Koopmans’ condition for density-functional theory

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In approximate Kohn-Sham density-functional theory, self-interaction manifests itself as the dependence of the energy of an orbital on its fractional occupation. This unphysical behavior translates into qualitative and quantitative errors that pervade many fundamental aspects of density-functional predictions. Here, we first examine self-interaction in terms of the discrepancy between total and partial electron removal energies, and then highlight the importance of imposing the generalized Koopmans’ condition—that identifies orbital energies as opposite total electron removal energies—to resolve this discrepancy. In the process, we derive a correction to approximate functionals that, in the frozen-orbital approximation, eliminates the unphysical occupation dependence of orbital energies up to the third order in the single-particle densities. This non-Koopmans correction brings physical meaning to single-particle energies; when applied to common local or semilocal density functionals it provides results that are in excellent agreement with experimental data—while providing an explicit total energy functional that preserves or improves on the description of established structural properties.

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I. INTRODUCTION

Density-functional approximations,1 which account for correlated electron interactions via an explicit functional $E_{xc}$ of the electronic density $\rho$, provide very good predictions of total energy differences for systems with nonfractional occupations.2 One of the notable successes of local and semilocal functionals overestimate or entire depletion of frontier orbitals. In quantitative terms, the local-spin-density (LSD) approximation and semilocal generalized-gradient approximations (GGAs) predict the energy differences of such reactions, namely, the electron affinity

$$A_N = E_N - E_{N+1}$$

and the first ionization potential

$$I_N = E_{N-1} - E_N$$

(1)

(2)

(where $E_N$ stands for the ground-state energy of the $N$-electron system) with a precision of a few tenths of an electron volt, which compares favorably to that of more expensive wave function methods.3

Considering the excellent performance of density-functional approximations in predicting total ionization energies, it is surprising to discover that the same theories fail in describing partial ionization processes. As a matter of fact, local and semilocal functionals overestimate (by as much as 40%) the absolute energy difference per electron

$$A_N = \left. \frac{dE_M}{dM} \right|_{M=N^+}$$

(3)

of an infinitesimal electron addition and underestimate with the same error differential energy changes

$$I_N = \left. \frac{dE_M}{dM} \right|_{M=N^+}$$

(4)

upon electron removal.

Analytically, the discrepancy between total and partial differential ionization energies manifests itself into the convexity of $E_{LSD}^N$ as a function of $N$ (Fig. 1) while the exact ground-state energy $E_N$ versus $N$ is known to be described by a series of straight-line segments with positive derivative dis-

![FIG. 1. (Color online) Convexity of the LSD ground-state energy as a function of the electron number, which results from the discrepancy between total and partial differential electron removal energies, i.e., $\delta^LSD_N < I_N < A_{N} < A_{N-1}$.](image-url)
continuities at integer values of \( N \).\(^4\) (For simplicity, we restrict the entire discussion to the case of the LSD functional; semilocal GGA functionals exhibit identical trends.)

It is important to note that the discrepancy described above can be related to the incorrect analytical behavior of the LSD chemical potential \( \mu_N^{\text{LSD}} \), i.e., the Lagrange multiplier associated to particle-number conservation in the near-zero temperature grand-canonical minimization of the total energy, \( \min_{\rho}(E[\rho]-\mu \int d\rho(r)) \),\(^4\) which can be interpreted physically as the opposite electronegativity of the system.\(^5,6\)

\[
\mu_N = -\chi_N = -\frac{A_N + \mathcal{I}_N}{2} \tag{5}
\]

and determines the direction and magnitude of electron transfer between separated molecular fragments.\(^4\) In fact, the exact dependence of the chemical potential \( \mu_N \) when \( N \) varies through an integer particle number \( Z \) is known to be

\[
\mu_N = \left\{ \begin{array}{ll}
-1 \quad & Z-1 < N < Z \\
-\frac{1}{2}(A_Z + I_Z) & N = Z \\
-A_Z & Z < N < Z + 1,
\end{array} \right.
\tag{6}
\]

where \( \chi_N \) denotes the Mulliken electronegativity\(^9\) of the \( N \)-electron system.\(^4,8\)

In related physical terms, the discrepancy between ionization energies is often interpreted as arising from electron self-interaction that causes the removal of a small fraction of an electron from a filled electronic state to be energetically less costly, in absolute value, than its addition to the corresponding empty state.\(^10\)

This self-interaction error is at the origin of important quantitative and qualitative failures that pervade crucial aspects of electronic-structure predictions.\(^11\) Consider, for example, the dissociation of a cation dimer \( X^+_2 \) in the infinite interatomic separation limit. Here, the energy cost of removing a small electron fraction from \( X \) is lower than the energy gained in adding an electron fraction to \( X^+ \). As a result, a portion of the electronic charge will transfer from \( X \) to \( X^+ \), eventually leading to the split-charge configuration \( 2X^{1/2} \), with a total energy that is correspondingly overstabilized with respect to the exact solution.

Related self-interaction artifacts explain other important failures of local and semilocal functionals in predicting electron-transfer processes,\(^12,13\) electronic transport,\(^14\) molecular adsorption,\(^15,16\) reaction barriers and energies,\(^17-19\) and electrical polarization in extended systems.\(^20-23\) Self-interaction is also connected to the underestimation of the differential energy gap \( E_{\text{ion}}^{\text{DOS}} = \mathcal{I}_N - A_N \) of molecules, semiconductors, and insulators within LSD and GGAs.\(^24-28\)

This work is organized as follows. First, we reexamine the Perdew-Zunger (PZ) one-electron self-interaction correction in terms of total and differential ionization energies. Then, after deriving an exact measure of the unphysical curvature of \( E_N \) as a function of \( N \) based on the generalized Koopmans’ theorem, we introduce a functional that minimizes self-interaction errors by enforcing Koopmans’ condition, thereby largely eliminating the discrepancy between total and differ-

\[ E^\text{PZ} = E^\text{LSD} + \sum_{i\sigma} (-E_{i\sigma}[\rho_{i\sigma}] - E_{xc}[\rho_{i\sigma}]), \tag{7} \]

\[ E^\text{PZ} = E^\text{LSD} + \sum_{i\sigma} \Xi_{i\sigma}, \tag{8} \]

\[ \tilde{h}^\text{PZ} = \tilde{h}^\text{LSD}[\rho] - \tilde{v}_{i\sigma}[\rho_{i\sigma}] - \tilde{u}_{xc,[\rho_{i\sigma}]], \tag{9} \]

where \( \rho_{i\sigma}(\mathbf{r}) \) denotes the orbital density (\( \sigma = \pm \frac{1}{2} \) represents the spin), \( \tilde{v}_{i\sigma}[\rho] = \delta E_{i\sigma}[\rho] / \delta \rho_{i\sigma}(\mathbf{r}) \) stands for the spin-dependent exchange-correlation potential, and the PZ one-electron correction contributions are defined as \( \Xi_{i\sigma} = -E_{i\sigma}[\rho_{i\sigma}] - E_{xc}[\rho_{i\sigma}] \).

The effect of the PZ self-interaction correction on the partial electron removal energies of an isolated carbon atom is depicted in Fig. 2. We observe that the LSD differential ionization potential \( \mathcal{I}_N^{\text{LSD}} / H^9262 \) and electron affinity \( A_N^{\text{LSD}} / H^9264 \) deviate by more than 5 eV from the experimental total removal energy, whereas their average is in close agreement with experiment. The PZ correction improves the precision of the predicted ionization potential \( \mathcal{I}_N^{\text{PZ}} / H^9262 \), reducing the error to less than 0.4 eV. The inaccuracy of the PZ electron affinity \( A_N^{\text{PZ}} / H^9261 \) remains almost unchanged due to the fact that the PZ one-electron correction vanishes for empty states (the slight variation of
the differential electron affinity from 17.79 to 17.65 eV is only due to the self-consistent reconfiguration of the occupied manifold). Consequently, the PZ functional achieves a substantial but only partial correction of self-interaction, reducing the unphysical convexity, i.e., the discrepancy \( A_{N-1} - \mathcal{I}_N \), by a factor of two.

Another notable feature in Fig. 2 is the overestimation of the average of the ionization potential \( \mathcal{PZ}_N \) and electron affinity \( \mathcal{A}_N \). Since the average \( \frac{1}{2} (\mathcal{A}_N + \mathcal{PZ}_N) \) roughly approximates \( I_N \), this deviation translates into overestimated total ionization energies and excessively negative ground-state energies \( E_N^{\text{PZ}} = -\sum_{\alpha=1}^N \mathcal{PZ}_N \). (Semiempirical relations between total energies and low-order ionization potentials have been also evidenced by Pucci and March in Ref. 40.) In the case of isolated atoms, the energy underestimation improves on the description of electron correlation, bringing PZ ground-state energies in remarkable agreement with experiment. However, in the more general case of many-electron polyatomic systems, PZ ground-state energies are found to be overcorrelated, yielding inaccurate dissociation energies and excessively short bond lengths.

Various downscaling methods have been proposed to correct the above trends. Nevertheless, the performance of such schemes is inherently limited by the fact that downscaling the PZ correction impairs the accuracy of differential ionization energies. For example, it is seen in Fig. 2 that a functional \( E_{\alpha}^{\text{PZ}} \), in which the self-interaction correction is linearly downscaled by a factor \( \alpha^{\text{PZ}} \)

\[
E_{\alpha}^{\text{PZ}} = E_{\text{LSD}} + \alpha^{\text{PZ}}(E^{\text{PZ}} - E^{\text{LSD}})
\]

cannot counterbalance the deviation of \( \mathcal{I}_N^{\text{PZ}} = \frac{1}{2} (\mathcal{A}_N^{\text{PZ}} + \mathcal{PZ}_N^{\text{PZ}}) \) without altering the precision of \( I_N^{\text{PZ}} \). More sophisticated downscaling methods exhibit identical trends.

In the next sections, we will derive a correction that cancels self-interaction errors for systems with fractional occupations without affecting the precision of total energy differences and equilibrium structural properties for systems with nonfractional occupations.

### B. Measure of self-interaction

The initial conceptual step in the construction of the self-interaction correction is to set forth a quantitative measure of self-interaction errors valid in the many-electron case—i.e., beyond the one-electron approximation of the Perdew-Zunger correction. With this self-interaction measure in hand, we will construct an improved self-interaction correction functional, working first in the simplified picture where electronic orbitals are kept frozen (Sec. II C), and then considering orbital relaxation as the next stage of refinement (Sec. II E).

Before doing so, to derive this measure, we start from the physical intuition that self-interaction relates to the unphysical variation of the energy of an orbital \( \varepsilon_{\alpha}(f) \) as a function of its own occupation \( f \). Hence, a necessary nonself-interaction condition can be written as

\[
\frac{d \varepsilon_{\alpha}(f)}{df} \bigg|_{f=0} = 0, \quad 0 \leq f \leq 1.
\]

(Here, \( f \) stands for a variable occupation while \( \varepsilon_{\alpha} \) denotes the orbital occupation that enters into the expression of the total energy functional.) Invoking Janak’s theorem, this necessary condition on orbital-energy derivatives can be restated as a criterion on the curvature of the total energy

\[
\frac{d^2 E_{\alpha}(f)}{df^2} \bigg|_{f=0} = 0, \quad 0 \leq f \leq 1,
\]

where \( E_{\alpha}(f) \) is the total energy \( E \) minimized under the constraint \( f_{\alpha}=f \) (leaving all other occupations unchanged). Thus, in the absence of self-interaction, the total energy does not display any curvature upon varying occupations. As a corollary, any self-interaction-free functional satisfies the linearity condition

\[
\Delta E_{\alpha} = -f_{\alpha} \frac{d E_{\alpha}(f)}{df} \bigg|_{f=0}, \quad 0 \leq f \leq 1,
\]

where

\[
\Delta E_{\alpha} = E_{\alpha}(0) - E_{\alpha}(f_{\alpha})
\]

denotes the removal energy of the orbital \( \psi_{\alpha} \).

Invoking once more Janak’s theorem, the above nonself-interaction condition yields the general Koopmans’ theorem

\[
\Delta E_{\alpha} = -f_{\alpha} \varepsilon_{\alpha}(f) + \Delta E_{\alpha}, \quad 0 \leq f \leq 1
\]

(The equivalence between Eq. (11) and Eq. (15) highlights the significance of Koopmans’ theorem for the quantitative assessment of self-interaction.) It is then convenient to measure self-interaction in terms of the energies \( \Pi_{\alpha} \) first introduced by Perdew and Zunger (see Sec. II D in Ref. 10) in analyzing discrepancies between orbital energies and vertical ionization energies.

\[
\Pi_{\alpha} = f_{\alpha} \varepsilon_{\alpha}(f) + \Delta E_{\alpha},
\]

Because the self-interaction energies \( \Pi_{\alpha} \) quantify deviations from the Koopmans’ linearity [Eq. (15)], they will be termed here non-Koopmans energies—the same terminology as that employed in Ref. 10 and 37.

Making then use of Slater’s theorem

\[
\Delta E_{\alpha} = -\int_{0}^{f_{\alpha}} df \varepsilon_{\alpha}(f)
\]

the non-Koopmans energy can be rewritten as

\[
\Pi_{\alpha} = \int_{0}^{f_{\alpha}} df \{ \varepsilon_{\alpha}(f) - \varepsilon_{\alpha}(f') \}.
\]

In this form, the energy \( \Pi_{\alpha}(f) \) is clearly seen to correspond to the integrated change of the orbital energy upon varying the orbital occupation—in particular, it is straightforward to verify that \( \Pi_{\alpha}(f)=0 \) if the orbital energy \( \varepsilon_{\alpha}(f') \) does not vary with the orbital occupation \( f' \).

Using the measure defined in Eq. (16), the nonself-
interaction criterion [Eq. (11)] can be restated exactly as a
\begin{equation}
\Pi_{\alpha\sigma}(f) = 0, \quad 0 \leq f \leq 1.
\end{equation}
Equation (19) is of central importance in this work; it pro-
vides a simple quantitative criterion in terms of a rigorous
nonlinearity measure to assess and correct self-interaction
errors. The suggestion of using non-Koopmans corrections to
minimize self-interaction has been recently introduced, in
preliminary form, by Dabo, Cococcioni, and Marzari in Ref.
49, and, in heuristic form, by Lany and Zunger in Ref. 36. In
the context of range-separated hybrid functionals, the impor-
tance of satisfying Koopmans’ equalities for frontier molecu-
lar orbitals has also been evidenced by Stein, Kronik and
Baer in Ref. 13. The consequences of Koopmans’ condition
are discussed in the next sections.

C. Bare non-Koopmans correction
On the basis of the above quantitative analysis, our objec-
tive now is to linearize the dependence of the total energy as
a function of orbital occupations by modifying the expres-
sion of the energy functional in order to cancel the non-
Koopmans terms \(\Pi_{\alpha\sigma}(f)\). To render this complex problem
tractable, we first consider the restricted case where all orbit-
als are frozen while the occupation of one of them is chang-
ing in the course of a fictitious ionization process (the
frozen-orbital approximation). Within this paradigm, Eq.
(19) becomes the restricted Koopmans’ condition
\begin{equation}
\Pi_{\alpha\sigma}^{\text{r}}(f) = 0, \quad 0 \leq f \leq 1,
\end{equation}
where the superscript \(\text{r}\) stands for unrelaxed. Here, the
frozen-orbital non-Koopmans energy \(\Pi_{\alpha\sigma}^{\text{r}}\) is defined as
\begin{equation}
\Pi_{\alpha\sigma}^{\text{r}}(f) = \int_{0}^{f} df'[\epsilon_{\alpha\sigma}^{\text{u}}(f') - \epsilon_{\alpha\sigma}^{\text{u}}(f)],
\end{equation}
where \(\epsilon_{\alpha\sigma}^{\text{u}}(f)\) is the unrelaxed orbital energy calculated keep-
ing all the orbitals frozen while setting \(f_{\alpha\sigma}\) to be \(f\). We un-
derscore that in the specific case of one-electron systems and
for \(f=0\), Eq. (20) yields the one-electron self-interaction con-
dition of Perdew and Zunger\(^{10}\)
\begin{equation}
\Xi_{\alpha\sigma} = 0.
\end{equation}
The restricted Koopmans’ condition is thus seen to encom-
pass the Perdew-Zunger condition, thereby representing a
more comprehensive criterion for assessing and correcting
self-interaction errors.

Expectedly, satisfying the restricted Koopmans’ condition
is exactly equivalent to fulfilling the restricted Koopmans’
theorem
\begin{equation}
\Delta E_{\alpha\sigma}^{\text{u}} = - f_{\alpha\sigma} \epsilon_{\alpha\sigma}^{\text{u}}(f), \quad 0 \leq f \leq 1,
\end{equation}
where
\begin{equation}
\Delta E_{\alpha\sigma} = E_{\alpha\sigma}^{\text{u}}(0) - E_{\alpha\sigma}^{\text{u}}(f_{\alpha\sigma})
\end{equation}
denotes the unrelaxed electron removal energy.

An example of a functional that exhibits a linear frozen-
orbital energy dependence is provided by the Hartree-Fock
(HF) theory, generalized to fractional occupations. Indeed, at
the HF level, one can verify that
\begin{equation}
\Pi_{\alpha\sigma}^{\text{HF}}(f) = 0, \quad 0 \leq f \leq 1
\end{equation}
due to the fact that the expectation value \(\epsilon_{\alpha\sigma}^{\text{HF}}(f)\) does not de-
pend on \(f\).

For functionals that do not satisfy the restricted Koop-
mans’ condition [Eq. (20)], the unrelaxed electron removal
can only be expressed in terms of the restricted Slater
integral
\begin{equation}
\Delta E_{\alpha\sigma}^{\text{u}} = - \int_{0}^{f_{\alpha\sigma}} df \epsilon_{\alpha\sigma}^{\text{u}}(f)
\end{equation}
(note that this relation is satisfied by any functional whether
it is subject or not to self-interaction errors).

These considerations allow us to introduce our corre-
correct function to satisfy Koopmans’ condition; at frozen
orbital, it is obtained by replacing Slater terms—Eq. (26),
where single-particle energies are function of their own
occupation—with Koopmans terms—Eq. (23), where single-
particle energies do not depend on occupations. In par-

icular, we evaluate here the Koopmans terms at a given orbital oc-
cupation \(f = f_{\text{ref}}\) that defines the reference transition state
the determination of the reference occupation \(f_{\text{ref}}\) shall be ex-
plained on the basis of Slater’s approximation and exchange-
correlation hole arguments in Sec. II F. Explicitly, in the
case of the LSD functional, the NK self-interaction correction
to the energy is defined as
\begin{equation}
E_{\alpha\sigma} = E_{\alpha\sigma}^{\text{LSD}} - \sum_{i\alpha} \left[ - f_{i\alpha} \epsilon_{i\alpha}^{\mu\text{LSD}}(f_{\text{ref}}) + \int_{0}^{f_{i\alpha}} df \epsilon_{i\alpha}^{\mu\text{LSD}}(f) \right],
\end{equation}
where the negative sign in front of the sum follows from the
convention that ionization energies are positive. Rewriting
Eq. (27) in terms of the frozen-orbital non-Koopmans ener-
gies of Eq. (21), we obtain
\begin{equation}
E_{\alpha\sigma} = E_{\alpha\sigma}^{\text{LSD}} + \sum_{i\alpha} \Pi_{i\alpha}^{\text{LSD}}(f_{\text{ref}}),
\end{equation}
where the non-Koopmans corrective terms can now be recast
into the explicit functional form
\begin{equation}
\Pi_{i\alpha}^{\text{LSD}}(f_{\text{ref}}) = f_{i\alpha}(2f_{\text{ref}} - f_{i\alpha}) E_{\alpha\sigma}^{\text{LSD}}[n_{i\sigma}] - E_{\alpha\sigma}^{\text{LSD}}[\rho] + E_{\alpha\sigma}^{\text{LSD}}[\rho] + \int d\rho_{i\alpha}(r) v_{\alpha\sigma}^{\text{LSD}}(r) \rho_{i\alpha}^{(\text{ref})}(r) dr,
\end{equation}
where \(n_{i\alpha}(r) = |\psi_{i\alpha}(r)|^{2}\) (\(\rho_{i\alpha}(r) = f_{i\alpha} n_{i\alpha}(r)\)). Here, the
electronic density \(\rho_{i\alpha}^{(\text{ref})}(r)\) stands for the reference transition-state
density
\begin{equation}
\rho_{i\alpha}^{(\text{ref})}(r) = f_{i\alpha} n_{i\alpha}(r) + \sum_{j\sigma'} f_{j\sigma'} n_{j\sigma'}(r),
\end{equation}
\begin{equation}
\rho(r) = (f_{\text{ref}} - f_{i\alpha}) n_{i\alpha}(r).
\end{equation}
We now derive the expression of the orbital-dependent NK
Hamiltonian. To this end, we first calculate the functional
derivative of the energy contribution $\Pi^\text{LSD}_{\text{ref}}(f_{\text{ref}})$ with respect to variations of the orbital density $\rho_{\text{ref}}$. The expression of the functional derivative reads

$$\frac{\delta \Pi^\text{LSD}_{\text{ref}}}{\delta \rho_{\text{ref}}(\mathbf{r})} = (f_{\text{ref}} - f_{\text{ref}}) \psi_{\text{H}}(\mathbf{r} ; [\rho_{\text{ref}}]) + \psi^\text{LSD}_{\text{xc,ref}}(\mathbf{r} ; [\rho_{\text{ref}}])$$

$$- \psi^\text{LSD}_{\text{xc,ref}}(\mathbf{r} ; [\rho]) + \psi_{\text{ref,ref}}(\mathbf{r}).$$

(32)

In Eq. (32), the potential $\psi_{\text{ref,ref}}$ in the LSD functional derivative reads

$$\psi_{\text{ref,ref}}(\mathbf{r}) = \int d\mathbf{r}' \psi^\text{LSD}_{\text{xc,ref}}(\mathbf{r}' , [\rho_{\text{ref}}]) n_{\text{ref}}(\mathbf{r}')$$

$$- \int d\mathbf{r}' d\mathbf{r}'' \psi^\text{LSD}_{\text{xc,ref}}(\mathbf{r}', \mathbf{r}'' ; [\rho_{\text{ref}}]) n_{\text{ref}}(\mathbf{r}') n_{\text{ref}}(\mathbf{r}''),$$

(33)

where $\psi^\text{LSD}_{\text{xc,ref}}(\mathbf{r}' , [\rho_{\text{ref}}])$ is the exchange-correlation contribution to $\psi^\text{LSD}_{\text{xc,ref}}(\mathbf{r} , \mathbf{r}')$. As a final result, the orbital-dependent NK Hamiltonian can be cast in the form

$$\hat{H}^\text{NK} = \hat{H}^\text{LSD}[\rho_{\text{ref}}] + \psi_{\text{ref,ref}} + \psi_{\text{ref,ref}},$$

(35)

where $\psi^\text{LSD}_{\text{ref,ref}}$ stands for the cross-derivative potential

$$\psi^\text{LSD}_{\text{ref,ref}}(\mathbf{r}) = \sum_{j\neq i} \frac{\delta \Pi^\text{LSD}_{\text{ref}}}{\delta \rho_{\text{ref}}(\mathbf{r})}.$$

(36)

In a nutshell, the NK Hamiltonian consists of the uncorrected LSD Hamiltonian calculated at the reference density $\hat{H}^\text{LSD}[\rho_{\text{ref}}]$ with the addition of two variational potentials. The first additional term $\psi_{\text{ref,ref}}$ results from the variation of the reference density as a function of $\rho_{\text{ref}}$ while the second term $\psi_{\text{ref,ref}}$ springs from the cross dependence of the non-Koopmans corrective terms. The effect of the $\psi_{\text{ref,ref}}$ and $\psi_{\text{ref,ref}}$ contributions that arise as byproducts of variationality is analyzed in the next section.

D. Comparative assessment

In this section, we assess the performance of the NK self-interaction correction, particularly focusing on the effect of variational terms on the accuracy of NK orbital predictions—i.e., on the cancellation of the unrelaxed frozen-orbital self-interaction measure $\Pi^\text{LSD}_{\text{ref,ref}}(f)$. One simple and probably the most direct way to evaluate the influence of $\psi_{\text{ref,ref}}$ and $\psi_{\text{ref,ref}}$ is to introduce a nonvariational orbital-energy scheme, the NK method, that consists of freezing the dependence of the reference transition-state densities and the cross dependence of corrective energy terms, thereby eliminating $\psi_{\text{ref,ref}}$ and $\psi_{\text{ref,ref}}$ contributions to the effective potential. Computed NK and NK0 orbital levels can then be compared for the direct assessment of $\psi_{\text{ref,ref}}$ and $\psi_{\text{ref,ref}}$ errors. Explicitly, the NK0 Hamiltonian can be written as

$$\hat{h}_{\text{NK}0} = \hat{h}_{\text{LSD}}[\rho] + \frac{\delta \Pi^\text{LSD}_{\text{ref}}}{\delta \rho_{\text{ref}}},$$

(37)

$$\hat{h}_{\text{NK}0} = \hat{h}_{\text{LSD}}[\rho_{\text{ref}}].$$

(38)

In the NK0 optimization scheme, the Hamiltonian given by Eq. (38) is employed to propagate orbital degrees of freedom at fixed $\rho_{\text{ref}}$. Reference transition-state densities are then updated according to Eq. (30). The procedure is iterated until self-consistency.

Due to the loss of variationality, the obvious practical limitation of the nonvariational NK0 orbital-energy method is that it cannot provide total energies and interatomic forces. However, NK0 is of great utility in evaluating the intrinsic performance of the NK correction. In itself, the NK0 formulation is also useful in determining orbital energy properties that are particularly affected by $\psi_{\text{ref,ref}}$ and $\psi_{\text{ref,ref}}$ errors.

Focusing now on computational predictions, the occupation dependencies of the LSD, HF, PZ, and NK unrelaxed orbital energies

$$e^i_{\text{ref}} = \left. \frac{dE^i_{\text{ref}}(f^i)}{df^i} \right|_{f^i_{\text{ref}}}$$

(39)

of the highest atomic orbital of carbon are depicted in Figs. 3 and 4(a). The salient feature of the LSD graph is the large variation of the orbital energy from $-19.40$ to $-6.15$ eV, reflecting the strong nonlinearity of the corresponding unrelaxed ionization curve. The PZ variation is found to be twice lower than for LSD, confirming the trends observed in Sec. II A. In contrast, the HF unrelaxed ionization curve exhibits a perfectly linear behavior (i.e., the unrelaxed orbital energy remains constant). This trend is closely reproduced by the NK functional [Fig. 4(a)]; on the scale of LSD residual non-Koopmans errors, the eye can barely distinguish any devia-
Furthermore, a Taylor series expansion of the error is of the fourth order in orbital densities.

The above observation is due to the fact that the variational contribution \( w_{\text{ref},\sigma} \) affects the orbital energy \( \epsilon_{\sigma} \) indirectly, i.e., only through the self-consistent response of the orbital densities since \( \langle \psi_{\sigma}\vert w_{\text{ref},\sigma}\vert \psi_{\sigma} \rangle = 0 \) at self-consistency. Furthermore, a Taylor series expansion of \( w_{\text{SD},\sigma} \) reveals that \( \langle \psi_{\sigma}\vert w_{\text{SD},\sigma}\vert \psi_{\sigma} \rangle \) does not cause notable departure from the linear Koopmans' behavior. In quantitative terms, the dominant term in the expansion of the residual NK self-interaction error is of the fourth order in orbital densities.

\[
\Pi_{\sigma\tau}^{\text{NK}}(f) = \frac{1}{4} \sum_{j \neq \sigma} f_{\sigma\tau}(2f_{\sigma\tau} - f_{j\sigma})(2f - f_{j\sigma})f_{\tau\sigma} \\
\times \int dr_{1234} f_{\text{SD}}(\rho_{\text{ref}})[1 - \rho_{\text{ref}}] n_{\sigma\tau}(r_1)n_{\sigma\tau}(r_2)n_{j\sigma}(r_3)n_{j\sigma}(r_4) + \cdots
\]

(40)

(41)

Despite the very good accuracy of the NK correction, direct confrontation with NK results—for which the non-Koopmans measure \( \Pi_{\sigma\tau}^{\text{NK}}(f) \) [Eq. (21)] is obviously canceled for any value of \( f \)—reveals that NK tends to underestimate orbital energies with deviations of 0.5–1.5 eV that gradually increase with \( f_{\text{ref}} \) [Figs. 4(a) and 4(b)]. The practical consequences of this observation will be discussed in Sec. III.

As a conclusion of this preliminary performance evaluation, the NK frozen-orbital correction results in a considerable reduction of residual errors \( \Pi_{\sigma\tau}^{\text{NK}}(f) \), bringing density-functional approximations in nearly exact agreement with the frozen-orbital linear trend while exhibiting a slight tendency to underestimate orbital energies. In the next sections, we present an extension of the NK correction beyond the frozen-orbital paradigm.

E. Screened non-Koopmans correction

Having derived and examined the bare non-Koopmans frozen-orbital correction, we now turn to the analysis of self-interaction for relaxed orbitals. The effect of the NK correction on relaxed partial ionization energies for carbon is illustrated in Fig. 5. The first important observation is that the NK correction decreases the unphysical convexity of \( E_N \), reducing the unphysical discrepancy \( A_{N-1} - \tilde{I}_N \) by a factor of 6 regardless of the reference occupation \( f_{\text{ref}} \).

The second notable feature is the fact that the NK correction reverts the convexity trend (i.e., \( A_N^{\text{NK}} > A_{N-1}^{\text{NK}} \)), transform-
KOOPMANS’ CONDITION FOR DENSITY-FUNCTIONAL THEORY

...ing the convex dependence of $E_N$ into a piecewise concave curve. In fact, bearing in mind that relaxation contributions to $E_N$ are always negative, it is relatively straightforward to show that any function that satisfies the restricted Koopmans’ condition [Eq. (20)] is piecewise concave. Consequently, in contrast to the $a$-$P$Z functional introduced in Sec. II A, there always exists a value of the coefficient $a_N^\text{NK}$ ($0 \leq a_N^\text{NK} \leq 1$) for which the $\alpha_N^\text{NK}$ functional

$$E_N^{\text{NK}} = E_N^{\text{LSD}} + a_N^\text{NK}(E_N - E_N^{\text{LSD}})$$

(42)

restores the agreement between $A_N^{\text{NK}}$ and $I_N^{\text{NK}}$. At this crossing point, the $\alpha_N^\text{NK}$ relaxed ionization curve is in close agreement with the exact linear trend described by the generalized Koopmans’ condition (see Sec. III A).

Making the approximation that ionization energies vary linearly with $\alpha_N^\text{NK}$, the value of the coefficient for which the crossing $A_N^{\text{NK}} = I_N^{\text{NK}}$ occurs can be estimated as

$$a_N^\text{NK} = \frac{A_N^{\text{LSD}} - I_N^{\text{LSD}}}{(A_N^{\text{LSD}} - I_N^{\text{LSD}}) - (A_N^{\text{NK}} - I_N^{\text{NK}})}.$$  

(43)

This initial estimate can then be refined using the secant-method recursion

$$a_{n+1} = a_n + \frac{1 - a_n)(A_N^{\text{NK}} - I_N^{\text{NK}})}{(A_N^{\text{NK}} - I_N^{\text{NK}}) - (A_N^{\text{NK}} - I_N^{\text{NK}})}.$$  

(44)

In practice, it is observed that only one or two iterations are sufficient to determine the value of the coefficient $a_N^\text{NK} = \lim_{n \to \infty} a_n$ and bring the difference $A_N^{\text{NK}} - I_N^{\text{NK}}$ below 0.2 eV.

Physically, the coefficient $a_N^\text{NK}$ is directly related to the magnitude of orbital relaxation upon electron removal. Indeed, $\alpha_N^\text{NK}$ can be viewed as a screening coefficient whose value is close to 1 for weakly relaxing ionized systems (since $E_N^{\text{NK}}$ is already piecewise linear in the absence of orbital relaxation) and is found to be small when relaxation is strong (see Sec. III A). The corrective factor $\alpha_N^\text{NK}$ introduced here is thus endowed with a clear physical interpretation.

As a final note, it should be emphasized that we have adopted here a simple picture of orbital relaxation through the coefficient $\alpha_N^\text{NK}$, which can be viewed as a uniform and isotropic screening factor. More elaborate screening functions could be employed (at the price of computational complexity). Such accurate screening approaches provide promising extensions of the $\alpha_N^\text{NK}$ method and represent an interesting subject for future studies.

F. Reference transition-state occupation

We now proceed to examine the influence of the reference occupation on the accuracy of calculated total electron removal energies. To this end, we compare the average of the $\alpha_N^\text{NK}$ differential electron removal energies (that closely approximates the total ionization potential) of carbon for $f_{\text{ref}}$ different from and equal to $\frac{1}{2}$ in Figs. 5(a) and 5(b), respectively. In the former case, the diagram indicates that the average

$$\frac{1}{2} (A_N^{\text{NK}} + I_N^{\text{NK}}) = I_N^{\text{NK}}$$

(45)

deviates significantly from its LSD counterpart, whereas such deviations do not occur in the case $f_{\text{ref}} = \frac{1}{2}$. In fact, from Eqs. (2), (28), and (42), it can be shown that

$$I_N^{\text{NK}}(f_{\text{ref}}) = I_N^{\text{LSD}} - \alpha_N^\text{NK} \Pi_N^{\text{LSD}}(f_{\text{ref}})$$

(46)

(neglecting orbital reconfiguration). Then, substituting the (restricted) Slater’s approximation

$$\Delta E_{\text{LSD}}(f_{\text{ref}}) = -\epsilon_{\text{LSD}}^{\text{ref}}(f_{\text{ref}} = 1)$$

(47)

into the definition of the non-Koopmans energy contributions [Eq. (21)], one can demonstrate that

$$\Pi_N^{\text{LSD}}(f_{\text{ref}} = 1) \approx 0$$

(48)

to arrive at the relation

$$I_N^{\text{NK}}(f_{\text{ref}} = 1) \approx I_N^{\text{SD}}.$$  

(49)

This result explains the accuracy of the $\alpha_N^\text{NK}$ correction in predicting total electron removal energies with the reference occupation $f_{\text{ref}} = \frac{1}{2}$.

It should be noted that in the particular case of one-electron systems, $f_{\text{ref}} = \frac{1}{2}$ is not the only possible choice for the reference occupation. Indeed, due to the fact the Hartree and exchange-correlation potentials vanish for empty solitary orbitals, the approximation

$$\Delta E_{\text{LSD}} = -\epsilon_{\text{LSD}}^{\text{ref}}(f = 0)$$

(50)

also holds; hence, the solution $f_{\text{ref}} = 0$ is equally valid. In fact, for one-electron systems, $f_{\text{ref}} = 0$ leads to the exact one-electron Hamiltonian and the exact solution to the one-electron Schrödinger problem.

As an alternative to transition-state arguments, the value of $f_{\text{ref}}$ can be justified by inspecting the sum rule satisfied by the exchange-correlation hole (xc hole).10,45,46 The xc hole $h_{\text{xc}}$ is defined by the relation

$$E_{\text{xc}} = \frac{1}{2} \int dr dr' \rho(r) h_{\text{xc}}(r, r') \frac{1}{|r - r'|}$$

(51)

and can be explicitly written through the adiabatic connection formalism as

$$\rho(r) h_{\text{xc}}(r, r') = \langle \tilde{\rho}_{2}(r, r') \rangle_{\lambda} - \rho(r) \rho(r')$$

(52)

with

$$\langle \tilde{\rho}_{2}(r, r') \rangle_{\lambda} = \int_{0}^{1} d\lambda \langle \Psi_{\lambda} | \tilde{\rho}_{2}(r, r') | \Psi_{\lambda} \rangle.$$  

(53)

In Eq. (53), $\tilde{\rho}_{2}(r, r') = \tilde{\psi}(r) \tilde{\psi}(r') \tilde{\psi}(r') \tilde{\psi}(r)$ denotes the pair density operator and $\Psi_{\lambda}$ stands for the ground state of a fictitious system where the Coulomb interaction is scaled down by $\lambda$ and the effective local single-electron potential $\nu_{\lambda}$.
is added to the Hamiltonian to keep the ground-state density constant with respect to \( \lambda \).

By definition, the exact xc hole corresponding to a system with a fractional number of electrons \( N+\omega \), satisfies the sum rule

\[
\int dr h_{xc}(r, r') = -1 + \omega (1 - \omega) \int_0^1 d\lambda \frac{\rho_{\lambda,N\omega+1}(r) - \rho_{\lambda,N}(r)}{\rho(r)},
\]

where the ground-state density \( \rho_{\lambda,M} \) of the system with \( M = N+\omega \) electrons depends on \( \lambda \) due to the fact that \( M \) differs from \( N+\omega \). It is important to note that the exact xc hole \( h_{xc} \) integrates to \(-1\) only for integer electron numbers at variance with the LSD xc hole \( h_{xc}^{\text{LSD}} \), which integrates to \(-1\) irrespective of the number of electrons.

Turning now to the \( \alpha \)N\text{K} xc hole \( h_{xc}^{\alpha \text{NK}} \), it is possible to derive a relation similar to Eq. (54)

\[
\int dr h_{xc}^{\alpha \text{NK}}(r, r') = -1 + \alpha \text{NK} \sum_{1a} f_{1a} [2 f_{1a} - f_{1\text{LSD}}] \frac{n_{1\text{LSD}}(r)}{\rho(r)}
\]

for \( N \geq 1 \) (the detailed derivation is presented in the Appendix). It is then clear that the occupation \( f_{1a} = \frac{1}{2} \) allows to satisfy the xc-hole sum rule exactly for integer number of electrons, and at least approximately for fractional numbers. (Note that in the case \( N = 1 \), the value \( f_{1a} = 0 \) corresponds also to the exact sum rule.) For PZ, it has been argued that the enforcement of a similar sum rule is critical to the quality of the self-interaction correction.

As a result, the explicit expression of the screened \( \alpha \)N\text{K} functional with \( f_{1a} = \frac{1}{2} \) reads

\[
E^{\alpha \text{NK}} = E^{\text{LSD}} + \alpha \text{NK} \sum_{1a} f_{1a} (1 - f_{1a}) E_{1a}[n_{1a}] + E_{xc}^{\text{LSD}} [\rho - \rho_{1a}]
\]

\[
+ \int dr u_{xc,1a}^{\text{LSD}}(r, \rho + \left( \frac{1}{2} - f_{1a} \right) n_{1a} - \rho_{1a}, - E_{xc}^{\text{LSD}} [\rho]).
\]

In the next section, we assess the predictive accuracy of the \( \alpha \)N\text{K} functional given by Eq. (56), first focusing on atoms then extending applications to molecular systems.

### III. APPLICATIONS

#### A. Atomic ionization

In order to probe the performance of the \( \alpha \)N\text{K} functional, we calculate the electron removal energies of a complete range of atomic elements, from hydrogen to xenon, using the all-electron LD1 code of the QUANTUM-ESPRESSO distribution. (Note that the atomic calculations that we report here do not account for relativistic effects; we have verified that the inclusion of scalar-relativistic contributions results in marginal deviations of \(<80\) meV in the absolute precision of \( \alpha \)N\text{K} differential electron removal energies.) The LD1 code proceeds by iterative integration of the spherically symmetric electronic-structure problem on logarithmic grids. The validity of this integration procedure, also employed by Perdew and Zunger in their original work, has been extensively discussed and carefully verified by Goedecker and Umrigar. More recently, Stengel and Spaldin have shown that the spherical approximation is actually required to prevent the appearance of unphysical hybrid states in the electronic structure of isolated atoms and predict physical atomic electron removal energies within orbital-dependent self-interaction corrections.

We first compute the screening coefficient \( \alpha \)N\text{K} [Eqs. (43) and (44)] for the elements of the five first periodic rows based on the electronic configurations tabulated in Ref. 55 (with the exception of Ni for which we used the slightly unfavored 4s\(^3\)3d\(^{10}\) configuration instead of 4s\(^2\)3d\(^{10}\) that overestimates the experimental ionization potential by more than 1 eV). The calculated \( \alpha \)N\text{K} are reported in Fig. 6. The graph confirms that the screening coefficient varies in a narrow range. Not unexpectedly, the value of \( \alpha \)N\text{K} is found to be maximal for the elements whose outermost electronic shell contains a single electron, such as hydrogen for which \( \alpha \)N\text{K} \( = 1 \) and alkali metals, e.g., \( \alpha \)N\text{K}(Na) = 0.99. In contrast, the coefficient \( \alpha \)N\text{K} saturates to low values for filled shell elements, namely, noble gases, e.g., \( \alpha \)N\text{K}(He)=0.66, alkaline earth metals, e.g., \( \alpha \)N\text{K}(Be)=0.72, and filled shell transition metals, e.g., \( \alpha \)N\text{K}(Pd)=0.69.
After calculating atomic screening coefficients, we compare $\alpha$NK differential electron affinity predictions with LSD and experiment in Fig. 7. The comparison demonstrates the predictive ability of the $\alpha$NK method, which brings partial electron removal energies $A_{N-1}$ in very close agreement with experimental total electron removal energies $A_{N-1}$, whereas LSD is found to considerably overestimate $A_{N-1}$. In quantitative terms, the differential LSD energy $\Delta A_{N-1}$ is overestimated by more than 4 eV with a standard deviation of 1.85 eV [Fig. 8(a)]. Comparable deviations are obtained with the PZ self-interaction correction. The HF energy $A_{N-1}$ are instead underestimated by a smaller margin of 1.48 eV. The $\alpha$NK correction results in substantial improvement in the calculation of partial electron removal energies, reducing the error to 0.31 eV. Here, it is quite interesting to note that the $\alpha$NK variational contributions counterbalance the slight tendency of the $\alpha$NK$_0$ correction to underestimate electron removal energies within LSD.

We now examine partial ionization potential predictions (Fig. 9). A marked difference with the above electron affinity results is the enhanced accuracy of the HF and PZ theories. The improved performance in predicting atomic ionization potentials results from the fact that orbital relaxation compensates the absence of correlation contributions in HF and cancels residual non-Koopmans errors in PZ. Nevertheless, even with beneficial error cancellation in favor of HF and PZ, the $\alpha$NK deviation is still the lowest, approximately equal to the PE mean absolute error of 0.34 eV [Fig. 8(b)].

The precision of $\alpha$NK differential ionization energies reflects the intrinsic accuracy of the underlying LSD total energy functional in reproducing subtle atomic ionization trends that are difficult to describe with, e.g., semiempirical approximations. To complement these observations, the full dependence of the LSD, PZ, and $\alpha$NK relaxed energies $\epsilon_{2p}$ of the highest occupied atomic orbital of carbon as a function of its occupation is depicted in Fig. 10. Confronting the LSD and PZ graphs with those presented in Fig. 3, it is seen that orbital relaxation causes a non-negligible decrease in the unphysical shift $\epsilon_{2p}(1)-\epsilon_{2p}(0)$ of 1.5 eV for both functionals. Additionally, the PZ orbital energy $\epsilon_{2p}$ becomes less curved at higher occupations, confirming that orbital relaxation enhances the performance of the PZ correction. Nevertheless, the inflexion of the curve remains important in the vicinity of $f_{2p}=0$ due to the fact that the PZ correction leaves the energy of the empty state almost unchanged. We observe that this unphysical trend is almost completely removed by the $\alpha$NK correction, clearly showing that the screened $\alpha$NK method is apt at imposing the generalized Koopmans’ condition for any fractional value of $f_{2p}$.

The above comparisons demonstrate the predictive performance of the non-Koopmans method in correcting atomic
differential electron affinities and first ionization potentials, placing \( \alpha_{N-1} \) and \( I_N \) predictions on the same level of accuracy with respect to experiment. The fact that \( \alpha_{NK} \) improves \( \alpha_{N-1} \) and \( I_N \) with the same precision ensures the accuracy of non-Koopmans total energy differences and related equilibrium properties. The results presented in the next sections provide further support to this conclusion.

B. Molecular ionization

In this section, we focus on the study of molecular systems. For this purpose, we have implemented the HF, PZ, and \( \alpha_{NK} \) methods in the plane-wave pseudopotential Car-Parrinello (CP) code of the QUANTUM-ESPRESSO distribution.\(^5\) In this code, orbital optimization proceeds via fictitious Newtonian damped electronic dynamics.

The main difficulty in the CP implementation of the HF, PZ, and \( \alpha_{NK} \) functionals is the correction of periodic-image errors that arise from the use of the supercell approximation.\(^1\) Such numerical errors preclude the accurate evaluation of exchange terms and orbital electrostatic potentials. To eliminate periodic-image errors in the plane-wave evaluation of exchange and electrostatic two-electron integrals, we employ countercharge correction techniques.\(^6\) In addition to this difficulty, explicit orthogonality constraints must be considered for the accurate calculation of the gradient of the orbital-dependent PZ and \( \alpha_{NK} \) functionals.\(^5\) To incorporate these additional constraints, we use the efficient iterative orthogonalization cycle implemented in the original CP code.\(^5\) In terms of computational performance, the cost of \( \alpha_{NK} \) calculations is here only 40% higher than that of PZ and lower than that of HF.

In Table I, we compare LSD, HF, PZ, and NK partial electron removal energy predictions for a representative set of molecules. In each case, molecular geometries are fully relaxed (the accuracy of equilibrium geometry predictions will be examined in Sec. III C). To perform our calculations, we employ LSD norm-conserving pseudopotentials\(^6\) with an energy cutoff of 60 Ry for the plane-wave expansion of the electronic wave functions. With this calculation parameter, we verify that \( \alpha_{N-1} \) and \( I_N \) are converged to within less than 50 meV.

It is frequently argued that substituting LSD pseudopotentials for their HF, PZ, and NK counterparts has minor effect on the predicted energy differences.\(^4\) Comparing our pseudopotential calculations with all-electron atomic results (see Sec. III A), we actually found that the use of LSD pseudopotentials yields HF, PZ, and NK electron removal energies with a typical error of 0.1–0.2 eV. However, since these moderate deviations affect HF, PZ, and NK predictions in identical manner, the pseudopotential substitution does not alter the validity of the present comparative analysis.

As expected, one conspicuous feature in Table I is the poor performance of LSD that predicts molecular partial electron removal energies with an average error of \( \pm 40\% \). As was the case for atoms, the PZ self-interaction correction reduces the error in predicting \( I_N \) to less than 14%, which corresponds to an average deviation of 1.68 eV, whereas \( \alpha_{N-1} \) predictions are not improved. In comparison, \( \alpha_{NK} \) partial ionization energies are predicted with a remarkable precision of 0.50 eV (4.1%) and 0.58 eV (4.8%) for \( \alpha_{N-1} \) and \( I_N \), respectively. The \( \alpha_{NK} \) accuracy in predicting molecular vertical ionization energies compares favorably (arguably, even more accurately) with that of recently published fully self-consistent GW many-body perturbation theory calculations.\(^6\)

It should also be noted that we perform here only one iteration [Eq. (43)] to determine the screening coefficient and that the calculated \( \alpha_{NK} \) vary in a very limited range of values, even narrower than that found in the case of atoms (Fig.
Similarly to the comparative analysis presented in Sec. II D, we use the reference orbital-energy formulation to evaluate the magnitude of $\tilde{w}_{\text{ref},i}$ and $\tilde{w}_{\text{ex},i}$ errors. In agreement with the trend already observed, $\alpha\text{NK}_0$ predictions for $A_{N-1}$ and $I_N$ are lower than $\alpha\text{NK}$ electron removal energies with negative shifts of 0.59 eV and 0.68 eV, respectively. (Thus, $\alpha\text{NK}_0$ results are found to be even closer to experiment than their $\alpha\text{NK}$ counterparts with mean absolute error margins of only 3.2% for $A_{N-1}$ and 2.5% for $I_N$.) This direct comparison indicates that $w_{\text{ref},i}^{\text{LSD}}$ and $w_{\text{ex},i}^{\text{LSD}}$ introduce non-negligible energy shifts in the calculation of frontier orbital levels. However, these errors are much smaller than typical self-interaction deviations of 4–5 eV, providing qualitative justifications of the excellent precision of the $\alpha\text{NK}$ method.

### C. Equilibrium structural properties

After analyzing partial electron removal energies, we now report on the accuracy of $\alpha\text{NK}$ equilibrium geometry calculations. We compare LSD, PZ, and $\alpha\text{NK}$ structural predictions to experimental bond lengths in Fig. 12 and we present LSD, HF, PZ, and $\alpha\text{NK}$ error bars in Fig. 13. The first important observation is the very good accuracy of LSD predictions with a mean absolute relative error of 1.1% for the seventeen molecules listed in Table I. PZ bond lengths are instead sensibly underestimated with a mean uncertainty of

---

**TABLE I.** LSD, HF, PZ, $\alpha\text{NK}$, and $\alpha\text{NK}_0$ differential molecular electron removal energies compared with experimental vertical electron removal energies. Mean deviations (MD), mean absolute deviations (MAD), and root mean squared deviations of the error (RMS) in absolute and relative terms are also reported. The adiabatic ionization lower bound is given when the experimental vertical ionization energy is not available. Energies are in eV.

<table>
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<tr>
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<th>LSD</th>
<th>HF</th>
<th>PZ</th>
<th>$\alpha\text{NK}$</th>
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<td>5.9%</td>
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*Reference 59.*

![Graph](image-url)
2.8%. In contrast with PZ calculations, αNK results deviate from experiment by a relative error margin of 0.8%, which is lower than that of LSD, demonstrating that the αNK self-interaction correction does not deteriorate and even improves LSD structural predictions, at variance with the conventional PZ self-interaction correction.

These results illustrate the tendency of PZ to overbind molecular structures, and confirm the systematic improvement brought about by the αNK correction. The promising potential of the αNK correction in predicting other thermodynamical properties (e.g., dissociation energies and vibrational frequencies) will be critically explored in a separate study.

D. Photoemission energies

Having validated the non-Koopmans self-interaction correction for the calculation of electron removal energies and equilibrium structures, we now evaluate the performance of the αNK and αNK₀ methods in predicting photoemission energies, for which LSD and GGAs exhibit notable failures.

From the theoretical point of view, the very poor performance of LSD and GGA is expected; Kohn-Sham density-functional theory eigenvalues are not meant to predict excited-state properties. In practice, total electron removal energies computed from constrained density-functional calculations (ΔSCF) are typically found to be in good agreement with experiment. This level of accuracy suggests that orbital-dependent self-interaction corrected functionals can provide orbital energies in accordance with spectroscopic results. This expectation is confirmed by PZ photoemission predictions for neon, argon, and krypton that we reproduce here using the LD1 code (Table II).

Nevertheless, similarly to the trend observed in Sec. III B, the predictive ability of PZ deteriorates in the case of molecular photoionization; this is at variance with the αNK method. To illustrate this fact, we compare LSD, HF, PZ, and non-Koopmans predictions for the photoemission spectrum (PES) of benzene in Table III and fullerene in Table IV. In the molecular photoemission calculations, we use the CP code with the computational procedure described in Sec. III B. We employ fully relaxed geometries for benzene and the LSD atomic structure of C₆₀, which is found to be in excellent agreement with the NMR experimental geometry.

Focusing first on benzene, we observe that LSD underestimates electron binding energies with errors as large as 4.35 eV for low-lying states. In contrast to LSD, the HF theory provides overestimated photoemission energies with absolute deviations that increase gradually from 0.12 to 3.19 eV when approaching the bottom of the PES. Similar trends are observed for PZ with the difference that the errors do not systematically increase with increasing photoemission energies, leading in particular to the incorrect ordering of the e₂g and a₂u, levels. In contrast, αNK restores the correct relative peak positions and yields slightly overestimated electron binding energies with an absolute precision of 4.9%. The slight tendency of αNK to overestimate electron binding energies is here again due to the influence of variational contributions, as directly confirmed by the performance of the αNK₀ orbital-energy method, which predicts photoemission energies in remarkable agreement with experiment.

Similarly to benzene, LSD energy predictions for fullerene are significantly underestimated. However, since the dispersion of the errors is much narrower than in the case of benzene, a simple shift of LSD photoemission bands, equal to the difference between the theoretical and experimental highest occupied molecular orbital levels, can bring the predicted PES in close agreement with experiment. Despite the excellent precision of HF in the top region of the spectrum, HF photoemission energies are largely overestimated for low-lying states. In addition, HF inverts the h₁ and g₂ states in the second photoemission band although it predicts the correct peak ordering in the third and fourth bands. The performance of PZ is found to be slightly worse than that of HF with significant qualitative errors in the grouping and ordering of the states. In contrast, αNK correctly shifts the spectrum and brings photoemission energies in very good agreement with experiment. Predicted αNK binding energies are also in excellent agreement with constrained LSD total energy differences providing a final validation of the performance of the αNK self-interaction correction.
correction in bringing physical meaning to orbital energies—i.e., in identifying orbital energies as opposite total electron removal energies.

**IV. SUMMARY AND OUTLOOK**

In summary, we have demonstrated that the correction of the nonlinearity of the ground-state energy $E_N$ as a function of the number of electrons $N$, at the origin of important discrepancies between total and differential electron removal energies, and related fundamental qualitative and quantitative self-interaction errors, can be achieved without altering the otherwise excellent performance of density-functional approximations in describing systems with nonfractional occupations. To construct the non-Koopmans self-interaction correction, we have first defined an exact non-Koopmans measure of self-interaction and adopted the frozen-orbital approximation, i.e., the framework of the restricted Koopmans’ theorem as a working alternative to the conventional one-electron paradigm. We have then accounted for orbital relaxation by introducing the screening coefficient $\alpha_{NK}$, which bears the physical significance of a uniform and isotropic screening factor that can be determined iteratively—thereby closely satisfying the generalized Koopmans’ condition. This self-interaction correction scheme can be applied to any lo-

**TABLE II.** LSD, HF, PZ, and $\alpha_{NK}$ orbital energies of neon, argon, and krypton compared with experimental photoemission energies. Relative MAD with respect to experimental photoionization results are also reported. The experimental photoemission energies of the spin-orbit doublets of $p$ and $d$ orbitals are indicated. Computational photoionization predictions do not include spin-orbit coupling. Energies are in eV.

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<th>$\alpha_{NK}$</th>
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<tr>
<td>Ar</td>
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<td>30.54</td>
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</tr>
<tr>
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<td>$2s$</td>
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<td>3218.88</td>
<td>3193.55</td>
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<td>Kr</td>
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<td>13.97</td>
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</tr>
<tr>
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<td>$4s$</td>
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</tr>
</tbody>
</table>

**TABLE III.** LSD, HF, PZ, $\alpha_{NK}$, and $\alpha_{NK_0}$ orbital energies of benzene compared with experimental photoemission energies. Relative MAD with respect to experimental photoionization results are also reported. Energies are in eV.

<table>
<thead>
<tr>
<th></th>
<th>LSD</th>
<th>HF</th>
<th>PZ</th>
<th>$\alpha_{NK}$</th>
<th>$\alpha_{NK_0}$</th>
<th>Expt. (^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$e_{1g}$</td>
<td>6.59</td>
<td>9.18</td>
<td>9.43</td>
<td>10.39</td>
<td>9.39</td>
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<td>$e_{2g}$</td>
<td>8.28</td>
<td>13.54</td>
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<td>12.48</td>
<td>11.8</td>
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<td>$a_{2u}$</td>
<td>9.43</td>
<td>13.64</td>
<td>12.99</td>
<td>13.25</td>
<td>12.60</td>
<td>12.5</td>
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<tr>
<td>$e_{1u}$</td>
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<td>16.02</td>
<td>17.67</td>
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<td>$b_{2u}$</td>
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<td>14.9</td>
</tr>
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<td>19.27</td>
<td>19.34</td>
<td>19.2</td>
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</tbody>
</table>

**References:**

\(^a\)Reference 62.

\(^a\)Reference 68.
TABLE IV. LSD, HF, PZ, and αNK orbital energies of fullerene C_{60} compared with constrained LSD total energy differences (Δ_{LSD}) and experimental photoemission energy bands. Energies are in eV.

<table>
<thead>
<tr>
<th>Band</th>
<th>LSD</th>
<th>HF</th>
<th>PZ</th>
<th>αNK</th>
<th>Δ_{LSD}</th>
<th>Expt</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
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<td>8.77</td>
<td>7.45</td>
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</tr>
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<td>9.64</td>
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<td>8.75</td>
<td>8.90</td>
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</tr>
<tr>
<td>IV</td>
<td>8.72</td>
<td>12.42</td>
<td>h_{g}</td>
<td>12.21</td>
<td>10.31</td>
<td>10.47</td>
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<tr>
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<td>12.69</td>
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<td>10.50</td>
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<td>10.79</td>
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<td>h_{g}</td>
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<td>g_{u}</td>
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<td></td>
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<td>h_{g}</td>
<td>15.66</td>
<td>h_{g}</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

aReference 65.
bReference 69.

cal, semilocal or hybrid density-functional approximation. The remarkable predictive performance of the non-Koopmans theory has been demonstrated for a range of atomic and molecular systems.

The theory developed here represents a significant step in the correction of electron self-interaction in electronic-structure theories. Nevertheless, interesting problems are left open. One central question is that similarly to the PZ approach, the αNK method leads to an orbital-dependent Hamiltonian, although it is always possible in principle to derive a consistent density-dependent formulation using, e.g., optimized effective potential mappings.27 It is a long held tenet that the orbital dependence of self-interaction functionals and the subsequent loss of invariance with respect to unitary transformation of the one-body density matrix precludes applications to periodic systems (e.g., conjugate polymers and crystalline materials). In future studies, we will explore solutions to this central conceptual difficulty without resorting to density-dependent unitary invariant mappings.

ACKNOWLEDGMENTS

The authors are indebted to X. Qian, N. Laachi, S. de Gironcoli, É. Cancès, and T. Körzdörfer for helpful suggestions and fruitful comments. This work has been performed using the QUANTUM-ESPRESSO package (http://www.quantum-espresso.org) (Ref. 54) and computational resources offered by the Minnesota Supercomputing Institute and DOE under Grant DE-FG02-05ER46253. I.D. and N.M. acknowledge support from MURI under Grant No. DAAD 19-03-1-016. I.D., Y.L., and M.C. acknowledge support from the Grant in Aid of the University of Minnesota and from Grant ANR 06-CIS6-014 of the French National Agency of Research. M.C. acknowledges partial support from NSF under Grant No. EAR-0810272 and from the Abu Dhabi-Minnesota Institute for Research Excellence (AD-MIRE). N.M., A.F., and N.P. acknowledge support from DOE under Grants No. SciDAC DE-FC02-06ER25794 and No. DE-FG02-05ER15728, and MIT-ISN.

APPENDIX: SCREENED NON-KOOPMANS EXCHANGE-CORRELATION HOLE SUM RULE

In this appendix, we derive the explicit expression of the αNK xc hole. Starting from the relation

$$E^{\alphaNK} = E^{LSD} + \alphaNK \sum_{i\sigma} \left[ E_{\text{Hxc}}^{\text{LSD}}[\rho - \rho_{i\sigma}] - E_{\text{Hxc}}^{\text{LSD}}[\rho] \right] + \int d\mathbf{r} \rho_{i\sigma}(\mathbf{r}) v_{\text{LSD}}^{\text{Hxc,\sigma}}(\mathbf{r};[\rho_{i\sigma}])$$

(A1)

and from the definition of the xc hole [Eq. (51)], the contributions to the total αNK xc hole arising from the three summation terms in Eq. (A1) can be worked out. Those terms will be labeled $h_{xc,i\sigma}^{(a)}$, $h_{xc,i\sigma}^{(b)}$, and $h_{xc,i\sigma}^{(c)}$, respectively. Focusing on the first term, it is straightforward to obtain

$$h_{xc,i\sigma}^{(a)}(\mathbf{r}, \mathbf{r}') = \frac{\rho(\mathbf{r}) - \rho_{i\sigma}(\mathbf{r})}{\rho(\mathbf{r})} \rho(\mathbf{r}') - \rho_{i\sigma}(\mathbf{r})$$

(A2)

Turning to the second term and including the appropriate sign, one obtains

$$h_{xc,i\sigma}^{(b)}(\mathbf{r}, \mathbf{r}') = -\rho(\mathbf{r}') - h_{xc}^{\text{LSD}}(\mathbf{r}, \mathbf{r}';[\rho]).$$

(A3)

The third term is more complicated to derive since its expression is based on the exchange-correlation potential. Making use of the relation
\[ v_{xc,\sigma}(r; \rho) = \frac{1}{2} \int dr' \frac{h_{xc}(r, r'; \rho)}{|r - r'|} + \frac{1}{2} \int dr' dr'' \rho(r') \delta h_{xc}(r', r''; \rho) \delta \rho_{\sigma}(r) \]

(A4)

the expression for \( h_{xc,\sigma}^{(c)}(r, r') \) becomes

\[ h_{xc,\sigma}^{(c)}(r, r') = \frac{2 \rho_{\sigma}(r)}{\rho(r)} \rho_{\sigma}^{ref}(r') + \frac{\rho_{\sigma}(r)}{\rho(r)} h_{xc}^{LSD}(r, r'; \{ \rho_{\sigma}^{ref} \}) + \rho_{\sigma}^{ref}(r) \int dr'' \rho_{\sigma}(r'') \delta h_{xc}^{LSD}(r, r''; \{ \rho_{\sigma}^{ref} \}) \delta \rho_{\sigma}(r'') \]

(A5)

Regrouping all the terms, the expression for the exchange-correlation hole of the αNK functional can be written as

\[ h_{xc,\sigma}^{\alpha NK}(r, r') = h_{xc}^{LSD}(r, r'; \{ \rho \}) + \frac{\alpha_{NK}(r)}{\rho(r)} \left( \sum_{i \sigma} \int dr'' \rho_{i \sigma}(r'') \delta h_{xc}^{LSD}(r, r''; \{ \rho_{i \sigma}^{ref} \}) \delta \rho_{\sigma}(r) \right) \]

\[ + f_{i \sigma} (2 f_{i \sigma} - 1) \int dr'' \rho_{i \sigma}(r'') \delta h_{xc}^{LSD}(r, r''; \{ \rho_{i \sigma}^{ref} \}) \delta \rho_{\sigma}(r') \]

\[ - h_{xc}^{LSD}(r, r'; \{ \rho \}) + h_{xc}^{LSD}(r, r'; \{ \rho_{i \sigma}^{ref} \}) \rho_{i \sigma}(r) \]

(A6)

The result given in Eq. (55) can be obtained from the above equation taking into account the xc-hole sum rule of the LSD functional, \( \int dr' h_{xc}^{LSD}(r, r') = -1 \) (valid for any electron number).
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50 Differential electron affinities can be directly calculated as the opposite energy of the lowest unoccupied orbital of the ionized atom since the derivative discontinuity contribution defined in Refs. 4 and 25 is absent for the LSD, HF, PZ, and non-Koopmans functionals.


