A Non-Local, Energy-Optimized Kernel for Structural Properties

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Motivation

To understand long-standing and fundamental problems in solid-state physics and materials chemistry by using the random phase approximation (RPA)

- structural phase transitions are extremely difficult (and extremely difficult to realize experimentally)
- interlayer binding (sometimes double minima is found)
- adsorption (binding and site preference)
- weak interactions in molecules, biomolecules
- reaction energies of certain kinds (self-interaction exposed)
RPA in the density functional context

- Adiabatic connection: density is constrained to physical ($\lambda=1$) value. $\Phi_0$ is a single-determinant of Kohn-Sham orbitals

$$E[n] = \langle \Phi_0[n]|\hat{H}_{\lambda=1}|\Phi_0[n]\rangle + E_c[n]$$

$$\hat{H}_{\lambda}[n] = \hat{T} + \hat{V}_{en} + \hat{V}_{nn} + \lambda \hat{V}_{ee}$$

- Zero-temperature fluctuation-dissipation theorem:

$$E_c = -\int_0^{1}\lambda d\lambda \text{Re} \int_0^{\infty} \frac{du}{2\pi} \langle V[\chi_\lambda(iu) - \chi_0(iu)]\rangle$$

$\chi_\lambda$: Density-density response function, $V$: bare Coulomb interaction

The total energy is computed as $\text{Exc} = \text{EXX} + \text{Ec}$, where $\text{EXX}$ is the Hartree-Fock exact-exchange energy evaluated using KS orbitals.

Ren et al., J. Mater. Sci. 47, 7447 (2012)
Density-density Response Function

\[
\chi^{-1}_\lambda(\omega) = \chi^{-1}_0(\omega) - \left[ V_\lambda + f_{xc}^{\lambda}(\omega) \right] \\
\chi_\lambda = \chi_0 + \chi_0 \left[ V_\lambda + f_{xc}^{\lambda} \right] \chi_\lambda
\]

- Poles of \( \chi_\lambda \) at excitations of interacting system
- Exact \( f_{xc}^{\lambda} \): spatially nonlocal complicated \( \omega \) behavior
- Approximate \( f_{xc} \) obtained from semilocal functionals or a model

**RPA:**

\[
f_{xc}(\vec{r}_1, \vec{r}_2, \omega) = 0
\]

Applications of RPA

• Atomization energies were too low by typically 10 kcal/mol, in comparison to experiment.

• vdW interactions
S22 data set, adsorption of organic molecules, i.e., benzene on metal surfaces, CO adsorption problem, graphene on metal surfaces

Although RPA is known for capturing vdW interactions, binding and cohesive energies are underestimated.

Question:

• How does the overall underestimation of binding by RPA affect structural properties and structural phase transitions?

• How much a beyond RPA treatment improves these properties?

Local field corrections

An intrinsic way to remedy the drawbacks of RPA is to restore the kernel within the ACFDT framework.

How to approach the kernel?

• derive the kernel from microscopic theory

• satisfy exact, known constraints

Most models use the paradigm of the uniform electron gas

ALDA kernel

• The ALDA is the next-simplest approximation for the kernel:

\[ f_{xc}^{ALDA}[n](\vec{r}, \vec{r}') = \delta(\vec{r} - \vec{r}') \frac{d^2[n \varepsilon_{xc}^{\text{unif}}(n)]}{dn^2} \bigg|_{n=n(\vec{r})} \]

\( \varepsilon_{xc}^{\text{unif}}(n) \) is the exchange-correlation energy per particle of an electron gas of uniform density

• ALDA is not exact for the homogeneous electron gas
• ALDA gives reasonable accuracy for low-frequency, long-wavelength excitations, but is not the right choice for a correction to RPA.
• Lein, Gross and Perdew studied the correlation energy per particle of the uniform gas using various kernels. They found that RPA and ALDA-corrected RPA make errors of about 0.5 eV, of opposite sign.
• ALDA is divergent in its correlation hole


What is known about the exchange-correlation kernel?

1. Why does the ALDA kernel fail even for the uniform electron gas?
   Is not exact even for the static exchange-correlation kernel of the uniform electron gas.
   Arbitrary small density variations around the uniform density requires a two-point density functional or double integral over the three-dimensional space, while LDA is only a one-point density functional or single integral.

2. Can a static kernel give accurate exchange-correlation energies?
   Ghosez, Gonze, and Godby had shown that the static kernel in an insulator at the exchange-only level has “ultranonlocality”, decaying at large \( u \) like a negative constant over \( u \).

3. Can we neglect the correlation part of the kernel?
   It is exact by construction in the high-density limit.

4. Error cancellation between exchange and correlation
   The kernel can be ultranonlocal for insulators and molecules as well.

Introducing the NEO (nonlocal energy-optimized) kernel

Three state-of-the-art features:

self-interaction correction

energy-optimization

ultranonlocality
NEO kernel: Short-range correction to the RPA correlation energy, for solid metals

Based on the knowledge of the uniform electron gas.

The exact $xc$ kernel in the uniform electron gas is known to be nonlocal but short-ranged, and our initial version of a NEO kernel also has these features:

$$f_{xc}^{\lambda,\text{NEO}}([n_{\uparrow}, n_{\downarrow}], \vec{R}, u) = -\frac{\lambda}{u} \sum_{\sigma} \left( \frac{n_{\sigma}}{n} \right)^2 \text{erfc} \left( -c(1 - z_{\sigma}^2)k_{F\sigma}^2 u^2 \right)$$

$$n = n_{\uparrow} + n_{\downarrow},$$
$$k_{F\sigma} = (6\pi^2 n_{\sigma})^{1/3}, z_{\sigma} = \tau_{\sigma}^w / \tau, \tau_{\sigma}^w = \left| \nabla n_{\sigma} \right|^2 / 8n_{\sigma}$$
$$\tau_{\sigma} = 1/2 \sum_{\alpha} \left| \nabla \phi_{\alpha\sigma} \right|.$$ 

The one-electron limit:
The singlet two-electron limit:
The uniform electron gas limit is:
The short-range limit:
The long-range limit:
Fitting to the second-order correlation energy

Correlation energy of the uniform electron gas:

\[ e^{RPA}_c = -\frac{12}{\pi} \frac{\hbar^2}{m\alpha_0^2 r_s a_0^2} \int_0^\infty dQ \int_0^\infty dWQ^2 [\alpha(Q,W) - \ln(1 + \alpha(Q,W))] \]

\[ Q = \frac{k}{2k_F}, \quad W = \frac{\Omega}{2k_F^2} \]

\[ \alpha(Q,W) \] is first-order in \( r_s \). \( \ln(1 + \alpha) = \alpha - \frac{1}{2} \alpha^2 + ... \)

Second-order exchange energy with the local-field factor:

\[ e^{2x}_c = \frac{3}{8\pi^3} \int_0^\infty dQ Q^2 \tilde{G}_x(Q) \int_0^\infty dWQ^2 \{2\beta\}^2 \]

\[ G(n; q, \omega) = -(q^2/4\pi)f_{xc}(n; q, \omega) \]

The “c” parameter in NEO provides a unique fit to the exact second-order correlation energy for the spin-unpolarized electron gas. It can be evaluated from explicit expressions given by von Barth and Hedin for RPA and by Langreth and Perdew beyond RPA. It is designed to yield the exact second-order exchange energy of the uniform gas, which is itself an average over frequencies.

Structural parameters

Average errors:

<table>
<thead>
<tr>
<th></th>
<th>PBE</th>
<th>RPA</th>
<th>NEO</th>
</tr>
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<tbody>
<tr>
<td>MD</td>
<td>$a_0$</td>
<td>0.025</td>
<td>0.008</td>
</tr>
<tr>
<td></td>
<td>$B$</td>
<td>$-9$</td>
<td>$-6$</td>
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<tr>
<td></td>
<td>$B$</td>
<td>9</td>
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</table>
RPA Renormalization

\[ \chi_\lambda = [1 - \chi_0 (V_\lambda + f_{xc}^\lambda)]^{-1} \chi_0 \]

\[ \chi_\lambda = \hat{\chi}_\lambda + \hat{\chi}_\lambda f_{xc}^\lambda \hat{\chi}_\lambda \]

Replacing \( \hat{\chi}_\lambda \) by \( \chi_0 \)

\[ \chi_\lambda = \hat{\chi}_\lambda + \chi_0 f_{xc}^\lambda \hat{\chi}_\lambda \]

RPA response function

First order approximation = RPAr1 with the NEO kernel

ACSOSEX with the NEO kernel


Both RPAr1 and ACSOSEX yield the exact second-order, unscreened perturbative correlation energy when the exact first-order kernel is used, but differ in their higher-order contributions.

ACSOSEX vs. RPAr1: homogeneous electron gas

The error from PW92 as a function of $r_s$ for $c = 0.264$.

- ACSOSEX overestimates, RPAr1 underestimates $\Delta E_C^{b\text{RPA}}$
- RPAr1 systematically recovers $> 90\%$ of $\infty$-order $\Delta E_C$
- structural properties show identical trends
ACSOSEX vs. RPA\textsubscript{Ar1}: inhomogeneous systems

Structural properties

<table>
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<tr>
<th></th>
<th>MD</th>
<th>RPA</th>
<th>ACSOSEX</th>
<th>RPA\textsubscript{Ar1}</th>
<th>NEO</th>
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<td>$B$</td>
<td>8</td>
<td>11</td>
<td>6</td>
<td>4</td>
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</table>
Phase Transition in Silicon from Diamond to $\beta$-tin

Equation of state:

$$E(V) = E_0 + \frac{9V_0B_0}{16} \left\{ \left( \frac{V_0}{V} \right)^2 - 1 \right\}^3 B_0' + \left( \frac{V_0}{V} \right)^2 \left[ 6 - 4 \left( \frac{V_0}{V} \right)^3 \right]$$

$E_0$: equilibrium energy  
$V_0$: equilibrium volume  
$B_0$: bulk modulus  
$B_0'$: the pressure derivative of the bulk modulus

<table>
<thead>
<tr>
<th>Method</th>
<th>$P_t$ (GPa)</th>
<th>$\Delta E$ (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PBE</td>
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<tr>
<td>RPA</td>
<td>13.4</td>
<td>0.38</td>
</tr>
<tr>
<td>ACSOSESEX</td>
<td>7.2</td>
<td>0.21</td>
</tr>
<tr>
<td>RPAr1</td>
<td>8.8</td>
<td>0.25</td>
</tr>
<tr>
<td>NEO</td>
<td>8.0</td>
<td>0.23</td>
</tr>
<tr>
<td>DMC</td>
<td>14 ± 1</td>
<td>0.42</td>
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<tr>
<td>Expt</td>
<td>11.3–12.6</td>
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RPAr1: NEO in RPAr1  
ACSOSEX: NEO in RPAr1-ACSOSEX

Higher order terms (HOT)

Finite-order RPAr:

\[ \chi_\lambda = \hat{\chi}_\lambda + \hat{\chi}_\lambda f_{xc}^\lambda \hat{\chi}_\lambda + \hat{\chi}_\lambda f_{xc}^\lambda \hat{\chi}_\lambda f_{xc}^\lambda \hat{\chi}_\lambda + \ldots \]

The n-th order RPAr correction: The n-th order RPAr correction is:

\[ E_c = E_c^{\text{RPA}} + \sum_{n=1}^{\infty} \Delta E_c^{\text{RPAr}^{-n}} [f_{xc}] \Delta E_c^{\text{RPAr}^{-n}} [f_{xc}] = -\int_0^1 d\lambda \int_0^{2\pi} \frac{du}{2\pi} \langle V(\hat{\chi}_\lambda (iu) f_{xc}^\lambda (iu))^n \hat{\chi}_\lambda (iu) \rangle \]

For the n-th order correction analytical integration wrt. \( \lambda \) is not feasible.

Solution: an approximation based on the AC formula

\[ \Delta E_c^{\text{bRPA}} [f_{xc}] = \sum_{n=2}^{\infty} \Delta E_c^{\text{RPAr}^{-n}} \approx (1 - \hat{b}) \Delta U_c^{\text{RPAr}2} \approx \frac{1}{2} \Delta U_c^{\text{RPAr}2} \]

\( \Delta U_c^{\text{RPAr}2} \) can be estimated from the coupling-strength integral at \( \lambda = 1 \)

\[ E_c^{\text{HOT}} [f_{xc}] = E_c^{\text{RPA}} + \Delta E_c^{\text{RPAr}1} [f_{xc}] + \frac{1}{2} \Delta U_c^{\text{RPAr}2} [f_{xc}] \]
RPA renormalization contributions to the RPA correlation energy with NEO, rALDA and CP07 kernels for the uniform electron gas at $r_s = 4$. The HOT correction is plotted for comparison with RPAr2-4.
Another paradigm in condensed matter physics: Jellium surface

Metal surface energies can be treated within the jellium model: valence electrons are neutralized by a positive background.

The ACFDT formalism was used for the jellium surface energy by Pitarke and Eguiluz.


In 2000, Yan, Perdew and Kurth argued that the correction to RPA, although large (about +0.5 eV per electron), would tend to cancel almost completely out of energy differences for processes that conserve electron number.


RPA can be expected to much better for the surface energy than for the bulk energy of a solid.
Jellium surface

The surface energy of jellium is the work invested to create a unit area of new surface:

$$\sigma_{xc} = \int_{-\infty}^{\infty} dz \{ n(z) \varepsilon_{xc} ([n], z) - \bar{n} \varepsilon_{xc}^{\text{unif}} (\bar{n}) \theta(z_0 - z) \}$$

semi-infinite surface, bulk contributions cancel

$n(z)$: non-unif. electron density

The jellium slab which is the finite version of the semi-infinite jellium

Wavevector decomposition

The xc energy contributions to the slab and surface

Jellium slab:  \( \varepsilon_{xc} = \int_{0}^{\infty} d \left( \frac{q_{\parallel}}{2k_F} \right) \varepsilon_{q_{\parallel}}^{xc} \)

Jellium surface:  \( \sigma_{xc} = \int_{0}^{\infty} d \left( \frac{q_{\parallel}}{2k_F} \right) \gamma_{q_{\parallel}}^{xc} \)

\( q_{\parallel} \): wave vector parallel to surface

\( z \): direction of the surface
Jellium surface energy within RPA

Invoke the xc hole integral generally:

$$E_{xc} = \frac{1}{2} \int dr \int dr' \frac{n(r)n_{xc}(r, r')}{|r - r'|}$$

The $q_{||}$ wavevector analysis:

$$\gamma_{q_{||}}^{xc} = \frac{k_f}{8\pi} \int dz \int dz' n(z)v_{q_{||}}(|z - z'|) \frac{1}{|z - z'|} d\lambda \left[n_{q_{||}, \lambda}^{xc}(z, z') - \bar{n}_{q_{||}, \lambda}^{xc}(|z - z'|)\right]$$

The xc hole density comes from the fluctuation dissipation theorem:

$$n_{q_{||}, \lambda}^{xc}(z, z') = -\frac{1}{\pi n(z)} \int_0^\infty du \chi_{q_{||}, \lambda}(z, z', iu) - \delta(z - z')$$

The density response function comes from the Dyson equation with or without the kernel:

$$\chi_{q_{||}, \lambda}(z, z', \omega) = \chi_{q_{||}, \lambda}^0(z, z', \omega) + \int dz_1 \int dz_2 \chi_{q_{||}, \lambda}^0(z, z_1, \omega) \times \left[\lambda v_{q_{||}}(|z_1 - z_2|) + f_{q_{||}, \lambda}^{xc}(z_1, z_2, \omega)\right] \chi_{q_{||}, \lambda}(z_2, z', \omega)$$

NEO kernel for jellium surface

NEO-I

\[
 f^{\alpha,\text{NEO}}_{xc}(n^\uparrow, n^\downarrow, \tilde{R}, u) = -\frac{\lambda}{u} \sum_{\sigma} \left( \frac{n_{\sigma}}{n} \right)^2 \text{erfc}(au) \quad a = \sqrt{\tilde{c} \left( 1 - Z_{\sigma}^2 \right) k_{F,\sigma}^2}
\]

Problem for jellium: it produces a long-ranged exchange kernel in the tail of the density of a jellium surface, since \( Z \to 1 \) and \( k_F \to 0 \) there.
The relevance of the meta-GGA ingredients

\[
Z = \tau / \tau^W \quad \alpha = (\tau - \tau^W) / \tau^{\text{unif}} \quad t^{-1} = \tau / \tau^{\text{unif}}
\]

1) Regions of one- and two-electron density:
\[
\tau = \tau^W \quad Z = 1 \quad t^{-1} = 5s^2 / 3 \quad \alpha = 0 \quad \text{(single bonds)}
\]

2) Regions of uniform density:
\[
\tau^W = 0 \quad Z = 0 \quad t^{-1} = 1 \quad \alpha = 1 \quad \text{(metallic bonds)}
\]

3) Regions of density overlap between closed shells:
\[
\tau^W \approx 0 \quad Z \approx 0 \quad t^{-1} \to \infty \quad \alpha \to \infty \quad \text{(weak bonds)}
\]

\[
\tau / \tau^{\text{unif}} \sim n / n^{5/3} \quad \text{as} \quad n \to 0 \quad t^{-1} \to \infty
\]

Only \( \alpha \) can recognize these three regions and treat them differently.

Solution: make use of our knowledge of recent meta-GGA’s

Replace \( (1 - Z^2_\sigma) \) by \( 3\alpha_\sigma - 3\alpha^2_\sigma + \alpha^3_\sigma \). This leads to: NEO-II

\[
f_{xc}^{\alpha,\text{NEO}}(n \uparrow, n \downarrow, \bar{R}, u) = -\frac{\gamma}{u} \sum_\sigma \left( \frac{n_\sigma}{n} \right)^2 \text{erfc}(au)
\]

\[
a = \sqrt{\tilde{c} \left( \alpha_\sigma - 3\alpha^2_\sigma + 3\alpha^3_\sigma \right) k^2_{F,\sigma}}
\]

\[
\alpha_\sigma = \left( \tau_\sigma - \tau^W_\sigma \right) / \tau^{unif}_\sigma
\]

properly diverges in the tail

We do not refit parameter “c” when switch from NEO-I to NEO-II

A. Ruzsinszky, L.A. Constantin, and J.M. Pitarke, to be submitted.
Correlation kernel: NEO-III

\[ f_{\sigma\sigma'}^{\text{xc}}(\vec{r}, \vec{r}') = -\delta_{\sigma\sigma'} \left[ \text{erfc}\left\{ \frac{c}{1+br_s} \left( 1 - \frac{Z_\sigma^2}{k_{F\sigma}} \right) \right\} \right] \]

- Parameters \( b \) and \( c \) come from fitting to the uniform electron gas over a wide range of densities.

- The expression correctly reduces to the high-density limit.
NEO-II leaves unchanged the correct NEO behaviors for one- and two-electron densities \((\alpha = 0)\) and slowly-varying densities \((\alpha = 1 + O(\nabla^2))\), but kills of the kernel in the tail of the surface density \((\alpha \to \infty)\)

Comparison between \(k_F^2 (1-Z^2)\) and \(k_F^2 (\alpha-3\alpha^2+\alpha^3)\) at a jellium surface of bulk parameter \(r_s = 6\). The surface is at \(z = 0\), the bulk is at \(z < 0\), and the vacuum is at \(z \to \infty\)
Wavevector resolution of the correlation energy for the jellium slab and surface

Two-dimensional wavevector analysis of the correlation energy per particle of the jellium slab with width $= 2.23\lambda_F$ ($\lambda_F=2\pi/k_F$), and $rs = 2.07$ for ISTLS, RPA, and the NEO kernels.

$rs = 2.07$ corresponds to Al(100) surface

Two-dimensional wavevector analysis of the jellium surface energy with $rs = 2.07$ for ISTLS, RPA, and the NEO kernels. Surface energy is the area below the curves.
Jellium correlation surface energies with different approaches at background parameter \( \text{rs} = 2.07 \)

<table>
<thead>
<tr>
<th>Method</th>
<th>Correlation surface energy (( \text{erg/cm}^2 ))</th>
</tr>
</thead>
<tbody>
<tr>
<td>ISTLS</td>
<td>714</td>
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<tr>
<td>RPA</td>
<td>743</td>
</tr>
<tr>
<td>NEO-I-Gaussian</td>
<td>687</td>
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<tr>
<td>NEO-I-erf</td>
<td>723</td>
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<td>NEO-II</td>
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<td>NEO-III</td>
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<tr>
<td>LDA</td>
<td>293</td>
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<tr>
<td>PBEsol</td>
<td>604</td>
</tr>
<tr>
<td>DMC</td>
<td>697±45</td>
</tr>
</tbody>
</table>
Exchange-correlation and correlation energies along the “z” direction in the jellium slab

![Graphs showing exchange-correlation and correlation energies along the “z” direction in the jellium slab.]}
Conclusions

• RPA energies require a substantial kernel correction.

• The standard ALDA kernel used in TDDFT is as bad as no kernel at all.

• The kernel can be modelled by satisfaction of exact constraints, e.g.; the NEO kernel.

• The kernel can be included exactly to all orders, but RPA renormalization is less expensive and almost as accurate.

• The kernel has a major effect on the correlation energy of bulk jellium, but much less effect on the jellium surface energy.
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