Symmetry-broken local-density approximation for one-dimensional systems

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I. INTRODUCTION

In 1965, Kohn and Sham [1] showed that the knowledge of an analytical parametrization of the uniform electron gas (UEG) correlation energy [2] allows one to perform approximate calculations for atoms, molecules, and solids [3]. This led to the development of various local-density approximation (LDA) correlation functionals (VWN [4], PZ [5], PW92 [6], etc.), all of which require information on the high- and low-density regimes of the UEG [7–35] and are parameterized using results from near-exact diffusion Monte Carlo (DMC) calculations [36–52].

The LDA is the simplest approximation within density-functional theory (DFT) [3]. It assumes that a real, nonuniform system (such as a molecule or a solid) may be treated as a collection of infinitesimally small UEGs of electronic density \( \rho \). In principle, if one knows the reduced (i.e., per electron) correlation energy \( e^{\text{FF}}_{\text{c}}(\rho) \) of the UEG for any density \( \rho \), by summing the individual contributions over all space, it is therefore possible to obtain the LDA correlation energy

\[
E^{\text{LDA}}_{\text{c}} = \int \rho(r) e^{\text{FF}}_{\text{c}}[\rho(r)] dr.
\]

Although it describes molecular bonding reasonably well compared to the Thomas-Fermi (TF) model [53,54] (which approximates the kinetic energy using the TF functional and ignores the exchange interaction between same-spin electrons), this rather crude approximation has managed only mixed success [55]. In particular, the functional consistently overestimates correlation energies, giving rise to errors up to some factor of 2 [3]. Fortunately, errors in the exchange and correlation energies approximately compensate each other, thus generating a total energy that is usually in good agreement with experimental results [56].

Since the emergent days of DFT, the UEG correlation energy \( e^{\text{FF}}_{\text{c}}(\rho) \) has always been defined as the difference between the exact energy \( e(\rho) \) and the Fermi fluid Hartree-Fock (FFHF) energy \( e^{\text{HF}}_{\text{c}}(\rho) \). However, in the early 1960s, Overhauser [57,58] showed that the FF state is never the Hartree-Fock (HF) ground state due to spin- and charge-density instabilities [59]. Therefore, for any density, one can find a symmetry-broken HF (SBHF) solution which has a lower energy than the FFHF solution. Unfortunately, the exact character of this SBHF solution is not given, nor is it necessary for it to remain the same over the full density range.

Zhang and Ceperley [60] have recently presented a computational “proof” of this statement. Performing unrestricted HF (UHF) calculations on the paramagnetic state of finite-size three-dimensional UEGs, they have succeeded in finding a ground state with broken spin symmetry in the high-density region. For lower densities, Trail et al. discovered that the Wigner crystal (WC) is more stable than the FF state for \( r_s > 1.44 \) in two dimensions and \( r_s > 4.5 \) in three dimensions [61], where the Wigner-Seitz radius \( r_s \) is the average distance between electrons. These calculations were recently refined by Holzmann and coworkers [62–67].

Physically, a WC represents a state whose energy is dominated by the potential energy term, resulting in the electrons localizing on lattice points. This situation typically occurs at low densities, where the WC becomes the ground state. At high densities the kinetic energy dominates, and the delocalized FF is the ground state. In addition to the usual FF and WC phases, they have also considered incommensurate crystals (a state in which the number of the charge-density maxima is higher than the number of electrons), showing that such a phase is always favored over the FF, independent of the imposed polarization and crystal symmetry and in agreement with the earlier prediction of Overhauser [57,58].

Here, we propose to construct a symmetry-broken version of the LDA, taking the one-dimensional (1D) UEG as an example and using SBHF energies instead of the usual FFHF expression. From an experimental point of view, 1D systems have recently attracted much attention due to their practical realization in carbon nanotubes [68–72], organic conductors [73–77], transition-metal oxides [78], edge states in quantum Hall liquids [79–81], semiconductor heterostructures [82–86], confined atomic gases [87–89], and atomic or semiconducting nanowires [90,91].
This paper is organized as follows. In Sec. II, we introduce the paradigm we have used to create a strict 1D UEG. Section III covers the acquisition of accurate SBHF energies at various densities, followed by the definition of the two correlation functionals in Sec. IV. These are compared in Sec. V for 1D systems, including atoms and molecules. Unless otherwise stated, atomic units are used throughout.

II. ONE-DIMENSIONAL UNIFORM ELECTRON GAS

A 1D UEG is constructed by confining a number $n$ of interacting electrons to a ring of radius $R$ of electronic density [7,9,92,93]

$$\rho = \frac{n}{2\pi R} = \frac{1}{2r_s},$$

(2)

where $r_s$ is the so-called Wigner-Seitz radius [3,59]. We refer the readers to Ref. [7] for more details about this paradigm, which has been shown to be equivalent to the more conventional “electrons-in-a-periodic-box” model in the thermodynamic limit (i.e., $n \rightarrow \infty$) [7,8,10,35] but mathematically simpler [35]. This can be qualitatively explained by the “shortsightedness” of the electronic matter [94,95]. Because the paramagnetic and ferromagnetic states are degenerate in strict 1D systems, we will consider only the spin-polarized case.

The expression of the (non-symmetry-broken) FFHF energy is [7]

$$e_{\text{FF}}^{\text{SB}}(r_s,n) = \frac{t_{\text{HH}}(n)}{r_s^2} + \frac{v_{\text{HH}}(n)}{r_s},$$

(3)

with

$$t_{\text{HH}}(n) = \frac{\pi^2 n^2 - 1}{24 n^2},$$

(4a)

$$v_{\text{HH}}(n) = \left(\frac{1}{2} - \frac{1}{8n^2}\right)\left[\psi\left(n + \frac{1}{2}\right) - \psi\left(\frac{1}{2}\right)\right] - \frac{1}{4},$$

(4b)

where $\psi(x)$ is the digamma function [98]. This corresponds to occupying the $n$ lowest plane waves

$$\phi_m(\theta) = \frac{\exp(i m \theta)}{\sqrt{2 \pi R}},$$

(5)

where $\theta$ is the angle of the electron around the ring and $m \in \mathbb{Z}$. The FFHF energy (3) diverges logarithmically for large numbers of electrons [7,99–101],

$$e_{\text{FF}}^{\text{SB}}(r_s,n) \sim \ln \frac{n}{r_s} + O(1),$$

(6)

but an identical divergence in the exact energy $e(r_s,n)$ results in a finite correlation energy

$$e_{\text{c}}^{\text{SB}}(r_s,n) = e(r_s,n) - e_{\text{FF}}^{\text{SB}}(r_s,n).$$

(7)

As stated in Sec. I, our goal here is to determine the SB correlation energy in the thermodynamic limit

$$e_{\text{c}}^{\text{SB}}(r_s) = \lim_{n \rightarrow \infty} e_{\text{c}}^{\text{SB}}(r_s,n)$$

via extrapolation, where

$$e_{\text{c}}^{\text{SB}}(r_s,n) = e(r_s,n) - e_{\text{FF}}^{\text{SB}}(r_s,n)$$

$$= e(r_s,n) - e_{\text{FF}}^{\text{SB}}(r_s) + \Delta e_{\text{c}}^{\text{SB}}(r_s,n)$$

$$= e_{\text{c}}^{\text{SB}}(r_s) + \Delta e_{\text{c}}^{\text{SB}}(r_s,n).$$

(9)

III. SYMMETRY-BROKEN HARTREE-FOCK CALCULATIONS

In order to obtain SBHF energies, we have written a self-consistent field program [102] using plane waves of the form (5) with

$$m = -\frac{M-1}{2}, \ldots, \frac{M-1}{2},$$

(10)

where we have used up to $M = 399$ to ensure that our energies are always converged within microhartree accuracy. As expected, large $r_s$ values require larger basis sets in order

<table>
<thead>
<tr>
<th>$n$</th>
<th>$r_s^{\text{SB}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>9</td>
<td>1.22</td>
</tr>
<tr>
<td>19</td>
<td>0.79</td>
</tr>
<tr>
<td>29</td>
<td>0.64</td>
</tr>
<tr>
<td>39</td>
<td>0.55</td>
</tr>
<tr>
<td>49</td>
<td>0.50</td>
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<tr>
<td>59</td>
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<tr>
<td>69</td>
<td>0.43</td>
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<tr>
<td>79</td>
<td>0.41</td>
</tr>
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<td>89</td>
<td>0.40</td>
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<tr>
<td>109</td>
<td>0.36</td>
</tr>
<tr>
<td>119</td>
<td>0.35</td>
</tr>
<tr>
<td>129</td>
<td>0.35</td>
</tr>
</tbody>
</table>

$\Delta e_{\text{c}}^{\text{SB}}$ (in millihartrees) for various $n$ and $r_s$, $r_s^{\text{SB}}$ is the lowest value of $r_s$ for which one can find a SBHF solution.
to converge the energy to the same accuracy due to the local character of the WC [103]. The required one- and two-electron integrals can be found in Ref. [7]. The results are reported in Table I, where we have also reported $r_s^{SB}$, the lowest $r_s$ value for which one can find a SBHF solution. It is interesting to note that $r_s^{SB}$ converges extremely slowly with respect to $n$, as shown in Fig. 1. We have found that the function (see the Appendix)
\begin{equation}
    r_s^{SB}(n) = \frac{a}{\ln n + b}
\end{equation}
(with $a = 1.13535$ and $b = -1.61346$) fits our data well [104]. Equation (11) reveals that, in order to observe a SBHF solution below $r_s = 0.2$, one needs at least 1500 electrons.

We have obtained the thermodynamic values by extrapolation via the following asymptotic form:
\begin{equation}
    \Delta \epsilon_{HF}^{SB}(r_s,n) = \Delta \epsilon_{HF}^{SB}(r_s) + An^{-2}.
\end{equation}
For each $r_s$ value we have used the six largest $n$ values of Table I, except for $r_s = 0.5$, where only the last three terms were considered. The quality of the fit is demonstrated in Fig. 2. A similar expression to (12) has been used by Lee and Drummond to extrapolate DMC calculations to the thermodynamic limit [36].

Physically, the appearance of the SBHF solution is characterized by the formation of a WC, i.e., where the electrons “crystallized” such that they are separated by an angle of $2\pi/n$ [7,99–101]. This phenomenon is easily understood in terms of the behavior of the kinetic and potential energies with respect to $r_s$, which scale as $r_s^{-2}$ and $r_s^{-1}$, respectively [see Eq. (3)]. Therefore, at small $r_s$, occupying the $n$ lowest plane waves is energetically favorable, as it minimizes the dominant kinetic energy contribution. However, there exists a critical density $r_s^{SB}$ for which it becomes compelling to break the spatial symmetry (by populating higher plane waves), albeit at the expense of a meagre increase in the kinetic energy. This has the effect of lowering the potential energy by localizing the electrons and, in doing so, effectively reducing the interelectronic interaction.

The formation of a Wigner crystal is illustrated in Fig. 3, where we have represented the HF and SBHF densities for 19 electrons at $r_s = 5$. One can see that, when the WC forms, a gap opens at the Fermi surface between the occupied and vacant orbitals. As shown in Table II, the symmetry-breaking stabilization has a very significant effect on the values of the correlation energy, especially at intermediate and low densities, where it represents a large fraction of the total correlation energy. These results confirm Overhauser’s prediction that in the thermodynamic limit, it is always favorable to break the spatial symmetry (see the Appendix). We have, however, observed that the stabilization becomes extremely small at high density.

IV. CORRELATION FUNCTIONALS

A. Local-density approximation

In this study, we used the LDA functional developed in Ref. [8], which has been constructed using the “robust” interpolation proposed by Cioslowski [105],

\begin{equation}
    \epsilon_c^{LDA}(r_s) = t^2 \sum_{j=0}^{3} c_j t^j (1-t)^{3-j},
\end{equation}
with

\begin{align*}
    c_0 &= k \eta_0, \quad c_1 = 4 k \eta_0 + k^{3/2} \eta_1, \\
    c_2 &= 5 \epsilon_0 + \epsilon_1 / k, \quad c_3 = \epsilon_0,
\end{align*}

and the high- and low-density expansions [8]

\begin{align}
    \epsilon_c(r_s) &= \epsilon_0 + \epsilon_1 r_s + O(r_s^2), \quad r_s \ll 1, \quad (15a) \\
    \epsilon_c(r_s) &= \frac{\eta_0}{r_s} + \frac{\eta_1}{r_s^{3/2}} + O(r_s^{-2}), \quad r_s \gg 1, \quad (15b)
\end{align}

where

\begin{align*}
    \epsilon_0 &= -\frac{\pi^2}{360}, \quad \epsilon_1 = +0.00845, \\
    \eta_0 &= -\ln(\sqrt{2\pi}) + 3/4, \quad \eta_1 = +0.359933,
\end{align*}
FIG. 3. Electronic density $\rho$ as a function of $\theta$ (left) and orbital energies $\epsilon$ as a function of $m$ (right) for the HF (solid line) and SBHF (dashed line) solutions for $n = 19$ at $r_s = 5$.

and $k = 0.418268$ is a scaling factor, determined by a least-squares fit of the DMC data given in Refs. [7,36]. As reported in Ref. [8], the LDA and DMC correlation energies agree to within 0.1 millihartree.

**B. Symmetry-broken local-density approximation**

We define the SBLDA functional as

$$e^{\text{SBLDA}}(r_s) = e^{\text{LDA}}(r_s) + \Delta e^{\text{SB}}_{\text{HF}}(r_s), \quad (16)$$

where $e^{\text{LDA}}(r_s)$ is given by (13). We propose to use the following expression for the symmetry-breaking stabilization:

$$\Delta e^{\text{SB}}_{\text{HF}}(r_s) = \frac{a_0}{b_0} r_s + \frac{a_1}{b_1} r_s^2 + \frac{a_2}{b_2} r_s^3 + \frac{\eta_0}{b_3} r_s^4 + \frac{\eta_1}{b_4} r_s^5 + \frac{\eta_2}{b_5} r_s^6, \quad (17)$$

where

$$a_0 = -0.0646228, \quad a_1 = 0.53562, \quad a_2 = -0.490719, \quad b_0 = 53.1171, \quad b_1 = 1.53114, \quad b_2 = 2.19606.$$ 

Expression (17) is illustrated in Fig. 4 alongside the corresponding data from Table II. The quality of (17) is excellent with a maximum error of 7 microhartrees compared to the values reported in Table I.

In the low-density limit, Eq. (17) behaves as

$$\Delta e^{\text{SB}}_{\text{HF}}(r_s) = -\frac{\eta_0}{r_s} + O(r_s^{-3/2}), \quad r_s \gg 1, \quad (18)$$

**TABLE II.** Correlation energies (in millihartrees) in the thermodynamic limit for various $r_s$.

<table>
<thead>
<tr>
<th>$r_s$</th>
<th>$-e^{\text{FF}}_c$</th>
<th>$-e^{\text{SB}}_c$</th>
<th>$1 - e^{\text{SB}}_c/e^{\text{FF}}_c$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>27.416</td>
<td>27.416</td>
<td>0%</td>
</tr>
<tr>
<td>0.5</td>
<td>23.962</td>
<td>23.486</td>
<td>2%</td>
</tr>
<tr>
<td>1</td>
<td>21.444</td>
<td>18.874</td>
<td>12%</td>
</tr>
<tr>
<td>2</td>
<td>17.922</td>
<td>11.984</td>
<td>33%</td>
</tr>
<tr>
<td>5</td>
<td>12.318</td>
<td>4.316</td>
<td>65%</td>
</tr>
<tr>
<td>10</td>
<td>8.292</td>
<td>1.525</td>
<td>82%</td>
</tr>
<tr>
<td>15</td>
<td>6.319</td>
<td>0.779</td>
<td>88%</td>
</tr>
<tr>
<td>20</td>
<td>5.133</td>
<td>0.478</td>
<td>91%</td>
</tr>
<tr>
<td>50</td>
<td>2.476</td>
<td>0.104</td>
<td>96%</td>
</tr>
<tr>
<td>100</td>
<td>1.358</td>
<td>0.034</td>
<td>97%</td>
</tr>
</tbody>
</table>

which, when combined with Eqs. (15b) and (16), gives

$$e^{\text{SBLDA}}_c(r_s) = O(r_s^{-3/2}), \quad r_s \gg 1. \quad (19)$$

This behavior is a direct consequence of the SBHF treatment allowing the electrons to localize at low densities, unlike the FFHF solution. It is therefore able to correctly identify the appearance of the WC phase, which results in the low-density expansion of the SBHF solution matching that of the exact energy up to $O(r_s^{-3/2})$. This difference in low-density behavior creates an important distinction between $e^{\text{LDA}}_c$ and $e^{\text{SBLDA}}_c$. For small $r_s$, we have imposed $\Delta e^{\text{SB}}_{\text{HF}}(r_s)$ to be quadratic in $r_s$, as shown in the Appendix.

**V. CORRELATION ENERGY IN ONE-DIMENSIONAL SYSTEMS**

In this section, we test the LDA and SBLDA functionals defined in Secs. IV A and IV B, respectively. The LDA and SBLDA correlation energies are obtained via

$$E^{\text{LDA/SBLDA}}_c = \int \rho(x) e^{\text{LDA/SBLDA}}_c[\rho(x)] dx, \quad (20)$$

which is computed by numerical quadrature. These quantities are calculated with the HF density; that is, they are not calculated self-consistently [106]. However, as expected [107], we have observed that the differences between self- and non-self-consistent densities are extremely small [9,10]. In some cases, we have also reported the exact, second-, and

FIG. 4. $\Delta e^{\text{SB}}_{\text{HF}}$ as a function of $r_s$ in the thermodynamic limit. The dashed curve represents the fit defined by Eq. (17).
the theoretical and computational details are provided. For additional information about these calculations, we refer the readers to Refs. [9,97], where theoretical and computational details are provided.

A. Two electrons in a box

As a simple example to illustrate the performance of the SBLDA functional, we have thoroughly studied the well-known system composed by two electrons in a box of length \( L \). In particular, we report variations in the correlation energy as a function of \( L \) obtained using the MP2, MP3, LDA, and SBLDA methods. In the interest of completion, we have also calculated the exact correlation energy of this system using a Hylleraas-type (Hy) expansion [109–113]. The results of this investigation are illustrated in Fig. 5, where the error in the correlation energy \( \Delta E_c = E_c - E_c^{\text{Hy}} \) is resolved as a function of \( L \). As expected, MP2 and MP3 yield reliable estimates of the correlation energy for this system. On the other hand, LDA returns a poor estimate of this energy. This discrepancy is especially clear in the high-density region (i.e., small \( L \)). By construction, LDA and SBLDA have similar performances at high densities. However, SBLDA is much more accurate than LDA at low densities, where stabilization returned from breaking the symmetry is most significant. Similar results are expected for different external potentials [114–116].

B. Atoms

We have calculated the ionization energies and electron affinities of 1D atoms [97] using the CHEMID software developed by one of the authors [117]. The values obtained with the LDA and SBLDA functionals are compared to the MP3 values in Table III, which has been observed to be an exceptionally accurate method in such systems [97]. Overall, LDA and SBLDA overestimate the ionization energies and electron affinities for these systems. It is interesting to note that, although the performance of the LDA and SBLDA functionals is quite poor compared to MP3 for small atoms, the results become rapidly more accurate for larger atoms. In particular, we observe that the accuracy of SBLDA improves faster than LDA. For example, although the deviation between MP3 and SBLDA is only 0.04 eV for the ionization energy of the F atom, the LDA is still 0.19 eV off. This effect is most easily seen by acknowledging that larger atoms have more diffuse orbitals which possess lower-density regions [97].

C. One-electron diatomics

As commonly reported, LDA-type functionals suffer from the self-interaction error (SIE) [118–122], i.e., the unphysical interaction of an electron with itself. This phenomenon is also known as the delocalization error and can be understood as the tendency of approximate functionals to artificially spread the electron density [123–126]. In Fig. 6, we have reported the SIE in the one-electron diatomic molecules \( \text{H}_2^+ \), \( \text{HeH}^2+ \), and \( \text{He}_2^{3+} \) as a function of the bond length. Although the SIE is obviously still present in SBLDA, one can see that it is less pronounced than it is in LDA. This statement is true at any bond length for the three molecules considered here. As an illustration, we have computed the energy of the H atom to be \(-0.5, -0.519\,054\), and \(-0.514\,971\) for HF, LDA, and SBLDA, respectively.

As reported in Table IV, the dissociation energy of \( \text{H}_2^+ \) is \(-0.3307, -0.3328\), and \(-0.3341\) for HF, LDA, and SBLDA. Table III. Ionization energy and electron affinity (in eV) of 1D atoms calculated with various methods.

\[
\begin{array}{cccccc}
\text{Atom} & \text{Ionization energy} & \text{Electron affinity} \\
 & A \rightarrow A^+ + e^- & A + e^- \rightarrow A^- \\
\hline
\text{He} & 33.895 & 34.393 & 34.325 & 0 & 0 & 0 \\
\text{Li} & 4.522 & 4.895 & 4.712 & 1.395 & 1.717 & 1.512 \\
\text{Be} & 10.408 & 10.822 & 10.669 & 0 & 0 & 0 \\
\text{B} & 2.099 & 2.386 & 2.190 & 0.638 & 0.875 & 0.688 \\
\text{C} & 4.730 & 5.056 & 4.865 & 0 & 0 & 0 \\
\text{N} & 1.14 & 1.38 & 1.20 & 0.34 & 0.51 & 0.37 \\
\text{O} & 2.56 & 2.83 & 2.63 & 0 & 0 & 0 \\
\text{F} & 0.68 & 0.87 & 0.72 & 0.2 & 0.3 & 0.2 \\
\text{Ne} & 1.5 & 1.7 & 1.5 & 0 & 0 & 0 \\
\end{array}
\]

FIG. 5. Error in correlation energy \( \Delta E_c \) for two electrons in a box of length \( L \) with MP2, LDA, and SBLDA.

FIG. 6. Self-interaction error in \( \text{H}_2^+ \) (green), \( \text{HeH}^2+ \) (red), and \( \text{He}_2^{3+} \) (blue) calculated with LDA (solid) and SBLDA (dashed) as a function of the bond length.
TABLE IV. Equilibrium bond length $R_{eq}$, transition structure bond length $R_s$, dissociation energy $E_{diss}$, and transition barrier $\Delta E_{ts}$ of 1D molecules calculated with various methods.

<table>
<thead>
<tr>
<th>Method</th>
<th>Molecules</th>
<th>$H_2^+$</th>
<th>HeH$^{2+}$</th>
<th>He$^{2+}$</th>
<th>$H_2$</th>
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<tr>
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<td>2.182</td>
<td>1.793</td>
<td>2.639</td>
</tr>
<tr>
<td></td>
<td>$R_s$</td>
<td>3.296</td>
<td>4.630</td>
<td>0.0209</td>
<td>0.0292</td>
</tr>
<tr>
<td></td>
<td>$E_{diss}$</td>
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<td>+0.0131</td>
<td>-0.1859</td>
</tr>
<tr>
<td></td>
<td>$\Delta E_{ts}$</td>
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<td>+0.2924</td>
<td>+0.045</td>
<td>-0.040</td>
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<tr>
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<td>2.581</td>
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</tr>
<tr>
<td></td>
<td>$R_s$</td>
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<td>4.630</td>
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<td>MP2</td>
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<td>+0.0131</td>
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<td>MP3</td>
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</tr>
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<td>4.630</td>
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<tr>
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<td>+0.045</td>
<td>-0.030</td>
</tr>
<tr>
<td>LDA</td>
<td>$R_{eq}$</td>
<td>2.573</td>
<td>2.176</td>
<td>1.790</td>
<td>2.627</td>
</tr>
<tr>
<td></td>
<td>$R_s$</td>
<td>3.291</td>
<td>4.630</td>
<td>0.0214</td>
<td>0.2964</td>
</tr>
<tr>
<td></td>
<td>$E_{diss}$</td>
<td>-0.3328</td>
<td>+0.1696</td>
<td>+0.0126</td>
<td>-0.1857</td>
</tr>
<tr>
<td></td>
<td>$\Delta E_{ts}$</td>
<td>+0.0214</td>
<td>+0.2964</td>
<td>+0.045</td>
<td>-0.030</td>
</tr>
<tr>
<td>SBLDA</td>
<td>$R_{eq}$</td>
<td>2.564</td>
<td>2.172</td>
<td>1.788</td>
<td>2.619</td>
</tr>
<tr>
<td></td>
<td>$R_s$</td>
<td>3.285</td>
<td>4.619</td>
<td>0.0216</td>
<td>0.2995</td>
</tr>
<tr>
<td></td>
<td>$E_{diss}$</td>
<td>-0.3341</td>
<td>+0.1697</td>
<td>+0.0124</td>
<td>-0.1864</td>
</tr>
<tr>
<td></td>
<td>$\Delta E_{ts}$</td>
<td>+0.0216</td>
<td>+0.2995</td>
<td>+0.045</td>
<td>-0.030</td>
</tr>
</tbody>
</table>

SBLDA, respectively. Like in three dimensions, LDA and SBLDA slightly overestimate the binding energy of $H_2^+$. The equilibrium bond lengths are 2.581, 2.573, and 2.564, showing that LDA and SBLDA predict bond lengths that are slightly too short.

As reported in Ref. [97], HeH$^{2+}$ and He$^{2+}$ are metastable, and it is instructive to know if LDA and SBLDA can predict this peculiar feature properly. Table IV reports the equilibrium bond length of these molecules as well as the transition-structure bond lengths and the height of the barrier. As we have observed in $H_2^+$, the bond lengths predicted by LDA and SBLDA are slightly too short, while the transition-state barriers are overestimated. It is interesting to note that both LDA and SBLDA predict (correctly) HeH$^{2+}$ and He$^{2+}$ as being metastable species.

D. Dissociating $H_2$

The apparently simple problem of stretching $H_2$ has been widely studied within DFT, as it reveals a common pitfall of approximate density functionals known as the static correlation error [123,126,127]. Figure 7 displays the error in the correlation energy $\Delta E_c$ of $H_2$, as computed by the LDA and SBLDA functionals, as a function of the bond length. The exact results have been obtained with a James-Coolidge-type (JC) expansion [97,128]. Although the error in the SBLDA correlation energy is still significant, we observe a clear improvement for all bond lengths compared to LDA. This result is encouraging given the simplicity of the SBLDA functional.

Table IV reports the equilibrium bond length of $H_2$ as well as its dissociation energy obtained with HF, MP2, MP3, LDA, and SBLDA. Compared to the exact results we observe, as previously reported [97], MP2 and MP3 are extremely accurate in one dimension. In contrast to the observations of Sec. V C, the LDA slightly underestimates the dissociation energy here (agreeing with MP3), while the SBLDA continues to overestimate the same value. Both functionals, however, continue to yield a shorter bond length.

VI. DISCUSSION AND CONCLUDING REMARKS

Inspired by Overhauser’s forecasts made some 50 years ago, we have constructed a symmetry-broken (SB) version of the commonly used local-density approximation for one-dimensional systems. The newly designed functional, which we have named SBLDA, has been shown to surpass the performance of its LDA parent in providing better estimates of the correlation energy. More importantly, we believe that this functional could be potentially useful as a superior starting point for more accurate approximations within density-functional theory, such as generalized gradient approximations or hybrid functionals [129]. The methodology presented here is completely general and can be applied to higher-dimensional systems, where SB Hartree-Fock calculations have already been performed [61,64–66]. The design of new exchange and correlation functionals for two- and three-dimensional systems based on the idea developed here is currently in progress in our group.

ACKNOWLEDGMENTS

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APPENDIX: BEHAVIOR OF $\Delta e^{SB}_{HF}$ NEAR $r^{SB}_s$

Near the critical density $r^{SB}_s$, it is possible to use a simple two-orbital model to study the symmetry-breaking process. When the symmetry breaking occurs, a small energy gap appears at the Fermi surface thanks to the mixing of the highest occupied molecular orbital $\phi_{s \uparrow}(\theta)$ and lowest unoccupied molecular orbital $\phi_{s \downarrow}(\theta)$. Therefore, to study the behavior of $\Delta e^{SB}_{HF}$ near $r^{SB}_s$ (see Sec. IV B), we consider the two orthonormalized molecular orbitals

$$\varphi_1(\theta) = \frac{\phi_{s \uparrow}(\theta) + c \phi_{s \downarrow}(\theta)}{\sqrt{1 + c^2}},$$

$$\varphi_2(\theta) = \frac{\phi_{s \uparrow}(\theta) - c \phi_{s \downarrow}(\theta)}{\sqrt{1 + c^2}}.$$  

This two-orbital model teaches us that there exists a critical density

$$r^{SB}_s(n) = \frac{\pi^2/4}{\gamma(n - 1/2) + 1/(\gamma n) + \gamma + 2(\ln 2 - 1)}$$

(A2)

where $\gamma$ is the Euler-Mascheroni constant [98] after which it is energetically favorable to mix these two orbitals, and the value

$$c^2 = \frac{1 - r^{SB}_s}{1 + r^{SB}_s/r_s + \frac{4}{\pi^2} \left(\frac{1}{2n+1} + \frac{1}{2n-1}\right) r^{SB}_s}$$

minimizes the energy for $r_s > r^{SB}_s$. Before $r^{SB}_s$, the solution is the usual FF state of energy

$$e^{FF}_{HF}(r_s, n) = \frac{t^{FF}_{HF}(n)}{r_s} + \frac{v^{FF}_{HF}(n)}{r_s},$$

(A4)

where

$$t^{FF}_{HF}(n) = \frac{\pi^2(n - 1)^2}{4n^2},$$

(A5a)

$$v^{FF}_{HF}(n) = \frac{\psi(n - 1/2) + \gamma + 2\ln 2}{n}$$

(A5b)

minimizes the kinetic energy. However, for $r_s > r^{SB}_s$, this is outweighed by negative contributions in the potential term that drive the symmetry-breaking process. The kinetic and potential parts of the symmetry-breaking stabilization are given by

$$e^{SB}_{HF}(r_s, n) = e^{FF}_{HF}(r_s, n) - \Delta e^{SB}_{HF}(r_s, n),$$

(A6a)

$$\Delta e^{SB}_{HF}(r_s, n) = \Delta e^{SB}_{HF}(r_s, n) + \Delta v^{SB}_{HF}(r_s, n),$$

(A6b)

with

$$\Delta e^{SB}_{HF}(r_s, n) = -\frac{\pi^2 c^2}{(1 + c^2)nr_s^2},$$

(A7)

$$\Delta v^{SB}_{HF}(r_s, n) = \left\{ \frac{\pi^2/4}{(1 + c^2)^2nr_s^2} - n c^2 \right\} \left( \ln 2 - \frac{1}{2n+1} - \frac{1}{2n-1} \right).$$

(A8)

In the thermodynamic limit, Eq. (A2) yields

$$r^{SB}_s \sim \frac{\pi^2/4}{\ln n + \gamma + 2(\ln 2 - 1)},$$

(A9)

which has motivated our use of a similar expression in Eq. (11). Because $\lim_{n \to \infty} r^{SB}_s(n) = 0$, it also proves that, in the thermodynamic limit, there must exist a SBHF solution for any $r_s > 0$ in agreement with Overhauser’s results [57,58].

Expanding $\Delta e^{SB}_{HF}$ at $r_s \sim r^{SB}_s$ yields

$$\Delta e^{SB}_{HF}(r_s, n) = \frac{\pi^2/4}{\ln n} \left[ \frac{\pi^2}{4} \left( \frac{1}{2n+1} + \frac{1}{2n-1} \right) + 1 \right] \left( r_s - r^{SB}_s \right)^2$$

$$+ O \left( (r_s - r^{SB}_s)^3 \right),$$

(A10)

showing that the behavior of $\Delta e^{SB}_{HF}$ is quadratic near $r^{SB}_s$.

[104] We have used the \( r_{\text{SB}} \) values for \( n = 49-99 \) from Table I to obtain the values of \( a \) and \( b \) in Eq. (11).