Density Functional Theory for The Many-Body Electron Problem

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OUTLINE

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- Hohenberg-Kohn Theorems
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- Results: H, He, H₂
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Many-Body Electron Problem

For fixed positions of the nuclei (adiabatic approximation) the electron Hamiltonian is

$$\hat{H} = \hat{T} + \hat{V}_{ext} + \hat{V}_{int} + E_{II}, \qquad (1)$$

where \hat{T} is the kinetic energy of the electrons, \hat{V}_{ext} is the potential acting on the electrons due to the nuclei,

$$\hat{V}_{ext} = \sum_{i,I} V_I(|\mathbf{r}_i - \mathbf{R}_I|), \qquad (2)$$

 \hat{V}_{int} is the many-body electron-electron interaction, and E_{II} is the classical ion-ion interaction.

The total energy is the expectation value

$$E = \frac{\langle \Psi | \hat{H} | \Psi \rangle}{\langle \Psi | \Psi \rangle} \equiv \langle \hat{H} \rangle = \langle \hat{T} \rangle + \langle \hat{V}_{int} \rangle + \int d^3 r V_{ext}(\mathbf{r}) n(\mathbf{r}). \quad (3)$$

The ground state wavefunction Ψ_0 is the state with lowest energy; that obeys the symmetries of the particles and all conservation laws.

Hohenberg-Kohn Theorems

• Theorem I: For any system of electrons in an external potential $V_{ext}(\mathbf{r})$, that potential is determined uniquely, except for a constant, by the ground state density $n(\mathbf{r})$.

Corollary I: Since the hamiltonian is thus fully determined, except for a constant shift of the energy, the full many-body wavefunction and all other properties of the system are also completely determined!

• Theorem II: A universal functional for the energy E[n] of the density $n(\mathbf{r})$ can be defined for all electron systems. The exact ground state energy is the global minimum for a given $V_{ext}(\mathbf{r})$, and the density $n(\mathbf{r})$ which minimizes this functional is the exact ground state density.

Corollary II: The functional E[n] alone is sufficient to determine the exact ground state energy and density. Excited states of the electrons must be determined by other means.

• Comment: Functionals may be arbitrarily complex!

Proof of Hohenberg-Kohn Theorems

Proof of Theorem I:

Suppose that there were two different external potentials $V_{ext}^{(1)}(\mathbf{r})$ and $V_{ext}^{(2)}(\mathbf{r})$ with the same ground state density $n(\mathbf{r})$. The two external potentials lead to two different hamiltonians, $\hat{H}^{(1)}$ and $\hat{H}^{(2)}$, which have different ground state wavefunctions, $\Psi^{(1)}$ and $\Psi^{(2)}$, which are hypothesized to have the same density $n(\mathbf{r})$. Then:

$$E^{(1)} = \langle \Psi^{(1)} | \hat{H}^{(1)} | \Psi^{(1)} \rangle < \langle \Psi^{(2)} | \hat{H}^{(1)} | \Psi^{(2)} \rangle.$$
(4)

which leads to

$$E^{(1)} < E^{(2)} + \int d^3r \{ V_{ext}^{(1)}(\mathbf{r}) - V_{ext}^{(2)}(\mathbf{r}) \} n(\mathbf{r}).$$
 (5)

But changing the labels leads to

$$E^{(2)} < E^{(1)} + \int d^3r \{ V_{ext}^{(2)}(\mathbf{r}) - V_{ext}^{(1)}(\mathbf{r}) \} n(\mathbf{r}).$$
(6)

which is a contradiction!

Theorem II (Not proved here) leads to

$$E_{HK}[n] = T[n] + V_{int}[n] + \int d^3 r V_{ext}(\mathbf{r}) n(\mathbf{r})$$

$$\equiv F_{HK}[n] + \int d^3 r V_{ext}(\mathbf{r}) n(\mathbf{r})$$
(7)

The Kohn-Sham Ansatz

The Kohn–Sham approach is to replace the original difficult interacting-particle system with an *auxiliary* system which can be solved more easily.

The widely used-version of the Kohn–Sham approach is to *as*sume one can find an auxiliary system of non-interacting "electrons" with same density as the true interacting system. The hamiltonian for this system can be written:

$$H_{eff} = -\frac{\hbar^2}{2m_e} \nabla^2 + V_{eff}(\mathbf{r}).$$
(8)

where

$$n_{eff}(\mathbf{r}) = \sum_{i=1}^{N} |\psi_i(\mathbf{r})|^2, \qquad (9)$$

and the kinetic energy T_{eff} is given by

$$T_{eff} = -\frac{\hbar^2}{2m} \sum_{i=1}^{N} \langle \psi_i | \nabla^2 | \psi_i \rangle, \qquad (10)$$

The Kohn-Sham energy is the sum

$$\tilde{E} = T_{eff} + \int V_{ext}(\mathbf{r}) n_{eff}(\mathbf{r}) dr + E_{Hartree}[n_{eff}] + E_{xc}[n_{eff}], \quad (11)$$

The Kohn-Sham Equations

The ground state can be found by minimizing with respect to the independent-electron wavefunctions $\psi_i(\mathbf{r})$

$$\frac{\partial \tilde{E}[n]}{\partial \psi_i(\mathbf{r})} = \frac{\partial T_{eff}}{\partial \psi_i(\mathbf{r})} + \frac{\partial E_{other}}{\partial n_{eff}(\mathbf{r})} \frac{\partial n_{eff}(\mathbf{r})}{\partial \psi_i(\mathbf{r})} = 0$$
(12)

subject to the orthonormalization constraints

$$\langle \psi_i | \psi_j \rangle = \delta_{i,j} \tag{13}$$

The Lagrange multiplier method constraints leads to the Kohn– Sham Schroedinger-like equations:

$$(H_{eff} - \epsilon_i)\psi_i(\mathbf{r}) = 0 \tag{14}$$

where the ϵ_i are the eigenvalues,

$$H_{eff}(\mathbf{r}) = -\frac{\hbar^2}{2m_e} \nabla^2 + V_{eff}(\mathbf{r}).$$
(15)

and

$$V_{eff}(\mathbf{r}) = V_{ext}(\mathbf{r}) + \frac{\partial E_{Hart}}{\partial n_{eff}(\mathbf{r})} + \frac{\partial E_{xc}}{\partial n_{eff}(\mathbf{r})}$$
(16)

$$= V_{ext}(\mathbf{r}) + V_{Hart}[n_{eff}] + V_{xc}[n_{eff}].$$
(17)

The total energy can be written

$$\tilde{E} = \sum_{i=1}^{N} \epsilon_i - \frac{1}{2} \int V_{Hart}(\mathbf{r}) n_{eff}(\mathbf{r}) dr + (E_{xc}[n] - \int (V_{xc} n_{eff}(\mathbf{r}) dr)$$
(18)

Exchange-Correlation Functional $E_{xc}[n]$

The Exchange-Correlation energy is defined by

$$F_{HK}[n] = T[n] + V_{int}[n] = \langle \hat{T} \rangle + \langle \hat{V}_{int} \rangle$$

$$\equiv T_{eff}[n] + E_{Hartree}[n] + E_{xc}[n]$$
(19)

where

$$E_{Hartree}[n] = \frac{1}{2} \int d^3r d^3r' \frac{n(\mathbf{r})n(\mathbf{r}')}{|\vec{\mathbf{r}} - \vec{\mathbf{r}}'|}$$
(20)

Thus the exchange-correlation functional is *defined* to be

$$E_{xc}[n] = \langle \hat{T} \rangle - T_{eff}[n] + \langle \hat{V}_{int} \rangle - E_{Hartree}[n]$$
(21)

Key points:

- T_{eff} is really calculated from the Kohn-Sham orbitals, even though it can be proven to be formally a functional of the density
- $E_{xc}[n]$ contains all the difficult terms is a functional of n by the Hohenberg-Kohn Theorem.
- $E_{xc}[n]$ is reasonably approximated as approximately local - unlike $T_{eff}[n]$ and $E_{Hartree}[n]$. This is the basis of the widespread use of this approach.

Exchange-Correlation Hole

(See, e.g., *Many-Particle Physics* by Mahan)

Around each electron at point \mathbf{r} other electrons are excluded to form a hole at points \mathbf{r}' , $n_{xc}(\mathbf{r}, \mathbf{r}')$.

- The Pauli principle (exchange) causes there to be a hole with *exactly one missing electron* compared to the average density of all electrons including the one under consideration.
- Correlation causes rearrangement but still *exactly one missing electron*.
- The energy is given by the interaction with the hole \overline{n}_{xc} averaged over all coupling constants e^2 (See notes for lecture 2.)

$$E_{xc}[n] = \int d^3r n(\mathbf{r}) \int d^3r' \frac{\overline{n}_{xc}(\mathbf{r}, \mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|}$$
(22)

LDA - Local Density Approximation

Assume $E_{xc}[n]$ is a sum of contributions from each point in space depending only upon the density at each point independent of other points. Then

$$E_{xc}[n] = \int d^3 r n(\mathbf{r}) \epsilon_{xc}(n(\mathbf{r}))$$
(23)

where $\epsilon_{xc}(n)$ is the x-c energy per electron

- Since $\epsilon_{xc}(n)$ is assumed to be universal, must be the same as for homogeneous electrons of density n.
- Exchange (e.g., Aschroft and Mermin, p. 411)

$$\epsilon_x(n) = -\frac{0.458}{r_s} \quad Hartree, \tag{24}$$

where r_s is the average distance between electrons given by $\frac{4\pi}{3}r_s^3 = \frac{1}{n}$.

- Correlation found by:
 - RPA approximation good at high density
 - Interpolation between low and high density Wigner (1934), Lindberg and Rosen (1970), ...
 - Essentially exact Monte Carlo Calculations done by Ceperley and Alder, 1980

Correlation Energy vs Density



Figure 1: Correlation energy of an unpolarized homogeneous electron gas as a function of the density parameter r_s . The most accurate results available are quantum Monte Carlo calculations; the curve labelled "Ceperley–Alder" is the work of those authors fitted to the interpolation formula of Vosko, Wilk, and Nusair; the Perdew–Zunger (PZ) fit is almost identical on this scale. In comparison are shown the Wigner interpolation formula, the RPA, and an improved many-body perturbation calculation taken from Mahan, where it is attributed to L. Lindgren and A. Rosen. Figure provided by H. Kim.

Kohn-Sham Equations in the LDA

In the LDA the potential in Kohn-Sham equations is found as a simple derivative:

$$H_{eff}(\mathbf{r}) = -\frac{\hbar^2}{2m_e} \nabla^2 + V_{eff}(\mathbf{r}).$$
(25)

and

$$V_{eff}(\mathbf{r}) = V_{ext}(\mathbf{r}) + \frac{\partial E_{Hart}}{\partial n_{eff}(\mathbf{r})} + \frac{\partial E_{xc}}{\partial n_{eff}(\mathbf{r})}$$
(26)

$$= V_{ext}(\mathbf{r}) + V_{Hart}[n_{eff}] + V_{xc}[n_{eff}].$$
(27)

where

$$V_{xc}(\mathbf{r}) = \frac{\partial E_{xc}}{\partial n_{eff}(\mathbf{r})} = \frac{\partial n_{eff}(\mathbf{r})\epsilon_{xc}(n_{eff}(\mathbf{r}))}{\partial n_{eff}(\mathbf{r})}$$
(28)

The equations are solved self-consistently with the density which results from the eigenfunctions of the independent electron equations

$$n_{eff}(\mathbf{r}) = \sum_{i=1}^{N} |\psi_i(\mathbf{r})|^2,$$
 (29)

Application to Jellium

The eigenvalues of Kohn-Sham Eq. are simply the noninteracting bands $\epsilon_k = \frac{\hbar^2}{2m}k^2$, shifted by the average value of V_{xc} . The total energy is the exact energy by definition.

LSD - Local Spin Density Approx.

- It is straightforward to generalize the Hohenberg-Kohn and Kohn-Sham approaches to functions of two densities n_{\uparrow} and n_{\downarrow} .
- The exchange energy is easily generalized since exchange is always a sum of terms for ↑ and ↓ spins.
- Correlation involves both spins, so it must be parametrized in terms of both n_{\uparrow} and n_{\downarrow} .
- Thus we are led to the *LSD* form $E_{xc}[n_{\uparrow}, n_{\downarrow}]$. All widely used forms are based upon fitting the energies found by Quantum Monte Carlo calculations for interacting electrons done by Ceperley and Alder.
- Parametrized forms given by Perdew and Zunger, 1981, and Vosko, Wilk, and Nusair, 1980.

GGA - Generalized Gradient Approx.

- Exchange and Correlation are *not* really local.
- Next approximation: consider $\epsilon_{xc}[n]$ a function of the density and the gradient of the density at each point
- The function is expressed in terms of the reduced density $s = \frac{|\nabla n|}{2k_F n}$, where $k_F = (3\pi^2 n)^{1/3}$.
- Now widely used especially in chemistry because the GGA improve the estimates of dissociation energies. The GGA lowers the energy of systems with larger gradients. It tends to lower the energy to dissociate a molecule into parts.
- Discussion of forms in paper: J. P. Perdew and Kieron Burke, "Comparison Shopping for a gradient-corrected density functional", Int. J. Quant.
- Programs for the LDA and various GGA's are available from:

http://www.phy.tulane.edu/ kieron/dft.html

Chem. 57, 309 (1996).

Typical Results

From Gunnarsson and Jones, Rev. Mod. Phys. 61, 689 (1989).

	HF	LSDA	Exp
Η	-1.0	-0.98	-1.0
Li	-14.866	-14.706	-14.957
$H_2(\text{binding})$	-3.64	-4.91	-4.75
$Li_2(\text{binding})$	-0.17	-1.01	-1.07

- Energies for solids like Si are in excellent agreement with experiment, e.g., lattice constant predicted: 5.40, Experiment: 5.43 Angstroms.
- Occupied bands typically very good
- Band gaps too low (Si: 0.5 eV, vs. 1.2 eV exp.)
- Failures: In strongly correlated systems like transition metal oxides, predicts metals when real material is a wide gap insulator.