Improved DFT Potential Energy Surfaces via Improved Densities

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Supporting Information

ABSTRACT: Density-corrected DFT is a method that cures several failures of self-consistent semilocal DFT calculations by using a more accurate density instead. A novel procedure employs the Hartree-Fock density to bonds that are more severely stretched than ever before. This substantially increases the range of accurate potential energy surfaces obtainable from semilocal DFT for many heteronuclear molecules. We show that this works for both neutral and charged molecules. We explain why and explore more difficult cases, for example, CH+, where density-corrected DFT results are even better than sophisticated methods like CCSD. We give a simple criterion for when DC-DFT should be more accurate than self-consistent DFT that can be applied for most cases.

Semilocal density functional theory (DFT) has become a popular workhorse for electronic structure calculations, despite its many known shortcomings. A major failure has always been the stretching of bonds into dissociated fragments. Prototypes of this failure are stretched H_2^+ and stretched H_2, for which semilocal DFT binding energies generically fail to vanish at large separations. The identification of self-interaction error for the former case and static correlation for the latter has been unified in the generic concept of delocalization error of semilocal functionals.

Our focus here is dissociation of heteronuclear diatomic molecular species. In many cases, semilocal DFT dissociates incorrectly into charged fragments, so that the potential energy curve tends to an incorrect value for large R, where R is the interatomic distance. Figure 1 shows the prototypical case, the binding energy curve of NaCl with standard approximations (PBE as a generic GGA functional, and B2PLYP, a double hybrid functional). Beyond about R = 4 Å, the PBE curve becomes highly inaccurate, as it tends to an incorrect dissociation limit, ~1 eV below the sum of Na and Cl atomic energies in PBE. Even a highly accurate double-hybrid functional such as B2PLYP goes to an incorrect dissociation limit of about ~0.7 eV. The work of ref 4 on density-corrected DFT (DC-DFT) showed that by using Hartree-Fock (HF) densities in DC-DFT (which is denoted as HF-DFT) accuracy could be maintained until ~5.6 Å. At that point the HF energy of neutral fragments dips below that of charged fragments, and the HF-DFT curves abruptly jump to zero. This is shown by the light-blue line for the PBE functional.

In the present work, we correct the procedure for applying HF-DFT to this problem and show that it substantially improves results. This procedure yields the solid blue line of Figure 1, which remains smooth and relatively accurate out to a remarkable 8.2 Å. We explain why this happens and that it is no accident. We show that a generalization applies to both molecular anions and cations. We also show some exceptions and how these can still be understood in terms of the basic concepts of DC-DFT. We give a simple criterion for when DC-DFT should improve results over self-consistent DFT. Thus, using our simple procedure, the potential energy surface of most heteronuclear diatomics can be greatly improved.

The theory behind DC-DFT is very straightforward. In a Kohn-Sham DFT (KS-DFT) calculation, the unknown piece of the energy, the exchange-correlation energy, $E_{XC}[\rho]$, is approximated, $E_{XC}[\rho]$. The self-consistent solution of the KS
equations then yields the density that minimizes the total energy, \( \rho^{\text{app}}(\mathbf{r}) \), and the calculated approximate total energy is \( E^{\text{app}}[\rho^{\text{app}}] \). We decompose the error in the energy by writing

\[
\Delta E = (E^{\text{app}}[\rho] - E) + (E^{\text{app}}[\rho^{\text{app}}] - E^{\text{app}}[\rho]) \tag{1}
\]

The first term is the functional error and is the error made in the energy even if we had used the exact density in our approximate functional. The second is the density-driven error and is the error in the energy due solely to the approximate density. In typical semilocal DFT calculations, the second term is much smaller than the first, and these are called normal. In unusual and interesting cases, the second-term becomes comparable or even dominates, and these are abnormal. In such cases, evaluation of the semilocal approximation on a more accurate density yields a significant reduction in the energy error. This is DC-DFT. For many density-driven errors of semilocal DFT calculations, HF densities are sufficient to make the density-driven error negligible.

These effects have been noticed since the start of KS-DFT, especially in the case of atomic anions, where the effect of the density-driven error is severe. Some of the earliest DFT calculations in quantum chemistry actually used HF densities to compare apples-to-apples when testing approximate functionals, but self-consistent densities rapidly became the norm. Pioneering works even showed that many difficult cases for semilocal DFT were improved by using HF densities. Other cases, such as the prototype of stretched H\(_2\), could not be This is a genuine error of the approximate energy functionals and is barely affected by using the exact density.

However, a quantitative and general analysis appeared only recently, showing that many cases of delocalization error could be greatly reduced (if not eliminated) by HF-DFT. This analysis has been used to explain how accurate electron affinities can be extracted from (formally) unconverged DFT calculations, why barrier heights are usually improved by evaluating functionals on HF densities, why the apparently very poor-quality potentials of semilocal approximations usually do not matter in the energy, and many other phenomena. A recent application of DC-DFT to ions and radicals in solution shows that the potential energy surfaces of chloride–water and hydroxide–water complexes are quite accurate with standard approximations such as B3LYP but are almost perfect when B3LYP is evaluated on HF densities. Moreover, while different approximations yielded significantly different potential energy surface when performed self-consistently, their agreement is so good in HF-DFT that there is no need to compare with sophisticated quantum-chemical methods like CCSD(T). We note that implementation of HF-DFT is particularly simple for evaluation of energies within any standard quantum chemical code. Simply solve the HF equations as usual, then use the output HF density as the initial guess for a DFT calculation, but stop the code before the density is updated, as described in a Web site. (Calculation details are given in Supporting Information.)

The present work concerns heteronuclear bond dissociation in semilocal DFT and how to improve it using DC-DFT. (Calculation details are given in Supporting Information.) From very early on, it was recognized that, in many cases, semilocal approximations cause molecules to dissociate into charged fragments. The linearity of the exact functional with particle number guarantees that all diatomics dissociate into fragments of integer charge. Semilocal approximations are smooth functions of particle number and lack the discontinuity in the derivative of the energy at integers. This, in turn, means they have an analytic minimum around some nonzero value of charge transfer between the fragments, \( q \), in the infinitely stretched limit. (An ensemble generalization cures this but only in the infinitely stretched limit.) Nowadays, we call this delocalization error. When \( q \) is finite, the energy of the molecule, stretched to \( R \to \infty \), does not match the sum of the energies of the fragments. In the classic case of NaCl, the stretched molecule is a full 1 eV lower in energy than the fragments in PBE. (In standard practice, this embarrassment is avoided by reporting the binding energy of a molecule as the difference between equilibrium energy and the sum of neutral atomic energies, not the difference between the equilibrium energy and infinitely stretched molecules.)

To explain the correct DC-DFT procedure for heteronuclear diatomics, in Figure 2, we plot several potential energy curves, using dashed lines for singlet and dotted lines for triplets. The self-consistent PBE singlet curve (brown) is smooth but goes to the incorrect limit. The HF curves (red) cross at 5.6 Å, where the minimizing HF density switches to the triplet. Thus, PBE, evaluated on the minimizing HF density, produces the cyan curve of Figure 1, but the correct procedure is to switch curves only when \( E^{\text{PBE}}[\rho^{\text{HF}}] > E^{\text{PBE}}[\rho] \). Thus, HF-DFT follows the blue dashed curve until \( R = 8.2 \) Å. Compared with CCSD(T) (black solid line), it is an almost constant shift prior to 8.2 Å.

In Figure 3, we plot Mulliken charges, \( Q(R) \), as a function of bond length for NaCl. The value of \( Q \) in self-consistent PBE differs only slightly from that of HF in the region \( R = 2.0–4.0 \) Å, but beyond 4 Å, in self-consistent PBE, \( Q \) drops slowly toward its asymptotic value of \( q = 0.4 \). The drop is the onset of the density-driven error, as the HOMO–LUMO gap becomes very small as \( R \) increases. This value of 8.2 Å is not much smaller than the value of 9.5 Å for the exact functional, in which region the charge localized on the ions suddenly drops to 0. On the contrary, the HF charge keeps increasing toward 1, until it suddenly drops to 0. This behavior is generic and can be traced back to the behavior of the approximations for fractionally charged fragments. The exact functional is linear between integer occupations. Semilocal functionals are concave, which causes the minimum of a stretched heteronuclear bond to differ from integer occupations, while HF is convex, producing downward cusps at the integer occupations. Usually this is overcome by using self-interaction-corrected DFT,
orbital-dependent DFT methods, hybrid functionals tuned to have correct linear behavior, and so on.

To practice this assumption, we show the results for several different approximations in Figure 4. The two purely semilocal curves, PBE and TPSS, are almost identical at large R, both self-consistently (dashed) and using the new version of HF-DFT (solid). The B3LYP curve, including $\sim 20\%$ exact exchange, does slightly better self-consistently because its density is $\sim 20\%$ better, but HF-B3LYP is almost identical to HF-PBE, and HF-TPSS is slightly better. They are each much closer to the CCSD(T) curve than their self-consistent analogs and tend toward the right value for large R. Moreover, their errors for large R are not much different from their errors at equilibrium.

Next, we apply our method to charged species to test its limitations. For anions, self-consistent DFT with semilocal approximations is notorious for not even binding the last electron, unless it is artificially (but accurately) confined by a limited basis set. In Figure 5, we plot the binding energy curve of CH$^-$ with several local approximations, both self-consistently and within HF-DFT. The differences between HF-DFT and self-consistent DFT are very small at equilibrium, but after $\sim 1.7$ Å, there is a huge difference. The self-consistent curves are too low and in fact plateau $\sim 1$ eV below the dissociated fragment energies of C$^-$ plus H. On the contrary, the HF-DFT curves are all in close agreement with one another and with the reference CCSD(T) curve. Again, the Mulliken charges of the various self-consistent calculations show the story. In the inset of Figure 5, we plot these for the H atom and see that the HF charge loosely agrees with the self-consistent local DFT charge out to $\sim 1.7$ Å but differs radically after this. Semilocal DFT allows about $q = |Q[H]| = 0.34$ of an electron to transfer from C$^-$ to the H atom, whereas HF localizes the extra electron entirely on the C atom.

We next consider a prototypical example of a molecular cation, CH$. We compare various binding energy curves of CH$^+$ in Figure 6 and Figure S1 (Supporting Information). It is well known that single-reference electronic structure calculations for the dissociation of CH$^+$ have difficulties, so we use a recent ground-state multireference configuration interaction (MRCI) result as reference. Because the atomic MRCI energies of C$^+$ are not given, we chose the MRCI energy of $R = 30$ au ($\sim 15.9$ Å) as the zero point. The pattern is very similar to the previous cases. All semilocal curves go to the incorrect dissociation limit $\sim 1.0$ eV below 0, while HF-DFT curves agree well with the MRCI curve. To emphasize this, in Figure 7, we

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Figure 3. Mulliken charge of Na in NaCl from HF and semilocal DFT calculations. The light-blue line is the HF charge following the previous procedure.

Figure 4. Binding energy curve of NaCl with various DFT approximations. Dashed lines are self-consistent and solid are HF-DFT.

Figure 5. Binding energy curve of CH$^-$ in self-consistent and HF-DFT. Inset: charge of H in CH$^-$ ion. The gray solid line is the HF charge.

Figure 6. Binding energy curves of CH$^+$ in self-consistent and HF-DFT. The black circled solid line is MRCI results from ref 22. Inset: charge of H in CH$^+$ ion. The gray solid line is the HF charge.

Figure 7. Binding energy curve of NaCl with various DFT approximations. Dashed lines are self-consistent and solid are HF-DFT.
plot the errors made by CCSD and the various HF-DFT calculations. (The self-consistent curves have errors of this scale.) Although all curves contain a kink, which is the point where curve crossing between singlet and triplet curve occurs, the magnitude of the error is typically smaller in HF-DFT than in CCSD, being particularly small for HF-TPSS.

So, should we expect all HF-DFT calculations to fix all dissociation problems, yielding accurate binding curves out to very large separations? If not, when do we expect HF-DFT to yield such good curves and when not? How would we know without having to run sophisticated calculations like CCSD(T), thereby defeating the purpose of these low-cost calculations? We next apply the analysis tools of DC-DFT to answer these questions.

The basic tool is to study the gap of the semilocal DFT calculation. An unusually small self-consistent gap (often 0 or below 0.5 eV) suggests a possible density-driven error, so a HF-DFT calculation should be done to see if energies change significantly. None of the neutral or cationic molecules studied here meet this criterion at equilibrium, implying that semilocal results are normal and barely changed by using HF-DFT. If near-equilibrium properties are the only ones needed, there is no reason to do HF-DFT (and substantial reasons to avoid it). Anions are abnormal at all distances and always improved by HF-DFT.

But in essentially every case, the gap shrinks to zero or near-zero at some finite value of \( R \), so that all stretched molecules could be abnormal in semilocal DFT. In fact, the gap of the molecular system in semilocal DFT can be somewhat ambiguous in a stretched molecular system. To find a more useful criterion, we study specifically the stretched limit, that is, where \( R \) is sufficiently large that all quantities are close to their limiting values. In this limit, a powerful result for approximate DFT for neutral molecules has long been known. If the isolated atoms satisfy

\[
\epsilon_{\text{HOMO}}(A) > \epsilon_{\text{LUMO}}(B)
\]

where \( A \) and \( B \) are either of the atoms, then this solution cannot be a minimum for the stretched molecule, and a self-consistent calculation will keep transferring charge until these values match one another. We denote that value by \( q = \lim_{R \to \infty} q(R) \). We have found that this criterion generalizes to molecular ions also. One first chooses the fragments with integer occupations that have lowest energy. Now one of them is an ion. In the case of anions, the LUMO eigenvalues of the semilocal functional are likely to be positive but well-defined with a carefully chosen basis set (as shown in ref 12). In Table S1 (Supporting Information), we give a table of atomic neutrals and ions showing the relevant levels.

Using the atomic inputs, we construct Table 1 for the molecules and molecular ions discussed here and other examples. The HOMO–LUMO criterion eq 2 applies in all cases for large \( R \): If it is satisfied, \( q \) is finite, and semilocal DFT tends to an incorrect energy at large \( R \). By incorrect, we mean different from the sum of the energies of the isolated fragments. The stretched molecule is abnormal and is usually improved by a HF-DFT calculation, as shown in our figures.

An important point is that this analysis applies only in the stretched limit. Even if a molecule is normal in this limit, that is, has \( q = 0 \), it still has a small or vanishing gap for finite \( R \) and must be checked for abnormality at finite \( R \). In such cases, HF-DFT might still yield an accurate binding energy curve out to larger \( R \) than self-consistent DFT, but we do not have a simple condition for predicting when this will be the case.

Figure 8 shows a typical case of a normal calculation. Semilocal functionals all go correctly to the zero dissociation limit, and HF-DFT worsens the result for all \( R \). The charges do not vary much and vanish at large \( R \) regardless of the functional you use. There are also some cases where the abnormality depends on the particular semilocal DFT calculation. Consider HF in Figure 9. From our table, PBE transfers a small charge at large \( R \), making it abnormal, and the PBE binding energy curve is improved by HF-PBE. This also holds for TPSS, where HF-TPSS slightly improves the results, but this is not true for B3LYP, so HF-B3LYP does not improve its result.

### Table 1. Various Properties of Molecular Species Calculated With PBE

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<th>( A )</th>
<th>( B )</th>
<th>( q )</th>
<th>( \epsilon_{\text{HOMO}}(A) )</th>
<th>( \epsilon_{\text{LUMO}}(B) )</th>
<th>( \Delta \epsilon )</th>
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"Amount of charge transfer in the dissociation limit, \( q \), is calculated at \( R = 5.0 \) Å and with the def2-QZVP basis.\(^{28}\) \( A \) is for electropositive atomic species, while \( B \) is for electronegative ones. If not specified, \( \epsilon \) is calculated with the aug-cc-pV5Z basis set.\(^{29}\)\(^{30}\) Calculated with the cc-pVSZ basis set.\(^{30}\)
We have commented only on heteronuclear diatomics, even though the classic cases of breakdown of DFT due to delocalization error are homogeneous (stretched H$_2^+$ and H$_3$). This is because we study only abnormal calculations and how to fix them. The classic examples do not have large density-driven errors.

To conclude, considerable evidence shows that, in many cases, semilocal DFT errors in heteronuclear binding energy curves are density-driven and can be reduced by using HF-DFT. We have significantly extended the range of DC-DFT beyond previous suggestions and have given a clear criterion for when DC-DFT will be better than self-consistent DFT. In the infinitely stretched limit, DC-DFT always yields the correct dissociation, which is the sum of the fragment energies (within DC-DFT). If a self-consistent DFT density incorrectly transfers charge, the HF density will be better. While this is not an ideal procedure in terms of model chemistries or a black box that can be applied unambiguously to every problem, DC-DFT provides a pragmatic approach for those who need dissociation curves at larger distances while retaining a DFT-level of computational cost as well as food for thought for more fundamental theory development.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpclett.5b01724.

Computational details and frontier orbital energies of atomic species. (PDF)

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**Notes**

The authors declare no competing financial interest.

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