Derivative Discontinuity in the Strong-Interaction Limit of Density-Functional Theory

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We generalize the exact strong-interaction limit of the exchange-correlation energy of Kohn-Sham density functional theory to open systems with fluctuating particle numbers. When used in the self-consistent Kohn-Sham procedure on strongly interacting systems, this functional yields exact features crucial for important applications such as quantum transport. In particular, the steplike structure of the highest-occupied Kohn-Sham eigenvalue is very well captured, with accurate quantitative agreement with exact many-body chemical potentials. While it can be shown that a sharp derivative discontinuity is present only in the infinitely strongly correlated limit, at finite correlation regimes we observe a slightly smoothened discontinuity, with qualitative and quantitative features that improve with increasing correlation. From the fundamental point of view, our results obtain the derivative discontinuity without making the assumptions used in its standard derivation, offering independent support for its existence.

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First-principles calculations of many-electron systems such as solids, molecules, and nanostructures are based, to a very large extent, on Kohn-Sham (KS) [1] density-functional theory (DFT) [2]. KS DFT is, in principle, an exact theory, in which the ground-state energy and density of an interacting many-electron system are mapped into a problem of noninteracting electrons moving in the effective one-body KS potential. In practice, KS DFT relies on approximations for the exchange-correlation energy that, although successful in very many cases, still have deficiencies that hamper its overall usefulness [3].

Exact KS DFT has many weird and counterintuitive features often missed by the available approximations. One of the weirdest and most elusive of these features is the derivative discontinuity of the exact exchangecorrelation energy functional at integer particle numbers N [4], which has been an incredibly long-debated issue [5–15] because its formal derivation relies on some (very reasonable) assumptions. This discontinuity makes the exact KS potential "jump" by a constant Δ_{xc} when we add to an N-electron system even a very tiny fraction η of an electron, aligning the chemical potential of the noninteracting KS system to the exact, interacting one, as schematically illustrated in Fig. 1. The derivative discontinuity has crucial physical consequences. For example, it accounts for the difference between the KS and the physical fundamental gaps [16–18], it allows a correct KS DFT description of molecular dissociation [4,19], and it should improve charge-transfer excitations in time-dependent DFT [20–22]. It also plays a fundamental role in quantum transport, especially to capture the physics of the Coulomb blockade and the Kondo effect [23–27]. These are all cases in which the standard approximations, which miss this discontinuity, work poorly. Corrections based on the explicit enforcement of the discontinuity have often been proposed as a practical solution (see, e.g., Refs. [28–31]).

Recently, it has been shown [32,33] that the exact strong-interaction limit of DFT [34,35] provides a nonlocal

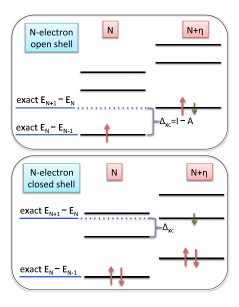


FIG. 1 (color online). Schematic illustration of the spin-restricted (same potential for spin- \uparrow and spin- \downarrow electrons) KS spectrum when adding a tiny fraction η of an electron to an N-electron system. Top: When the KS N-electron system is open shell, the whole KS spectrum "jumps" by a positive constant $\Delta_{\rm xc} = I_N - A_N$, which aligns the KS HOMO to minus the electron affinity $-A_N = E_{N+1} - E_N$. Bottom: When the KS system is closed shell, the positive constant $\Delta_{\rm xc}$ aligns the KS N-electron LUMO to minus the exact affinity $-A_N$. In both cases, in the exact KS system the HOMO is always equal to the many-body chemical potential.

density functional, called "strictly correlated electrons" (SCE) [34–36], which can be used to approximate the exchange-correlation energy of KS DFT, capturing key features of strong correlation such as charge localization in low-density quantum wires without any symmetry breaking [32,33]. This approximation becomes asymptotically exact in the very low-density (or infinitely strong correlation) limit [32,33]. The purpose of this Letter is to generalize the SCE functional to the case of fractional particle numbers, yielding an exchange-correlation functional for open systems that becomes more and more accurate as correlation increases, and that can already be used for modeling nanotransport through low-density quantum wires and dots. Our results also support the assumptions that were made to derive the existence of the derivative discontinuity [4] and provide new insight for the construction of approximate functionals.

The Letter is organized as follows. After briefly reviewing the essential background material, we present the rigorous generalization of the SCE functional to fractional electron numbers. We then show that, without breaking the spin symmetry, the self-consistent KS results with the SCE functional display the right discontinuities at integer electron numbers when correlation is very strong. This is obtained without making any additional assumption or imposing any *ad hoc* constraint. Finally, we draw our conclusions and discuss some perspectives. Hartree (effective) atomic units are used throughout.

Fractional particle numbers in KS DFT.—At zero temperature, open systems with fluctuating particle numbers have been first analyzed in the KS framework by considering statistical mixtures [4]. When dealing with DFT for quantum mechanical systems, we can work with pure states in which a degenerate system is composed by well separated fragments, and one focuses on the energy and density of one of the fragments alone [37–39]. A very simple example is a stretched H₂⁺ molecule, in which on each proton we find, on average, 1/2 electron [13,19]. A key point to prove the existence of the derivative discontinuity is the (empirical) observation that, for integer particle numbers N, the energy E_N of an N-electron system in a given external potential is a concave-up function, i.e., $E_N \leq (1/2)(E_{N+1} + E_{N-1})$. This implies that for fractional electron numbers $Q = N + \eta$ (with $0 \le \eta \le 1$) the exact many-electron ground-state energy E_O and density $\rho_O(\mathbf{r})$ lie on the line connecting the values at the two adjacent integers:

$$E_{N+\eta} = (1-\eta)E_N + \eta E_{N+1}, \tag{1}$$

$$\rho_{N+\eta}(\mathbf{r}) = (1-\eta)\rho_N(\mathbf{r}) + \eta \rho_{N+1}(\mathbf{r}). \tag{2}$$

In KS DFT, one usually aims at obtaining the exact quantities of Eqs. (1) and (2) by means of a noninteracting system of $Q = N + \eta$ electrons in the effective single-particle KS potential $v_s(\mathbf{r}; [\rho_Q])$, which necessarily

changes as we change Q. The energy is then minimized by giving integer occupation to the N single particle KS spin orbitals with lowest eigenvalues and fractional occupation η to the frontier orbital(s), often called HOMO [40,41].

In the exact KS theory, the HOMO eigenvalue ϵ_{HOMO} is the derivative of the total energy of Eq. (1) with respect to the particle number Q, $(\partial E_O/\partial Q) = \epsilon_{HOMO}$ [4,40]. Thus, the exact ϵ_{HOMO} is constant between any two adjacent integers (say, N and N + 1) and equal to the interacting chemical potential $E_{N+1} - E_N$, jumping to a different value when crossing an integer. This "steplike" behavior of the KS ϵ_{HOMO} is not captured by the standard approximate functionals (see, e.g., Refs. [38,41,42]), unless imposed a priori via additional constraints in a spin-unrestricted framework, as, e.g., in Refs. [28,29,31]. The alignment of the exact KS HOMO eigenvalue with the interacting chemical potential implies that the exact KS one-body potential must jump by a constant Δ_{xc} (the derivative discontinuity) when crossing an integer, $v_s(\mathbf{r}; [\rho_{N^+}]) - v_s(\mathbf{r}; [\rho_{N^-}]) =$ $\Delta_{\rm xc}$ (see Fig. 1).

Strong-interaction limit.—The strong-interaction limit of DFT is given by the SCE functional $V_{ee}^{\rm SCE}[\rho]$, defined as the minimum of the electronic interaction alone over all the wave functions yielding the density ρ [34,36,43],

$$V_{ee}^{\text{SCE}}[\rho] = \min_{\Psi \to \rho} \langle \Psi | \hat{V}_{ee} | \Psi \rangle. \tag{3}$$

It can be shown [32,44,45] that in the low-density (or strong-interaction) limit the exact Hartree and exchange-correlation functional $E_{\rm Hxc}[\rho]$ of KS theory tends asymptotically to $V_{ee}^{\rm SCE}[\rho]$.

Physically, the functional $V_{ee}^{\rm SCE}[\rho]$ portrays the *strict* correlation regime, where the position ${\bf r}$ of one electron determines all the other N-1 electronic positions ${\bf r}_i$ through the so-called co-motion functions, ${\bf r}_i={\bf f}_i[\rho]({\bf r})$, some nonlocal functionals of the density [34,35,46,47]. Therefore, the net repulsion on an electron at position ${\bf r}$ due to the other N-1 electrons depends on ${\bf r}$ alone and can be *exactly* represented [33,34,46,47] by a local one-body potential,

$$\nabla \tilde{v}_{\text{SCE}}[\rho](\mathbf{r}) = -\sum_{i=2}^{N} \frac{\mathbf{r} - \mathbf{f}_{i}[\rho](\mathbf{r})}{|\mathbf{r} - \mathbf{f}_{i}[\rho](\mathbf{r})|^{3}},$$
 (4)

which is also the functional derivative of $V_{ee}^{\text{SCE}}[\rho]$, $\delta V_{ee}^{\text{SCE}}[\rho]/\delta \rho(\mathbf{r}) = \tilde{v}_{\text{SCE}}[\rho](\mathbf{r})$ [32,33]. In terms of the co-motion functions, we have [34]

$$V_{ee}^{\text{SCE}}[\rho] = \sum_{i=1}^{N-1} \sum_{j=i+1}^{N} \int d\mathbf{r} \frac{\rho(\mathbf{r})}{N} \frac{1}{|\mathbf{f}_{i}[\rho](\mathbf{r}) - \mathbf{f}_{j}[\rho](\mathbf{r})|}.$$
 (5)

SCE for fractional particle numbers.—The generalization of the SCE formalism to noninteger particle numbers Q is not obvious, because in Eqs. (4) and (5) the sum runs over the integer number of electrons N. To proceed in a rigorous way [3,13,19,37,39], we first analyze a well-separated Q-electron fragment inside a degenerate system

with total integer electron number M, as schematically illustrated in Fig. 2.

Actually, from Eq. (5) it is not evident that $V_{ee}^{\text{SCE}}[\rho]$ separates into the sum of contributions of the isolated fragments, because the interaction $|\mathbf{f}_i[\rho](\mathbf{r}) - \mathbf{f}_j[\rho](\mathbf{r})|^{-1}$ between two electrons on a given fragment may contribute significantly to the integral when \mathbf{r} spans the region of another fragment. However, we have recently shown [48] that another exact expression for $V_{ee}^{\text{SCE}}[\rho]$ is

$$V_{ee}^{\text{SCE}}[\rho] = \frac{1}{2} \int d\mathbf{r} \rho(\mathbf{r}) \sum_{k=2}^{N} \frac{1}{|\mathbf{r} - \mathbf{f}_{k}[\rho](\mathbf{r})|}, \quad (6)$$

which separates into a sum of fragment contributions, showing explicitly that $V_{ee}^{\rm SCE}[\rho]$ is size consistent.

We start from (quasi-)1D systems, for which we can construct the SCE solution analytically. The main findings and conclusions, however, should hold also for 2D and 3D systems [49]. Indeed, we have also performed a three-dimensional self-consistent calculation for a spherically symmetric density, for which we can deduce the fractional SCE solution from our 1D construction, finding similar results.

Figure 2 depicts the reasoning behind the construction of the SCE functional for Q electrons. In the illustrated example, we have solved the SCE problem for M=5 electrons, for a density made of two well-separated identical fragments, each integrating to Q=2.5=N+0.5 particles. We have then studied the "local" SCE solution on the fragments, indicated in Fig. 2 by black circles. Here, the positions $f_i(x)$ of the other four electrons are shown as

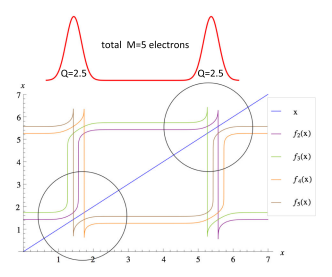


FIG. 2 (color online). Simple example of the analysis used to deduce the SCE solution for fractional electron numbers: we considered a density with M=5 electrons, made of two separated identical fragments (modeled here with two Gaussians), and we have studied the exact SCE solution on each fragment. The graphic shows the positions $f_i(x)$ of the other four electrons as a function of the position x of the first electron. The two black circles represent the "local" SCE solution on each fragment.

a function of the position x of the first electron. We find that two adjacent strictly correlated positions $f_i(x)$ and $f_{i+1}(x)$ on the fragment always satisfy the condition of total suppression of fluctuations [36],

$$\int_{f_i(x)}^{f_{i+1}(x)} \rho(x') dx' = 1, \tag{7}$$

so that there are values of x for which we have three electrons in the fragment, and values of x for which we find only two particles (the third electron is in the other fragment).

Similarly, we find that the general SCE solution for a density integrating to $Q = N + \eta$ electrons can be easily obtained from Eq. (7) (see the Supplemental Material [50] for a detailed derivation). From now on, we work with the fragment alone, considering that on our x axis only the density $\rho_O(x)$ is present. The co-motion functions read

$$f_i(x) = \begin{cases} N_e^{-1}[N_e(x) + i - 1] & x < a_{N+1+\eta-i} \\ N_e^{-1}[N_e(x) + i - N - 2] & x > a_{N+2-i} \\ \infty & \text{otherwise,} \end{cases}$$
(8)

where the function $N_e(x)$ is defined via the density $\rho_O(x)$,

$$N_e(x) = \int_{-\infty}^x \rho_Q(x') dx', \tag{9}$$

 $a_k = N_e^{-1}(k)$, and i = 2, ..., N+1. Notice that even if we have N+1 co-motion functions, when $x \in [a_{N+1+\eta-i}, a_{N+2-i}]$ one of the electrons stays at infinity, so that there are x intervals for which we find N+1 electrons in the density, and x intervals for which we have only N electrons (one of the electrons cannot "enter" in the density).

Self-consistent KS SCE for $N + \eta$ electrons.—From Eqs. (8), (6), and (4) we can now construct the SCE functional $V_{ee}^{\rm SCE}[\rho_Q]$ and its functional derivative $\tilde{v}_{\rm SCE}[\rho_Q]({f r})$ for any one-dimensional density integrating to a noninteger particle number Q. [For $Q = N + \eta$ electrons, the sum in Eqs. (6) and (4) runs up to N+1.] We then consider Q electrons in the quasi-one-dimensional model quantum wire of Refs. [51,52], in which the effective electron-electron interaction is obtained by integrating the Coulomb repulsion on the lateral degrees of freedom [51,53], and is given by $w_b(x) = (\sqrt{\pi}/2b) \exp(x^2/4b^2) \operatorname{erfc}(x/2b)$. The parameter b fixes the thickness of the wire, set to b = 0.1 throughout this work, and erfc(x) is the complementary error function. As in Ref. [52], we consider an external harmonic confinement $v_{\rm ext}(x) = (1/2)\omega^2 x^2$ in the direction of motion of the electrons, which for small ω yields bound states with very low density, where the functional $V_{ee}^{\text{SCE}}[\rho]$ becomes closer and closer to the exact $E_{\rm Hxc}[\rho]$. This is crucial to show that when $E_{\rm Hxc}[\rho] \to V_{ee}^{\rm SCE}[\rho]$ we recover the exact

We have then performed self-consistent, spin-restricted (same KS potential for \uparrow and \downarrow spins) KS calculations with

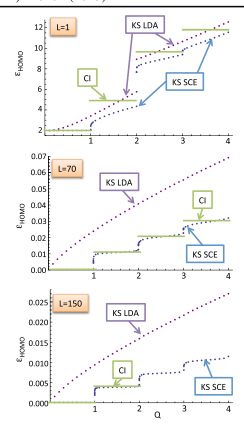


FIG. 3 (color online). Self-consistent results for the spin-restricted KS HOMO eigenvalue (in Hartrees) as a function of the particle number Q for a quasi-1D quantum wire with harmonic confinement along the direction of motion of the electrons, $v_{\rm ext}(x) = (1/2)\omega^2 x^2$, and $\omega = 4/L^2$. The KS SCE results are compared with the standard KS LDA and with the exact chemical potential $E_{N+1} - E_N$ from full CI calculations [33].

the SCE potential, as in Refs. [32,33], this time by giving fractional occupation to the KS HOMO orbital [40,41,54]. The SCE potential $\tilde{v}_{\text{SCE}}[\rho_Q](x)$, representing the approximate Hartree plus exchange-correlation potentials in our KS SCE calculations, is obtained by integrating the 1D analog of Eq. (4), with boundary condition $\tilde{v}_{\text{SCE}}[\rho_Q] \times (|x| \to \infty) = 0$, using the co-motion functions of Eq. (8).

In Fig. 3 we show our self-consistent KS SCE results for the HOMO eigenvalue as a function of the particle number, comparing with the KS local density approximation (LDA) values (the 1D LDA functional is from Ref. [55]), and with the full configuration-interaction (CI) results [33] for the chemical potential $E_{N+1} - E_N$. We have considered a moderately correlated case, L = 1, and two strongly correlated cases, L = 70 and L = 150, where L is an effective confinement length such that $\omega = 4/L^2$. For moderate correlation, L = 1, we see that LDA, having $\Delta_{xc} = 0$, shows a discontinuity in the HOMO value only when filling a new shell (even N, case illustrated in the second panel of Fig. 1), while KS SCE shows a small vertical change also when the N system is open shell (at odd N). When correlation becomes stronger (L = 70 and 150), the KS

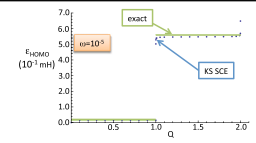


FIG. 4 (color online). Self-consistent results for the spin-restricted KS HOMO eigenvalue as a function of the particle number Q for a three-dimensional electronic system in the external harmonic potential, $v_{\rm ext}(\mathbf{r}) = (1/2)\omega^2 r^2$, with $\omega = 10^{-5}$. The KS SCE results are compared with the exact chemical potential $E_{N+1} - E_N$ [60,61].

SCE self-consistent HOMO approaches more and more the exact step structure, with very good quantitative agreement with the full CI chemical potentials. For such cases, KS LDA yields essentially a continuous curve, since the single particle energies are all very close.

In Fig. 4, we show the result for a 3D system, in which the electron-electron interaction is Coulombic and the external potential is harmonic ("Hooke's atom"). The co-motion functions can be deduced from Eq. (8), as detailed in the Supplemental Material [50]. For small ω , we find that the self-consistent KS SCE HOMO again approaches the exact step structure, which becomes sharper and sharper as correlation increases (ω decreases) [56], similarly to the 1D results of Fig. 3. Notice, again, that for $0 \le Q \le 2$, we have one KS orbital with occupancy Q.

Concluding remarks and perspectives.—The discontinuity in the HOMO for open shell systems (case in the upper panel of Fig. 1) from the self-consistent KS SCE is not just a unique result, but also an independent proof that the exact KS formalism should have this feature. In fact, we have only used the exact $V_{ee}^{\rm SCE}[\rho_Q]$ in the KS self-consistent procedure, without imposing any other condition on our functional. Until now, this feature has only been captured in the context of lattice Hamiltonians [57] or by imposing it a priori in a spin-unrestricted framework [58], as, e.g., in Refs. [28,29,31].

From a practical point of view, our results could already be used to model transport through a correlated quantum wire or quantum dot, going beyond the lattice calculations of Refs. [25,27]. Our findings also provide novel insight for the construction of approximate exchange-correlation functionals: the challenge is to transfer this exact behavior into approximations for less extreme correlation regimes relevant for solid-state physics and chemistry (see Supplemental Material [50] for a test on the H atom). Another challenge is the construction of the SCE functional for general 3D geometry, for which a promising route (that can be generalized to open systems along the lines of this work) is given in Refs. [47,59].

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- [1] W. Kohn and L. J. Sham, Phys. Rev. 140, A1133 (1965).
- [2] P. Hohenberg and W. Kohn, Phys. Rev. 136, B864 (1964).
- [3] A. J. Cohen, P. Mori-Sánchez, and W. Yang, Chem. Rev. 112, 289 (2012).
- [4] J. P. Perdew, R. G. Parr, M. Levy, and J. L. Balduz, Phys. Rev. Lett. 49, 1691 (1982).
- [5] O. Gunnarsson and K. Schönhammer, Phys. Rev. Lett. 56, 1968 (1986).
- [6] R. W. Godby, M. Schlüter, and L. J. Sham, Phys. Rev. Lett. 56, 2415 (1986).
- [7] L. Kleinman, Phys. Rev. B 56, 12 042 (1997).
- [8] J. P. Perdew and M. Levy, Phys. Rev. B **56**, 16 021 (1997).
- [9] L. Kleinman, Phys. Rev. B **56**, 16029 (1997).
- [10] F.E. Zahariev and Y.A. Wang, Phys. Rev. A 70, 042503 (2004).
- [11] M. Grüning, A. Marini, and A. Rubio, Phys. Rev. B 74, 161103 (2006).
- [12] E. Sagvolden and J. P. Perdew, Phys. Rev. A 77, 012517 (2008).
- [13] P. Mori-Sanchez, A. J. Cohen, and W. Yang, Phys. Rev. Lett. 102, 066403 (2009).
- [14] P. Gori-Giorgi and A. Savin, Int. J. Quantum Chem. 109, 2410 (2009).
- [15] V. Brosco, Z. Ying, and J. Lorenzana, Sci. Rep. 3, 2172 (2013).
- [16] J. P. Perdew and M. Levy, Phys. Rev. Lett. **51**, 1884 (1983).
- [17] L. J. Sham and M. Schlüter, Phys. Rev. Lett. 51, 1888 (1983).
- [18] J. P. Perdew, Int. J. Quantum Chem., Quantum Chem. Symp. 19, 497 (1986).
- [19] A. J. Cohen, P. Mori-Sanchez, and W. Yang, Science 321, 792 (2008).
- [20] O. Gritsenko and E. J. Baerends, J. Chem. Phys. 121, 655 (2004).
- [21] D. Vieira, K. Capelle, and C. A. Ullrich, Phys. Chem. Chem. Phys. 11, 4647 (2009).
- [22] M. Hellgren and E. K. U. Gross, Phys. Rev. A 85, 022514 (2012).
- [23] C. Toher, A. Filippetti, S. Sanvito, and K. Burke, Phys. Rev. Lett. 95, 146402 (2005).
- [24] K. Capelle, M. Borgh, K. Kärkkäinen, and S. M. Reimann, Phys. Rev. Lett. **99**, 010402 (2007).
- [25] S. Kurth, G. Stefanucci, E. Khosravi, C. Verdozzi, and E. K. U. Gross, Phys. Rev. Lett. 104, 236801 (2010).
- [26] G. Stefanucci and S. Kurth, Phys. Rev. Lett. **107**, 216401 (2011).
- [27] S. Kurth and G. Stefanucci, Phys. Rev. Lett. 111, 030601 (2013).
- [28] I. Dabo, A. Ferretti, N. Poilvert, Y. Li, N. Marzari, and M. Cococcioni, Phys. Rev. B 82, 115121 (2010).
- [29] S. Refaely-Abramson, S. Sharifzadeh, N. Govind, J. Autschbach, J. B. Neaton, R. Baer, and L. Kronik, Phys. Rev. Lett. 109, 226405 (2012).
- [30] L. Kronik, T. Stein, S. Refaely-Abramson, and R. Baer, J. Chem. Theory Comput. **8**, 1515 (2012).

- [31] E. Kraisler and L. Kronik, Phys. Rev. Lett. 110, 126403 (2013).
- [32] F. Malet and P. Gori-Giorgi, Phys. Rev. Lett. 109, 246402 (2012).
- [33] F. Malet, A. Mirtschink, J. C. Cremon, S. M. Reimann, and P. Gori-Giorgi, Phys. Rev. B 87, 115146 (2013).
- [34] M. Seidl, P. Gori-Giorgi, and A. Savin, Phys. Rev. A 75, 042511 (2007).
- [35] P. Gori-Giorgi, G. Vignale, and M. Seidl, J. Chem. Theory Comput. 5, 743 (2009).
- [36] M. Seidl, Phys. Rev. A 60, 4387 (1999).
- [37] W. Yang, Y. Zhang, and P. W. Ayers, Phys. Rev. Lett. 84, 5172 (2000).
- [38] J. P. Perdew, A. Ruzsinszky, G. I. Csonka, O. A. Vydrov, G. E. Scuseria, V. N. Staroverov, and J. Tao, Phys. Rev. A 76, 040501 (2007).
- [39] P. W. Ayers, J. Math. Chem. 43, 285 (2008).
- [40] J. F. Janak, Phys. Rev. B 18, 7165 (1978).
- [41] O. A. Vydrov, G. E. Scuseria, and J. P. Perdew, J. Chem. Phys. 126, 154109 (2007).
- [42] A. J. Cohen, P. Mori-Sanchez, and W. Yang, Phys. Rev. B 77, 115123 (2008).
- [43] M. Seidl, J. P. Perdew, and M. Levy, Phys. Rev. A 59, 51 (1999).
- [44] M. Levy and J.P. Perdew, Phys. Rev. A **32**, 2010 (1985).
- [45] P. Gori-Giorgi and M. Seidl, Phys. Chem. Chem. Phys. 12, 14 405 (2010).
- [46] P. Gori-Giorgi, M. Seidl, and G. Vignale, Phys. Rev. Lett. **103**, 166402 (2009).
- [47] G. Buttazzo, L. De Pascale, and P. Gori-Giorgi, Phys. Rev. A 85, 062502 (2012).
- [48] A. Mirtschink, M. Seidl, and P. Gori-Giorgi, J. Chem. Theory Comput. **8**, 3097 (2012).
- [49] L.O. Wagner, E.M. Stoudenmire, K. Burke, and S.R. White, Phys. Chem. Chem. Phys. 14, 8581 (2012).
- [50] See Supplemental Material at http://link.aps.org/supplemental/10.1103/PhysRevLett.111.126402 for additional data and details of the derivation of Eq. (8).
- [51] S. Bednarek, B. Szafran, T. Chwiej, and J. Adamowski, Phys. Rev. B 68, 045328 (2003).
- [52] S. H. Abedinpour, M. Polini, G. Xianlong, and M. P. Tosi, Eur. Phys. J. B 56, 127 (2007).
- [53] L. Calmels and A. Gold, Phys. Rev. B 56, 1762 (1997).
- [54] A.P. Gaiduk, D.S. Firaha, and V.N. Staroverov, Phys. Rev. Lett. 108, 253005 (2012).
- [55] M. Casula, S. Sorella, and G. Senatore, Phys. Rev. B 74, 245427 (2006).
- [56] For the failure of standard functionals at this correlation regime see, e.g., M. Taut, A. Ernst, and H. Eschrig, J. Phys. B **31**, 2689 (1998).
- [57] N. A. Lima, M. F. Silva, L. N. Oliveira, and K. Capelle, Phys. Rev. Lett. 90, 146402 (2003).
- [58] An exception to the spin-unrestricted construction is Ref. [13], which attempts to construct, in a spin-restricted framework, a functional showing the linear behavior of Eq. (1).
- [59] C. B. Mendl and L. Lin, Phys. Rev. B 87, 125106 (2013).
- [60] M. Taut, Phys. Rev. A 48, 3561 (1993).
- [61] J. Cioslowski and K. Pernal, J. Chem. Phys. **113**, 8434 (2000).