Quantum Molecular Dynamics: Representation & Solution

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How to represent & solve Kohn-Sham equations in QMD?





Representation: Plane-Wave Basis

• Pseudopotentials result in slowly varying wave functions that can be represented on a regular grid, which in turn can be represented as a linear combination of plane waves, *i.e.*, Fourier transform



See "Fourier transform as a resolution of unity in a vector space" https://aiichironakano.github.io/phys516/03QD-slide.pdf

Numerics: Fast Fourier Transform

• O(NlogN) fast Fourier-transform (FFT) algorithm is typically used to perform Fourier transform



Periodic Solid

• Consider a periodic solid with the unit cell spanned by vectors a, b & c



• Fourier transform of a periodic function

$$u(\mathbf{r}) = \sum_{\mathbf{G}} u_{\mathbf{G}} \exp(i\mathbf{G} \cdot \mathbf{r})$$
$$\mathbf{G} = \frac{2\pi}{\mathbf{a} \cdot (\mathbf{b} \times \mathbf{c})} [l(\mathbf{b} \times \mathbf{c}), m(\mathbf{c} \times \mathbf{a}), n(\mathbf{a} \times \mathbf{b})] \quad (l, m, n \in \mathbb{Z})$$

Bloch's theorem

n: band index

$$\psi_{n\mathbf{k}}(\mathbf{r}) = \exp(i\mathbf{k} \cdot \mathbf{r}) u_{n,\mathbf{k}}(\mathbf{r})$$

 $= \sum_{\mathbf{G}} u_{n,\mathbf{k}}(\mathbf{G}) \exp(i(\mathbf{k} + \mathbf{G}) \cdot \mathbf{r})$

 $\mathbf{k} \in$ first Brillouin zone in the reciprocal space

Electronic Bands: Infinite Lattice



J. R. Chelikowsky & M. L. Cohen, Phys. Rev. B 10, 5095 ('74)

See notes on (1) plane-wave basis & (2) supercell

QMD Algorithm



Molecular Dynamics Modes

QXMD software supports various **MD** modes

Use gradient & Hessian information

- Structural optimization, *e.g.*, quasi-Newton method, see https://aiichironakano.github.io/phys760/MNK.pdf
 - > Relax atomic positions {R_I} to minimize the energy {R_I^{*}} = argmin_{{R_I}}(min_{{ $\psi_n(\mathbf{r})$ }} E[{ $\psi_n(\mathbf{r})$ }, {R_I}])
- Molecular dynamics
 - > Follow atomic trajectories by numerically integrating Newton's second law of motion

$$M \frac{d^2}{dt^2} \mathbf{R}_I = -\left\langle \Psi_0 \right| \frac{\partial h(\mathbf{r}, \mathbf{R}(t), t)}{\partial \mathbf{R}(t)} \left| \Psi_0 \right\rangle$$

> Microcanonical (NVE), canonical (NVT) & isobaric (NPT) ensembles are supported

Martyna et al., Mol. Phys. 87, 1117 ('96)

For classical molecular dynamics (*e.g.***, velocity Verlet algorithm), see** PHYS 516 (*Methods of Computational Physics*, <u>https://aiichironakano.github.io/phys516/02MD-slide.pdf</u>)</u> & MASC 575 (*Basics of Atomistic Simulation of Materials*)

Self-Consistent Field Iteration

$$\begin{pmatrix} -\frac{\hbar^2}{2m} \frac{\partial^2}{\partial \mathbf{r}^2} + \hat{V}_{\text{ion}} + \hat{V}_{\text{H,xc}}[\rho(\mathbf{r})] \end{pmatrix} \psi_n(\mathbf{r}) = \epsilon_n \psi_n(\mathbf{r})$$
Given $\rho(\mathbf{r})$,
iteratively obtain
 $\{\psi_n, \epsilon_n\}, e.g., \text{by}$
preconditioned
conjugate gradient
$$\rho(\mathbf{r}) = \sum_n |\psi_n(\mathbf{r})|^2 \Theta(\mu - \epsilon_n)$$
Chemical potential
 $N = \int d\mathbf{r}\rho(\mathbf{r})$

Self-Consistent Field Iteration



Orthogonalization by Matrix Decomposition

• Gram-Schmidt orthonormalization: The orthonormal basis set $Q = [q_1...q_m]$ is obtained starting from an arbitrary set of *m* vectors, $S = [s_1...s_m]$ as

$$\mathbf{q}_{1} = \mathbf{s}_{1} / |\mathbf{s}_{1}|$$
for $i = 2$ to m

$$\mathbf{q}'_{i} = \mathbf{s}_{i} - \sum_{j=1}^{i-1} \mathbf{q}_{j} (\mathbf{q}_{j} \cdot \mathbf{s}_{i}) \quad \frac{\text{Projection!}}{\widehat{P}} |\mathbf{s}_{i}\rangle$$

$$\mathbf{q}_{i} = \mathbf{q}'_{i} / |\mathbf{q}'_{i}| \quad \sum_{j=1}^{i-1} |\mathbf{q}_{j}\rangle \langle q_{j}|$$
endfor

• The Gram-Schmidt procedure amounts to QR decomposition, S = QR, where R is an $m \times m$ right-triangle matrix m

$$n \begin{bmatrix} m \\ s_{1} & s_{2} & s_{3} & s_{4} \end{bmatrix} = n \begin{bmatrix} q_{1} & q_{2} & q_{3} & q_{4} \end{bmatrix} \begin{bmatrix} |q_{1}'| & q_{1} \cdot s_{2} & q_{1} \cdot s_{3} & q_{1} \cdot s_{4} \\ 0 & |q_{2}'| & q_{2} \cdot s_{3} & q_{2} \cdot s_{4} \\ 0 & 0 & |q_{3}'| & q_{3} \cdot s_{4} \\ 0 & 0 & 0 & |q_{4}'| \end{bmatrix} m$$

$$\therefore \mathbf{s}_{i} = |\mathbf{q}_{i}'|\mathbf{q}_{i} + \sum_{j=1}^{i-1} \mathbf{q}_{j}(\mathbf{q}_{j} \cdot \mathbf{s}_{i})$$
Hasegawa *et al.*, *SC* ('11)

• For higher parallelization, Cholesky decomposition (BLAS3) is used instead https://aiichironakano.github.io/phys516/Cholesky.pdf

Charge Mixing

- Fixed-point charge mapping in self-consistent field iteration $\rho_{in}(\mathbf{r}) \mapsto v_{Hxc}(\mathbf{r}) \mapsto \{\psi_n(\mathbf{r})\} \mapsto \rho_{out}(\mathbf{r})$
- Directly using $\rho_{out}(\mathbf{r})$ as $\rho_{in}(\mathbf{r})$ in the next iteration step often destabilizes numerical iteration
- Charge mixing $\rho_{in}^{i} \leftarrow \sum_{j=1}^{n} \alpha_{i} \rho_{in}^{i-j}$
- Determine the mixing coefficients α_i in order to minimize the residual $R[\rho_{in}(\mathbf{r})] \equiv \rho_{out}[\rho_{in}] \rho_{in}$
- See note on Pulay charge mixing

Conjugate-Gradient Minimization of Energy Functional

i: iteration index; *n*: band index

"gradient"
$$g_n^{(i)} = -\frac{\delta E\left[\left\{\psi_n^{(i)}\right\}, \left\{\vec{R}_I(t)\right\}\right]}{\delta\psi_n^{(i)}} + \varepsilon_n^{(i)}\psi_n^{(i)} \equiv -H\psi_n^{(i)} + \varepsilon_n^{(i)}\psi_n^{(i)}}{\varepsilon_n^{(i)}} = \int d^3r\psi_n^{(i)*}H\psi_n^{(i)}}$$

"preconditioning" $\tilde{g}_n^{(i)} = \hat{P}g_n^{(i)}$
"conjugate gradient" $h_n^{(i)} = \tilde{g}_n^{(i)} + \beta h_n^{(i-1)}, \beta = \int d^3rg_n^{(i)} \cdot g_n^{(i)} / \int d^3rg_n^{(i-1)} \cdot g_n^{(i-1)}}$
"new wave function" $\psi_n^{(i+1)} = C(\lambda) \left(\psi_n^{(i)} + \lambda h_n^{(i)}\right)$
with constraint $\int d^3r\psi_n^{(i+1)*}\psi_m = 0 \quad (m \le n)$
 $i \leftarrow i + 1$ if $|\varepsilon_n^{(i+1)} - \varepsilon_n^{(i)}| > \varepsilon$

See lecture on iterative minimization (https://aiichironakano.github.io/phys516/QD2CG.pdf) & notes on (1) conjugate-gradient (CG) method, (2) CG electronic-state solver, (3) CG DFT solver & (4) 2D electron example

Real-Space Grid as a Basis

- Wave functions & electron density are represented by numerical values on real-space grid points
- Finite difference expansion for the kinetic-energy operator

$$\frac{\partial^2 \psi_n}{\partial x^2} \bigg|_{\mathbf{r}_{ijk} = (x_i, y_j, z_k)} = \sum_{l=-L}^{L} C_l \psi_n(x_i + lh, y_j, z_k) + O(h^{2L+2})$$

(short-ranged operation)

The calculations are performed completely in "real space"

- Suitable for systems with vacuum (e.g., clusters, surfaces)
- Efficient implementation on parallel computers

J.R. Chelikowsky, N. Troullier & Y.Saad, Phys. Rev. Lett. 72, 1240 ('94)

Acceleration of Convergence

Preconditioning $(H - \varepsilon)(\psi + \delta) = 0$ approximately **Enhanced convergence rate of short** $\frac{-)(H-\varepsilon)\psi = -g}{(H-\varepsilon)\delta = g} \stackrel{\text{invert?}}{\Rightarrow} \delta \leftarrow (H-\varepsilon)^{-1}g$ wavelength components of the residual $\tilde{g}_n(x_i, y_i, z_k) = \hat{P}g_n^{(i)}$ $= \sum_{l_1=-1}^{1} \sum_{l_2=-1}^{1} \sum_{l_2=-1}^{1} c_{l_1 l_2 l_3} g_n (x_i + l_1 h, y_j + l_2 h, z_k + l_3 h)$ Multigrid method [Brandt '77, Bernholc et al. '96, Beck, '00] To reduce long wavelength components of the residual, l = 2

$$\begin{pmatrix} -\frac{\hbar^2}{2m}\nabla^2 + V \end{pmatrix} \varphi = g_n^{(i)} \\ \psi_n^{(i)} \leftarrow \psi_n^{(i)} + \varphi$$

on a coarse grid



/= 1

Iterative Solution of Linear Systems



Multigrid Method



Parallel Multigrid Method

Domain decomposition with boundary-layer caching



2D computational & communication costs (isogranular or weak scaling)

 $N \times N$ grids each on $P \times P$ processors: $T(N^2 P^2, P^2) = a \log NP + bN + cN^2$ Speedup $S_{P^2} = \frac{N^2 P^2 T (N^2, 1)}{N^2 T (N^2, P^2)} = \frac{P^2 (cN^2)}{a \log NP + bN + cN^2} = \frac{P^2}{1 + \frac{b}{cN} + \frac{a}{cN} \log NP}$ Parallel efficiency $E_{P^2} = \frac{S_{P^2}}{P^2} = \frac{1}{1 + \frac{b}{cN} + \frac{a}{cN} \log NP}$ Highly scalable!

Nakano et al., Comput. Phys. Commun. 83, 181 ('94)

For the definition of parallel efficiency, see https://aiichironakano.github.io/cs596/MPI-Pi.pdf

Global Communications



See note on multigrid preconditioned CG

Real-Space DFT on Hierarchical Grids

Efficient parallelization of DFT: real-space approaches

- High-order finite difference [Chelikowsky, Troullier, Saad, '94]
- Multigrid acceleration [Bernholc et al., '96; Beck, '00]
- Double-grid method [Ono, Hirose, '99] ~ obsolete, with PAW
- Spatial decomposition/divide-&-conquer

Multigrid acceleration of preconditioned conjugate gradient

F. Shimojo *et al.*, "Embedded divide-and-conquer algorithm on hierarchicalreal-space grids," *Comput. Phys. Commun.* **167**, 151 ('05)

Adaptive high-resolution grid for atomic pseudopotentials

Hierarchical Computing



cf. globally- sparse-yet-locally-dense eigensolver [J. H. Lam et al., Nature Commun. 15, 3479 ('24)]

• Hierarchical band (*i.e.*, Kohn-Sham orbital) + space + domain (BSD) decomposition



Divide-&-conquer domains

Globally-Sparse Yet Locally-Dense Eigensolver



Iterative Krylov-subspace eigensolver

• 250-fold speed-up over state-of-the-art for 2.4M atom molecular vibrational modes

J. H. Lam et al., Nature Commun. 15, 3479 ('24)

Finite-Element DFT

- DFT calculation using a higher-order adaptive spectral finite-element (FE) basis outperforms that with the plane-wave basis for larger (*e.g.*, > 10,000 electrons) systems: see DFT-FE code [S. Das *et al.*, *Comput. Phys. Commun.* 280, 108473 ('22); https://github.com/dftfeDevelopers/dftfe]
- 2023 Gordon-Bell award: 659.7 Pflop/s (43.1% of the peak) by the DFT-FE code for 619,124 electrons on 8,000 GPU nodes of the Frontier supercomputer [S. Das *et al.*, *Proc. Supercomputing*, SC ('23)]

