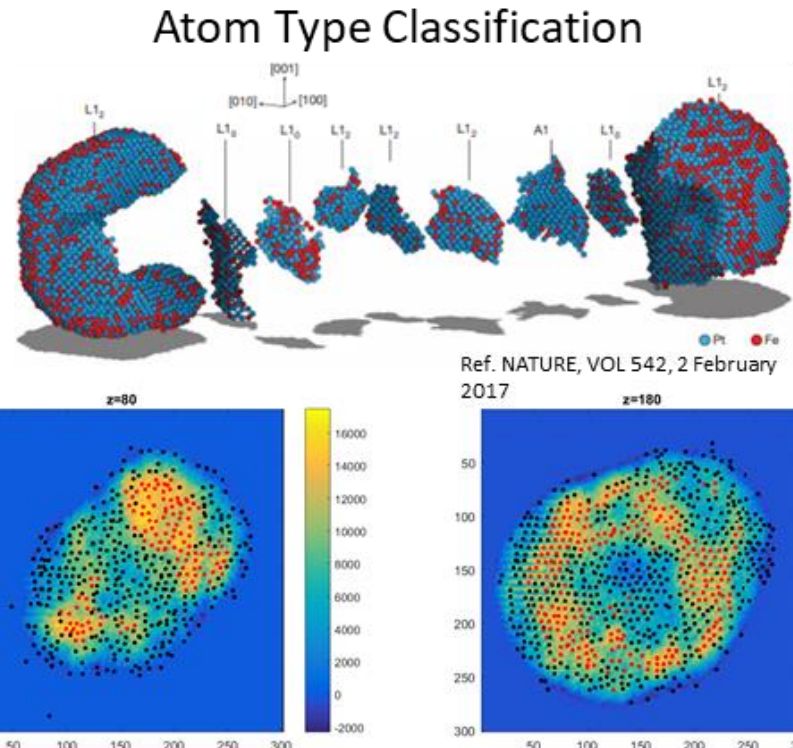


合作計畫	無
合作團隊	交大材料鄒年棣實驗室

案例	透過非監督式學習，完成晶粒邊界辨識、結構分群
客戶目標	晶粒邊界辨識，達成結構分群，以進行區塊結構相似度分析，並與人為經驗公式相互驗證
問題困難描述	人工觀察大量模擬結果資料耗工費時，傳統分群方法需要先前知識才能協助分群。
訓練資料來源	分子動力學模擬結果
機器學習引擎	Modularity
結果	針對晶粒邊界辨識與其結構達成自動分群，與人工經驗法則結果一致。
Status	會議論文已發表 論文撰寫中

Case2-奈米粒子影像重建

種類與缺陷型態深度辨識機器

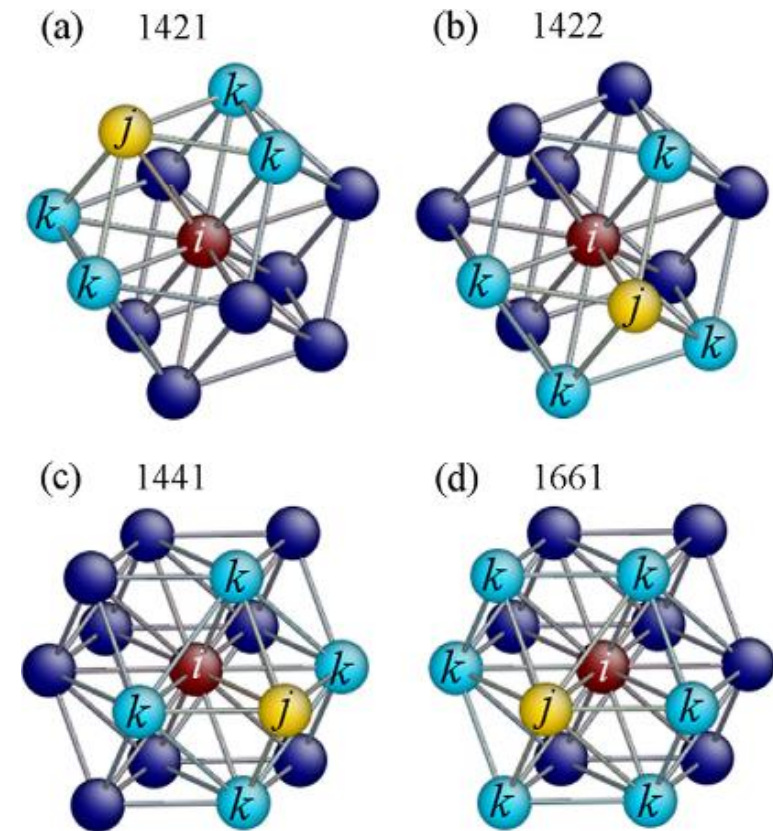
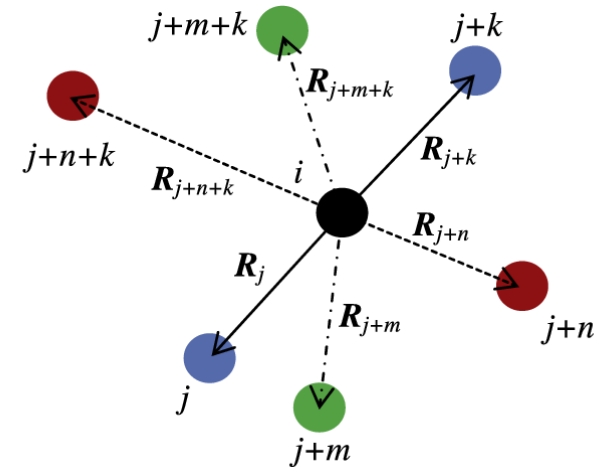
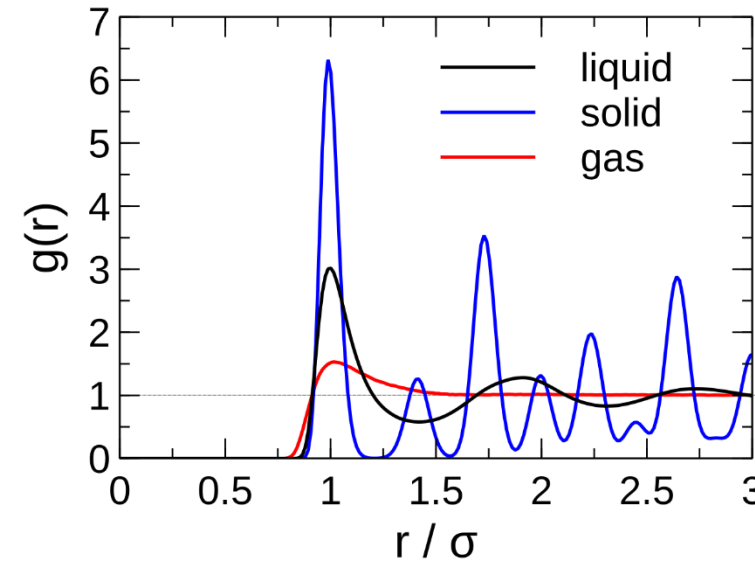


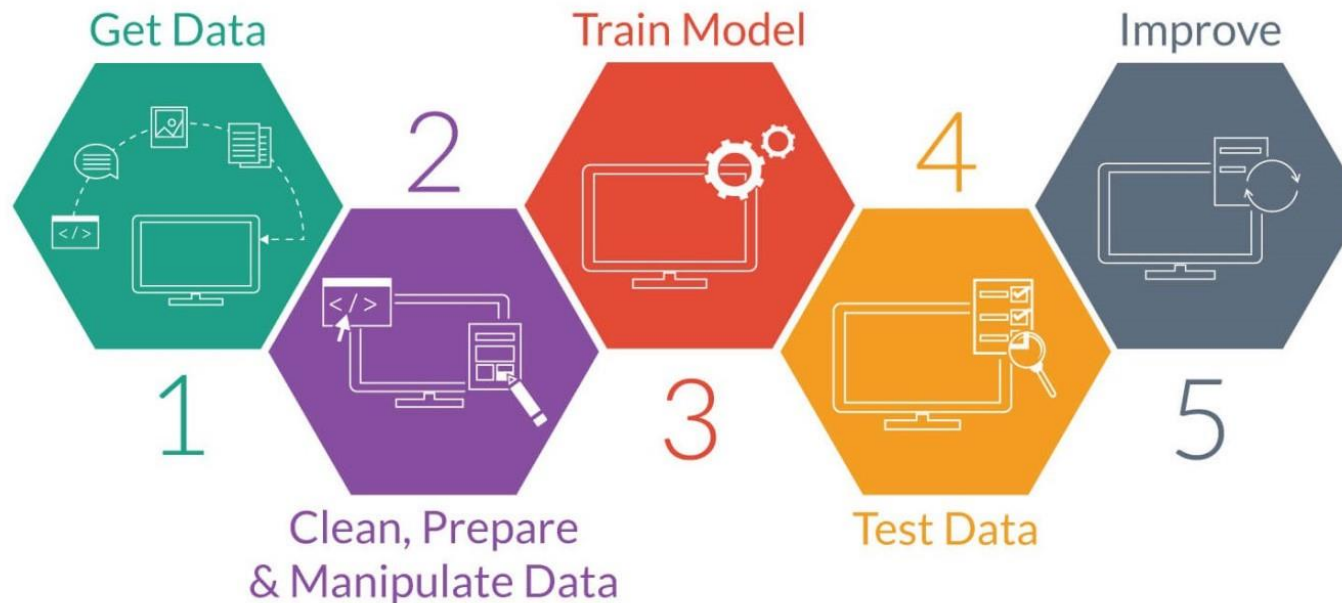
合作計畫	國研院創新計畫
合作團隊	清大工科陳健群實驗室

案例	三維斷層造影之原子種類與缺陷型態辨識
客戶目標	透過深度學習技術，定位原子座標、辨識原子種類、分析缺陷型態
問題困難描述	需要人工判斷原子種類與缺陷邊界，會因人為偏見造成不一致性的誤判。
訓練資料來源	原子級三維斷層造影顯微技術之實驗資料
機器學習引擎	Convolutional Neural Network, Ensemble Learning, Active Learning
結果	成功完成原子座標定位、種類辨識、缺陷邊界型態與相似度分析。降低人為誤判機會。
Status	通過實驗組驗證 論文撰寫中(Nature Materials)

Classification of Crystallographic Groups by Machine Learning

A new powerful approach helping us to identify the microstructure.





使用軟體產
生資料

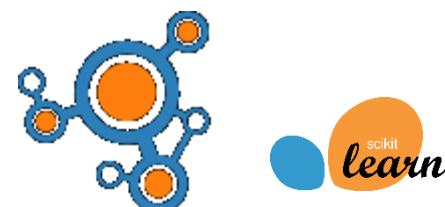
檔案處理

特徵萃取



建立網路

分群演算法



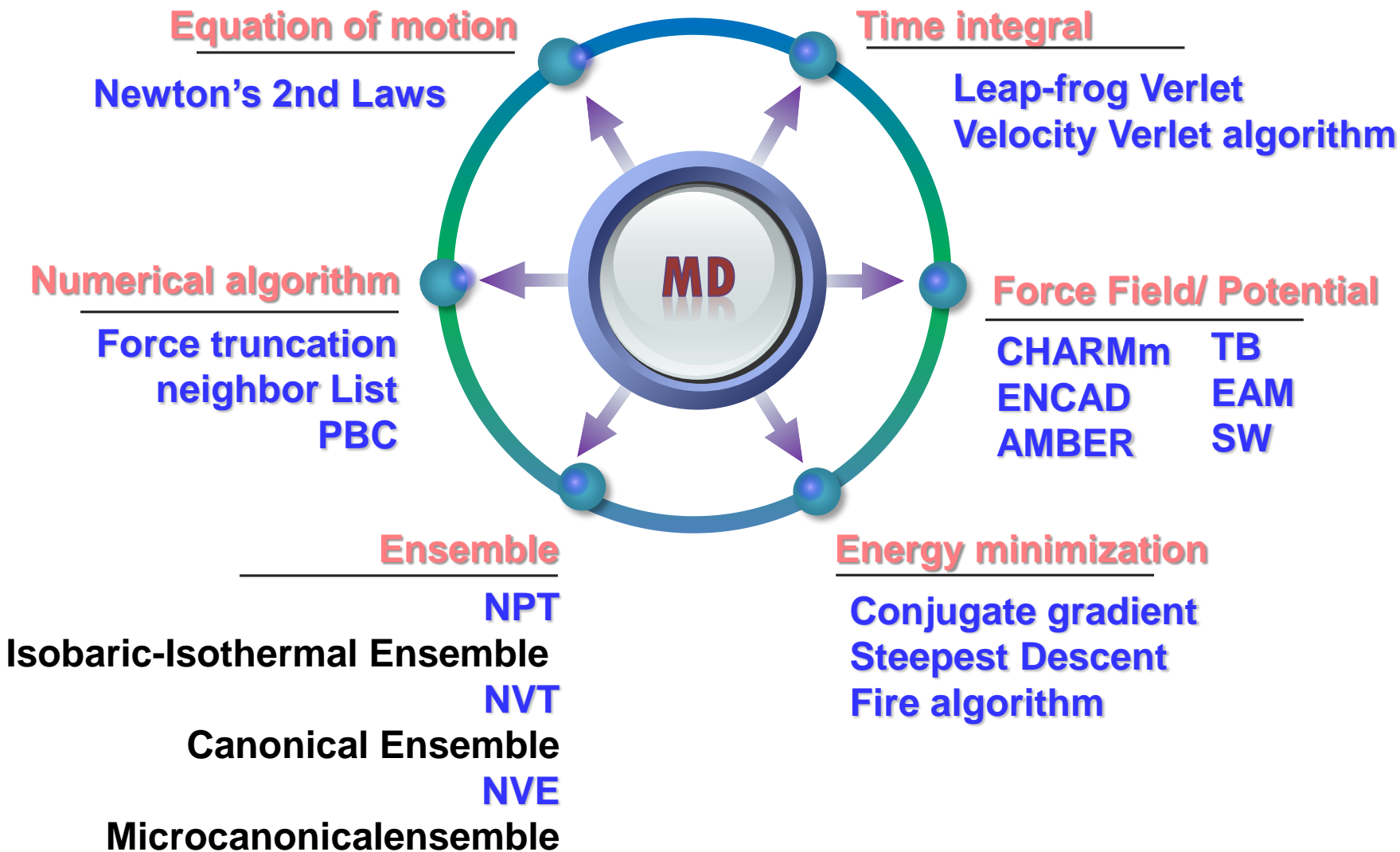
用測試資料
檢驗演算法

調整萃取特徵
方法

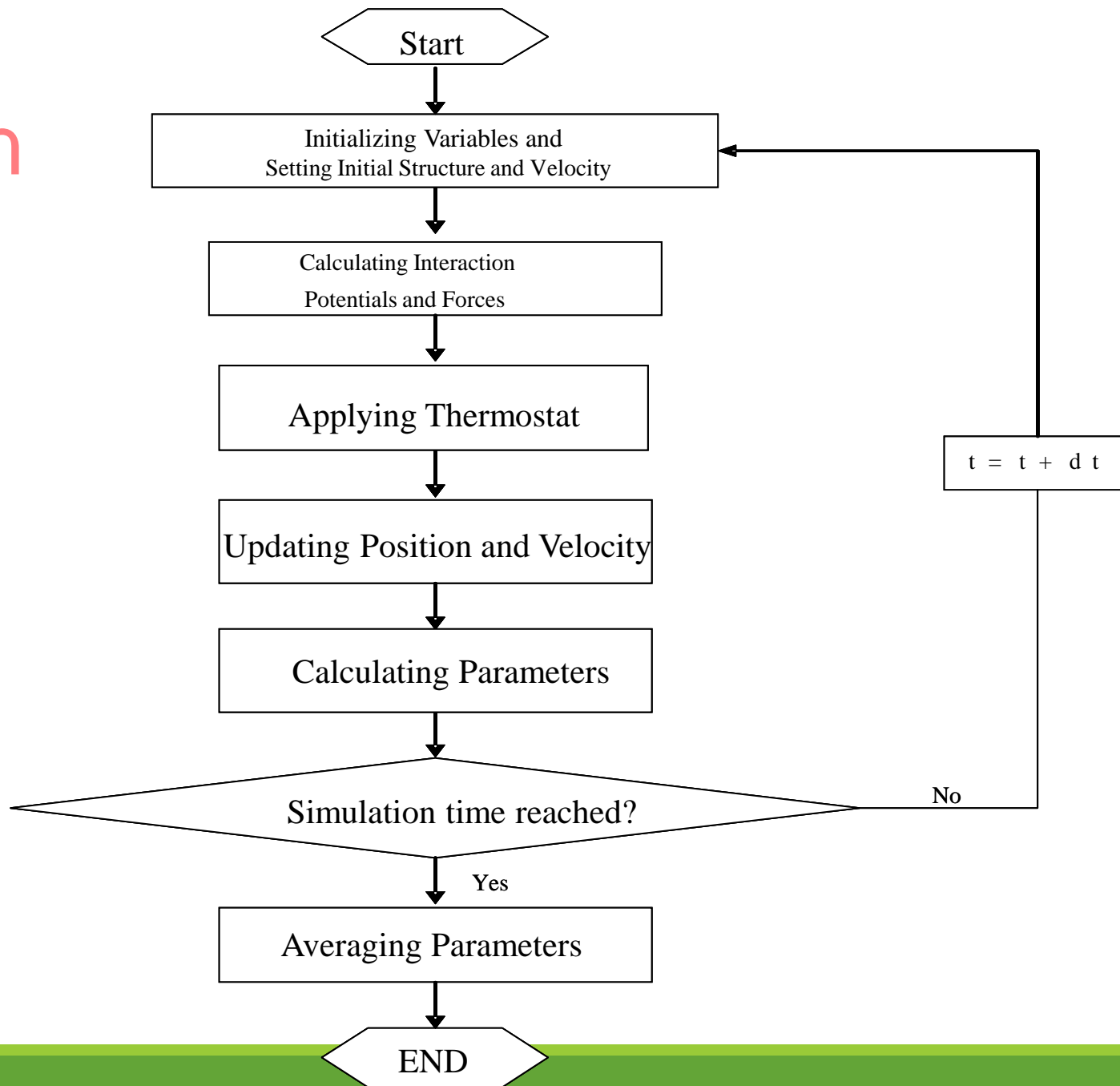
Molecular Dynamics (MD)

- A computer simulation technique that allows one to predict the time evolution of a system of interacting particles (atoms, molecules,.. etc.)
- N-body simulation
- 利用一個描述原子間作用關係來描述一個系統。
- 系統內的所有原子遵循著牛頓運動定律。

Quantum Molecular Dynamics; Classical Molecular Dynamics



Flow Chart of MD simulation



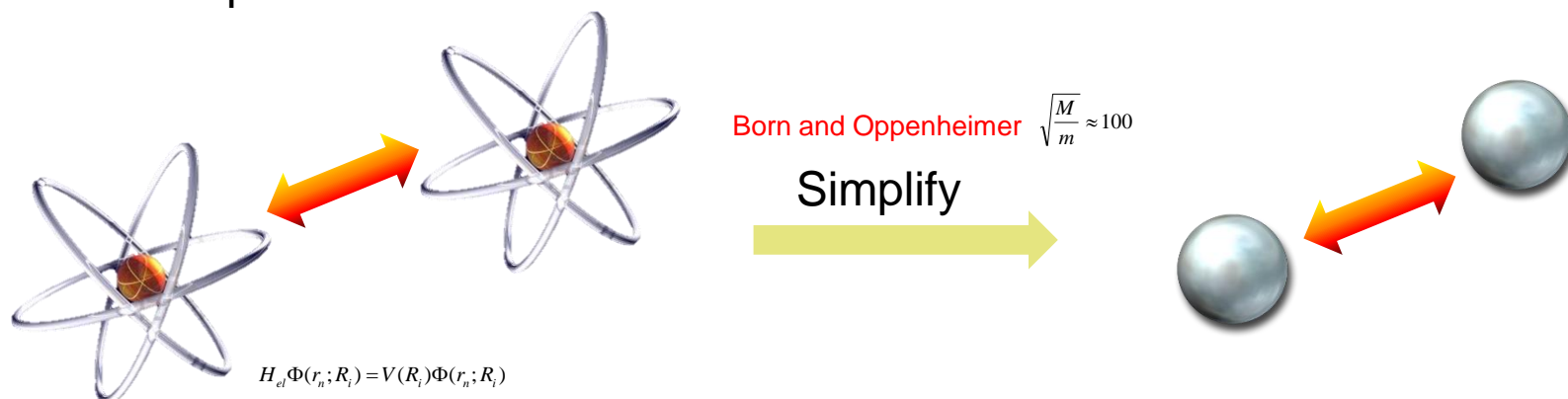
Initial velocity and energy conservation

- Velocities are randomly assigned to the atoms. To achieve faster equilibration atoms can be assigned velocities with the expected equilibrium velocity, i.e. Maxwell distribution.
- Momentum and energy are to be conserved throughout the simulation period.
- Energy conservation is sensitive to the choice of integration method and size of the time step.
- Angular momentum conservation.

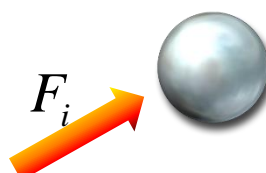
Equation of motion

How Does Molecular Dynamics Work?

In the molecular dynamics method, each atom is treated as a point mass in space



Once the force on each atom is computed, atomic motion is determined through application of Newton's Laws



$$F_i(t) = m_i \frac{d^2 r_i(t)}{dt^2} = -\frac{\partial \Phi(r_i)}{\partial r_i}$$

Second-order ordinary differential equation which can be numerically integrated to find new atomic positions!

Time integration

First-order

Second-order

High-order

Leap-frog Verlet algorithm

$$r(t + \delta t) = r(t) + v(t + \frac{1}{2} \delta t) \delta t$$

$$v(t + \frac{1}{2} \delta t) = v(t - \frac{1}{2} \delta t) + a(t) \delta t$$

Runge–Kutta methods

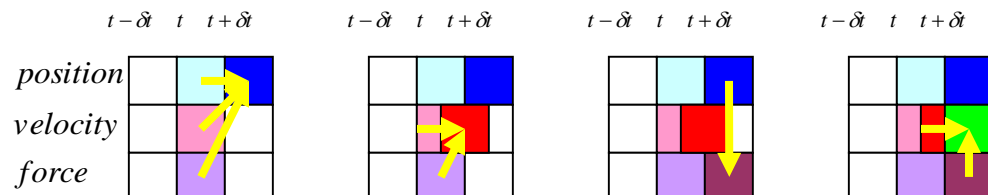
SSPRK3

...

Velocity Verlet algorithm

$$r(t + \delta t) = r(t) + v(t) \delta t + \frac{1}{2} a(t) \delta t^2$$

$$v(t + \delta t) = v(t) + \frac{1}{2} (a(t) + a(t + \delta t)) \delta t$$



Ensembles

An ensemble is a collection of all possible systems which have different microscopic states but have an identical macroscopic or thermodynamic states. There exist different ensembles with different characteristics.

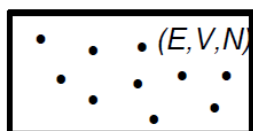
- **Microcanonical ensemble** (NVE) : The thermodynamic state characterized by a fixed number of atoms, N, a fixed volume, V, and a fixed energy, E. ([isolated system](#))

- **Canonical Ensemble** (NVT): This is a collection of all systems whose thermodynamic state is characterized by a fixed number of atoms, N, a fixed volume, V, and a fixed temperature, T. ([system in thermostat](#))

- **Isobaric-Isothermal Ensemble** (NPT): This ensemble is characterized by a fixed number of atoms, N, a fixed pressure, P, and a fixed temperature, T. ([system in barostat](#))

- Microcanonical ensemble

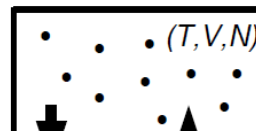
isolated system



*energy transfer not allowed
E constant*

- Canonical ensemble

heat reservoir T



*energy transfer allowed
T constant*

Temperature Control

- Velocity Rescaling

This method is appropriate and efficient for a **homogeneous** system or an inhomogeneous system with **slight difference of mass among atoms**. In the latter case, this method is only appropriate in initial stages of equilibrating the system to relax the random noise of initial velocity.

- Weak coupling method :Berendsen Thermostat / Brownian motion

This method does generate a fluctuation at about the desired temperature, but has disadvantages similar to the rescaling method.

- Stochastic collision method : Andersen thermostat

The drawback of this method is that it cannot generate a smooth trajectory and has **less realistic dynamics**. Therefore, it has been rarely used for molecular dynamics simulations.

- Extended system coupling (Extended Lagrangian) method :

Although the addition of the extended Lagrangian has definitely carried out a correct canonical distribution in property and produced a fluctuation of energy, this assumption of an extended system still cannot make a connection to statistical mechanics.

Temperature Control

- Velocity Rescaling $V_i^{\text{new}} = V_i^{\text{old}} \lambda$ $\lambda = \sqrt{T_d / T_\alpha}$
 - Weak coupling method : Berendsen Thermostat / Brownian motion
-

$$m\dot{v} = F_i - m_i \gamma_i v_i + R(t) \quad (\text{Langevin's equation})$$

- Stochastic collision method : Andersen thermostat
- Extended system coupling (Extended Lagrangian) method :
 - The Nose-Hoover extended system method

$$\boxed{\begin{aligned}\dot{\mathbf{r}}_i &= \frac{\mathbf{p}_i}{m_i} \\ \dot{\mathbf{p}}_i &= -\nabla_i V - \zeta \mathbf{p}_i \\ \zeta &= \frac{\left(\sum_i \frac{\mathbf{p}_i^2}{m_i} - g k_B T \right)}{Q} = \mathbf{V}_T \left[\frac{\sum_i \frac{\mathbf{p}_i^2}{m_i}}{g k_B T} - 1 \right] = \mathbf{V}_T \left[\frac{T^*}{T} - 1 \right]\end{aligned}}$$

Notice that: (1) for $T^* > T$ (system too hot), ζ will increase (if it is positive the system cools down)
(2) for $T^* < T$ (system too cold), ζ will decrease (if it is negative the system heat up)

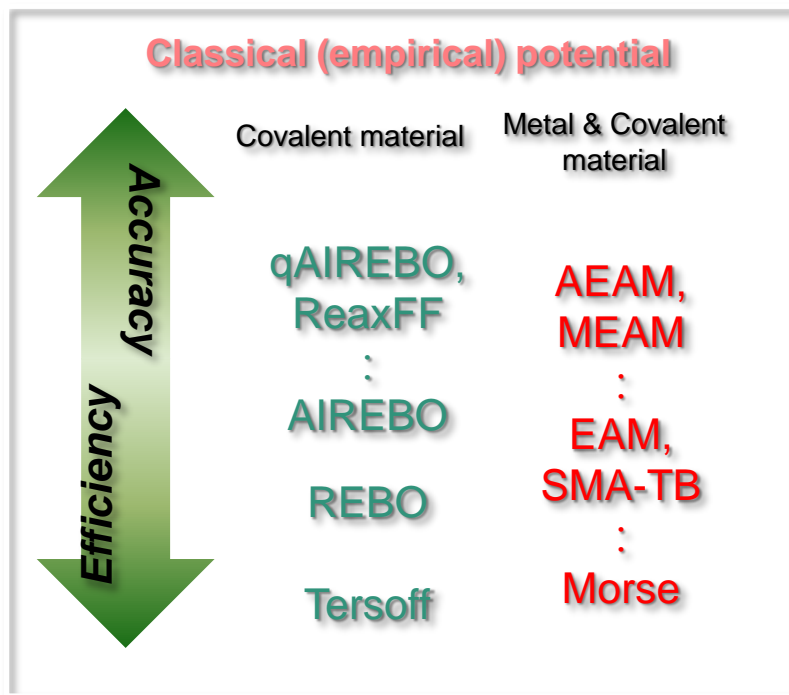
Force Field

In the context of molecular modeling, a force field refers to the form and parameters of mathematical functions used to describe the potential energy of a system of particles (typically molecules and atoms). Force field functions and parameter sets are derived from both experimental work and high-level quantum mechanical calculations. "All-atom" force fields provide parameters for every type of atom in a system.

The usage of the term "force field" in chemistry and computational biology differs from the standard usage in physics. In chemistry it is a system of potential energy functions rather than the gradient of a scalar potential.

To be effective, an analytic Force Field (potential) must possess the following critical properties:

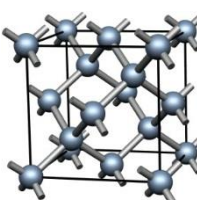
- Accuracy (reproduce properties of interest as closely as possible)
- Computational speed (calculations are fast with simple potentials)
- Transferability (can be used to study a variety of properties for which it was not fit)
- Flexibility (accommodate a wide range of structures in a fitting database)



The potential parameters in the force field are derived from

- ab-initio quantum mechanics
- Crystallography.
- Spectroscopy
-

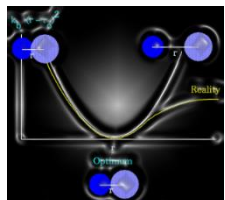
$$\nu = \frac{1}{2\pi} \sqrt{\frac{K}{\mu}}$$



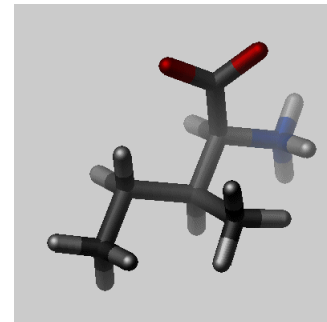
Nonreactive force field (AMBER,CHARMM...)

$$U_{total} = U_{bond} + U_{bending} + U_{dihedral} + U_{vdw} + U_{col}$$

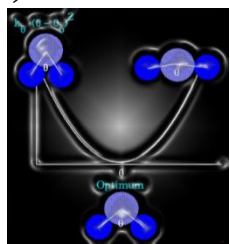
(a) Bond strength



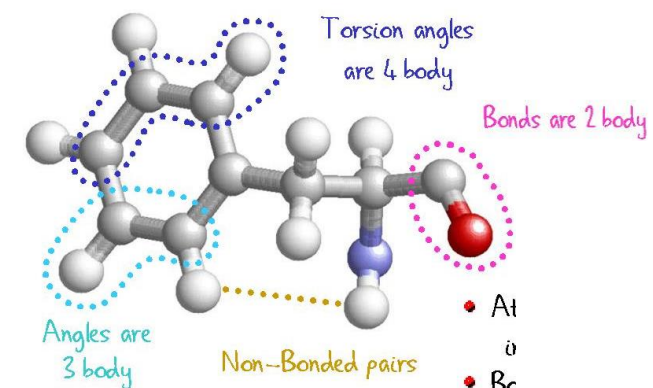
$$U_{bond\ length} = \sum_{N=1}^{N_0} \frac{1}{2} K_b^i (b_i - b_0^i)^2$$



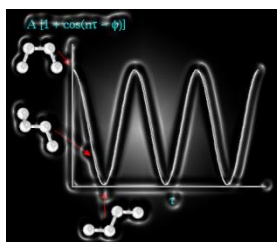
(b) Bond bending



$$U_{bending} = \sum_{N=1}^{N_0} \frac{1}{2} K_\theta^i (\theta_i - \theta_0^i)^2$$



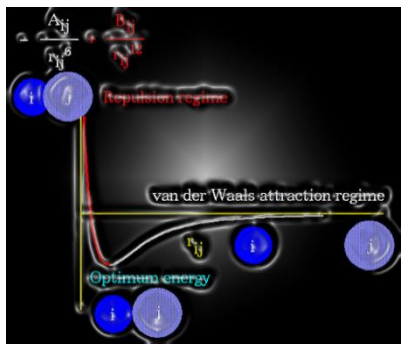
(c) Dihedral Angle



$$U_{dihedral} = \sum_{N=1}^{N_\phi} K_\phi^i \left\{ 1 - \cos[N^i(\phi_i - \phi_0^i)] \right\}$$

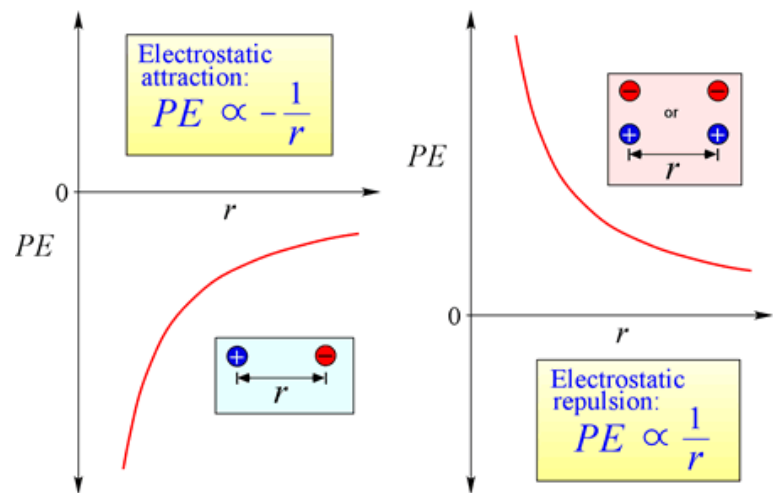
- Atoms are connected in a simple graph.
- Bonded Interactions: 2, 3, 4 body.
- Non-Bonded Interactions.

(d) Van der Waal



$$U_{\text{vdw}} = \left[\epsilon^{ij} \left(\frac{r_0^{ij}}{r_{ij}} \right)^{12} - 2 \epsilon^{ij} \left(\frac{r_0^{ij}}{r_{ij}} \right)^6 \right]$$

(e) Coulomb force



$$U_{\text{col}} = \sum_{\text{partialcharges}} \frac{q^i q^j}{r_{ij} \epsilon}$$

Treating Long-Range Electrostatic Interactions
particle-mesh Ewald (PME); Ewald sum

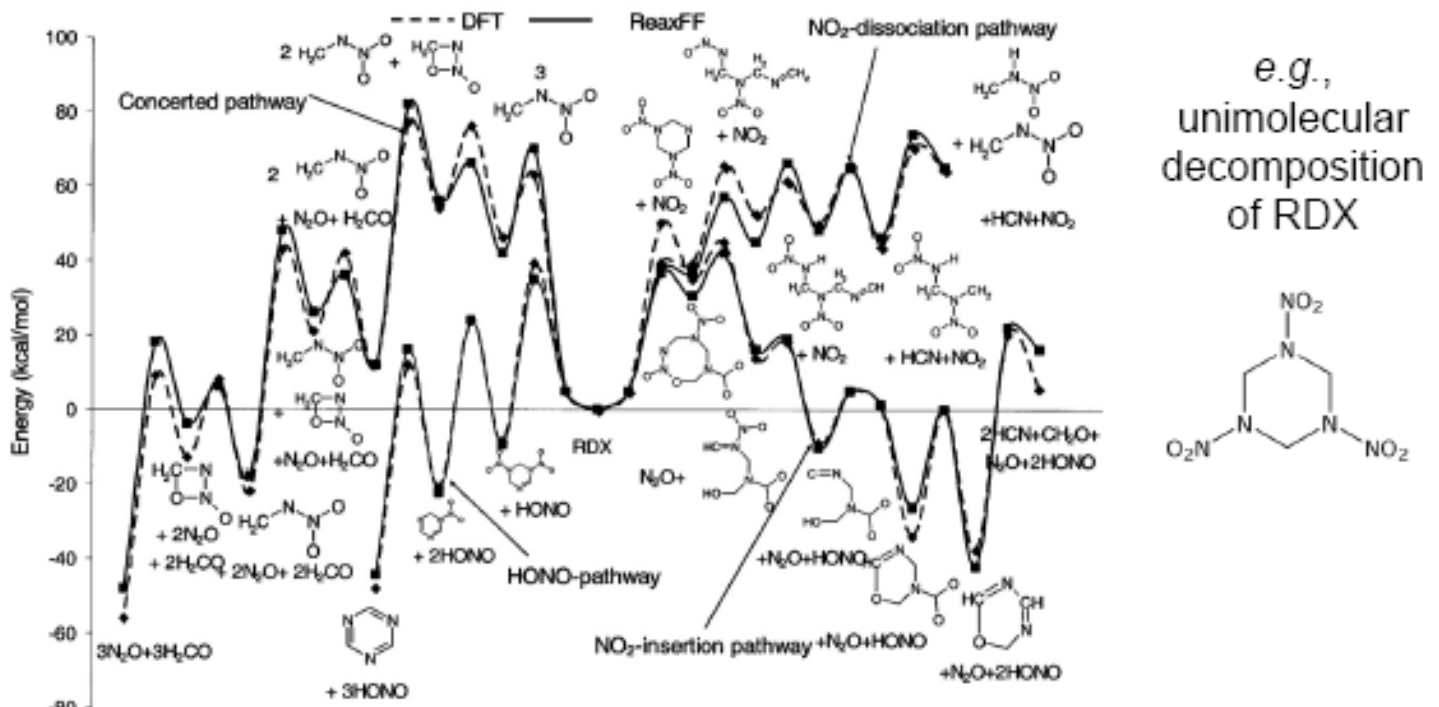
ReaxFF Potential Energy Function

$$E_{\text{system}} = E_{\text{bond}} + E_{\text{lp}} + E_{\text{over}} + E_{\text{under}} + E_{\text{val}} + E_{\text{pen}} + E_{\text{coa}} + E_{\text{C2}} + E_{\text{triple}} + E_{\text{tors}} + \\ E_{\text{conj}} + E_{\text{H-bond}} + E_{\text{vdWaals}} + E_{\text{Coulomb}}$$

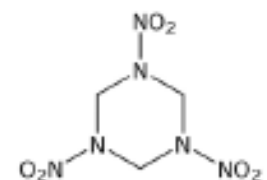
- E_{bond} : bond energy; attractive term, directly derived from bond orders
- E_{lp} : Lone pair energy; penalty for breaking up lone pairs in O, N
- E_{over} : Overcoordination energy: penalty for overcoordinating atoms
- E_{under} : Undercoordination energy: stabilizes undercoordinated atoms
- E_{val} : Angle strain; equilibrium angle depends on bond order central atom
- E_{pen} : Penalty for 'allene'-type molecules (H2C=C=CH2)
- E_{coa} : Angle conjugation; stabilizes -NO2 groups
- E_{C2} : C2 correction: destabilizes C=C
- E_{triple} : triple bond related, first mentioned in publications from 2008
- E_{tors} : Torsion energy: bond-order dependent V2-term
- E_{conj} : Torsion conjugation: general conjugation stability
- $E_{\text{H-bond}}$: Hydrogen bond
- E_{vdWaals} : van der Waals: calculated between every atom
- E_{Coulomb} : Coulomb interaction: calculated between every atom; polarizable charges get updated every iteration

ReaxFF/Ab Initio Comparison

ReaxFF can describe a wide variety of chemical reactions.



e.g.,
unimolecular
decomposition
of RDX



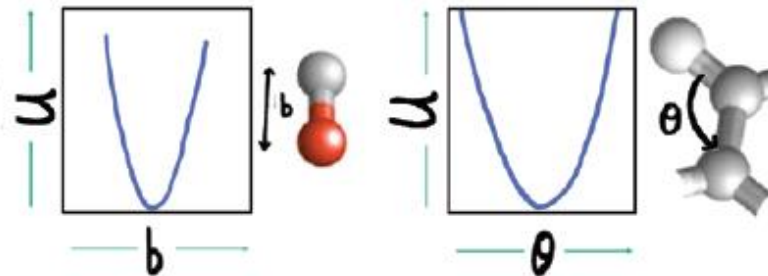
Strachan, et al, JCP, 122, 054502 ('05).

(Original slide from Quenneville presentation)

MOLECULAR POTENTIAL ENERGY

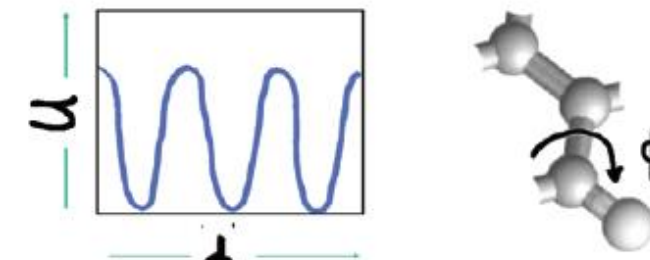
$$U = \sum_{\text{All}}^{\frac{1}{2}K_b(b-b_0)^2} + \sum_{\text{All}}^{\frac{1}{2}K_\theta(\theta-\theta_0)^2}$$

Hooke 1635



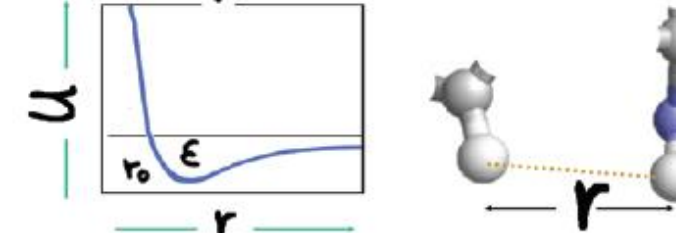
$$+ \sum K_\phi [1 - \cos(n\phi + \delta)]$$

All Torsion



$$+ \sum \epsilon \left[\left(\frac{r_0}{r} \right)^{12} - 2 \left(\frac{r_0}{r} \right)^6 \right]$$

All Nonbonded

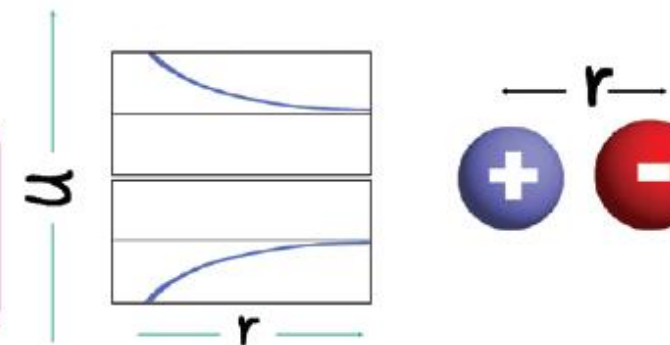


$$+ \sum 332q_i q_j / r$$

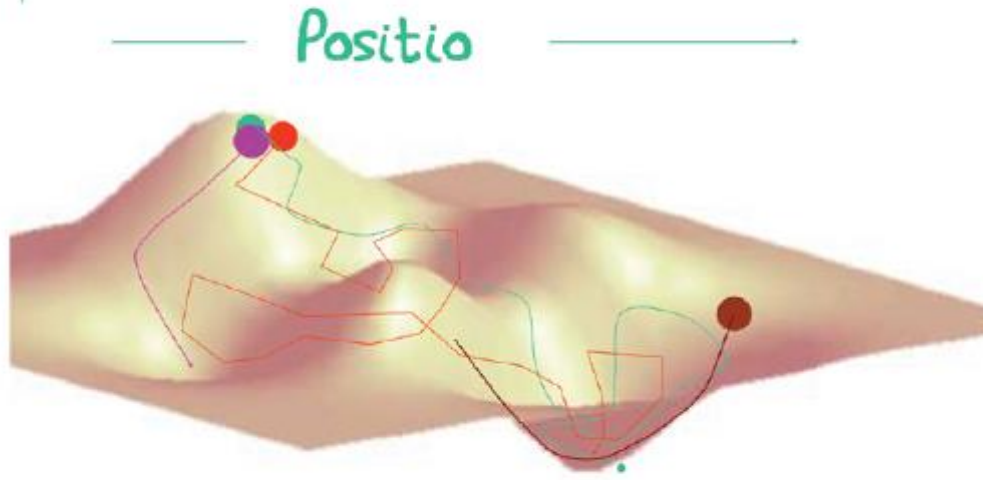
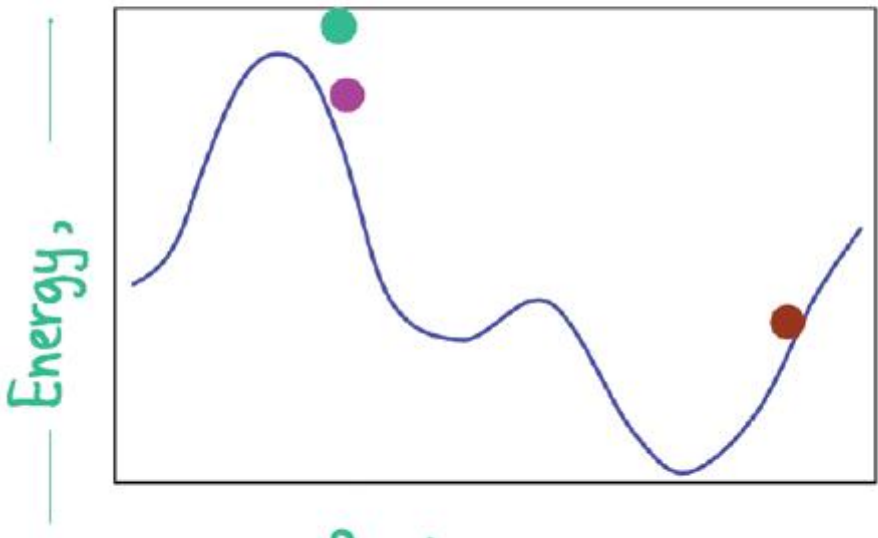
All partial

Coulomb 1736

Simple sum
over many
terms

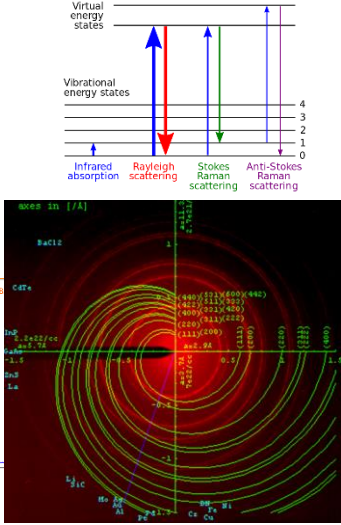
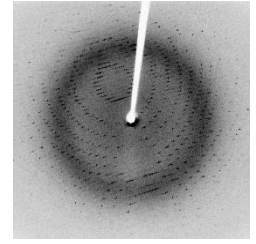


MOVING OVER ENERGY SURFACE

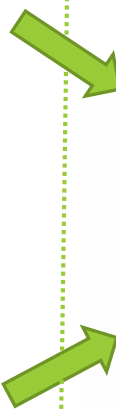


- EM: Energy Minimization drops into local minimum.
Euclid 325 BC
- NMD: Normal Mode Dynamics vibrates about minimum.
Galileo 1564
- MD: Molecular Dynamics uses thermal energy to move smoothly over surface.
Newton 1643
- MC: Monte Carlo Moves are random. Accept with probability $\exp(-\Delta U/kT)$.
Metropolis 1915

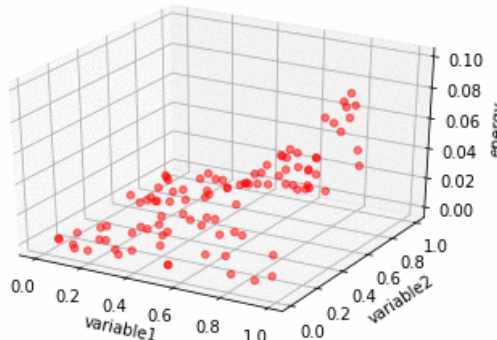
Fitting energy surface in high dimensional data space with Machine Learning



Experimental measurement



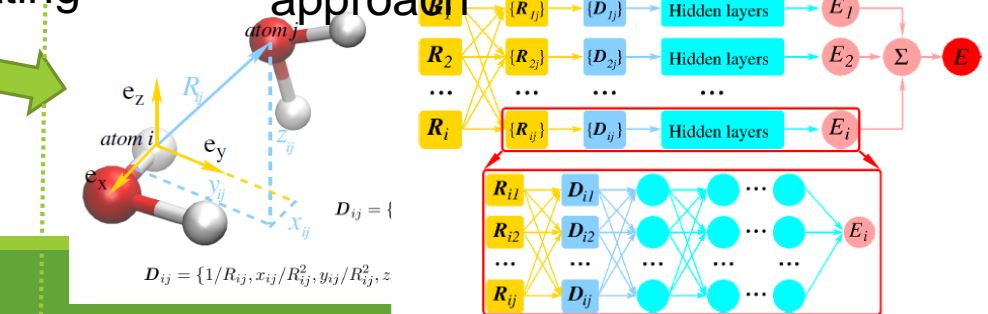
Real P.E.S.



Human fitting



NN fitting

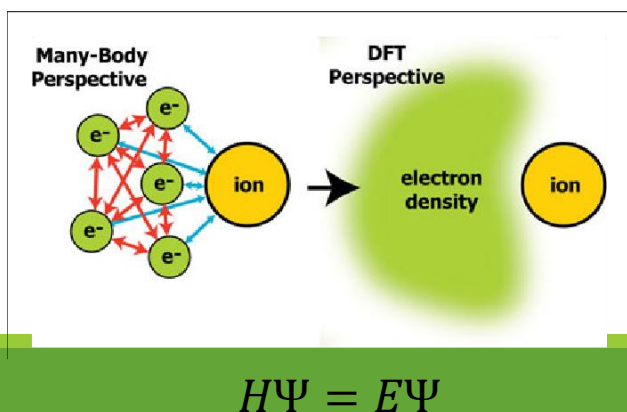


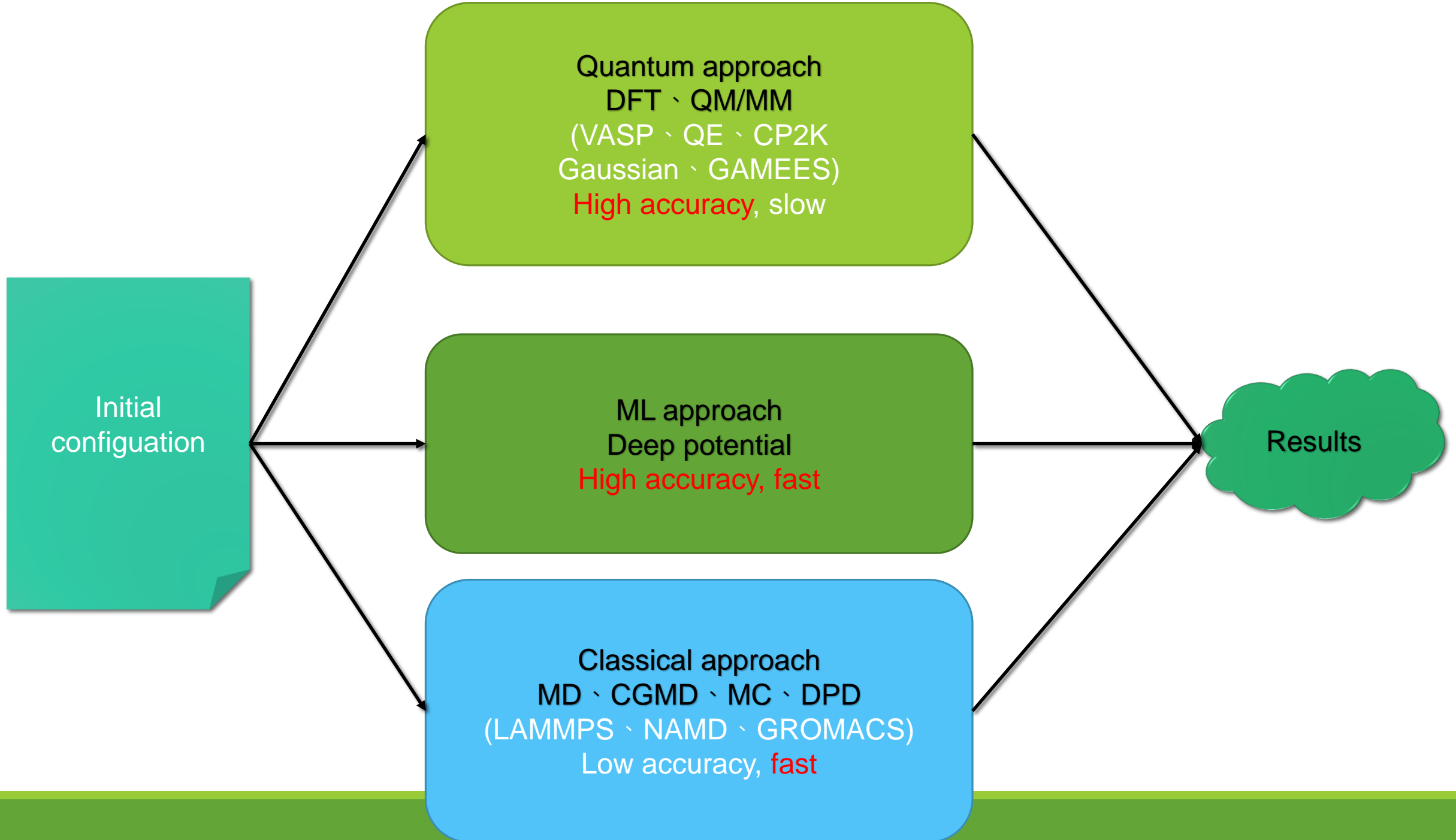
Ordinary approach

MOLECULAR POTENTIAL ENERGY

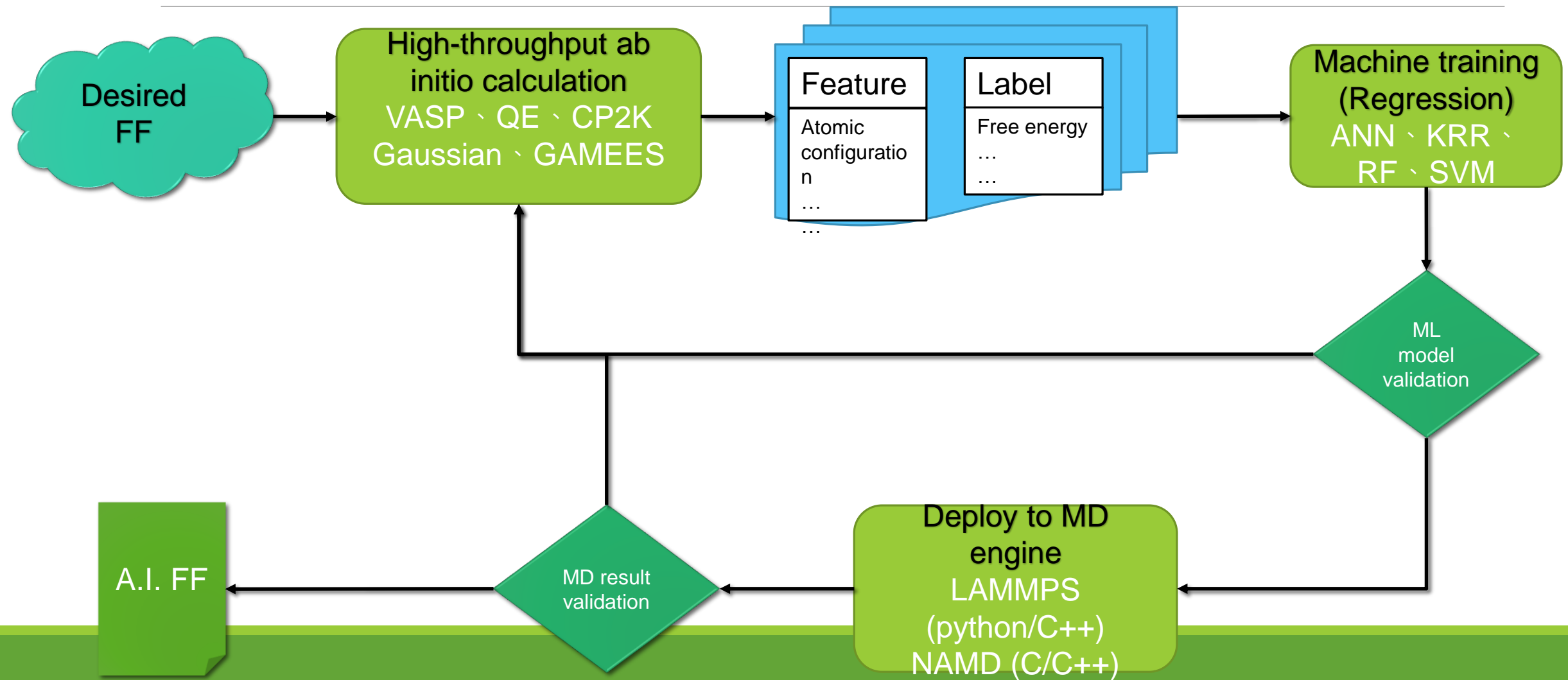
$$U = \sum_{\text{All}} \frac{1}{2} K_b (b - b_0)^2 + \sum_{\text{All}} \frac{1}{2} K_\theta (\theta - \theta_0)^2 + \sum_{\text{All Torsion}} K_\phi [1 - \cos(n\phi + \delta)] + \sum_{\text{All Nonbonded}} \epsilon [(r_0/r)^12 - 2(r_0/r)^6] + \sum_{\text{All partial}} 332 q_i q_j / r$$

Hooke 1635
Fourier 1768
Van der Waals 1837
Coulomb 1736
Simple sum over many terms





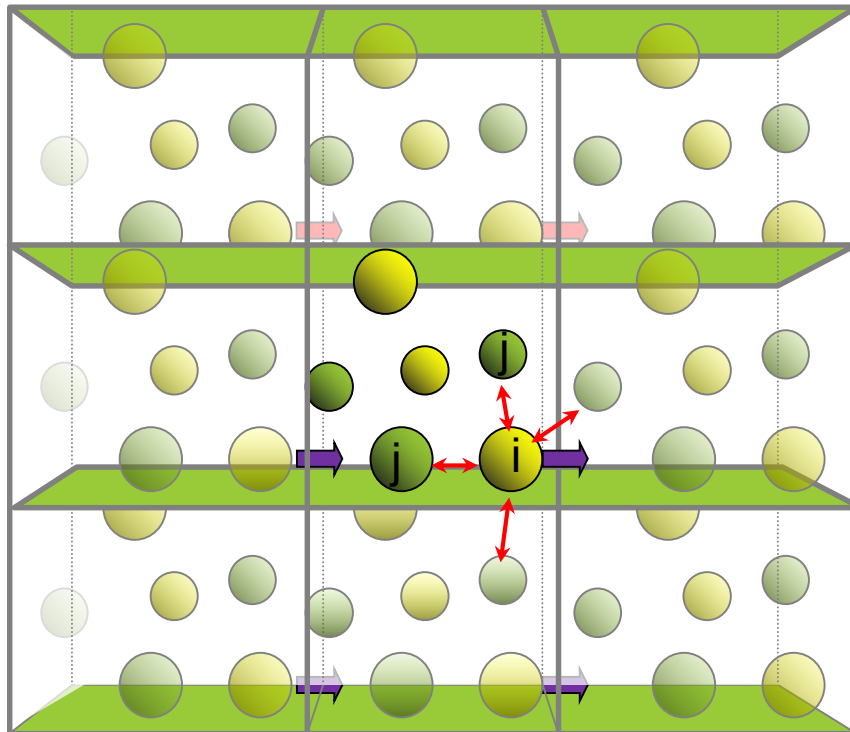
Flow chart for Deep Potential



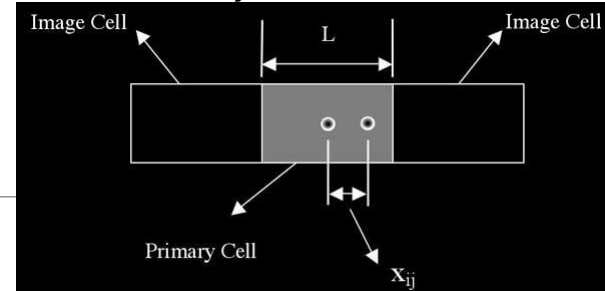
Numerical algorithm

- Periodic Boundary Condition
- Minimum Image Criterion

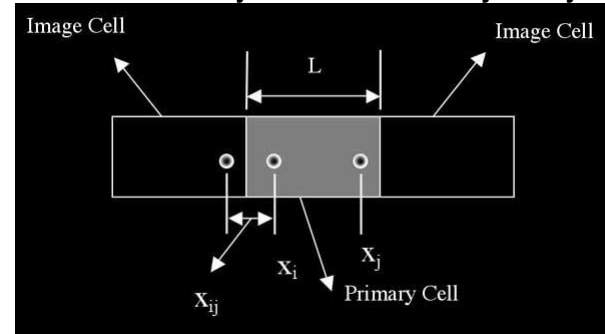
Case B: $x_{ij} < -0.5L \rightarrow x_{ij} = x_{ij} + L$



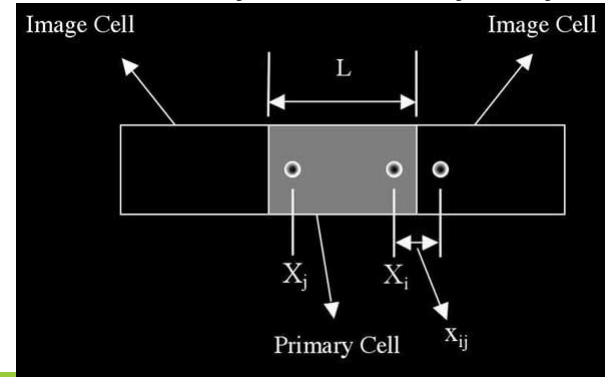
Case A: x_{ij} is less than $0.5L$



Case B: $x_{ij} < -0.5L \rightarrow x_{ij} = x_{ij} + L$



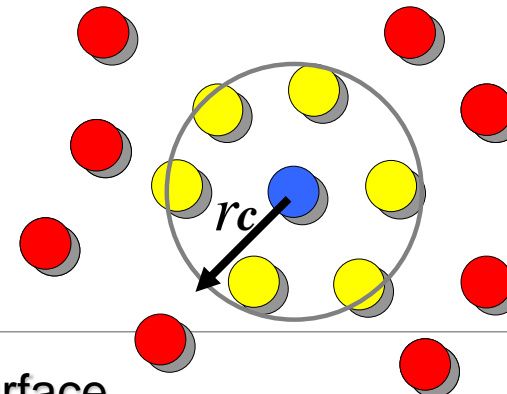
Case C: $x_{ij} > 0.5L \rightarrow x_{ij} = x_{ij} - L$



Force Truncation

● Truncation

$$U(r) = \begin{cases} U(r) & r < R_c \\ 0 & r \geq R_c \end{cases}$$



- **Shift function:** modifies the entire potential energy surface

$$U_{nb}(r) = \begin{cases} U_{nb}(r) \left(1 - \left(\frac{r}{R_c}\right)^2\right)^2 & r < R_c \\ 0 & r \geq R_c \end{cases} \quad \dots \quad \text{Brook et. al.}$$

$$U_{nb}(r) = \begin{cases} U_{nb}(r) - \left[U_{nb}(R_C) + (r - R_C) \left[\frac{dU_{nb}(R_C)}{dr} \right] \right] & r < R_C \\ 0 & r \geq R_C \end{cases} \dots \text{Levitt et. al.}$$

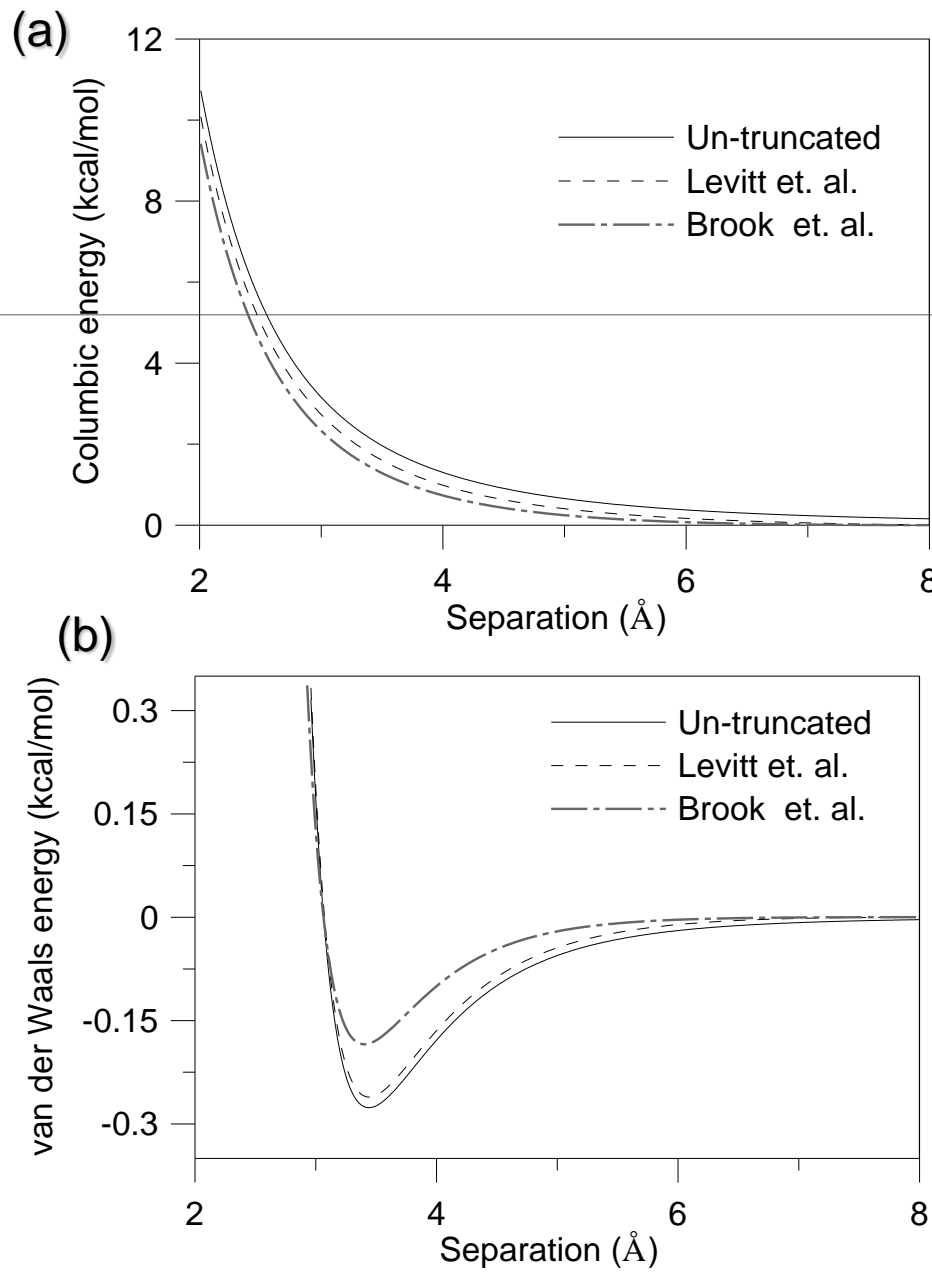
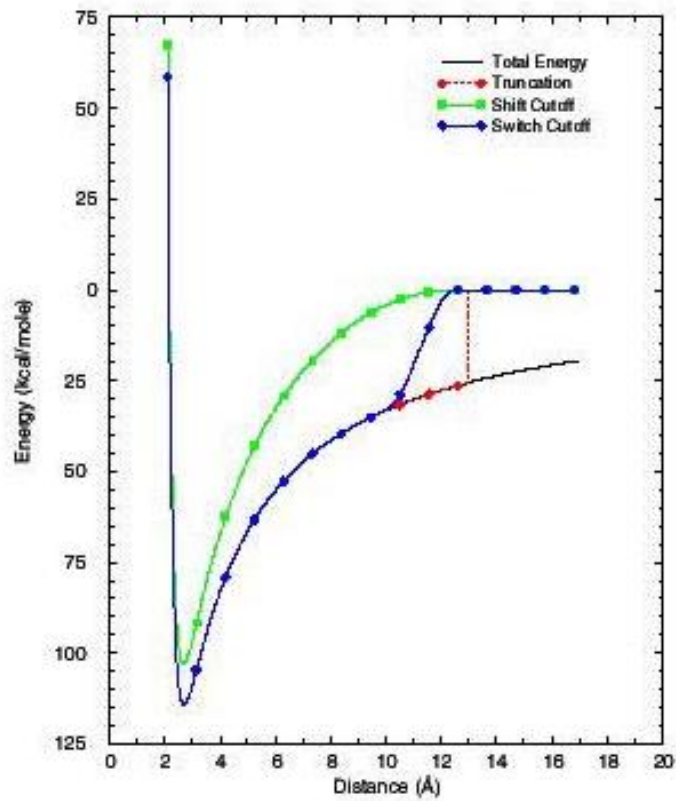
- **Switch function:** modifies the interaction potential over a predefined range of distances

$$U_{nb}(r) = \begin{cases} U_{nb}(r) & r < r_l \\ U_{nb}(r) \left\{ \frac{(r_c - r)^2(r_c + 2r - 3r_l)}{(r_c - r_l)^3} \right\} & r_l \leq r < r_c \\ 0 & r_c \leq r \end{cases}$$

CHARMm

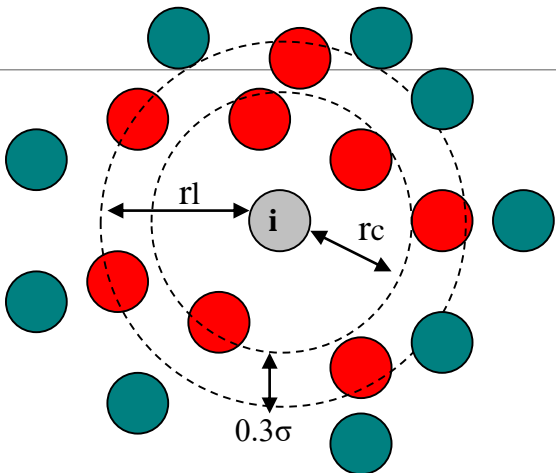
● The Ewald summation technique

(a) Columbic energy profile between two water molecules and (b) the van der Waals energy profile between oxygen atom in C=O group and nitrogen atom in N-H group with and without truncation methods.

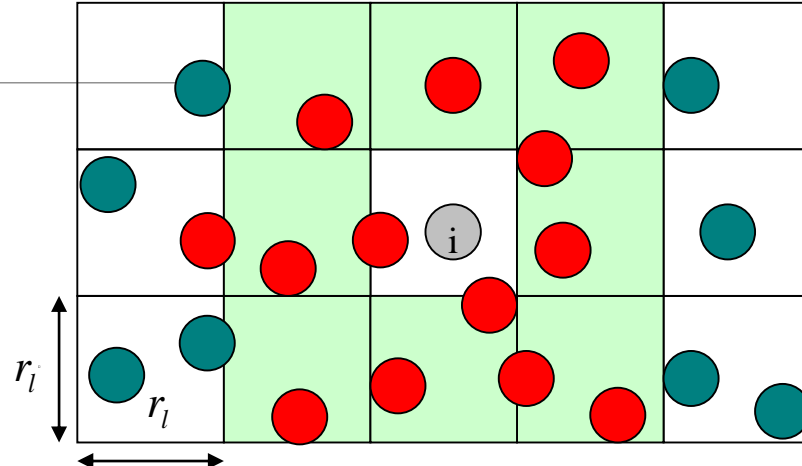


Efficient Neighbours List

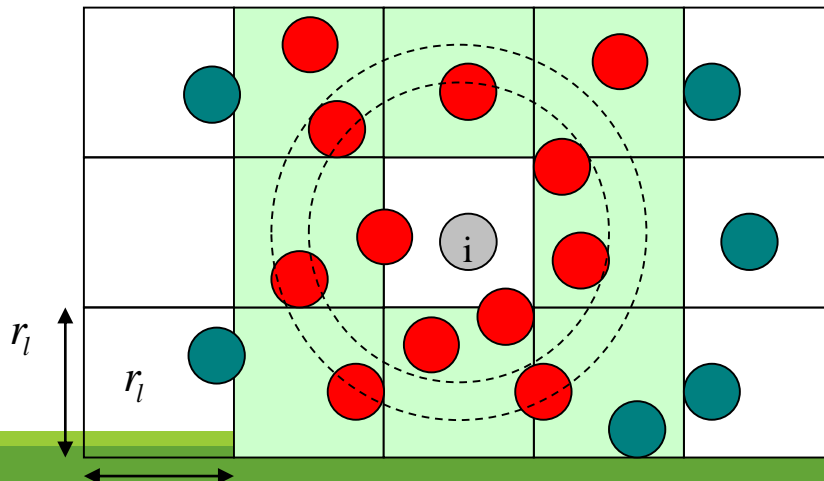
● Verlet List



● Cell Link



● Verlet List + Cell Link



Packages for molecular modeling

Package Name	Developer	Features
LAMMPS	Sandia National Labs	atomic, polymeric, biological, metallic, granular, and coarse-grained systems
NAMD	Parallel Programming Labs	structural biology
GROMACS	Science for Life Labs	general purpose
DL_POLY	Daresbury Labs	general purpose
Desmond	D. E. Shaw Research	biological and chemical systems
AMBER	Peter Kollman et. Al.	biomolecules
SPaSM	Los Alamos National Labs	Large scale simulation

Installation of Windows pre-built binary (ICMS version)

weppage Screenshot

LAMMPS-ICMS Windows Installer Repository

This repository is hosting pre-compiled Windows installers of the [LAMMPS](#) molecular dynamics simulation software package. The binaries are built semi-automatically with MinGW64 Linux to Windows cross compilers using up-to-date snapshots of the [LAMMPS-ICMS git repository](#) hosted at the [Institute for Computational Molecular Science](#) at Temple University. The LAMMPS binaries contain **all** optional packages included in the source distribution **except**: KIM (license is not GPL compatible), USER-CUDA (CUDA does not support cross compilation), KOKKOS (does not support cross-compilation), and REAX (superseded by the USER-REAXC package).



Some Notes on GPU Support

These Windows binaries include (experimental on Windows) GPU acceleration via the [GPU](#) package. This is achieved through compiling the GPU package in OpenCL mode and linking to an [OpenCL v1.2](#) compatible ICD loader. This means the executables do not contain any vendor provided code and should be compatible with GPUs from both [AMD](#) and [Nvidia](#). The GPU package has been compiled for mixed precision computation and is currently somewhat tuned for Nvidia (Fermi generation) GPUs. It does not yet work with OpenCL drivers for CPUs (like those included in the Intel and AMD OpenCL SDKs). Work is under way to improve and expand OpenCL support in LAMMPS and particularly provide means so that the individual kernels are optimized for specific platforms at runtime. Please watch [Mike Brown's LAMMPS accelerator library web page](#) for status updates.

When reporting problems, please always include the exact [version](#) of the installer and the output of the `ocl_get_devices` tool.

Looking for pre-compiled Linux binary RPMs? They are just a [mouse click away](#).

Installing LAMMPS on Windows

There are installer packages for 32-bit and 64-bit versions of Windows available.

- [Latest version for 32-bit Windows](#)
- [Latest version for 64-bit Windows](#)

[32-bit Windows download area](#)
[64-bit Windows download area](#)

The respective download directory will contain installer packages that are labeled with the date they were compiled on and one package labeled as *latest*. It is usually recommended to download and install the latest package via the link above. The other packages are provided in case there is a problem with it. Download the installer executable suitable for your machine, execute it, and follow the instructions in the dialogs. Each version will install into a different directory, so it is possible to have multiple versions installed at the same time. Both kinds of packages contain:

- A regular multi-threaded LAMMPS executable called `lmp_serial`. This should always work.
- A multi-threaded LAMMPS executable that also supports parallel execution via MPI message passing. This executable is called `lmp_mp1` and requires installation of a suitable MPICH2 package to work.
- the LAMMPS manual in PDF format
- the [colvirs](#) reference manual in PDF format
- the potential files bundled with the LAMMPS source code
- most of the example inputs, reference outputs and related files
- the benchmark inputs and reference outputs
- the tools `restart2data`, `binary2txt`, `chain`, `msi2lmp`, `ocl_get_devices`

Both executables will run in serial when executed directly. Please see below for instructions on how to perform parallel runs. To use the MPI based parallelism, you also need to install [MPICH2 from Argonne lab](#). For 32-bit Windows you have to download and install [mpich2-1.4.1p1-win-ia32.msi](#) or any compatible version. Correspondingly, for 64-bit Windows you have to download and install [mpich2-1.4.1p1-win-x86-64.msi](#) or any compatible version.

Running LAMMPS on Windows

General Comments

<http://rpm.lammps.org/windows.html> Fri Aug 15 2014 10:35:07 GMT+0800 (台北標準時間)

Choose executable file based on the architecture of your PC, then double click on it to install. If you want to run LAMMPS in parallel mode, [mpich2-1.4.1p1-win-x86-64.msi](#) is also needed.

LAMMPS Windows Installer Repository

<https://packages.lammps.org/windows.html>



↑

Run LAMMPS on Windows

1. Install lammps on your windows system, put the path of lammps exe file(Ex. `lmp_serial.exe`) in the PATH environment variable (Ex. `C:\Program Files\LAMMPS 64-bit 16Aug2018\bin`).
2. Put input file(Ex. `in.xxx`), potential file or coordinate file (optional)in project folder (`X:\project_folder`).
3. Open cmd , and change directory to project folder("X:" then "cd project_folder").
4. Or press shift and right click to open cmd in the project_folder.
5. Use the following command to run lammps "`C:\Program Files\LAMMPS 64-bit 16Aug2018\bin\lmp_serial.exe -in in.xxx`"
6. Use the following command to run lammps with multi-threading parallelization "`C:\Program Files\LAMMPS 64-bit 16Aug2018\bin\lmp_serial.exe -in in.xxx -pk omp 4 -sf omp`"

LAMMPS Command-line options

Imp_serial.exe -in in.xxx -pk omp 4 -sf omp -v rseed 46

- -i or -in Specify a file to use as an input script.
- -pk or -package Invoke the package command with style and args
- -sf or -suffix Use variants of various styles if they exist. The specified style can be gpu, intel, kk, omp, opt, or hybrid.
- -v or -var Specify a variable that will be defined for substitution purposes when the input script is read.
(referenced as \$abc or \${abc} in the input script)
- ...

LAMMPS input script

- LAMMPS executes by reading commands from a input script (text file).
- One line at a time.
- The input script is read one line at a time and each command takes effect when it is read.
- Some commands are only valid when they follow other commands.

```
timestep 0.5  
run    100  
run    100
```

```
run    100  
timestep 0.5  
run    100
```

LAMMPS input script

- All characters from the first “#” character onward are treated as comment and discarded.
- If the last printable character on the line is a “&” character, the command is assumed to continue on the next line.
- The input script is read one line at a time and each command takes effect when it is read.
- The line is broken into “words” separated by white-space (tabs, spaces).
- The first word is the command name. Other words in the line are arguments.
- If you want text with spaces to be treated as a single argument, it can be enclosed in either single or double or triple quotes.
- If you want multiple lines of an argument to retain their line breaks, the text can be enclosed in triple quotes.

LAMMPS input script structure

設定檔可區分為4個部分

Initialization

Atom Definition

Settings

Run a Simulation

Settings

Run a Simulation

...

命令可以分類成9群

1. Initialization
2. Atom Definition
3. Force Fields
4. Settings
5. Fixes
6. Computes
7. Outputs
8. Actions
9. Miscellaneous

LAMMPS

INITIALIZATION

ATOM DEFINITION

SETTINGS

Force fields

Parameters

Fixes

Computes

Output

RUN

Actions

Miscellaneous

Commands

[atom_modify](#), [atom_style](#), [boundary](#), [dimension](#), [newton](#), [processors](#), [units](#)

[create_atoms](#), [create_box](#), [lattice](#), [read_data](#), [read_dump](#), [read_restart](#), [region](#), [replicate](#)

[angle_coeff](#), [angle_style](#), [bond_coeff](#), [bond_style](#), [dielectric](#), [dihedral_coeff](#), [dihedral_style](#), [improper_coeff](#), [improper_style](#), [kspace_modify](#), [kspace_style](#), [pair_coeff](#), [pair_modify](#), [pair_style](#), [pair_write](#), [special_bonds](#)

[neighbor](#), [neigh_modify](#), [group](#), [timestep](#), [reset_timestep](#), [run_style](#), [min_style](#), [min_modify](#).

[fix](#), [fix_modify](#), [unfix](#)

[compute](#), [compute_modify](#), [uncompute](#)

[dump](#), [dump_image](#), [dump_modify](#), [dump_movie](#), [restart](#), [thermo](#), [thermo_modify](#), [thermo_style](#), [undump](#), [write_data](#), [write_dump](#), [write_restart](#)

[run](#), [temper](#), [minimize](#), [neb_prd](#), [rerun](#)

[delete_atoms](#), [delete_bonds](#), [displace_atoms](#), [change_box](#)

[clear](#), [echo](#), [if](#), [include](#), [jump](#), [label](#), [log](#), [next](#), [print](#), [shell](#), [variable](#)

Command Format

- ✓ **COMMAND** (+ group-ID) (+**args...**) (style (+**args...**))(+**其它(+ values ...)**) (+ keyword (+ values ...))

Example:

neb etol ftol N1 N2 Nevery file-style arg

- etol = stopping tolerance for energy (energy units)
- ftol = stopping tolerance for force (force units)
- N1 = max # of iterations (timesteps) to run initial NEB
- N2 = max # of iterations (timesteps) to run barrier-climbing NEB
- Nevery = print replica energies and reaction coordinates every this many timesteps
- file-style=*final* or *each* or *none*

final arg = filename

filename = file with initial coords for final replica coords for intermediate replicas are linearly interpolated between first and last replica

each arg = filename

filename = unique filename for each replica (except first) with its initial coords

none arg = no argument

all replicas assumed to already have their initial coords

- ✓ fix +ID + group-ID + style (+**args...**) (+ keyword (+ values ...))
- ✓ compute +ID + group-ID + style (+**args...**) (+ keyword (+ values ...))
- ✓ variable+name + style_name +args...

※若**arg** 或**value**為斜字體即為字元(*yes, no, final ...*)，若非則為數字

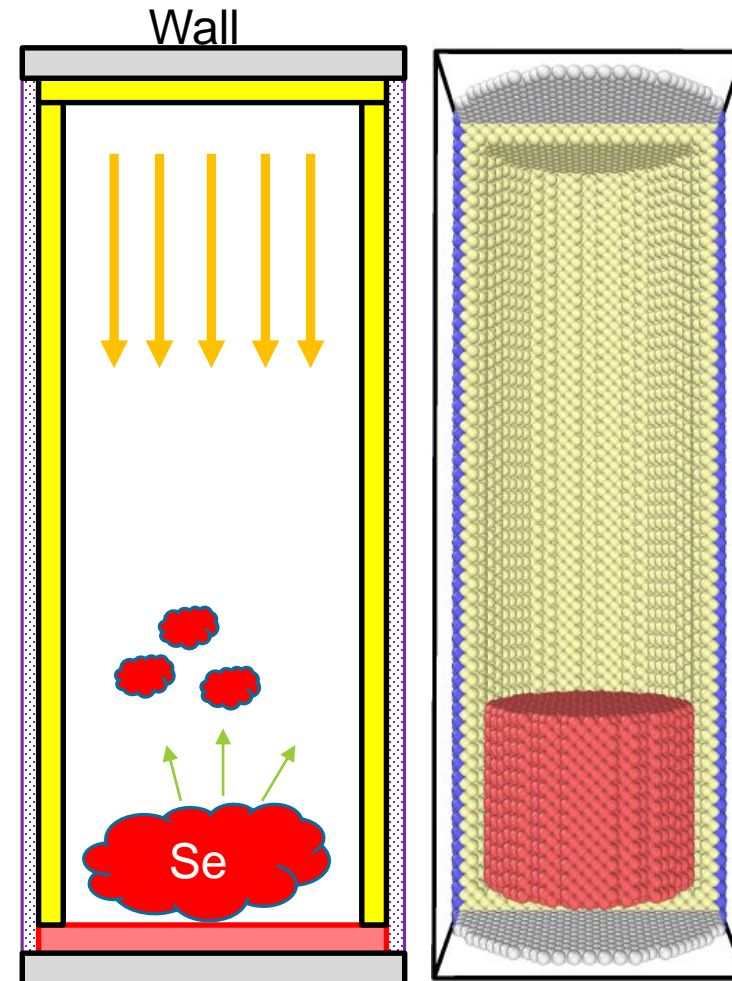
command參數	說明
style(+arg)	<p>擁有多重功能參數之command 會有style參數</p> <ul style="list-style-type: none"> ✓ 功能參數 -style · 名稱有特定 -arg · 部份名稱為特定，部份由使用者給數據，可為數字或字元 ✓ 無指定群組 · e.g.: <ul style="list-style-type: none"> min_style (選擇能量最小化方法) lattice (選擇晶格) pair_style (選擇勢能函數) ✓ 指定群組 · e.g.: <ul style="list-style-type: none"> fix (+ID + group - ID + style_name) compute (+ID + group - ID + style_name)
ID, group-ID, region-ID, fix-ID,	參數中的指定代號,指定群組代號,指定區間代號 , fix代號 (使用者給定)
keyword (+value)	<p>可額外選用之子功能/子條件之參數(可給可不給)</p> <ul style="list-style-type: none"> -keyword · 名稱有特定 -value · 部份名稱為特定，部份由使用者給數據，可為數字或字元
file	<p>檔案名稱 · e.g.:</p> <p>dump ID group-ID style N file args</p>
name	<p>名稱 · e.g.:</p> <p>variable name</p>
其它	以上參數以外的名稱通常為供特定功能所設置之參數名稱，其定義請閱讀manual

You should know about this

region	area
region-ID	name of region
group	a cluster of atoms
group-ID	Name of group
type	atom type
molecule	molecule id #
id	atom id #

Addforce zone
→

Heating zone



compute command --diagnostics

centro/atom - centro-symmetry parameter for each atom

cna/atom - common neighbor analysis (CNA) for each atom

com - center-of-mass of group of atoms

com/molecule - center-of-mass for each molecule

coord/atom - coordination number for each atom

group/group - energy/force between two groups of atoms

gyration/molecule - radius of gyration for each molecule

heat/flux - heat flux through a group of atoms

inertia/molecule - inertia tensor for each molecule

ke - translational kinetic energy

msd - mean-squared displacement of group of atoms

msd/molecule - mean-squared displacement for each molecule

pe - potential energy

pe/atom - potential energy for each atom

pressure - total pressure and pressure tensor

rdf - radial distribution function $g(r)$ histogram of group of atoms

reduce - combine per-atom quantities into a single global value

reduce/region - same as compute reduce, within a region

stress/atom - stress tensor for each atom

temp - temperature of group of atoms

voronoi/atom - Voronoi volume and neighbors for each atom

fix command (1/2) --Operations

- addforce - add a force to each atom
- ave/atom - compute per-atom time-averaged quantities
- ave/time - compute/output global time-averaged quantities
- deform - change the simulation box size/shape
- deposit - add new atoms above a surface
- efield - impose electric field on system
- freeze - freeze atoms in a granular simulation
- gravity - add gravity to atoms in a granular simulation
- gcmc - grand canonical insertions/deletions
- heat - add/subtract momentum-conserving heat
- indent - impose force due to an indenter
- langevin - Langevin temperature control
- momentum - zero the linear and/or angular momentum of a group of atoms
- move - move atoms in a prescribed fashion
- neb - nudged elastic band (NEB) spring forces
- npt - constant NPT time integration via Nose/Hoover
- npt/asphere - NPT for aspherical particles
- npt/sphere - NPT for spherical particles
- nve - constant NVE time integration
- nvt - constant NVT time integration via Nose/Hoover

fix command (2/2) --Operations

nvt - constant NVT time integration via Nose/Hoover

press/berendsen - pressure control by Berendsen barostat

print - print text and variables during a simulation

restrain - constrain a bond, angle, dihedral

rigid - constrain many small clusters of atoms to move as a rigid body with NVE integration

setforce - set the force on each atom

shake - SHAKE constraints on bonds and/or angles

srd - stochastic rotation dynamics (SRD)

temp/berendsen - temperature control by Berendsen thermostat

temp/rescale - temperature control by velocity rescaling

thermal/conductivity - Muller-Plathe kinetic energy exchange for thermal conductivity calculation

viscosity - Muller-Plathe momentum exchange for viscosity calculation

viscous - viscous damping for granular simulations

wall/lj93 - Lennard-Jones 9-3 wall

wall/reflect - reflecting wall(s)

Data Visualization for Atomistic/Molecul- ar Simulations

These are high-quality visualization packages we have used and recommend.

LAMMPS can either write output files directly in an input format recognized by these programs

[VMD](#)

[AtomEye](#)

[OVITO](#)

[ParaView](#)

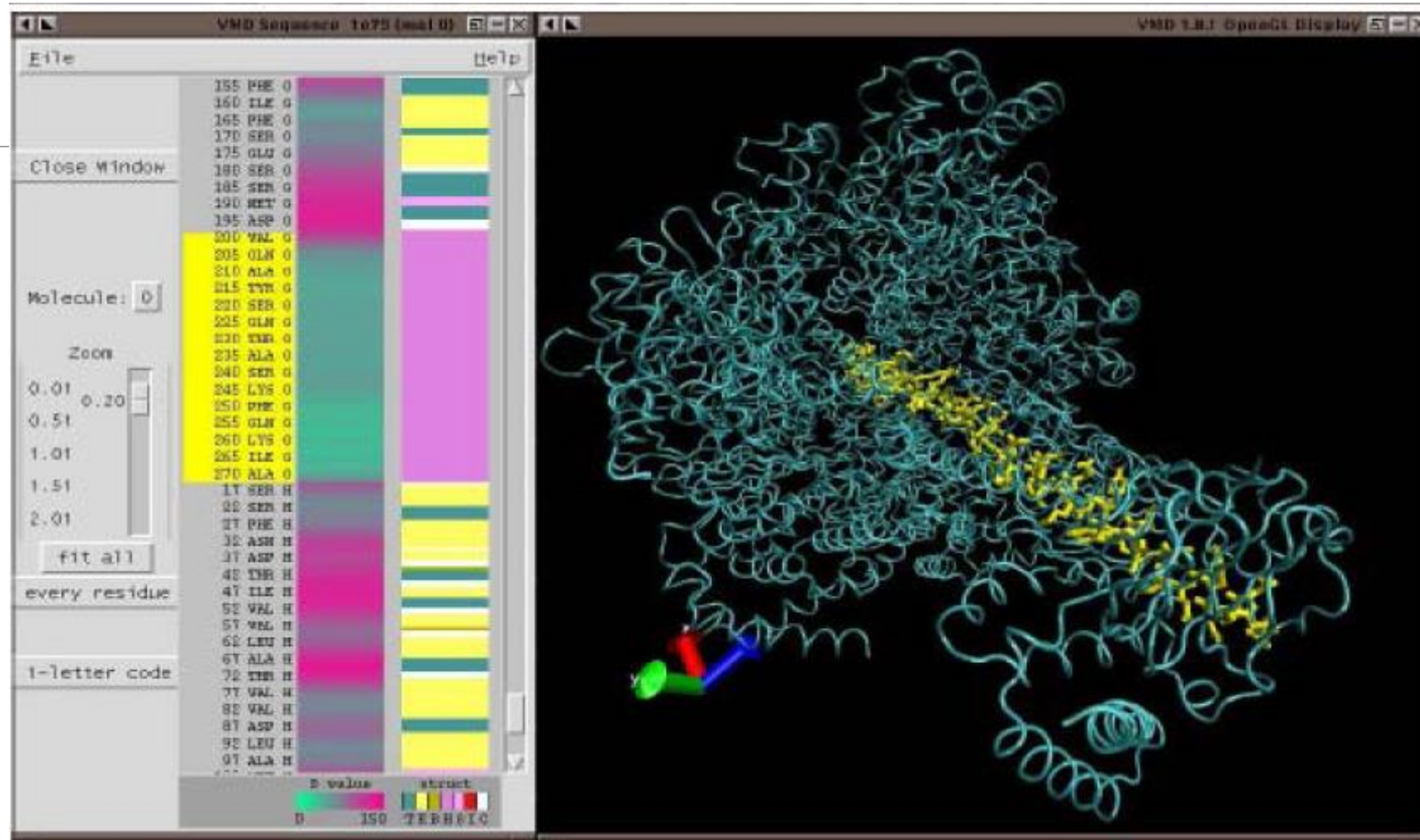
[PyMol](#)

[Raster3d](#)

[RasMol](#)

VMD

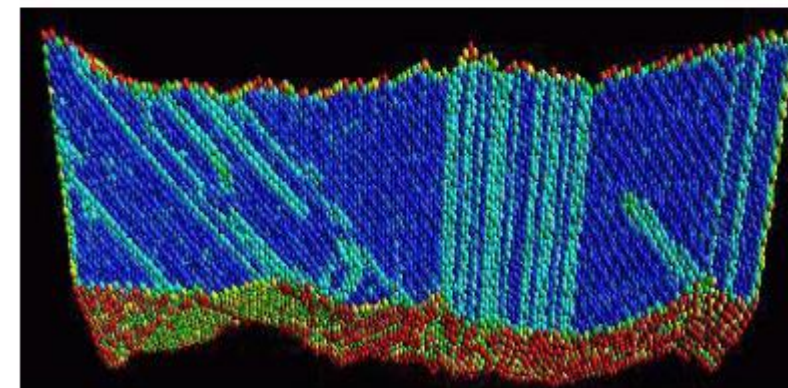
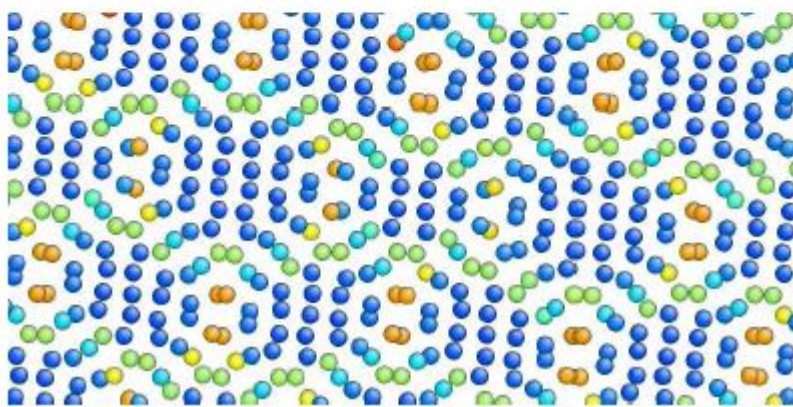
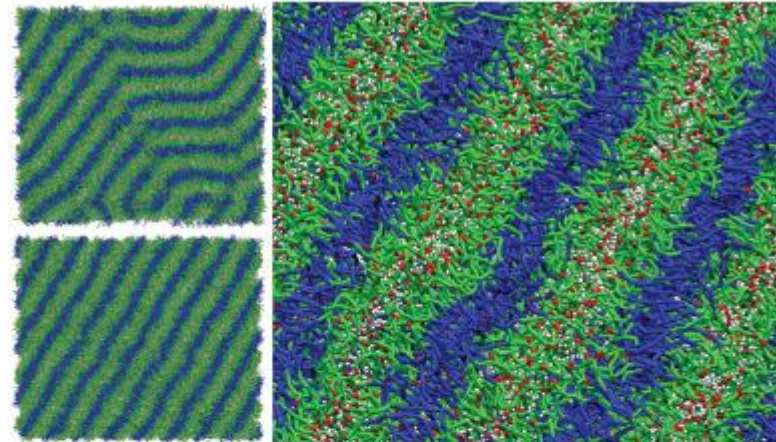
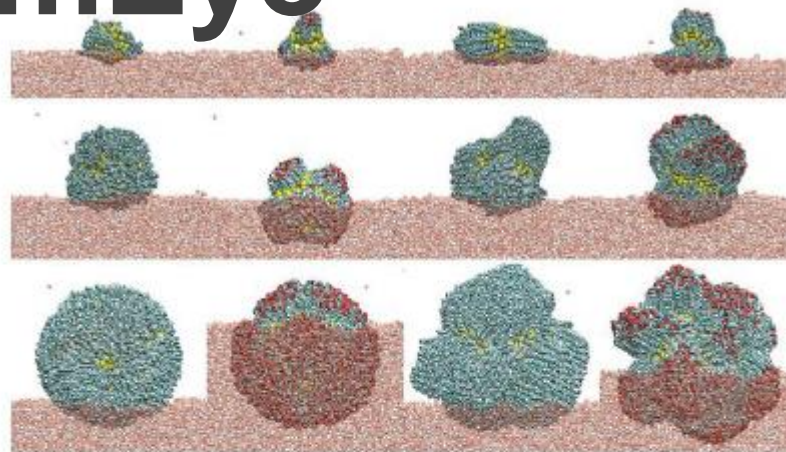
<http://www.ks.uiuc.edu/Research/vmd>



- Feature : surface, electronic density, hydrogen bond...
- Format : .pdb, mol, xyz, POSCAR, gjf ...

AtomEye

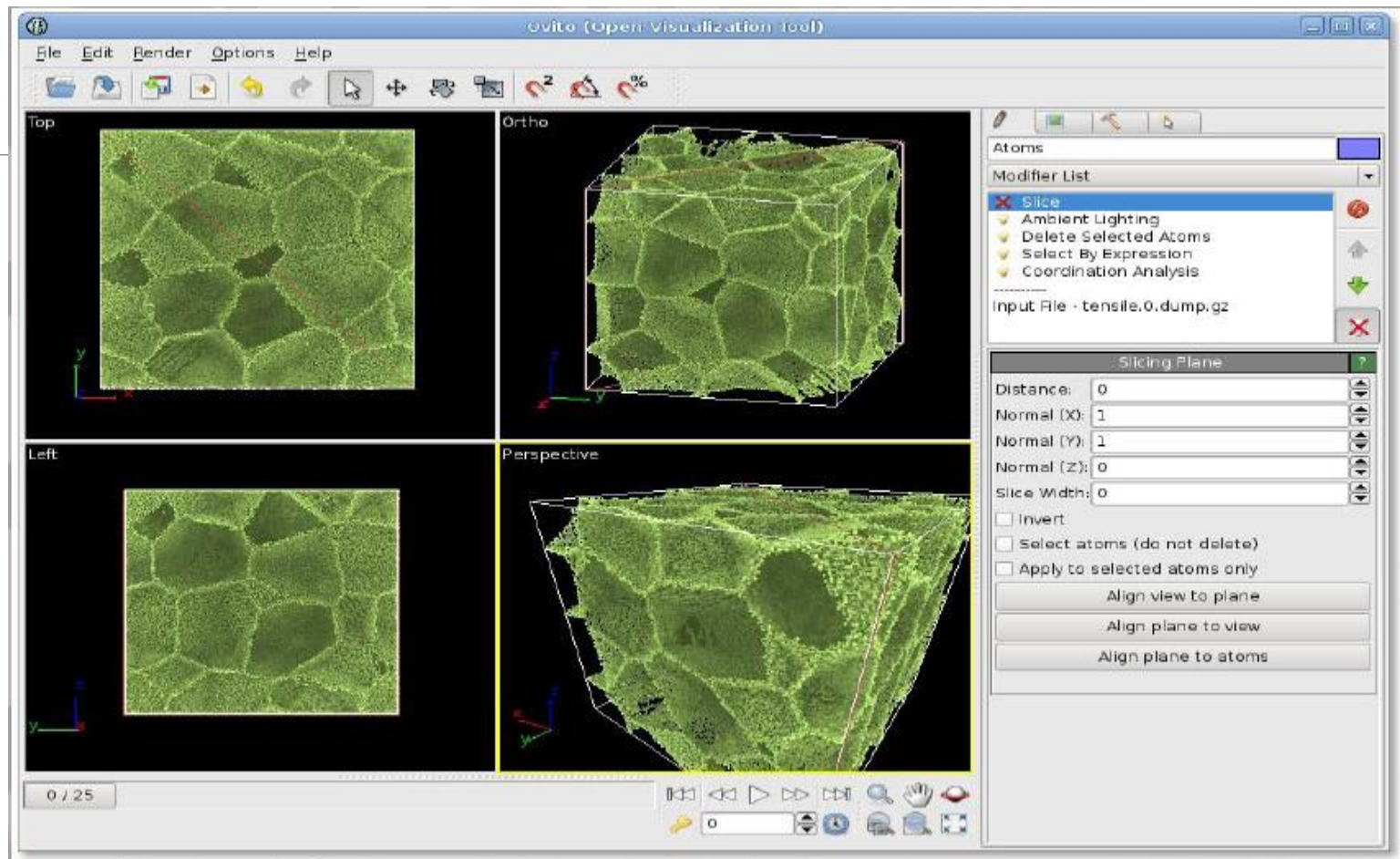
<http://li.mit.edu/Archive/Graphics/A/>



- Feature : default → coordination number, central symmetry parameter, local von Mises shear strain
- Format : cfg

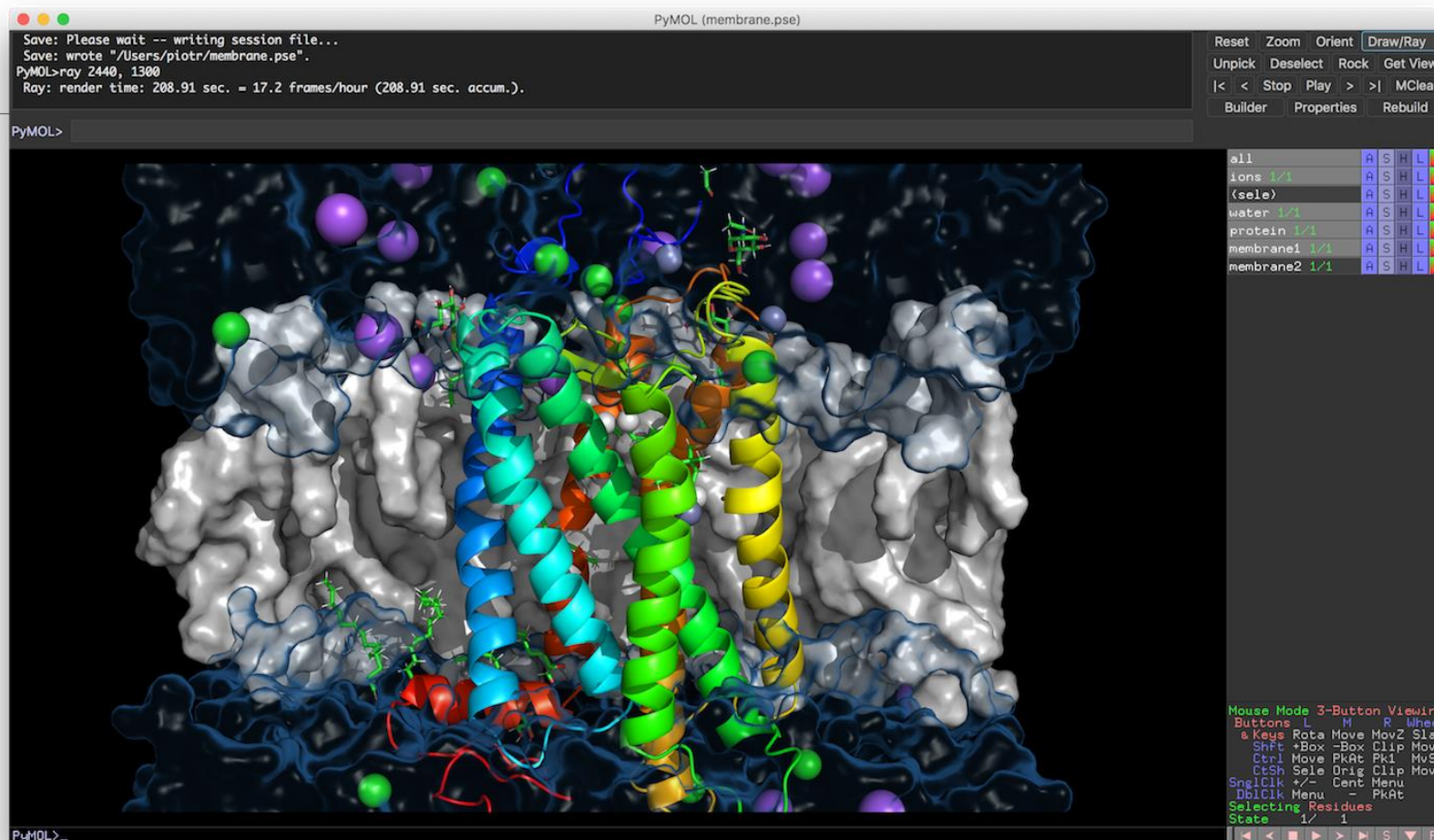
Ovito

<http://www.ovito.org/>



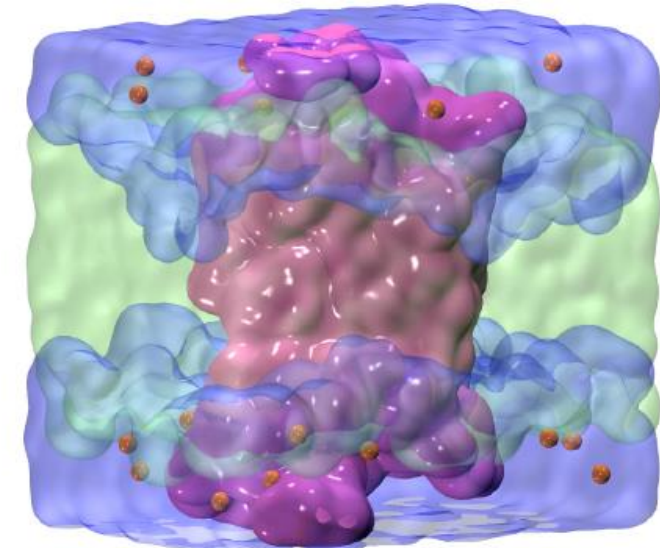
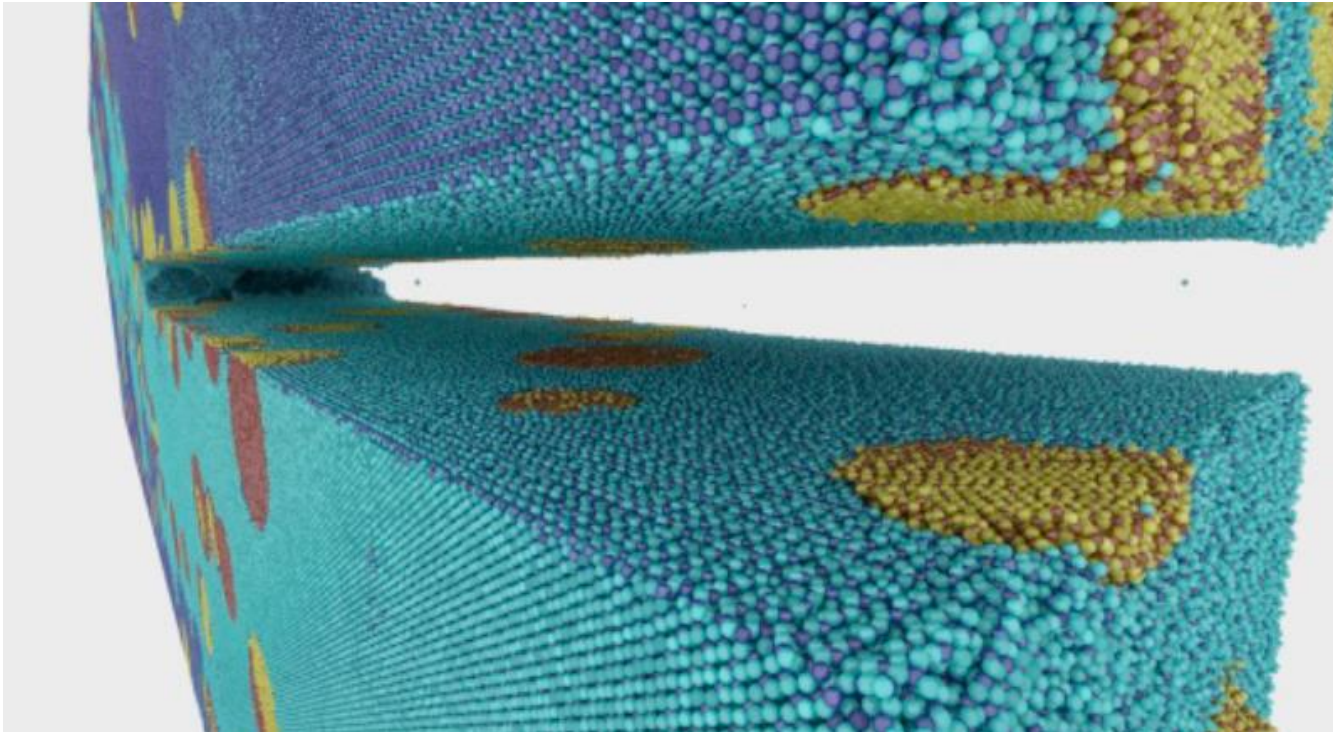
- Feature : A fast analyzing configuration viewer for crystal structures → FCC BCC HCP
- Format : xyz, cfg

Pymol



- Feature : surface, electronic density, hydrogen bond...
- Format : .pdb, mol, xyz, POSCAR, gjf ...

paraview



- Feature : Highly customized analyze tool and large scale visualization.
- Format : xyz, cfg, vtk,...

Melting-Cu

Model system Cu

Initial conditions T, P...

Supercell number of atoms, PBCs ...

Interatomic potentials EAM

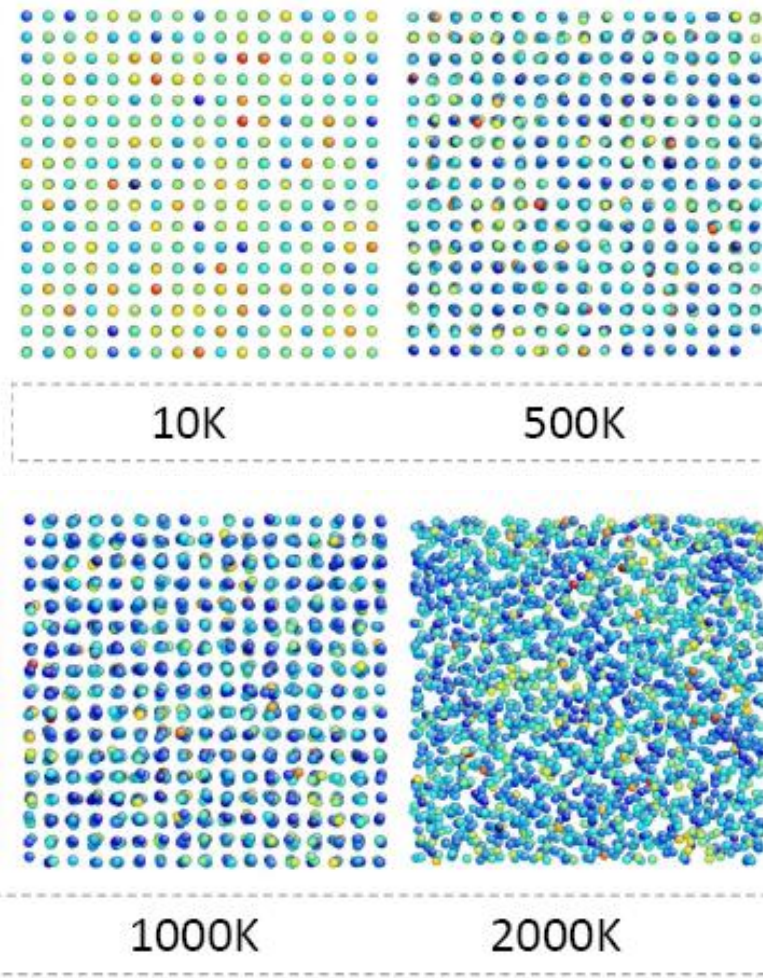
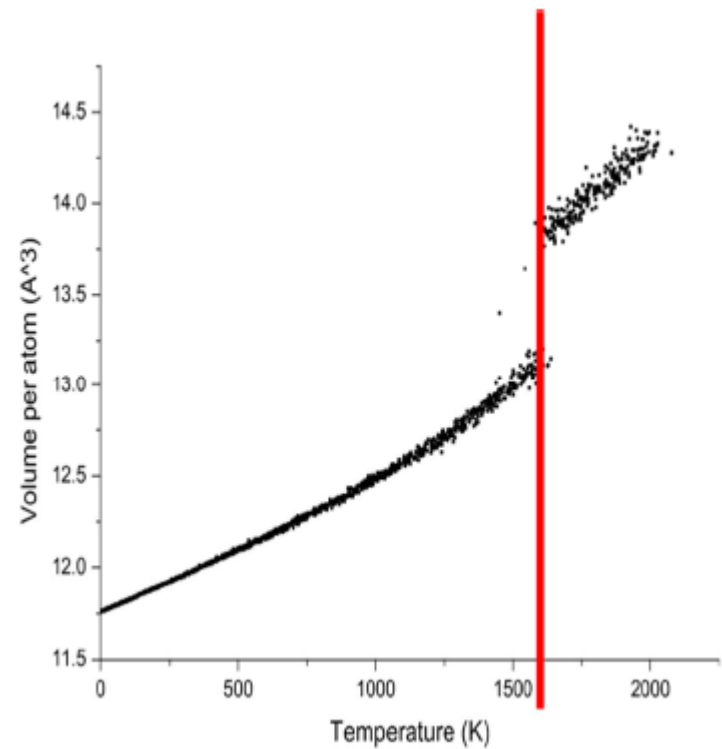
Ensembles NPT (Nosé-Hoover thermostat)

input script

Melting

```
units          metal
boundary      p p p
atom_style    atomic
lattice       fcc 3.61
region        box block 0 4 0 4 0 4
create_box    1 box
create_atoms  1 box
pair_style    eam
pair_coeff   * * Cu_u3.eam
timestep     0.01
thermo       1000
variable      N equal step
variable      pote equal pe
variable      T equal temp
variable      Press equal press
variable      V equal vol
velocity      all create 10 825577 dist gaussian
fix           extra all print 100 "${N} ${T} ${V} ${pote} ${Press}" file data
dump          1 all cfg 10000 a*.cfg id type xs ys zs c_ke
dump_modify   1 element Cu
fix           1 all npt temp 10 2000 1 iso 0 0 10
run           120000
```

若超過三組輸出參數 需使用{}



Initialization

□ units metal

設定模擬使用的單位，有metal, **lj(default)**, real, si, cgs, electron, micro, nano等選擇。

□ boundary p p p

設定模擬空間三個方向週期邊界條件，p 代表週期邊界，f 代表固定邊界，

□ atom_style atomic

設定原子的型態，有angle or **atomic(default)** or body or bond or charge or dipole or dpd or edpd or mdpd or tdpd or electron or ellipsoid or full or line or meso or molecular or peri or smd or sphere or spin or tri or template or hybrid，需要跟據模擬的需求自己選擇。

Atom definition

- lattice fcc 3.615
定義長度，方便在模擬中做度量，我們要件晶格結構，所以用lattice constant作單位很方便。
- region cubic block 0 10 0 10 0 10
定義區域，後面可以用來做其他的操作，像是選取或是建立範圍。
- create_box 1 cubic
跟據前一行建立的區域建立模擬空間，還有其他的建立方式。
- create_atoms 1 box
在前一行的模擬空間中(晶格點上)，建立原子，1代表原子種類的數目。
- mass 1 63.55
定義原子的質量，偶而在勢能函數也會有定義，但明確寫出比較好。

Atom definition

- **print \${rseed}**

從command-line option 的-v rseed xxx讀入值，作為亂數種子。

- **velocity all create 300.0 \${rseed} mom yes rot yes dist gaussian**

設定所有原子的速度，使速度滿足300K時的高斯分布，並且動量、角動量都為0。

- **pair_style eam**

設定使用eam勢能函數來描述原子間作用力，為了移植性，不明確寫出suffix，改用command-line option 的-pk -sf來控制是否使用加速套件。

- **pair_coeff 1 1 "C:\Program Files\LAMMPS 64-bit
16Aug2018\Potentials\Cu_u3.eam"**

明確指出勢能函數檔案的位置，沒有指明路徑就會在兩個地方依序搜尋1. in.xxx同一個資料夾。2. LAMMPS_POTENTIALS這個變數定義的位置。

Setting

neighbor 2.0 bin

設定鄰近表列的建立方式，2.0表示以比截斷半徑多2.0(長度單位)的長度建立表列，bin代表用長度來建立表列，nsq代表用數量來建立表列。

neigh_modify every 2 delay 20 check yes ## 20+1 step=>check

設定更新表列的頻率，every 2代表每隔2 step更新一次表列，delay 20 代表離上次更新表列超過20 steps後也強制更新，check yes 代表檢查每個原子的位移有無超過0.5 skin厚度。

timestep 0.001 #1fs

設定每次疊代的時間長度，單位是跟據先前unit的選擇。

thermo 1000

輸出熱力性質的頻率。

thermo_style custom step time temp press vol pe ke etotal enthalpy

調整想要監控的性質。

thermo_modify flush yes

強迫從記憶體中傾印到螢幕。

Setting

```
variable    a1 equal step  
variable    a2 equal time  
variable    a3 equal temp  
variable    a4 equal press  
variable    a5 equal vol  
variable    a6 equal "pe/atoms"  
variable    a7 equal "ke/atoms"  
variable    a8 equal etotal  
variable    a9 equal enthalpy  
  
fix        tdo all print 1000 "${a1} ${a2} ${a3} ${a4} ${a5} ${a6} ${a7} ${a8} ${a9}" file tdo.txt title "Step  
Time Temp Press Volume PotEng KinEng TotEng Enthalpy" screen no  
  
min_style   cg  
minimize    0.0 1.0e-6 20000 200000
```

因為thermo_style的輸出在螢幕以及
log.lammps檔案，避免手動複製處理的手續，
所以客制化輸出自己想要監控的性質。

消除一開始人為設定初始結構的artifact，
使用共軛梯度法cg，鬆弛結構。

Run a simulation

fix 1 all nve

設定nve，絕熱的thermal state。

fix 2 all temp/rescale 10 300.0 300.0 10.0 0.8

但因為數值疊代的誤差，會造成系統的能量會偏離初始狀態，使用rescale方法保持。

Run a simulation

write_dump all cfg initial.cfg mass type xs ys zs vx vy vz modify element Cu
 輸出起始的結構

dump 1 all cfg 10000 dump.*.cfg mass type xs ys zs vx vy vz fx fy fz

dump_modify 1 element Cu pad 6

在模擬過程中標準輸出檔案，ascii檔，人可以直接讀和編輯。（MDANALYSIS不能讀，需要用ovito轉成data檔）

dump 2 all xyz 10000 b*.xyz

輸出另一種檔案形式，ascii檔，人可以直接讀和編輯。（MDANALYSIS可以讀，但是缺了mass info。）

dump 3 all dcd 10000 c.dcd

輸出另一種檔案形式，binary檔，人不能直接解讀。（MDANALYSIS可以讀，ovito讀不了。）

run 100000

執行模擬

Run a simulation

unfix 1

unfix 2

解除上一段模擬所施加的operation。

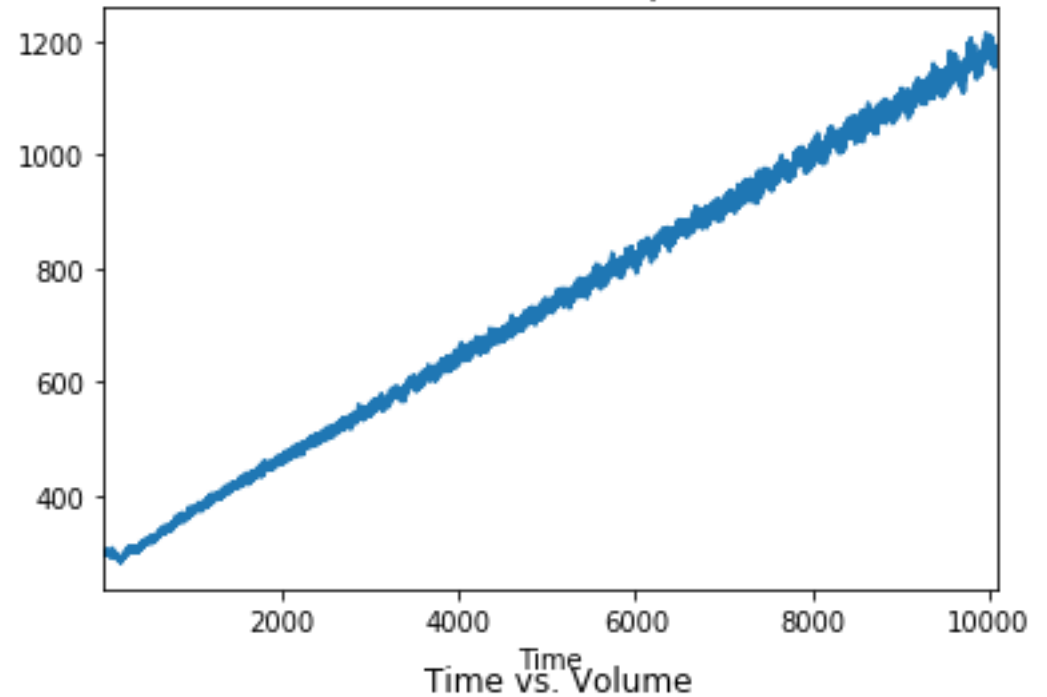
fix 3 all npt temp 300.0 1200.0 100.0 iso 1.01325 1.01325 1000.0

重新賦予系統npt的thermal state，並且從300K升溫至1200K

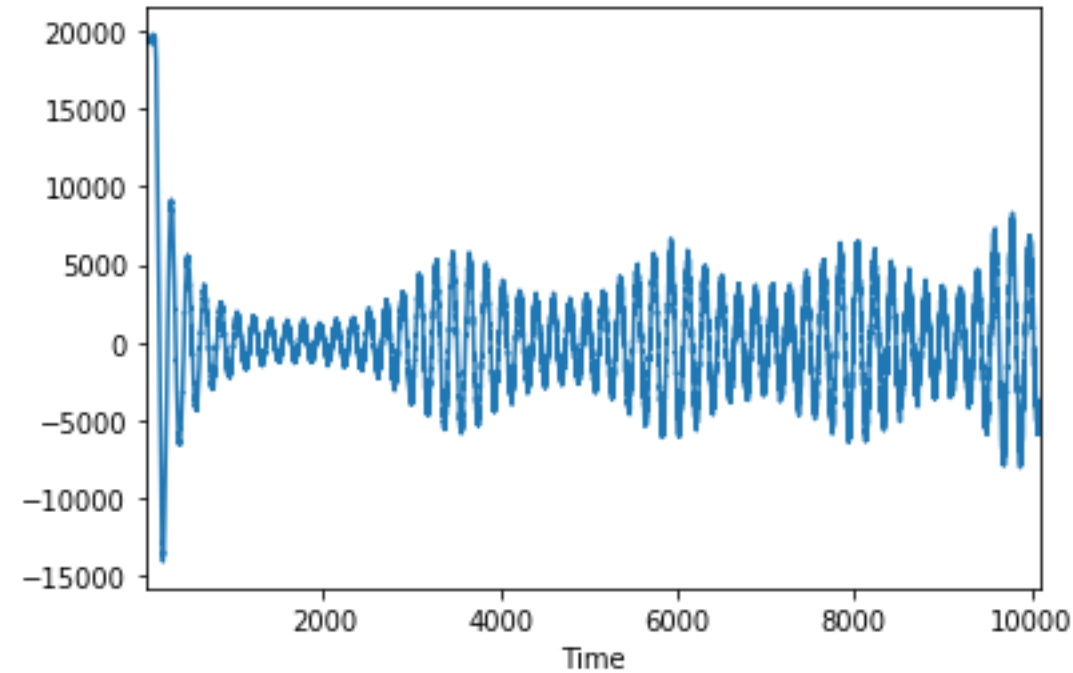
run 100000

再次執行模擬

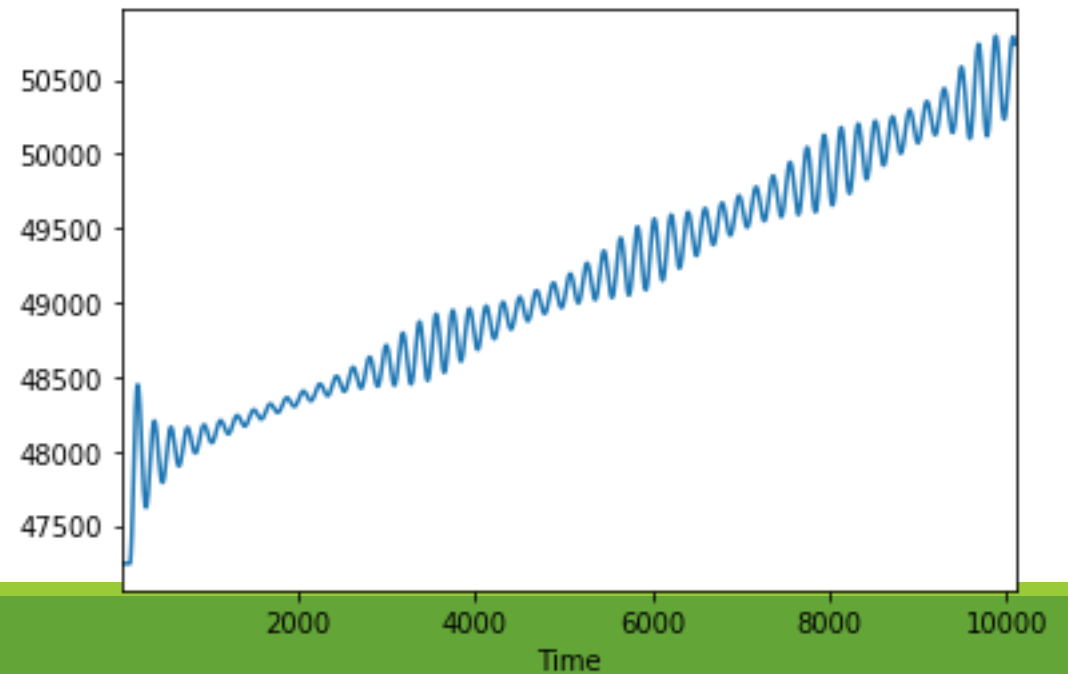
Time vs. Temp



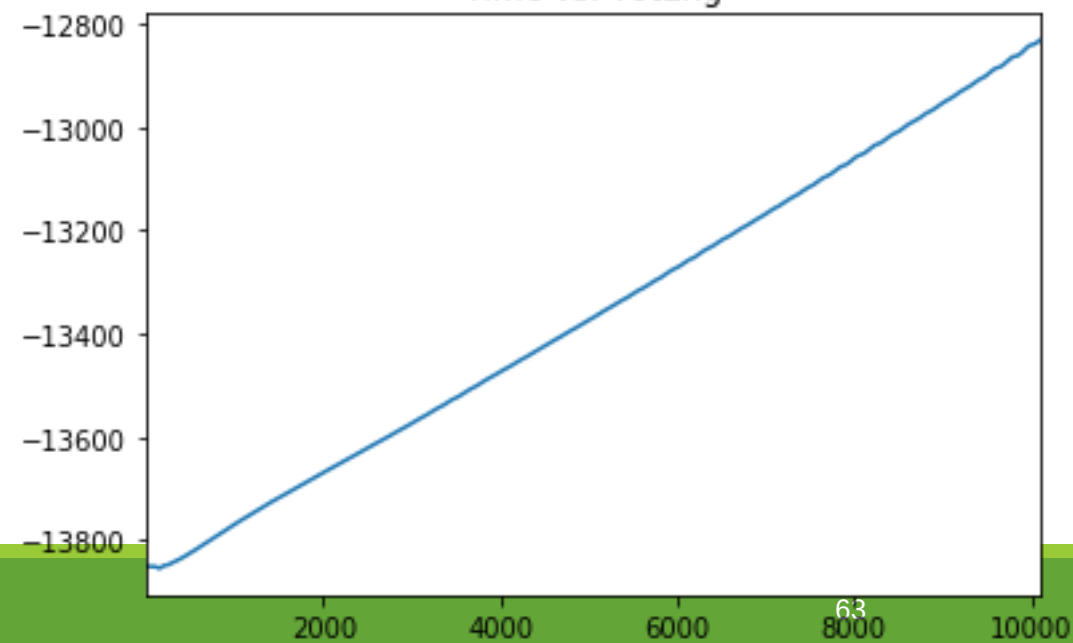
Time vs. Press



Time vs. Volume



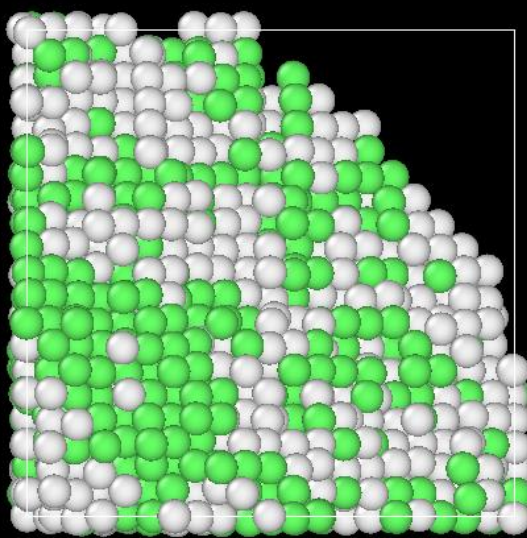
Time vs. TotEng



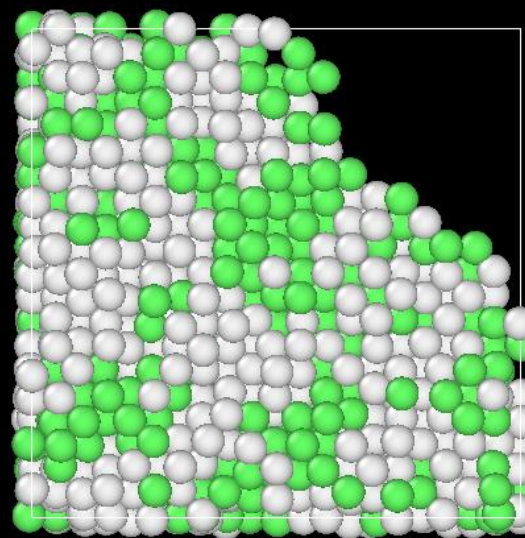
File Edit Scripting Options Help



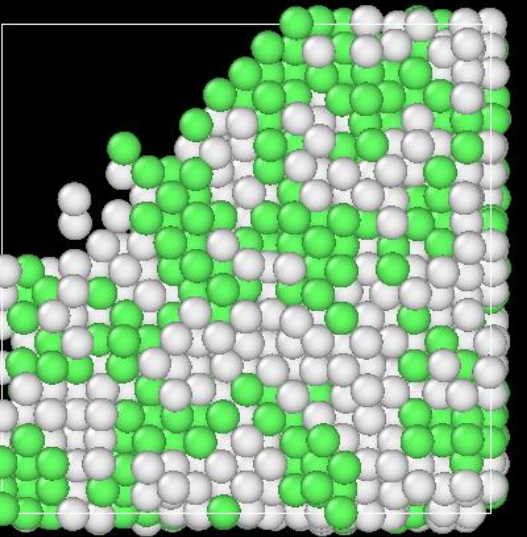
Top



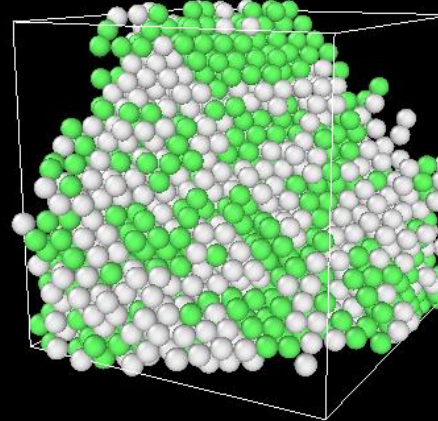
Front



Left



Perspective



0

50

100

150

200

250

300

350

400

450

500

550

600

680 / 1009

750

800

850

900

950

1000

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Add modification...

Display

 Simulation cell Particles Forces

Modifications

 Slice Common neighbor analysis Coordination analysis

Input

dump.6810000.cfg [CFG]

Simulation cell

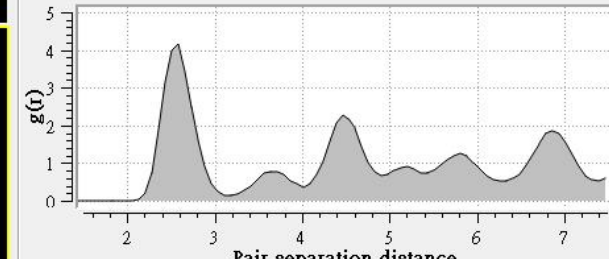
Particle types

Coordination analysis

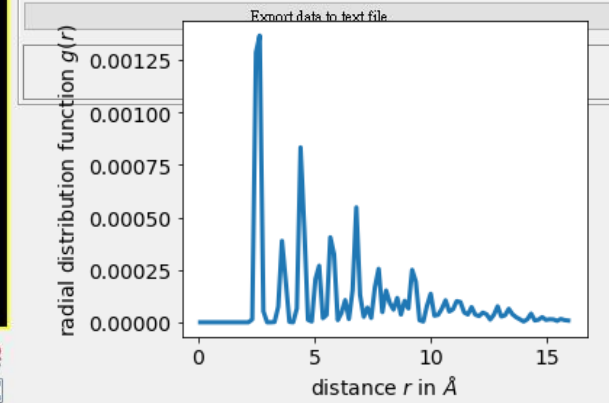
Cutoff radius: 7.5

Number of histogram bins: 100

Radial distribution function:



Pair separation distance

distance r in Å

Melting-Si

Model system Si

Initial conditions T, P...

Supercell number of atoms, PBCs ...

Interatomic potentials Tersoff

Ensembles NPT (Nosé-Hoover thermostat)

input script-1

```
units          metal
atom_style    atomic

lattice        diamond 5.431
region         cubic block 0 3 0 3 0 5
create_box     1 cubic
create_atoms   1 box
mass           1 28.0855
pair_style     tersoff
pair_coeff    * * Si.tersoff Si

velocity       all create 300.0 456236 sum yes mom
yes dist gaussian
neighbor       1 bin
neigh_modify   every 1 delay 20 check yes ## 20+1
step=>check
thermo         1000
timestep      0.002
```

input script-2

```
variable          Pot equal " pe/atoms"
variable          kee equal "ke/atoms"
variable          tstep equal "step*dt"
fix              eng all print 200 "${tstep} ${Pot} ${kee}" file Energy.dat screen no
variable          vv atom "(vx*vx+vy*vy+vz*vz)^{1/2} "
#####compute average energy/atom#####
#compute          pe_a    all pe/atom
#fix              vva all ave/atom 10 70 1000 vx vy vz
#fix              pe_a all ave/atom 10 70 1000 c_pe_a
#dump             id all custom 1000 dump.melt x y z mass f_pe_a
#####compute average energy/atom#####

fix              1 all nvt temp 300 300 1
run              500
unfix            1

#fix              1 all nve
#fix              2 all temp/rescale 10 300.0 300.0 10.0 1.0
fix              2 all npt temp 2000 4000 1 x 1 1 1 y 1 1 1
#####deform#####
#fix              3 all deform 1 z erate 0.005 units box

run              200000
```