第35回CMDワークショップ

第一原理計算の基礎

大阪大学 大学院基礎工学研究科 物質創成専攻 未来物質領域

草部 浩一

Concept: Motion of electrons

An electron moves as a wave, found to be a particle.

Interference of electron wave

Why are these pictures different?

An imaginative picture of the electron

r

Position : Momentum :

4

Both determined at a time *t*

An electron as a wave, which is in 'a scattering state'.

Particle position at the time *t* is not determined to be any specific state at *r*. The state with 'a group velocity' is composed of a superposition of '*p* states'.

Direct imaging of "electron wave"

A. Hashimoto, K. Suenaga, A. Gloter, K. Urita & S. Iijima, *Nature* **430**, 870 - 873 (2004).

 0.74 nm 2 nm 0.21 nm

Cf. M. Ziatdinov, et al., *Phys. Rev. B* **89**, 155405-1-15 (2014).

Therefore, we recognize "existence of the electron wave". > 5

Concepts in QED

- I am rather delighted that we must resort to such peculiar rules and strange reasoning in order to understand Nature, and I enjoy telling people about it.
- There is no "wheels and gears" beneath this analysis of Nature; if you want to understand *Her*, this is what you have to take.

"QED THE STRANGE THEORY OF LIGHT AND MATTER" Richard P. Feynman (Princeton Univ. Press, 1985.)

- The coordinates $(x, y, z), \qquad \sigma = \begin{pmatrix} 1 \\ 0 \end{pmatrix}, \text{ or } \begin{pmatrix} 0 \\ 1 \end{pmatrix}.$ $\begin{array}{c} \hline \hline \hline \hline \hline \end{array}$ α , or $0'$ 1) WWW $(\sigma = \begin{array}{cc} \cdot & \cdot & \cdot \\ \cdot & \cdot & \cdot \\ \cdot & \cdot & \cdot \end{array})$, or $\begin{array}{cc} \cdot & \cdot & \cdot \\ \cdot & \cdot & \cdot \\ \cdot & \cdot & \cdot \end{array}$. **SANA SANA** 1999999 88888 (1) (0) \vert , or **SANA SANA** 18888 $\begin{pmatrix} 0 \\ 0 \end{pmatrix}$, c (0) $\langle 1 \rangle$ $\mathbf{r} = (x, y, z), \qquad \sigma = \begin{bmatrix} 1 \\ 2 \end{bmatrix}, \text{ or }$ $(\mathbf{r}_1, \sigma_1, \mathbf{r}_2, \sigma_2, \ldots, \mathbf{r}_N, \sigma_N) = (x_1, \ldots, x_N).$
- Wavefunction : an amplitude by projection $\Psi(\mathbf{r}) = \langle \mathbf{r} | \cdot | \Psi \rangle = \langle \mathbf{r} | \Psi \rangle,$ $\Psi(\mathbf{r}_{1}, \sigma_{1}, \mathbf{r}_{2}, \sigma_{2}, \ldots, \mathbf{r}_{N}, \sigma_{N})$ $=\langle \mathbf{r}_{1}, \sigma_{1}, \mathbf{r}_{2}, \sigma_{2}, \ldots, \mathbf{r}_{N}, \sigma_{N} | \Psi \rangle.$ $\ket{\Psi}$ State vector

7

 $\vert{\bf r}\rangle$

The state may be measured by the coordinates.

What is $|\Psi\rangle$? $1 \sim$ 1 $\left[\uparrow\uparrow\downarrow\downarrow\right]$ $\langle \Psi_{S=0} \rangle = \frac{1}{\sqrt{2}} |\!| \uparrow \rangle_{1} \otimes |\downarrow \rangle_{2} - |\downarrow \rangle_{1} \otimes |\uparrow \rangle_{2} = \frac{1}{\sqrt{2}} |\!| \uparrow \downarrow \rangle - |\downarrow \uparrow \rangle.$ $\left|\left|\uparrow\right\rangle_{1}\otimes\left|\downarrow\right\rangle_{2}-\left|\downarrow\right\rangle_{1}\otimes\left|\uparrow\right\rangle_{2}=-\frac{1}{\sqrt{2}}\left|\left|\uparrow\downarrow\right\rangle-\left|\downarrow\uparrow\right\rangle\right].$ $2^{\frac{1}{2}}$

$\Psi_{S=0}$ by

Prof. P.A.M. Dirac

"A vector" !

is with ^a name "Ψ" added to the symbol.

Experimentally, we only see 'the action' as amplitudes.

When a time coordinate is fixed for a system, a Hamiltonian operator is determined by 'the gauge fixing'.

Separation of degrees of freedom

Based on difference in the mass, separation of the quantum mechanical motion is done. (the error scales as (*m*/*M*) 6/4.) *Cf. S. Takahashi, K. Takatsuka, J. Chem. Phys. 124, 144101 (2006).*

$$
\{\hat{H}_{el} + \hat{H}_{el-ion} + \hat{H}_{ion-ion}\} \Psi \rangle \otimes |\Phi_{ion}(\{\mathbf{R}_{I}\})\rangle
$$
\n
$$
= \{E + E_{ion-ion}\} \Psi \rangle \otimes |\Phi_{ion}(\{\mathbf{R}_{I}\})\rangle.
$$
\n
$$
= \{E + E_{ion-ion}\} \Psi \rangle \otimes |\Phi_{ion}(\{\mathbf{R}_{I}\})\rangle.
$$
\n
$$
m: electron mass
$$

• Existence of stable materials allows us to fix positions (or "distribution") of the mass center. Then, we may also make use of "existence of the order" in the material *a priori*. $\langle H_{el} + H_{el-ion} + H_{ion-ion} | \Psi \rangle \otimes | \Phi_{ion} (\{ \mathbf{R}_I \}) \rangle$

= { $E + E_{ion-ion} | \Psi \rangle \otimes | \Phi_{ion} (\{ \mathbf{R}_I \}) \rangle$. muclear mass

Existence of stable materials allows us to fix positions (or

"distribution") of the mass center. Then, we may also m

The order is given by "the charge distribution of each species". In relativistic form, it is by "distribution of the 4-current".

Born-Oppenheimer approximation

 $\mathbf{(r)} = \sum_{I=1}^{\infty} \frac{\mathbf{Z}_I}{|\mathbf{r} - \mathbf{r}|}$ *Nion* $I=1$ **I** \uparrow **I** I *I ext* $Z_i e$ *v* 1 2 **r R** and define the external potential $v_{ext}(\mathbf{r}) = \sum_{k=1}^{\infty} \frac{Z_{ik}}{|\mathbf{r}-\mathbf{R}|^2}$ for the electrons, we have, Multiply $\langle \{ \mathbf{R}_I \} |$ to the Schrödinger equation from the left,

$$
\hat{H}|\Psi\rangle
$$
\nHere, this operator is a two-component Fermion field operator.
\n
$$
= \left[-\sum_{\sigma} \int d^3r \left(\frac{\hbar^2}{2m} \right) \hat{\psi}_{\sigma}^{\dagger}(\mathbf{r}) \nabla^2 \hat{\psi}_{\sigma}(\mathbf{r}) + \int \int d^3r d^3r' \frac{e^2 : \hat{n}(\mathbf{r}) \hat{n}(\mathbf{r})}{|\mathbf{r} - \mathbf{r}|} + \int d^3r v_{ext}(\mathbf{r}) \hat{n}(\mathbf{r}) \right] |\Psi\rangle
$$
\n
$$
= E|\Psi\rangle.
$$
\nThe Coulomb Kernel is fixed as the photon propagator in a steady state.

Therefore, in a "well-defined" precise description of the quantum mechanical motion of electron, we may start from the electron Hamiltonian in the equation above.

Hamiltonian

- Static state of electron system:
	- Represented by potential:

$$
v_{ext}(\mathbf{r}_i) \qquad V(\mathbf{r}_i, \mathbf{r}_j) = \frac{e^2}{|\mathbf{r}_i - \mathbf{r}_j|}.
$$

$$
\frac{1}{\frac{1}{r_i}}
$$

- Cf: Ehrenfest's theorem $\langle \Psi | \mathbf{p} | \Psi \rangle$ = $\langle \Psi | -\nabla v_{ext}(\mathbf{r}) | \Psi \rangle$ *dt d*
- The Hamiltonian
	- Velocities of electrons < the velocity of light,
	- In a steady material, internal gauge field behaves as the Coulomb potential.

$$
H=-\sum_{i=1}^N\left(\frac{\hbar^2}{2m}\right)\nabla^2_{\mathbf{r}_i}+\sum_{\langle i,j\rangle}\frac{e^2}{\left|\mathbf{r}_i-\mathbf{r}_j\right|}+\sum_{i=1}^Nv_{ext}(\mathbf{r}_i).
$$

Electrons in a potential

Therefore, charge density should be determined.

Schrödinger equation : partial differential equation

Quantum model		
Quantum model	mechanical	calculation
$H\Psi(\{\mathbf{r}_i, \sigma_i\}) = -\sum_{i=1}^{N} \left(\frac{\hbar^2}{2m}\right) \nabla_x^2 \Psi + \sum_{i=1} \frac{e^2}{ \mathbf{r}_i - \mathbf{r}_j } \Psi + \sum_{i=1}^{N} \nu_{\text{ext}}(\mathbf{r}_i) \Psi = E\Psi(\{\mathbf{r}_i, \sigma_i\}).$		
<i>We</i> need to employ "an efficient (fast) calculation method" equivalent to solver of the differential equation.		
From Schrödinger picture (Wave equation)		
From Schrödinger picture (Matrix equation or operator formalism)		
From Differential equation (Function of coordinates)		
from Differential equation (Function of coordinates)		
to Green's function method (Inverse operator or resolvent)		
or realize one of them, however, we need, at least and at any time,		
a set of well-converged basis functions " to have an efficient expansion!		
This part consists from purely mathematical and practically computational problems.		

We need to employ "an efficient (fast) calculation method" equivalent to a solver of the differential equation.

From Schrödinger picture (Wave equation) to Heisenberg picture (Matrix equation or operator formalism)

From Differential equation (Function of coordinates) to Green's function method (Inverse operator or resolvent)

To realize one of them, however, we need, at least and at any time, "a set of well-converged basis functions" to have an efficient expansion!

> This part consists from purely mathematical and practically computational problems.

Modern calculation methods

The density determining "order"

- Energy as a functional of v_{ext} : $E[v_{\text{ext}}]$.
- A functional derivative:

$$
E[v_{ext} + \delta v_{ext}] - E[v_{ext}] = \int d^3r \frac{\delta E[v_{ext}]}{\delta v_{ext}(\mathbf{r})} \delta v_{ext}(\mathbf{r}) + O(\delta^2)
$$

= $\int d^3r \delta v_{ext}(\mathbf{r}) n(\mathbf{r}) + O(\delta^2)$ (1)
• The order parameter : $n(\mathbf{r})$.
K. Kusakabe and I. Maruyama, J. Phys. A: Math. Theor. 44 (2011) 135305.

• The order parameter : *n*(**r**).

A theorem in quantum mechanics

- Consider two potentials, $v_1(r)$ and $v_2(r)$, which are different more than a constant from each other.
- Let $|\Psi_1\rangle$ and $|\Psi_2\rangle$ be an eigen state of a system by v_1 (*r*) and that by v_2 (*r*), respectively.
- Then, $|\Psi_1\rangle$ cannot be an eigen state of $v_2(r)$ and vice versa.

Proof: The unique continuation theorem for the Poisson equation tells us that the statement above is correct. See the next page.

Note: you can find that the next proof is easily extended for a wider class of Hamiltonians appearing in a model space.

- When our system is treated by the Born-Oppenheimer approximation, nuclei is treated as fixed point charges.
- v_1 and v_2 follows the Poisson equation.
- Suppose that ψ is a solution of two Schrödinger equation given by v_1 and v_2 , where v_1 and v_2 are different from each other more than a constant.
- We have $v_1 \psi = (v_2 + const.) \psi$.
- Let *N* be the number of electrons.
- We may find an open set *S* in \mathbb{R}^3 , where

$$
\psi(\mathbf{r}_1+\mathbf{r},\sigma_1,\mathbf{r}_2,\sigma_2,\cdots,\mathbf{r}_N,\sigma_N)\neq 0
$$

for $r \in S$.

- In *S*, we have $v_1(r) = v_2(r) + \text{const.}$
- By the unique continuation theorem, we conclude that $v_1(r) = v_2(r)$ $+$ const. in \mathbb{R}^3 .
- This conclusion contradicts to the assumption.

Hohenberg-Kohn's theorem

• By the last theorem, we have a next inequality which is a contradiction.

$$
E_1 = \langle \Psi_1 | T + V_{ee} | \Psi_1 \rangle + \int d^3 r n v_1
$$

\n
$$
= \langle \Psi_1 | H_1 | \Psi_1 \rangle < \langle \Psi_2 | H_1 | \Psi_2 \rangle
$$

\n
$$
= \langle \Psi_2 | T + V_{ee} | \Psi_2 \rangle + \int d^3 r n v_1
$$

\n
$$
= \langle \Psi_2 | T + V_{ee} | \Psi_2 \rangle + \int d^3 r n v_2 + \int d^3 r n (v_1 - v_2)
$$

\n
$$
= \langle \Psi_2 | H_2 | \Psi_2 \rangle + \int d^3 r n (v_1 - v_2)
$$

\n
$$
< \langle \Psi_1 | H_2 | \Psi_1 \rangle + \int d^3 r n (v_1 - v_2)
$$

\n
$$
= \langle \Psi_1 | T + V_{ee} | \Psi_1 \rangle + \int d^3 r n v_2 + \int d^3 r n (v_1 - v_2)
$$

\n
$$
= \langle \Psi_1 | H_1 | \Psi_1 \rangle = E_1.
$$

• This implies existence of "one-to-one correspondence between the potential and the electron density". (by Hohenberg and Kohn (1964))

A logic tells physics

- Fixed positions of nuclei give a scalar potential *vⁱ* .
- When $v_2 \neq v_1 + \text{const.}, \Psi_1 \neq \Psi_2$, where Ψ_i is a ground state for a problem by v_i . (by the unique continuation theorem.)
- In this case, $n_1 \neq n_2$. (Proof by Hohenberg & Kohn.) Namely,
	- If " $n_1 = n_2 = n$ " and " $\Psi_1 \neq \Psi_2$ ", then $(v_1 - v_2) = \langle \Psi_2 | H_2 | \Psi_2 \rangle + | d^3 r n (v_1 - v_2)$ $(v_1 - v_2) = \langle \Psi_1 | T + V_{ee} | \Psi_1 \rangle + | d^3 r n v_2 + | d^3 r n (v_1 - v_2) = \langle \Psi_1 | H_1 | \Psi_1 \rangle = E_1.$ *Thus*, $E_1 < E_1$, which *is a contradiction.* . $3 \cdot \sqrt{2}$ $2\sqrt{u}$ $3 \cdot \cdot \cdot$ 1 v_2 $\langle 1_1 | 1_$ $v_{ee} | 1_1 / 1_$ u_{ee} $3 \cdot \sqrt{2}$ $1 < \langle \Psi_1 | H_2 | \Psi_1 \rangle + \int d^3 r n (v_1 - v_2) = \langle \Psi_1 | T + V_{ee} | \Psi_1 \rangle + \int d^3 r n v_2 + \int d^3 r n (v_1 - v_2) = \langle \Psi_1 | H_1 | \Psi_1 \rangle = E_1.$ $3 \cdot \sqrt{1}$ $1 \quad \sqrt{2}$ $\sqrt{2}$ $\sqrt{2}$ $\sqrt{2}$ $\sqrt{2}$ $3 \cdot \sqrt{2}$ $2 \sqrt{u}$ $3 \cdot \cdot \cdot$ $=\langle \Psi_2 | T + V_{ee} | \Psi_2 \rangle + \int d^3 r n v_2 + \int d^3 r n (v_1 - v_2) = \langle \Psi_2 | H_2 | \Psi_2 \rangle + \int d^3 r n (v_1 - v_2)$ 1/1/1/1/1 $3 \cdot \cdot \cdot$ $1 - \left(\frac{1}{1} |I_1| \right) \left(\frac{1}{1} \right) \left(\frac{1}{2} |I_1| \right) \left(\frac{1}{2} - \left(\frac{1}{2} |I_1| \right) \right)$ $3 \dots$ $\mathbf{1}$ $\mathbf{1}$ $\mathbf{1}$ $\mathbf{1}$ \mathbf{v}_{ee} $\mathbf{1}$ $\mathbf{1}$ \mathbf{u} $E_1 = \langle \Psi_1 | T + V_{ee} | \Psi_1 \rangle + \int d^3 r n v_1 = \langle \Psi_1 | H_1 | \Psi_1 \rangle < \langle \Psi_2 | H_1 | \Psi_2 \rangle = \langle \Psi_2 | T + V_{ee} | \Psi_2 \rangle + \int d^3 r n v_1$
= $\langle \Psi_2 | T + V_{ee} | \Psi_2 \rangle + \int d^3 r n v_2 + \int d^3 r n (v_1 - v_2) = \langle \Psi_2 | H_2 | \Psi_2 \rangle + \int d^3 r n (v_1 - v_2)$ $=\langle \Psi_1|T+V_{ee}|\Psi_1\rangle+\int d^3r n v_1=\langle \Psi_1|H_1|\Psi_1\rangle<\langle \Psi_2|H_1|\Psi_2\rangle=\langle \Psi_2|T+V_{ee}|\Psi_2\rangle+\int d^3r n v_1$

- Thus, we have a one to one correspondence between *v* and *n*.
- Namely, *n* gives *v*, the Hamiltonian, and the wave function of the ground state Ψ with energy $E[n]$.

Idea by Kohn and Sham

• Density as a variable

- Search optimum *n*(*r*) by referencing "another wave function $\phi(r)$ ". (by Kohn & Sham (1965))
- Caution 1 by Harriman $:\;\;\forall n(r)\!\in I_{_N}, \exists \, \phi_{_i}(r) \!\! \S. t.\{\phi_{_i}(r)\}\!\rightarrow\! n(r).$
- Caution 2 by Lieb : $A_N \subset I_N$, $A_N \neq A_N$ '.

We need other energy functionals than the Hohenberg-Kohn functional.

A set of density

⚫ V-representable density : *AN*

● Density of a ground state, which is an eigenstate of an interacting Hamiltonian operator for an external potential $V_{ext}(r)$.

⚫ N-representable density : *IN*

⚫ Density satisfying next three conditions.

$$
n(\mathbf{r}) \geq 0, \quad \int n(\mathbf{r}) d\mathbf{r} = N, \quad \int |\nabla n(\mathbf{r})^{1/2}|^2 d\mathbf{r} < \infty.
$$

Energy as a functional of density

- The Levy-Lieb universal energy density functional $[n] = min\langle \Psi | \hat{T} + \hat{V}_{ee} | \Psi \rangle.$ $\Psi \rightarrow n$ *ee n* $F[n] = min\langle \Psi | \hat{T} + \hat{V}_{ee} | \Psi \rangle$. \hat{T} : kinetic energy operator, \hat{V}_{ee} : e : kinetic energy operator, V_{ee} : electron-electron interaction. Since $\hat{T} + \hat{V_{ee}}$ is a positive quadratic form,
- The Lieb energy functional $[n] = \inf_{\mathbf{r}} \operatorname{tr} \hat{\Gamma}(\hat{T} + \hat{V}_{ee}),$ $\hat{r} \rightarrow n$ $(1 \mid V_{ee})$ $F_L[n] = \inf_{\hat{r}} \text{tr} \hat{\Gamma}(\hat{T} + \hat{V}_{ee}),$ $\hat{\Gamma}$: density matrix. For a density, $n({\bf r}){\ge}0,$ which is ensemble-V (EV) representable,

$$
\begin{aligned}\n\mathbf{F}_{0} &= \langle \Psi_{GS} | \hat{T} + \hat{V}_{ce} | \Psi_{GS} \rangle + \int n_{GS}(\mathbf{r}) v_{ext}(\mathbf{r}) d^{3}r \\
&= \min_{n} \left\{ \min_{\Psi \to n(\mathbf{r})} \langle \Psi | \hat{T} + \hat{V}_{ce} | \Psi \rangle + \int n(\mathbf{r}) v_{ext}(\mathbf{r}) d^{3}r \right\} \\
&= \min_{n} \left\{ \min_{\Psi' \to n(\mathbf{r})} \langle \Psi' | \hat{T} | \Psi' \rangle + F[n] - F_{T}[n] + \int n(\mathbf{r}) v_{ext}(\mathbf{r}) d^{3}r \right\} \\
&= \min_{n} \left\{ \min_{\Psi' \to n(\mathbf{r})} \left\{ \langle \Psi' | \hat{T} | \Psi' \rangle + F[n_{\Psi'}] - F_{T}[n_{\Psi'}] \right. \\
&\quad \left. + \int n_{\Psi}(\mathbf{r}) v_{ext}(\mathbf{r}) d^{3}r \right\} \right\} \\
&= \min_{\Psi'} \left\{ \langle \Psi' | \hat{T} | \Psi' \rangle + \frac{e^{2}}{2} \int \frac{n_{\Psi'}(\mathbf{r}) n_{\Psi'}(\mathbf{r})}{|\mathbf{r} - \mathbf{r}'|} d^{3}r d^{3}r' + F[n_{\Psi'}] \\
&= \frac{e^{2}}{2} \int \frac{n_{\Psi'}(\mathbf{r}) n_{\Psi'}(\mathbf{r})}{|\mathbf{r} - \mathbf{r}'|} d^{3}r d^{3}r' - F_{T}[n_{\Psi'}] + \int n(\mathbf{r}) v_{ext}(\mathbf{r}) d^{3}r \right\} \\
&= \left[\min_{\Psi'} \left\{ \langle \Psi' | \hat{T} | \Psi' \rangle + \frac{e^{2}}{2} \int \frac{n_{\Psi'}(\mathbf{r}) n_{\Psi'}(\mathbf{r})}{|\mathbf{r} - \mathbf{r}'|} d^{3}r d^{3}r' + E_{xc}[n_{\Psi'}] \right\} \right] \mathbf{Definition of } E_{xc}[\mathbf{n}].\n\end{aligned}
$$

Two cautions on Kohn-Sham scheme

Modified energy functional and phase transition points.

$$
F_{\lambda}[n] = \min_{\Psi \hookrightarrow n} \langle \Psi' | \hat{T} + \lambda \hat{V}_{ee} | \Psi' \rangle.
$$

K. Kusakabe and I. Maruyama, J. Phys. A: Math. Theor. **44**, 135305 (2011).

Ex. Metal \leftrightarrow Mott insulator, Kramers doublet \leftrightarrow local Kondo singlet

See also contents of 13th lecture.

of "Theory in Materials Science @ ES-OU".

Differentiability of the Lieb functional is shown in the following conditions, only.

- *n*(**r**) is an EV-representable,
- Direction $n_1(r)$ is an EV-rep.,
- $n(r) > 0$, and $n_1(r) > 0$,
- Second order derivatives of *n*1 (**r**), *n*(**r**), and G.S. w.f., are square integrable.
- Special directional derivative of *F*^L [*n*] is given.

P.E. Lammert, Int. J. Quant. Chem. **107**, 1943 (2007).

Note that functional derivative is considered with variation out of H^1 in general sense, if one do not make a caution on the domain.

DFT & LDA/GGA

Definition of energy functional Model functional introduced,

Hohenberg-Kohn theorem

Harriman-Lieb construction Levy-Lieb constrained min.

Lieb functional with definition of the domain **then, we have the steps !**

Kohn-Sham variational method

Many-body perturbation theory using K-S orbitals

Effective many-body models to search "exact results"

Caution: On the searching path, definition of the Kohn-Sham equation by using functional derivative of the Lieb functional is not always possible. While directional derivative is given for limited directions, not only interpolation but extrapolation of *n*(*r*) is requested to determine effective potential.

Functional derivative

Variation w.r.t. density

- General variation of *n*(*r*) includes, *r n*(*r*) $\delta n(r)$ *r*
- Variation maintaining *N* has to be non-local.

Non-linear functional

Fréchet derivative w.r.t. $\delta n(r)$ is accessible for linear functional. But, $F_{L}[n]$ is not a linear functional. *n*(*r*)

Since Ψ giving "non-*N*-rep. $n(r)$ " is not found, "ensemble *v*-rep. *n*(*r*)" is requested. The Gâteaux derivative of the Lieb functional has directional dependence on $\delta n(r)$ as a function of r.

Classification of density

 v -representable $v - \overline{a}$ density distribution given by a quantum eigen state in a potential

- $\mathcal{A}(N)$: by an eigen state (a ground state) of an interacting N-body electron system
- $\mathcal{A}(N)'$: by an eigen state (a ground state) of a non-interacting N-body Fermion system

N-representable by an anti-symmetric wave function of an *N*-particle system

- **•** $\mathcal{I}(N)$: positive, integrable, thus normalizable to give *N*, (∇*n*(*r*)^{1/2})² is also square integrable
- The Harriman-Lieb construction gives a many-body wave function giving *n*(*r*). Thus existence of the wave functions for constrained minimization is certified.

Linear combination of two or more v-representable density distributions (Ensemble average as statistical average) Ensemble v-representable

- Defined as an interpolating point of two well-defined end points
- Two end points are v-representable (non-negative, without diverging points)
- A statistical average found on the way of the search is not pure v-representable.

The rigorous DFT method requires "huge amount of calculation steps" far bigger than the statistical quantum mechanics method for N-body problem.

We do not use the Lieb functional.

$$
G_{\mathcal{T}}[\Psi'] = \langle \Psi' | \hat{T} | \Psi' \rangle - F_{\mathcal{L},\mathcal{T}}[n_{\Psi'}] + F_{\mathcal{L}}[n_{\Psi'}] + \int \nu_{\text{ext}}(\mathbf{r})n_{\Psi'}(\mathbf{r})d^{3}r.
$$

\n
$$
G_{\mathcal{T}}[\Psi'] = \langle \Psi' | \hat{T} | \Psi' \rangle + \frac{e^{2}}{2} \int \frac{n_{\Psi'}(\mathbf{r})n_{\Psi'}(\mathbf{r})}{|\mathbf{r} - \mathbf{r}|} d^{3}rd^{3}r' + E_{\mathcal{X}}[n_{\Psi'}] + \int \nu_{\text{ext}}(\mathbf{r})n_{\Psi'}(\mathbf{r})d^{3}r.
$$

\nLet's assume that $|\Psi'\rangle$ is a minimizer w.r.t. variation: $\frac{\delta}{\delta\langle \Psi'|} \delta G_{\mathcal{T}}[\Psi'] - E(\langle \Psi' | \Psi' \rangle - 1) \rangle = 0$, where density variation is assumed to be given as,
\n
$$
n(\mathbf{r}) = \langle \Psi' | \hat{n}(\mathbf{r}) | \Psi' \rangle,
$$
\n
$$
\delta n(\mathbf{r}) = \langle \Psi' | \hat{n}(\mathbf{r}) | \Psi' \rangle + \delta |\Psi' \rangle - n(\mathbf{r}) : \text{two end points are positive. (Good. But,})
$$
\n
$$
= \{\delta \langle \Psi' | \hat{n}(\mathbf{r}) | \Psi' \rangle\} + \langle \Psi' | \hat{n}(\mathbf{r}) \delta | \Psi' \rangle\} + \delta \langle \Psi' | \hat{n}(\mathbf{r}) \delta | \Psi' \rangle\}.
$$
\n
$$
\{\hat{\mathcal{T}} + \int d^{3}r v_{\text{eff}}[n, \delta n[\delta \langle \Psi' | \hat{n}(\mathbf{r}) | \Psi' \rangle] + \delta \langle \Psi' | \delta n(\mathbf{r}) \delta | \Psi' \rangle\}.
$$
\n
$$
\{\hat{\mathcal{T}} + \int d^{3}r v_{\text{eff}}[n, \delta n[\delta \langle \Psi' | \hat{n}(\mathbf{r}) | \Psi' \rangle] + \delta \langle \Psi' | \delta n(\mathbf{r}) \delta | \Psi' \rangle\}.
$$
\n
$$
\{\hat{\mathcal{T}} + \int d^{3}r v_{\text{eff}}
$$

Why do we use DFT?

- Degree of freedom is separated into electrons and nuclei by the Born-Oppenheimer approximation.
	- Electron density characterizes the system!
- Let's determine the free energy (total energy at T=0K) by the electron density.

Hamiltonian

• Born-Oppenheimer approx.

$$
H = -\sum_{i=1}^{N} \left(\frac{\hbar^2}{2m} \right) \nabla_{\mathbf{r}_i}^2 + \sum_{\langle i,j \rangle} \frac{e^2}{|\mathbf{r}_i - \mathbf{r}_j|} + \sum_{i=1}^{N} v_{ext}(\mathbf{r}_i).
$$
\n
$$
E[v_{ext} + \delta v_{ext}] - E[v_{ext}]
$$
\n
$$
- \int d^3 r \frac{\delta E[v_{ext}]}{\delta v_{ext}} \delta v_{ext}(\mathbf{r}) + O(\delta^2).
$$

- ➢ Classical configuration of nuclei,
- ➢ Determination of static potential.

Order parameter

• Funct. deriv. w.r.t. ext. pot.

$$
E[v_{ext} + \delta v_{ext}] - E[v_{ext}]
$$

= $\int d^3 r \frac{\delta E[v_{ext}]}{\delta v_{ext}(\mathbf{r})} \delta v_{ext}(\mathbf{r}) + O(\delta^2)$
= $\int d^3 r \delta v_{ext}(\mathbf{r}) n(\mathbf{r}) + O(\delta^2)$

Statistical mechanics offers a solution.

- L.D. Landau had predicted relevance of order parameters.
	- Energy of condensed matter is given by "a function of physical variables" i.e. a functional of a function.
		- Landau considered analytic functions at first.
		- Phenomenological, but "exact", since the idea comes from observation of phenomena.
- Present DFT is interpreted as a realization. $E_{\nu_{\text{ext1}}}[n]$
	- Order parameter = Electron density
	- Energy functional = Landau functional

 $E_{_{\nu_{ext2}}}[n]$

n : density

The idea goes back to the original Thomas-Fermi arguments. As such, the history of the quantum statistical mechanics is long.

Derive differential equations

- First, set up a differentiable energy functional.
	- $E_{xc}[n] \rightarrow \overline{E}_{xc}[n]$ $G_{T}[\Psi] \rightarrow \overline{G}_{T}[\Psi]$
- Derive a determining equations, which are to be self-consistent equations.

 $\frac{\partial}{\partial \Psi} \left\{ \overline{G}_T \left[\Psi \right] - E \left(\left\langle \Psi \right| \Psi \right\rangle - 1 \right) = 0, \quad \Box$ $\delta \Psi$ $\frac{\partial}{\partial \Psi} \Big\{ \overline{G}_T \big[\Psi \big] - E \big(\! \big\langle \Psi \big| \Psi \big\rangle \! - \! 1 \big) \! \big\} \! = 0, \quad \blacktriangleright \quad \Big\{ \!\hat{T} + \int d^3r v_{\mathit{eff}} \big(\mathbf{r} \big) \! \hat{n} \big(\mathbf{r} \big) \! \Big\} \Psi \Big\rangle \! = E \big| \Psi \big\rangle,$ $({\bf r}) = |d^3r'|$ $(\mathbf{r}') \quad \delta E_{rc} |n|$ () $\left(\mathbf{r} \right)$ and $\left(\mathbf{r} \right)$ $v_{\text{av}} + \frac{\partial \mathbf{E}_{xc}[\mu]}{\partial \mathbf{E}_{ox}} + v_{ext}(\mathbf{r}),$ $n(\mathbf{r}) = \sum \langle \Psi | \hat{n}(\mathbf{r}) | \Psi \rangle.$ ' ' $^3r'$ $\frac{n(\mathbf{I} + \mathbf{I})}{r}$ + $\frac{\partial E_{xc}[\mu]}{\partial \mathbf{I}}$ + $v_{ext}(\mathbf{r}),$ $-\mathbf{r}'$ $\delta n(\mathbf{r})$ $=\int d^3r' \frac{n(\mathbf{r})}{|\mathbf{r}-\mathbf{r}'|} + \frac{\partial L_{xc}l''l}{\partial n(\mathbf{r})} + v_e$ σ δE |n| **r** $\mathbf{r} - \mathbf{r}'$ $\delta n(\mathbf{r})$ $\cos n$ \mathbf{r}') δE _r $\mathbf{r} = \int d^3 r' \frac{n(\mathbf{r})}{|\mathbf{r} - \mathbf{r}'|} + \frac{\partial \mathcal{L}_{xc}[\mathbf{r}]}{\partial n(\mathbf{r})} + v_{ext}(\mathbf{r}),$ $n(\mathbf{r}')$ $\delta E_{rc} |n|$ ($v_{\text{eff}}(\mathbf{r}) = \int d^3 r' \frac{n(\mathbf{r})}{\ln |\mathbf{r}|} + \frac{\partial E_{xc}[\mathbf{r}]}{\partial |\mathbf{r}|} + v_{ext}(\mathbf{r})$

This strategy gives us "realistic approaches" of DFT. We had developed "systematic improvement" of any approximation in DFT.

● The exchange-correlation energy functional may be written

$$
E_{xc}[n] = \frac{e^2}{2} \int d\mathbf{r} n(\mathbf{r}) \int d\mathbf{r}' \frac{1}{|\mathbf{r} - \mathbf{r}'|} n_{xc}(\mathbf{r}, \mathbf{r}')
$$

with

as,

$$
n_{xc}(\mathbf{r}, \mathbf{r}') = n(\mathbf{r}') \int_0^1 [g_n(\mathbf{r}, \mathbf{r}', \lambda) - 1] d\lambda
$$

=
$$
\int_0^1 \left[\langle \Psi_\lambda | (\hat{n}(\mathbf{r}) - n(\mathbf{r})) (\hat{n}(\mathbf{r}') - n(\mathbf{r}')) \Psi_\lambda \rangle / n(\mathbf{r}) - \delta(\mathbf{r} - \mathbf{r}') \right] d\lambda.
$$

Quantum Fluctuation!

● In the local density approximation (LDA), we approximate $n_{\rm xc}$ by that of the homogeneous electron gas as,

$$
n_{\scriptscriptstyle xc}(\mathbf{r},\mathbf{r}')=n(\mathbf{r})\int_0^1\Big[g^{\rm hom}{}_n(\mathbf{r},\mathbf{r}',\lambda)-1\Big]d\lambda.
$$

'Density-functional theory of atoms and molecules,' by R.G. Parr, **Same B. W. Yang, Springer, §8.5.**

LDA exchange-correlation energy can be written as,

 $(\mathbf{r}, \mathbf{r}') = n(\mathbf{r}) \left[\int g^{\text{hom}}_{n}(\mathbf{r}, \mathbf{r}', \lambda) \right]$

Why local density approximations?
\nDA exchange-correlation energy can be written as,
\n
$$
\overline{E}_{xc}[n] = \frac{e^{2}}{2} \int d^{3}rn(\mathbf{r}) \int d^{3}r' \frac{1}{|\mathbf{r} - \mathbf{r}'|} \overline{n}_{xc}(\mathbf{r}, \mathbf{r}') = \int d^{3}rn(\mathbf{r}) \varepsilon_{xc}(n(\mathbf{r})).
$$
\n
$$
\overline{n}_{xc}(\mathbf{r}, \mathbf{r}') = n(\mathbf{r}) \int_{0}^{1} \left[g^{hom}{}_{n}(\mathbf{r}, \mathbf{r}', \lambda) - 1\right] d\lambda.
$$
 This LDA exchange-
\ncorrelation hole satisfies a
\nsum rule. (See next page.)
\nThe value is given for any $n(\mathbf{r})$.
\nWhen we choose LDA, the Kohn-Sham equation reads,
\n
$$
\left\{-\frac{\hbar^{2}}{2m} \Delta_{\mathbf{r}} + v_{eff}(\mathbf{r})\right] \phi_{i}(\mathbf{r}) = \varepsilon_{i} \phi_{i}(\mathbf{r}),
$$
For this set of equations, we have
\ncomputer codes as numerical
\nsolvers of quantum mechanical
\n
$$
v_{eff}(\mathbf{r}) = \int d^{3}r' \frac{n(\mathbf{r}')}{\vert \mathbf{r} - \mathbf{r}' \vert} + \frac{\partial \overline{E}_{xc}[n]}{\partial \overline{E}_{xc}(\mathbf{r})} + v_{ext}(\mathbf{r}),
$$
 problems for electron systems.

- $\int_{0}^{1} [g^{\hom}{}_n(\mathbf{r},\mathbf{r}',\lambda)-1] d\lambda.$ Thi $\overline{n}_{xc}(\mathbf{r}, \mathbf{r}') = n(\mathbf{r}) \int_0^1 \left[g^{\text{hom}}{}_n(\mathbf{r}, \mathbf{r}', \lambda) - 1\right] d\lambda.$ corre • This LDA exchangecorrelation hole satisfies a sum rule. (See next page.) **oximation?**
 d³*rn***(r**) $\varepsilon_{xc}(n(\mathbf{r}))$.

s LDA exchange-

relation hole satisfies a

n rule. (See next page.)

e value is given for any $n(\mathbf{r})$. **roximation?**
 $\int d^3r n(\mathbf{r}) \varepsilon_{xc}(n(\mathbf{r})).$
his LDA exchange-
prrelation hole satisfies a
um rule. (See next page.)
he value is given for any $n(\mathbf{r})$.
	- The value is given for any $n(\mathbf{r})$.

When we choose LDA, the Kohn-Sham equation reads,

$$
\begin{cases}\n-\frac{\hbar^2}{2m}\Delta_{\mathbf{r}} + v_{\text{eff}}(\mathbf{r})\phi_i(\mathbf{r}) = \varepsilon_i \phi_i(\mathbf{r}), & \text{For the complex solution} \\
v_{\text{eff}}(\mathbf{r}) = \int d^3 r' \frac{n(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} + \frac{\partial \overline{E}_{xc}[n]}{\partial n(\mathbf{r})} + v_{\text{ext}}(\mathbf{r}), & \text{probe} \\
n(\mathbf{r}) = \sum_{\sigma} \sum_{\varepsilon_i \le E_F} |\phi_i(\mathbf{r})|^2. & \text{Re}\n\end{cases}
$$

For this set of equations, we have computer codes as numerical solvers of quantum mechanical problems for electron systems.

Reasonable accuracy !!!

If you need, you may always make a correction by introducing "multi-reference representation of a model many-body system".

A sum rule in the image:
$$
\begin{array}{ll}\n\mathbf{A} & \text{sum rule } \text{in } \mathbf{D} \text{ is given by the following equation.} \\
\mathbf{A} & \text{sum value } \sum_{n(\mathbf{r})n(\mathbf{r})} \frac{\partial(\mathbf{r}-\mathbf{r}')}{n(\mathbf{r})} \\
&= \frac{\left\langle \hat{n}(\mathbf{r})\hat{n}(\mathbf{r}') \right\rangle_{\mathbf{r}_{n}} - \partial(\mathbf{r}-\mathbf{r}')}{n(\mathbf{r})}}{\frac{\partial(\mathbf{r})\hat{n}(\mathbf{r}) - \partial(\mathbf{r}') \right\rangle_{\mathbf{r}_{n}} - \partial(\mathbf{r}-\mathbf{r}')}{n(\mathbf{r})}} \\
&= \frac{\left\langle \hat{n}(\mathbf{r})\hat{n}(\mathbf{r}') \right\rangle_{\mathbf{r}_{n}} - \partial(\mathbf{r}-\mathbf{r}')}{n(\mathbf{r})}}{\frac{\partial(\mathbf{r})\hat{n}(\mathbf{r}) - \partial(\mathbf{r}') \right\rangle_{\mathbf{r}_{n}} - \partial(\mathbf{r}-\mathbf{r}')}{n(\mathbf{r})}} \\
&= \int d^{3}r' \left| \int_{0}^{1} \frac{\hat{n}(\mathbf{r})\hat{n}(\mathbf{r}') \right\rangle_{\mathbf{r}_{n}} \cdot d\lambda - n(\mathbf{r}')\partial(\mathbf{r}-\mathbf{r}')}{n(\mathbf{r})} \\
&= \int d^{3}r' \left| \int_{0}^{1} \frac{\hat{n}(\mathbf{r})\hat{n}(\mathbf{r}') \right\rangle_{\mathbf{r}_{n}} \cdot d\lambda - n(\mathbf{r}')\partial(\mathbf{r}-\mathbf{r}')}{n(\mathbf{r})} \\
&= \int d^{3}r' \left| \int_{0}^{1} \frac{\hat{n}(\mathbf{r})\hat{n}(\mathbf{r}') \partial(\mathbf{r}\cdot\mathbf{r}')}{n(\mathbf{r})}\right\rangle_{\mathbf{r}_{n}} \cdot d\lambda - n(\mathbf{r}')\partial(\mathbf{r}-\mathbf{r}')}{n(\mathbf{r})}\n\end{array}
$$
\n
$$
\begin{array}{ll}\n\mathbf{A} & \text{sum value } \hat{n}(\mathbf{r})\hat{n}(\mathbf{r}')\partial(\mathbf{r})\hat{n}^{(3)}\hat{n}^{(3)}
$$

Thus, although we have completely different formulae for DFT and LDA, the same sum rule for the exchange-correlation hole is satisfied.

Correlation energy density of LDA

Let's introduce r_s by $\frac{r_e}{V} = 1/\left(\frac{m}{2}r_s^3\right)$ we may divide $\varepsilon_{\rm xc}$ as, $\varepsilon_{\rm xc}(r_s) = \varepsilon_{\rm x}(r_s) + \varepsilon_c(r_s)$. $\bigg)$ \setminus Ķ. \setminus $\sqrt{}$ $=1/\frac{4\pi}{\pi}r_s^3$ 3 4 $1/\frac{1}{2}r_s$ $\frac{e}{r} = 1 / \frac{4\pi}{\pi} r$ *V* N_e ¹ $(4\pi$

How to solve the Kohn-Sham egs?

$$
\begin{aligned}\n&\left\{-\frac{\hbar^2}{2m}\Delta_{\mathbf{r}} + v_{\text{eff}}(\mathbf{r})\right\}\phi_i(\mathbf{r}) = \varepsilon_i\phi_i(\mathbf{r}),\\
v_{\text{eff}}(\mathbf{r}) &= \int d^3r' \frac{n(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} + \frac{\partial \overline{E}_{xc}[n]}{\partial n(\mathbf{r})} + v_{\text{ext}}(\mathbf{r}),\\
n(\mathbf{r}) &= \sum_{\sigma} \sum_{\varepsilon_i \leq E_F} |\phi_i(\mathbf{r})|^2. \quad &\text{A good point: A complete basis set giving the expansion of the many-body state vector.}\n\end{aligned}
$$

We need to find a solution $\{\phi_i(r)\}$ satisfying

- 1. Symmetry of the system (Crystal symmetry, molecular symmetry)
- 2. Self-consistency

This is a least version for DFT. It is corrected by "quantum charge fluctuation".

Band structure of cubic diamond

An energy gap appears and the system is a wide-gap semiconductor. LDA by PW91.

Plane-wave expansion with ultra-soft PP.

Bonding charge in hex-diamond

- In a covalent crystal, we can see charge density of electrons at each bond connection.
- Yellow object represents charge density and white spheres are carbons.

The density map is rather accurate, because it is determined self -consistently.

Band structure of graphite

The π -band is half-filled and there are small Fermi pockets both for electrons and holes. (Semimetal)

Bonding charge in graphite

- Bonding charge comes from σ electrons.
- This system is a semimetal where the Fermi surface is made of π -bands.

When you see a localized state mediating quantum information process, you may introduce "a solver for correlated electron systems" at any time.

Band structure of Graphene

We want to see "what is seen in the right figure"!!

Idea to use gradient of density

- To overcome limitation of LDA, Kohn & Sham had considered gradient corrections.
- Let's first introduce another order parameter,

or density for two spins, $n_{\uparrow}(r)$, $n_{\downarrow}(r)$, or density of multiple species (electrons & nuclei). $|n, j|$ (n, j) $|\mathbf{y}| = \min \langle a, b | | H(\mathbf{r}) d\mathbf{r} | a, b \rangle,$ $,b\rangle \rightarrow (n,$ $F[n, j] = \min_{a,b} \left| \frac{a,b}{b} \right| H(r) dr | a,b$ $a,b\rangle \rightarrow \! (n$ $\bm{\mathrm{j}}$ = $\min_{[a,b)\to (n,\bm{\mathrm{j}})}\langle a,b|\int H(\mathbf{r})d\mathbf{r}|a,b\rangle,$ $F[n,\mathbf{m}]$ = $\min_{[a,b)\to (n,\bm{\mathrm{j}})}$ (n,m) \mathbf{m}] = $\min_{|a,b\rangle \rightarrow (n,\mathbf{m})} \langle a,b|\hat{T}+\hat{V}_{ee}|a,b\rangle.$ $F|n,m| = \min \langle a,b|T+\hat{V}_{eq}|a,b\rangle.$ $a,b\rangle \rightarrow (n,m)$ $= \min \langle a,b|T+\hat{V}_{\text{ee}}|a,b\rangle.$ $\rightarrow (n,m)$ **m**

• Naively,

$$
E_{xc}^{GEA}[n_{\uparrow},n_{\downarrow}] = E_{xc}^{LSDA}[n_{\uparrow},n_{\downarrow}] + \sum_{\sigma,\sigma'} \int d^3r C_{\sigma,\sigma'}(n_{\uparrow},n_{\downarrow}) \frac{\nabla n_{\sigma}}{n_{\sigma}^{2/3}} \cdot \frac{\nabla n_{\sigma'}}{n_{\sigma'}^{2/3}}.
$$

Generalized gradient approximation

- GEA fails due to violation of sum rules.
- Introduce a GGA exchange-correlation energy functional, whose ex.-corr. hole satisfies desired sum rules.

$$
E_{xc}^{GGA}[n_{\uparrow}, n_{\downarrow}] = \int d^3r f(n_{\uparrow}, n_{\downarrow}, \nabla n_{\uparrow}, \nabla n_{\downarrow}).
$$

For the case of exchange hole,

$$
\tilde{n}_x^{GGA} = -\frac{1}{2}n(\mathbf{r})\tilde{y}(\mathbf{r},\mathbf{u})\theta(\tilde{y}(\mathbf{r},\mathbf{u}))\theta(u_x(\mathbf{r})-u),
$$

where $\theta(x)$ is the step function.

'Electronic Structure – Basic Theory and Practical Methods,' by R.M. Martin, Cambridge University Press.

Which do we use, LDA or GGA?

• 3d ferromagnetic metals:

- LSDA does not reproduce stability of ferromagnetism.
- Spin-GGA reproduces magnetism even quantitatively. (This is practical understanding.)

Internal field considered

The result suggests existence of *m*(*r*) in a solution of DFT simulation.

 $\hat{\mathbf{m}}(\mathbf{r})\rangle, \quad \big<\hat{\mathbf{j}}(\mathbf{r})\big>, \quad \text{or}$ $\hat{j}_{\mu}(x)$ = $\langle \overline{\psi}(x) \gamma_{\mu} \psi(x) \rangle$ $\hat{i}(x)$

may be used to determine the internal vector photon field to derive an effective potential problem determining an L^2 basis.

An effective action and the resulting effective Hamiltonian contain all the electron-electron interaction processes in manybody description.

Decision by empirical rules

- Good 3d metals,
- Strong metallic ferromagnets,
- 3d semiconductors,
- Molecules,
- Molecular solids without π - π stacking.

GGA OK GGA not in use

- Graphite, h-BN, CF,
- Weak ferromagnets,
- Molecular solids with π - π stacking.

For these materials, LDA gives rather reasonable solution.

We may use GGA to obtain a starting (tentative, or initial) solution for consideration of

- Van-der-Waals crystals,
- Materials showing strong-correlation effects.

If both LDA and GGA fails?

We might be able to solve the problem using

- For the van-der-Waals crystals,
	- GGA + van-der-Waals correction,
	- EXX + RPA correlation.

• For the strong-correlation problems,

- GGA + short-range correlation,
- EXX + (RPA + vertex corrected) correlation.

Please consult experts!!!

The bulk modulus

$$
B_0 = -\frac{Vdp}{dV} = \frac{Vd^2E}{dV^2},
$$

\n
$$
E(V) = E_0 + \frac{B_0V}{B_0} \left(\frac{(V_0/V)^{B_0}}{B_0 - 1} + 1 \right) - \frac{B_0V_0}{B_0 - 1}.
$$

We can compare the methods for c-BN.

- LDA is known to give reasonable estimation.
- People noticed that GGA gives a shifted value.
- Interpretation became possible after BS & PUS have been known.
- Correction of GGA is known to be possible.
- Then, evaluation of TB or force field may be allowed.

Cf. Nagakubo, et al. Appl. Phys. Lett. 102, 241909 (2013).

A tight-binding model by DFT

Ref. K. Kuroki, et al. Phys. Rev. Lett. 101, 087004 (2008).

The Graduate School of Engineering Science

GW+beyond as "a many-body PT

Evaluation of semiconductor gap by self-consistent GW + vertex correction

Ref. M. Shishkin, et al. Phys. Rev. Lett. 99, 246403 (2007).

The Graduate School of Engineering Science

Diffusion Monte-Carlo method

Electronic state and charge distribution of a Hydrogen molecule adsorbed on benzene

Ref. Y. Kanai and J.C. Grossman, Phys. Rev. A 80, 032504 (2009).

An almighty method

• Space of DFT models

• Distance:
$$
\|n_i - n_{i+1}\|.
$$

• Variational principle: DFVT by K.K. (2009).

Q: Why? A: Its because "axiom for materials" tells this strategy.

-vicinity around

exact solution

CASE I

Super processes

To have an initial state of correlated electron system, we may utilize

The upconversion Hamiltonian

$$
\left(H^1+P_AV_{ee}P_A+H^1_{C,counter}+P_AV_{ee}(1-P_A)\frac{1}{H^1+H^1_{C,counter}-E}(1-P_A)V_{ee}P_A\right)\vert\Psi_A\rangle=E\vert\Psi_A\rangle.
$$

which leads us to the convergent model series in the DFT model space. Cf. K. Kusakabe, I. Maruyama, "Electronic state calculation method, electronic state calculation device, computer program", PCT filed No. PCT/JP2011/068589, date 2011.8.18.

Processes by "Feynman Diagram

Symmetrized bare interaction

The full super-process for an electron-hole interaction

A polarization bubble diagram with a vertex correction in the above scattering channel

A third-order ladder diagram for the scattering channel

First principles calculation

- When an effective Hamiltonian is derived by renormalization of the phase space of $L^2(R^{3N})$, it is in a form of "a many-body correlation problem", which is tractable in a simulation.
- Its solver is always faster than the encapsulation in $L^2(R^3)$ by the KS scheme.
- This conclusion is a FP.

Why is the KS-DFT not FP? Why do "first principles" exist?

- There are virtual photons.
- There is no real photon.
- Thus any theory for any system in nature needs to solve a many-body correlation problem.
- There are non-Abelian gauge fields.
- Determination of their source matter fields requires to describe the propagator of the gauge bosons, which is far beyond the U(1) theory.

A comment on U_{eff} and t

A narrow band system becomes 'the strongly correlated electron system'

For a metallic system, we have variety of phases including

- An anomalous metallic state for a heavy Fermion system,
- A high-temperature superconducting state of cuprates, where U_{eff}/t becomes large.

But, a further difficult question is to conclude a Fermi liquid state following the renormalization group algorithm by starting from a DFT basis set.

Better description tells physics.

K. Nishiguchi, et al. PRB (2018).

Stronger superconductivity in OP.

Dynamical problems

- Properties are to be given by
	- Symmetry,
	- Self-consistency,
	- Time evolution path.
	- So, dynamics is determined by
		- Initial conditions,
		- Consistency,
		- Conservation laws.
- There, quantum correlation is cut, because we are in the space after the primary observation in our universe.

Cautions:

- 1. Separation into sub-systems is dangerous without consideration.
- 2. We have a different heat-bath & environment for a history of the sub-system different from an observed one.

The density functional theory is no more than a method to describe a closed system in equilibrium. Once the time evolution starts, it means that we need the quantum-statistical mechanics for non-equilibrium systems.

When an initial state is created,

Consider a semiconductor having conduction and valence bands with a band width around 1eV. Assume that the material (each domain of poly-crystal sample) is in a nanometer scale.

When DFT is applied to the steady initial state of the electron system, the band structure is given by a solution of the Kohn-Sham equation. Wannier orbitals are given by a unitary transformation of the Kohn-Sham Bloch orbitals. Therefore,

In this process of theoretical description, we need to generalize DFT from single-referenced to multi-referenced DFT.

The trend in theory of DFT

Growing numbers of APS papers

Survey in APS journals

① Correlated electron model & DFT ② Many-body perturbation & DFT ③ Quantum Monte Carlo & DFT

Thank you for your attention.

K. Kusakabe 63

