

量子物理学・ナノサイエンス第211回セミナー

## Development of correlation energy functional based on the transcorrelated density functional theory

講師	:	Dr. Naoto Umezawa*
		National Institute for Materials Science, Tsukuba
日程	:	12月27日(水)16:00-17:00
場所	:	本館1階 H156 物理学系輪講室

## 既要

We propose a new formulation of the correlation energy functional derived from the transcorrelated method in use in density functional theory (TC-DFT). An effective Hamiltonian, H<sub>TC</sub>, is introduced by a similarity transformation of a many-body Hamiltonian, H, with respect to a complex function F:  $H_{TC} = 1/F$  H F. It is proved that an expectation value of  $H_{TC}$  for a normalized single Slater determinant,  $D^n$ , corresponds to the total energy:  $E[n] = \langle \Psi^n | H | \Psi^n \rangle / \langle \Psi^n | \Psi^n \rangle = \langle D^n | H_{TC} | D^n \rangle$ under the two assumptions: (1) The electron density  $n(\mathbf{r})$  associated with a trial wave function  $\Psi^n =$  $D^n F$  is v-representable and (2)  $\Psi^n$  and  $D^n$  give rise to the same electron density n (r). This formulation, therefore, provides an alternative expression of the total energy that is useful for the development of novel correlation energy functionals. By substituting a specific function for F, we successfully derived a model correlation energy functional, which resembles the functional form of the screened exchange method. The proposed functional, named the extended screened exchange (ESX) functional, is described within two-body integrals and is parametrized for a numerically exact correlation energy of the homogeneous electron gas. The ESX functional does not contain any ingredients of (semi-) local functionals and thus is totally free from self-interactions. The computational cost for solving the self-consistent-field equation is comparable to that of the Hartree-Fock method. We apply the ESX functional to electronic structure calculations for a solid silicon, Hion, and small atoms. The results demonstrate that the TC-DFT formulation is promising for the systematic improvement of the correlation energy functional.

\*Present affiliation : Semiconductor R&D Center, Samsung Electronics, Korea

**Reference:** N. Umezawa, "Extended screened exchange functional derived from transcorrelated density functional theory" *J. Chem. Phys.* **147**, 104104 (2017)

連絡教員 物理学系 斎藤 晋(内線 2070)

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