Self-consistent density-based basis-set correction: How much do we lower total energies and improve dipole moments?

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ABSTRACT
This work provides a self-consistent extension of the recently proposed density-based basis-set correction method for wave function electronic-structure calculations [E. Giner et al., J. Chem. Phys. 149, 194301 (2018)]. In contrast to the previously used approximation where the basis-set correction density functional was a posteriori added to the energy from a wave-function calculation, here the energy minimization is performed including the basis-set correction. Compared to the non-self-consistent approximation, this allows one to lower the total energy and change the wave function under the effect of the basis-set correction. This work addresses two main questions: (i) What is the change in total energy compared to the non-self-consistent approximation and (ii) can we obtain better properties, namely, dipole moments, with the basis-set corrected wave functions. We implement the present formalism with two different basis-set correction functionals and test it on different molecular systems. The main results of the study are that (i) the total energy lowering obtained by the self-consistent approach is extremely small, which justifies the use of the non-self-consistent approximation, and (ii) the dipole moments obtained from the basis-set corrected wave functions are improved, being already close to their complete basis-set values with triple-zeta basis sets. Thus, the present study further confirms the soundness of the density-based basis-set correction scheme.

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I. INTRODUCTION

One of the main limitations of electronic-structure calculations based on wave-function theory (WFT) is the slow convergence of the results with respect to the size of the one-electron basis set. This aspect is of fundamental importance in quantum chemistry as WFT methods have otherwise many interesting features. In particular, in a given basis set \( B \), WFT methods can usually be systematically improved toward the exact solution provided by full configuration interaction (FCI). At the root of the slow convergence of WFT lies the singularity of the Coulomb interaction: Because it becomes infinite at electron–electron coalescence points, it creates a derivative discontinuity of the wave function at these points, the so-called electron–electron cusp, which is not representable by the usual incomplete basis sets employed. To cure the slow convergence problem of WFT, two main approaches have emerged: basis-set extrapolation techniques and explicitly correlated F12 methods. Basis-set extrapolation techniques consist in exploiting the known asymptotic behavior of WFT properties as a function of the size of the basis set in order to estimate the complete basis-set (CBS) limit based on several calculations with basis sets of increasing sizes. Explicitly correlated F12 methods consist in supplementing to the usual basis sets a correlation factor that restores the electron–electron cusp and accelerates the convergence toward the CBS limit. Although these F12 methods are increasingly popular in quantum chemistry, they have the drawback of needing rather complex three- and four-electron integrals and, more generally, of involving a relatively complex mathematical formalism, which makes the adaptation of a WFT method to its F12 version a non-trivial task.
An alternative path to speed up the convergence of WFT calculations with respect to the size of the basis set has been recently proposed by some of the present authors by exploiting the ability of range-separated density-functional theory (RSDFT) to recover the short-range correlation effects missing from an incomplete basis set. The central idea developed in Ref. 13 is to define a mapping between the electron–electron Coulomb interaction projected into an incomplete basis set \( \mathcal{B} \) and the non-diverging long-range electron–electron interaction \( \text{erf}(\mu r_{12})/r_{12} \) used in RSDFT. The connection is done through the definition of a range-separation parameter \( \mu \), which varies in space and automatically adapts to the basis set \( \mathcal{B} \). Once this adaptive range-separation parameter is defined, one can use a special flavor of short-range correlation density functionals used in RSDFT for the estimation of the correlation energy missing in the considered basis set \( \mathcal{B} \). An important property of this RSDFT-based approach is that the basis-set energy correction properly vanishes in the CBS limit. This strategy was successfully validated for the calculations of ionization energies. Once this adaptive range-separation parameter is considered as the best variational approximation to the exact ground-state energy \( E_0 \) and energy \( E^\mathcal{B}_0 \) converge to the exact ground-state energy \( E_0 \) and energy \( E^\mathcal{B}_0 \), respectively.

The paper is organized as follows: In Sec. II A, we present the exact theory of the self-consistent basis-set correction scheme using either a function of density only or a function of both density and on-top pair density and we recall the non-self-consistent approximation previously employed. In Sec. II B, we introduce our approximations of the unknown exact basis-set correction functional by short-range functionals. In Sec. II C, we explain how we solve the self-consistent basis-set correction equations by a selected configuration interaction algorithm. In Sec. III, we report and discuss the results on the total energies of the Be atom and the BH molecule and on the dipole moments of the BH, FH, H\(_2\)O, and CH\(_2\) molecules. Section IV contains our conclusions. Unless otherwise specified, Hartree (Ha) atomic units (a.u.) are used throughout the paper.

II. THEORY

A. Self-consistent basis-set correction

1. Basis-set correction as a function of density

We start by reviewing the scheme where the basis-set correction is written as a function of density, which was first developed in Ref. 13. Given a basis set \( \mathcal{B} \), the exact ground-state energy \( E_0 \) of an electronic system can be approximated by the energy \( E^\mathcal{B}_0 \) obtained by the following minimization over \( \mathcal{B} \)-representable one-electron densities \( n^\mathcal{B}_0 \), i.e., densities that can be obtained from a wave function \( \Psi^\mathcal{B}_0 \) belonging to the \( N \)-electron Hilbert space generated by the basis set \( \mathcal{B} \),

\[
E^\mathcal{B}_0 = \min_{n^\mathcal{B}_0} \left\{ F[n^\mathcal{B}_0] + \int \text{d}r \, v_{\text{ne}}(r) \, n^\mathcal{B}_0(r) \right\},
\]

where \( v_{\text{ne}}(r) \) is the nuclei–electron interaction potential. In this expression, \( F[n^\mathcal{B}_0] \) is the standard Levy–Lieb constrained-search universal density functional evaluated at \( n^\mathcal{B}_0 \),

\[
F[n^\mathcal{B}_0] = \min_{\Psi^\mathcal{B}} \left\{ \langle \Psi^\mathcal{B} | \hat{T} + \hat{W}_{\text{ee}} | \Psi^\mathcal{B} \rangle \right\},
\]

where \( \hat{T} \) and \( \hat{W}_{\text{ee}} \) are the kinetic-energy operator and the Coulomb electron–electron interaction operator, respectively, and the notation \( \Psi^\mathcal{B} \rightarrow n^\mathcal{B}_0 \) means a \( N \)-electron wave function yielding the density \( n^\mathcal{B}_0 \). It is important to notice that the wave functions \( \Psi^\mathcal{B} \) used in the definition of \( F[n^\mathcal{B}_0] \) in Eq. (2) are not restricted to be expandable in the basis set \( \mathcal{B} \) but should instead be thought of as expanded on a complete basis set. The minimizing density \( n^\mathcal{B}_0 \) in Eq. (1) can be considered as the best variational approximation to the exact ground-state density \( n_0 \). Importantly, when the basis set \( \mathcal{B} \) is increased up to the CBS limit, the density \( n^\mathcal{B}_0 \) and energy \( E^\mathcal{B}_0 \) converge to the exact ground-state density \( n_0 \) and energy \( E_0 \), respectively.

\[
\lim_{\mathcal{B} \rightarrow \text{CBS}} n^\mathcal{B}_0 = n_0 \quad \text{and} \quad \lim_{\mathcal{B} \rightarrow \text{CBS}} E^\mathcal{B}_0 = E_0.
\]

Since the \( \mathcal{B} \)-representability restriction is only applied to densities and not to wave functions, the basis-set convergence of \( E^\mathcal{B}_0 \) to \( E_0 \) is much faster than in a usual WFT calculation.

We then decompose the universal Levy–Lieb density functional as

\[
F[n^\mathcal{B}_0] = \min_{\Psi^\mathcal{B}} \left\{ \langle \Psi^\mathcal{B} | \hat{T} + \hat{W}_{\text{ee}} | \Psi^\mathcal{B} \rangle + \hat{E}[n^\mathcal{B}_0] \right\},
\]

where \( \Psi^\mathcal{B} \) designates wave functions restricted to the \( N \)-electron Hilbert space generated by the basis set \( \mathcal{B} \) and \( \hat{E}[n^\mathcal{B}_0] \) is the complementary basis-set correction density functional required to make Eq. (4) exact. This basis-set correction functional \( \hat{E}[n^\mathcal{B}_0] \) recovers the part of the energy that is missing in the first term of the right-hand side of Eq. (4) due to the basis-set restriction of the wave functions \( \Psi^\mathcal{B} \). Inserting Eq. (4) into Eq. (1) and recombinining the two minimizations, we can obtain \( E^\mathcal{B}_0 \) by the following minimization over \( \mathcal{B} \)-restricted wave functions \( \Psi^\mathcal{B} \):

\[
E^\mathcal{B}_0 = \min_{\Psi^\mathcal{B}} \left\{ \langle \Psi^\mathcal{B} | \hat{T} + \hat{W}_{\text{ee}} + \hat{V}_{\text{ne}} | \Psi^\mathcal{B} \rangle + \hat{E}[n^\mathcal{B}_0] \right\},
\]

where \( \hat{V}_{\text{ne}} \) is the nuclei–electron interaction operator and \( n^\mathcal{B}_0(r) = \langle \Psi^\mathcal{B} | \hat{n}(r) | \Psi^\mathcal{B} \rangle \) is the density of the wave function \( \Psi^\mathcal{B} \), where we have introduced the density operator \( \hat{n}(r) = \sum_{\sigma \in \{1, \ldots N\}} \hat{\psi}_\sigma^\dagger(r) \hat{\psi}_\sigma(r) \) written in real-space second quantization. The minimizing wave function \( \Psi^\mathcal{B}_0 \) in Eq. (5) satisfies the following self-consistent Schrödinger-like equation:

\[
\hat{H}^\mathcal{B}_0[n^\mathcal{B}_0] | \Psi^\mathcal{B}_0 \rangle = E^\mathcal{B}_0 | \Psi^\mathcal{B}_0 \rangle,
\]

where \( \hat{H}^\mathcal{B}_0 \) is the Lagrange multiplier associated with the normalization constraint of the wave function \( \Psi^\mathcal{B}_0 \) and the effective Hamiltonian is defined for a given density \( n^\mathcal{B}_0 \) as

\[
\hat{H}^\mathcal{B}_0[n^\mathcal{B}_0] = \hat{T} + \hat{W}_{\text{ee}} + \hat{V}_{\text{ne}} + \hat{V}_{\text{ext}}[n^\mathcal{B}_0].
\]

In this expression, \( \hat{T} \), \( \hat{W}_{\text{ee}} \), and \( \hat{V}_{\text{ne}} \) are the kinetic, electron–electron, and electron–nuclei operators projected in
the \( N \)-electron Hilbert space generated by the basis set \( B \), and \( \hat{V}^B[n^B] \) is the one-electron effective potential operator,

\[
\hat{V}^B[n^B] = \int \mathrm{d}r \, \hat{v}^B(r) \, n^B(r),
\]

where \( \hat{v}^B(r) = \delta \hat{E}^B[n^B]/\delta \hat{n}^B(r) \) and \( \hat{n}^B(r) \) is the density operator projected in the basis set \( B \).

Using real-valued spatial orthonormal orbitals \( \{\phi_n \} \) spanning the same space as the basis set \( B \), the expression of the effective Hamiltonian in second quantization is

\[
\hat{H}^B_{\text{eff}}[n^B] = \sum_{pq} \left( \hbar_{pq} + \hat{v}_{pq}^B \right) \hat{c}_{pq} + \frac{1}{2} \sum_{pqrs} \hat{w}_{pqrs} \hat{c}_{pq} \hat{c}_{rs},
\]

where \( \hat{c}_{pq} = \hat{d}^+_p \hat{d}_q \) and \( \hat{w}_{pqrs} = \hat{E}_{pq} - \delta_{pq} \hat{E}_{ps} \) are the spin-singlet one- and two-particle elementary operators, \( \hbar_{pq} \) are the usual one-electron integrals, \( \hat{w}_{pqrs} = \langle pq|rs \rangle \) are the usual two-electron integrals, and \( \hat{v}_{pq}^B \) are the one-electron integrals associated with the effective potential \( \hat{v}^B(r) \).

\[
\hat{v}_{pq}^B = \int \mathrm{d}\mathbf{r} \hat{\phi}_p(r) \hat{v}^B(r) \hat{\phi}_q(r).
\]

Note that in Eq. (9), we put \( B \) on top of the sum symbols to indicate that the sums run over all orbitals generated by the basis set \( B \).

Finally, note that we have considered the total density \( n^B \) for simplicity, but the theory can be trivially extended to spin densities \( n^B_{\uparrow} \) and \( n^B_{\downarrow} \).

2. Basis-set correction as a function of density and on-top pair density

We now extend the theory to allow for a basis-set correction functional depending on both the density \( n_{\uparrow\uparrow}(\mathbf{r}) \) and the on-top pair density \( n_{\uparrow\downarrow}(\mathbf{r}) \) of a wave function \( \Psi^B \), where we have introduced the on-top pair density operator \( \hat{n}_{\uparrow\downarrow}(\mathbf{r}) = \sum_{\{\uparrow\downarrow\}} \langle \uparrow\downarrow|\hat{\psi}(\mathbf{r})\hat{\psi}(\mathbf{r})\rangle \). In the spirit of the generalized Kohn–Sham scheme (see also Ref. 22), we write the universal Levy–Lieb density functional as

\[
\begin{align*}
F[n^B] &= \min_{\Psi^B \in \mathcal{H}} \left\{ \langle \Psi^B|\hat{T} + \hat{W}_{\text{xc}}|\Psi^B \rangle + \hat{E}^B[n_{\uparrow\uparrow}, n_{\uparrow\downarrow}] \right\},
\end{align*}
\]

where \( \hat{E}^B[n_{\uparrow\uparrow}, n_{\uparrow\downarrow}] \) can be any functional of \( n_{\uparrow\uparrow} \) and \( n_{\uparrow\downarrow} \) such that the minimization in Eq. (11) exactly gives \( F[n^B] \). Insertion into Eq. (1) leads to

\[
\hat{H}^B_{\text{eff}}[n^B_{\uparrow\uparrow}, n^B_{\uparrow\downarrow}][\Psi_0^B] = \hat{E}_0^B[\Psi_0^B],
\]

and the minimizing wave function \( \Psi_0^B \) satisfies the following self-consistent Schrödinger-like equation:

\[
\hat{H}^B_{\text{eff}}[n^B_{\uparrow\uparrow}, n^B_{\uparrow\downarrow}][\Psi_0^B] = \hat{E}_0^B[\Psi_0^B],
\]

where the effective Hamiltonian is defined for a \( B \)-representable density \( n^B \) and on-top pair density \( n^B_{\uparrow\downarrow} \) as

\[
\hat{H}^B_{\text{eff}}[n^B, n^B_{\uparrow\downarrow}] = \hat{T}^B + \hat{W}_{\text{xc}} + \hat{V}_{\text{eff}} + \hat{\tilde{V}}^B[n^B, n^B_{\uparrow\downarrow}] + \hat{\tilde{W}}^B[n^B, n^B_{\uparrow\downarrow}].
\]

In this expression, \( \hat{\tilde{V}}^B[n^B, n^B_{\uparrow\downarrow}] \) is the one-electron effective potential operator,

\[
\hat{\tilde{V}}^B[n^B, n^B_{\uparrow\downarrow}] = \int \mathrm{d}\mathbf{r} \, \hat{\tilde{v}}^B(r) \, n^B(r),
\]

where \( \hat{\tilde{v}}^B(r) = \delta \hat{E}^B[n^B]/\delta \hat{n}^B(r) \) and \( \hat{\tilde{w}}^B[n^B, n^B_{\uparrow\downarrow}] \) is the two-electron effective interaction operator,

\[
\hat{\tilde{W}}^B[n^B, n^B_{\uparrow\downarrow}] = \frac{1}{2} \int \mathrm{d}\mathbf{r} \, \hat{\tilde{w}}^B(r) \, n^B_{\uparrow\downarrow}(r),
\]

where \( \hat{\tilde{w}}^B(r) = 2\delta \hat{E}^B[n^B, n^B_{\uparrow\downarrow}]/\delta n^B_{\uparrow\downarrow}(r) \) and \( n^B_{\uparrow\downarrow}(r) \) is the on-top pair density operator projected in the basis set \( B \). The second-quantized expression of the effective Hamiltonian is

\[
\hat{H}^B_{\text{eff}}[n^B, n^B_{\uparrow\downarrow}] = \sum_{pq} \left( \hbar_{pq} + \hat{v}_{pq}^B \right) \hat{c}_{pq} + \frac{1}{2} \sum_{pqrs} \left( \hat{w}_{pqrs} + \hat{\tilde{w}}_{pqrs} \right) \hat{c}_{pq} \hat{c}_{rs},
\]

leading to the following approximation for \( E_0^B \), for the basic theory of Sec. II A 1:

\[
E_0^B \approx E_{\text{FCI}}^B + \hat{E}^B[\Psi_{\text{FCI}}^B],
\]

where \( E_{\text{FCI}}^B \) is the standard FCI energy (or an estimate of it) in the basis set \( B \), and for the extended theory of Sec. II A 2,

\[
E_0^B \approx E_{\text{FCI}}^B + \hat{E}^B[\Psi_{\text{FCI}}^B, n^B_{\uparrow\downarrow}],
\]

The approximation in Eq. (19) is in fact equivalent to approximating the minimizing density \( n_0^B \) in Eq. (1) by the standard FCI ground-state density \( n_{\text{FCI}}^B \),

\[
n_0^B(\mathbf{r}) \approx n_{\text{FCI}}^B(\mathbf{r}),
\]

which seems intuitively a reasonable approximation as one expects \( n_{\text{FCI}}^B \) and \( n_0^B \) to be both close to the exact density \( n_0 \), and the encouraging numerical results obtained for energies with this non-self-consistent approximation tend to confirm the validity of Eq. (22). Nevertheless, in the present study, we will investigate the quantitative effect on energies and dipole moments of performing the minimization in Eq. (5) or Eq. (12).
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ARTICLE

B. Approximations for the basis-set correction functional \( E^B \)

1. Local range-separation parameter

As originally proposed in Ref. 13, the basis-set correction functional \( E^B \) can be mapped to the so-called short-range correlation functional with multideterminant reference introduced in Ref. 23 in the context of RSDFT. This mapping relies on the definition of a local range-separation parameter \( \mu \).

\[
\mu^B(\mathbf{r}) = \frac{\sqrt{\pi}}{2} W^B(\mathbf{r}),
\]

which provides a local measure of the incompleteness of the basis set \( B \). It is defined such that the long-range electron–electron interaction of RSDFT, \( W^B(\mathbf{r}) = \text{erf}(\mu \mathbf{r})/r_\mathbf{r} \), coincides at coalescence (i.e., at \( r_\mathbf{r} = 0 \)) with an effective interaction representing the Coulomb electron–electron interaction projected in the basis set \( B \). The expression of this effective interaction at coalescence is

\[
\bar{W}^B(\mathbf{r}) = \left\{ \begin{array}{ll}
\frac{f_n^B(\mathbf{r})}{n_n^B}\psi^B_\mathbf{r}(\mathbf{r}) & \text{if } n_{n_n^B}\psi^B_\mathbf{r}(\mathbf{r}) \neq 0,
\infty & \text{otherwise},
\end{array} \right.
\]

with

\[
f_n^B(\mathbf{r}) = \sum_{\mu \gamma \rho \nu} w_{\rho \gamma \nu \mu} g_{\rho \gamma \nu \mu}(\mathbf{r}) \phi_{\rho}(\mathbf{r}) \phi_{\gamma}(\mathbf{r}) \phi_{\nu}(\mathbf{r}),
\]

where \( g_{\rho \gamma \nu \mu}(\mathbf{r}) = 2\langle \Psi_{\text{loc}}^B|d^\dagger_\rho d^\gamma d^\nu d^\mu|\Psi_{\text{loc}}^B \rangle \) is the opposite-spin two-electron density matrix of some “localizing” wave function \( \Psi_{\text{loc}}^B \), and \( n_{n_n^B}\psi^B_\mathbf{r}(\mathbf{r}) \) is its associated on-top pair density.

\[
n_{n_n^B}\psi^B_\mathbf{r}(\mathbf{r}) = \sum_{\mu \gamma \rho \nu} \Gamma_{\rho \gamma \nu \mu} \phi_{\rho}(\mathbf{r}) \phi_{\gamma}(\mathbf{r}) \phi_{\nu}(\mathbf{r}).
\]

The wave function \( \Psi_{\text{loc}}^B \) is only used to localize the effective interaction projected in the basis set \( B \). The local range-separation parameter is very weakly dependent on this wave function \( \Psi_{\text{loc}}^B \). It should be thought of as essentially dependent on the basis set \( B \). Importantly, in the CBS limit, the effective interaction goes to the Coulomb interaction that diverges at coalescence, and consequently, the local range-separation parameter goes to infinity,

\[
\lim_{B \to \text{CBS}} \mu^B(\mathbf{r}) = \infty,
\]

independently of \( \Psi_{\text{loc}}^B \), which is fundamental to guarantee the correct behavior of the theory in the CBS limit.

2. Approximate basis-set correction functionals from short-range functionals

Approximations for the basis-set correction functional \( E^B \) are obtained by using the previously defined local range-separation parameter in short-range correlation functionals. Specifically, for the basis-set correction functional in Eq. (5), we use the so-called PBE-UEG basis-set correction functional (UEG stands for “uniform electron gas”),

\[
E_{\text{PBE-UEG}}^B[n] = \int d\mathbf{r} \epsilon_{\text{c,md}}^\text{at} (\mathbf{r}, \nabla n(\mathbf{r}), n_2^\text{UEG}(\mathbf{r}), \mu(\mathbf{r})),
\]

and for the basis-set correction functional in Eq. (12), we use the so-called PBE-OT basis-set correction functional (OT stands for “on-top”),

\[
E_{\text{PBE-OT}}^B[n, n_2] = \int d\mathbf{r} \epsilon_{\text{c,md}}^\text{at}(\mathbf{r}, \nabla n(\mathbf{r}), n_2(\mathbf{r}), \mu(\mathbf{r})),
\]

where we have dropped the superscript \( B \) in the density, in the on-top pair density, and in the local range-separation parameter for simplicity. In these expressions, the short-range (sr) correlation energy density with multideterminant (md) reference \( \epsilon_{\text{c,md}}^\text{at} \) has the following generic form in terms of the density \( n \), the density gradient \( \nabla n \), the on-top pair density \( n_2 \), and the range-separation parameter \( \mu \).

\[
\epsilon_{\text{c,md}}^\text{at}(\mathbf{r}, \nabla n(\mathbf{r}), n_2(\mathbf{r}), \mu(\mathbf{r}))) = \frac{\epsilon_{\text{c,PBE}}^\text{at}(\mathbf{r}, \nabla n(\mathbf{r}))}{1 + \beta(n_s, n_2, \mu) \mu^3},
\]

where \( \epsilon_{\text{c,PBE}}^\text{at}(\mathbf{r}, \nabla n(\mathbf{r})) \) is the usual Perdew–Burke–Ernzerhof (PBE) correlation energy density\(^\text{\textsuperscript{24}}\) and \( \beta(n_s, n_2, \mu) = (2\sqrt{\pi}(1 - \sqrt{2})/3)\).

The key difference between the PBE-UEG and PBE-OT functionals is the on-top pair density used. The PBE-UEG functional uses \( n_2^\text{UEG}(\mathbf{r}) \), which is an estimate of the exact on-top pair density using a parametrization of the on-top pair density of the uniform electron gas (UEG) at density \( n(\mathbf{r}) \),

\[
n_2^\text{UEG}(\mathbf{r}) = n(\mathbf{r})^2 g_0(n(\mathbf{r})),
\]

where the on-top pair-distribution function \( g_0(n) \) is taken from Eq. (46) of Ref. 26. By contrast, the PBE-OT functional uses \( n_2(\mathbf{r}) \), which is an estimate of the exact on-top pair density obtained from extrapolating the input on-top pair density \( n_2^\text{at}(\mathbf{r}) \) of the wave function \( \Psi^B \) to the limit \( \mu \to \infty \) (see Ref. 26),

\[
n_2(\mathbf{r}) = \left(1 + \frac{2}{\sqrt{\mu} r_\mathbf{r}} \right)^{-1} n_2^\text{at}(\mathbf{r}).
\]

As shown in Ref. 16, the difference between the two flavors of on-top pair densities comes from the treatment of strong correlation. While \( n_2^\text{UEG}(\mathbf{r}) \) is a good approximation of the exact on-top pair density for weakly correlated situations, when strong-correlation effects are present, it fails to represent the large depletion of the exact on-top pair density, and in this case, \( n_2(\mathbf{r}) \) provides a much better approximation of the exact on-top pair density.

The explicit expression of the one-electron effective potential associated with the PBE-UEG functional,

\[
\tilde{v}_{\text{PBE-UEG}}^B(\mathbf{r}) = \frac{\delta E_{\text{PBE-UEG}}^B[n]}{\delta n(\mathbf{r})},
\]

was already given in a previous study.\(^\text{\textsuperscript{24}}\) The corresponding potential for the PBE-OT functional,

\[
\tilde{v}_{\text{PBE-OT}}^B(\mathbf{r}) = \frac{\delta E_{\text{PBE-OT}}^B[n, n_2]}{\delta n(\mathbf{r})},
\]

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has a very similar expression, with the simplification that the on-
top pair density \( n_2 \) in Eq. (29) is taken as independent of the den-
sity, whereas \( n_2^{\text{loc}} \) in Eq. (28) is a function of density. For the
PBE-OT functional, we have in addition the two-electron effective
interaction,

\[
\omega_{\text{PBE-OT}}^\theta = \frac{\delta E_{\text{PBE-OT}}^\theta [n, n_2]}{\delta n_2 (r)}. \tag{36}
\]

Its explicit expression

\[
\omega_{\text{PBE-OT}}^\theta (r) = \frac{\partial^2 E_{\text{PBE-OT}}^\theta (n, n_2)}{\partial n_2 (r)} \cdot \nabla n_2 (r), \tag{37}
\]

where

\[
\frac{\partial n_2 (r)}{\partial n_2 (r)} = \left( 1 + \frac{2}{\sqrt{n_2 (r)}} \right)^{-1}, \tag{38}
\]

and

\[
\frac{\partial^2 E_{\text{PBE-OT}}^\theta (n, n_2, \mu)}{\partial n_2 (r)} = \frac{\varepsilon_{\text{PBE-OT}}^\theta (n, \nabla n_2, \mu)}{2} \frac{\mu^3 c (n_2)^2}{n_2}. \tag{39}
\]

Note that the localizing wave function \( \Psi_{\text{loc}}^\theta \) used in the
definition of the local range-separation parameter (see Sec. II B 1) is taken
as fixed, i.e., we do not consider variations of \( \Psi_{\text{loc}}^\theta \). Importantly,
since the local range-separation parameter diverges in the CBS
limit [Eq. (27)] and since the correlation energy per particle \( \mu^\theta \in
\Psi_{\text{loc}}^\theta \) in Eq. (30) vanishes for \( \mu \to \infty \), both basis-set
correction functionals correctly vanish in the CBS limit,

\[
\lim_{\mu \to \infty} E_{\text{PBE-UEG}}^\theta [n] = 0, \tag{40}
\]

\[
\lim_{\mu \to \infty} \omega_{\text{PBE-OT}}^\theta [n, n_2] = 0, \tag{41}
\]

i.e., the CBS limit is unaltered by the correction.

3. Frozen-core version of the basis-set correction

functionals

When the wave function \( \Psi^\theta \) is calculated in the frozen-core
approximation, we use the frozen-core version of the basis-set
correction functionals introduced in Ref. 15. The basis-set correction
functionals become

\[
\hat{E}^\theta [n_{\text{loc}}^\theta] \rightarrow \hat{E}^\theta [n_{\text{loc}}^\theta], \tag{42}
\]

and

\[
\hat{E}^\theta [n_{\text{loc}}^\theta, n_{\text{val}}^\theta] \rightarrow \hat{E}^\theta [n_{\text{loc}}^\theta, n_{\text{val}}^\theta], \tag{43}
\]

where \( \Psi_{\text{val}}^\theta \) is the wave function \( \Psi^\theta \) truncated to the "valence"
orbital space (i.e., with all core orbitals removed). Accordingly, the
local range-separation parameter becomes

\[
\mu^\theta (r) \rightarrow \mu_{\text{val}}^\theta (r), \tag{44}
\]

where

\[
\mu_{\text{val}}^\theta (r) = \frac{\sqrt{\pi}}{2} W_{\text{val}}^\theta (r). \tag{45}
\]

The valence-only effective interaction at coalescence is

\[
W_{\text{val}}^\theta (r) = \begin{cases} 
\frac{f_{\Psi_{\text{loc}}^\theta}^\theta (r)}{n_{\Psi_{\text{loc}}^\theta}^\theta (r)} & \text{if } n_{\Psi_{\text{loc}}^\theta}^\theta (r) \neq 0, \\
\infty & \text{otherwise},
\end{cases} \tag{46}
\]

where

\[
f_{\Psi_{\text{loc}}^\theta}^\theta (r) = \sum_{pqrs} w_{pqrs} \Gamma_{pqrs} \phi_{pq}^\theta (r) \phi_{rs}^\theta (r) \phi_{pq}^\theta (r) \phi_{rs}^\theta (r), \tag{47}
\]

where \( \Gamma_{pqrs} = 2 \langle \Psi_{\text{loc},\text{val}}^\theta | \hat{\alpha}_{pq}^\theta \hat{\alpha}_{rs}^\theta | \Psi_{\text{loc},\text{val}}^\theta \rangle \) is the opposite-spin two-electron density matrix of the localization wave function truncated to the valence orbital space \( \Psi_{\text{loc},\text{val}}^\theta \) and \( n_{\Psi_{\text{loc}}^\theta}^\theta (r) \) is its associated on-
top pair density,

\[
n_{\Psi_{\text{loc}}^\theta}^\theta (r) = \sum_{pqrs} \Gamma_{pqrs} \phi_{pq}^\theta (r) \phi_{rs}^\theta (r) \phi_{pq}^\theta (r) \phi_{rs}^\theta (r). \tag{48}
\]

In Eqs. (47) and (48), the indication "\( \text{loc} \)" on top of the sum symbols
means that the sums are only over valence (i.e., non-core) orbitals. It
is noteworthy that \( \mu_{\text{val}}^\theta (r) \) still fulfills Eq. (27), and thus, the frozen-
core versions of the basis-set correction functionals still correctly
vanish in the CBS limit.

Correspondingly, the frozen-core versions of the effective
Hamiltonians in Eqs. (7) and (14) are simply obtained by setting to
zero all the one-electron effective integrals \( \varepsilon_{pq}^\theta \) and the two-electron
effective integrals \( \omega_{pq}^\theta \), if at least one orbital involved in the integral
is a core orbital.

C. Selected configuration interaction to solve
the self-consistent eigenvalue equations

To solve the self-consistent basis-set correction eigenvalue

\[
E_{\text{eff}}^\theta (k) = \frac{1}{\mathcal{R}(k)} \sum_{I \in \mathcal{R}(k)} c_I^k |I\rangle, \tag{49}
\]

where \( \mathcal{R}(k) \) denotes a set of Slater determinants. We then use the
density of this wave function to form the following effective
Hamiltonian at the next iteration \( k \):

\[
E_{\text{eff}}^\theta (k) = \frac{1}{\mathcal{R}(k)} \sum_{I \in \mathcal{R}(k)} c_I^k |I\rangle, \tag{50}
\]

and we want to find the associated ground-state wave function

\[
\Psi_{\text{eff}}^\theta (k) \equiv E_{\text{eff}}^\theta (k) |\Psi_{\text{eff}}^\theta (k)\rangle. \tag{51}
\]

This wave function is obtained by the CIPSI algorithm as

\[
|\Psi_{\text{eff}}^\theta (k)\rangle = \sum_{I \in \mathcal{R}(k)} c_I^k |I\rangle, \tag{52}
\]
where \( R^{(k)} \) is the new set of Slater determinants at iteration \( k \). According to the CIPSI algorithm, the set \( R^{(k)} \) is obtained by repeatedly adding to a reference wave function \( \Psi_{\text{ref}}^{R^{(k)}} \) the determinants \( K \) having the largest second-order perturbation theory (PT2) contributions \( |c_{K,\text{PT2}}^{(k)}| \) with

\[
|c_{K,\text{PT2}}^{(k)}| = \frac{\left( \frac{\partial^2}{\partial r^2} \left( \Psi_{\text{ref}}^{\Psi_{\text{ref}}^{R^{(k)}}} \right) - \Psi_{\text{ref}}^{\Psi_{\text{ref}}^{R^{(k)}}} \right) \right)^2}{\left( K \right) H_{\text{eff}}^{(k)} \left( K \right) - \left( \Psi_{\text{ref}}^{\Psi_{\text{ref}}^{R^{(k)}}} \right) H_{\text{eff}}^{(k)} \left( \Psi_{\text{ref}}^{\Psi_{\text{ref}}^{R^{(k)}}} \right)},
\]

iteratively doubling the number of determinants in \( \Psi_{\text{ref}}^{R^{(k)}} \) until the absolute value of the total PT2 energy correction due to the missing determinants,

\[
|\varepsilon_{\text{PT2}}^{(k)}| = \sum_{K \in \mathcal{G}^{(k)}} |c_{K,\text{PT2}}^{(k)}|,
\]

is smaller than a given threshold. To reduce the cost of the evaluation of the PT2 contribution, the semi-stochastic multi-reference approach of Garniron et al.\(^{30}\) is used, adopting the technical specifications recommended in that work.

This determines the set of determinants \( R^{(k)} \), which is then fixed for the rest of the iteration \( k \). The energy \( E_0^{R^{(k)}} \) for this iteration is then determined according to the minimization in Eq. (5),

\[
E_0^{R^{(k)}} = \min_{\Psi_0^{R^{(k)}}} \left\{ \sum_{I} \left[ T + W_{\text{ee}} + V_{\text{ex}} \Psi_0^{R^{(k)}} \right] + E_{\text{loc}}^{R^{(k)}} \right\},
\]

which amounts to solving the iterative equation,

\[
H_{\text{eff}}^{(k)} \left( \Psi_0^{R^{(k)}} \right) = E_0^{R^{(k)}} \left( \Psi_0^{R^{(k)}} \right),
\]

for the optimal coefficients of the determinants \( \{c_I^{(k)}\} \) leading to the minimizing wave function at iteration \( k \),

\[
|\Psi_0^{R^{(k)}}| = \sum_{I \in \mathcal{G}^{(k)}} c_I^{(k)} |I|.
\]

The iterations over \( k \) are repeated until the variation of \( E_0^{R^{(k)}} \) is smaller than a given threshold. The evaluation of the dipole moment is obtained as the expectation value of the dipole operator over the converged wave function \( \Psi_0^{R^{(k)}} \).

The same approach is used for solving Eq. (13), which involves the on-top pair density in addition to the density.

III. COMPUTATION OF TOTAL ENERGIES AND DIPOLE MOMENTS

A. Computational details

We study the total ground-state energies of the Be atom and BH molecule together with the dipole moments of the BH, FH, and \( \text{H}_2\text{O} \) molecules in their ground states and of the \( \text{CH}_2 \) molecule in its lowest spin-singlet state. We report the standard CIPSI (i.e., near FCI) results without the basis-set correction (referred to as “CIPSI”) as well as CIPSI results including the basis-set correction using the PBE-UEG and PBE-OT functionals with or without self-consistency. The non-self-consistent calculations are referred to as “CIPSI-PBE-UEG” and “CIPSI-PBE-OT,” whereas the self-consistent calculations are referred to as “SC CIPSI-PBE-UEG” and “SC CIPSI-PBE-OT” where SC stands for self-consistent. The orbitals used for all converged CIPSI calculations are the natural orbitals obtained from a standard CIPSI calculation stopped at a total PT2 energy correction smaller in absolute value than 0.001 hartree. For the localizing wave function \( \Psi_{\text{loc}}^{\Psi_{\text{ref}}^{R^{(k)}}} \) involved in the definition of the local range-separation parameter \( \mu_c^{\Psi_{\text{ref}}^{R^{(k)}}} \) (see Sec. II B 1), we choose either a single Slater determinant (SD) built from the natural orbitals of the largest CIPSI wave function (which we refer to as \( \mu_{\text{CIPSI}}^{\Psi_{\text{ref}}^{R^{(k)}}} \)) or the largest CIPSI wave function (which we refer to as \( \mu_{\text{CIPSI}}^{\Psi_{\text{ref}}^{R^{(k)}}} \)). We use the Dunning correlation-consistent basis-set family.\(^{31-35}\) We perform both non-frozen-core calculations using the core-valence aug-cc-pCVXZ basis sets and frozen-core calculations (with the 1s orbitals of non-hydrogen atoms frozen) using the valence aug-cc-pVXZ basis sets and the corresponding frozen-core version of the basis-set correction (see Sec. II B 3). All the CIPSI calculations have been performed with QUANTUM PACKAGE.\(^{37}\)

We also report the dipole moment at the coupled cluster singles doubles perturbative triples [CCSD(T)] level, which was taken from Ref. 37 for the BH and FH molecule and obtained using linear-response calculations from the DALTON software.\(^{38,39}\) For the BH and \( \text{H}_2\text{O} \) molecules. The molecular geometries are taken from Ref. 37 for BH and FH and from Ref. 40 for \( \text{H}_2\text{O} \) and \( \text{CH}_2 \).

B. Total energies of the Be atom and the BH molecule

Tables I and II report the total energies of the Be atom and the BH molecule, respectively, calculated using the aug-cc-pCVXZ basis sets with CIPSI without any basis-set correction and with different basis-set corrections. It can be observed that the total energies obtained with the basis-set corrections converge much faster toward the estimated exact total energies than the total energies obtained without any basis-set correction. For the Be atom, all the basis-set corrected total energies from the aug-cc-pCVXZ to the aug-cc-pCVQZ basis sets have errors below 1.6 mhartree \( \approx 1 \) kcal/mol compared to the estimated exact total energy, whereas without basis-set correction such an accuracy is not even reached with the aug-cc-pCVQZ basis set. Similar trends are observed for the BH molecule: All the basis-set corrected total energies have errors below 1 kcal/mol already from the aug-cc-pCVTZ basis set, whereas without basis-set correction such an accuracy is reached only with the aug-cc-pCV5Z basis set.

Focusing now on the differences between the various basis-set corrections, we can notice that (i) using the local range-separation parameter \( \mu_{\text{CIPSI}}^{\Psi_{\text{ref}}^{R^{(k)}}} \) gives larger basis-set corrections than using \( \mu_{\text{CIPSI}}^{\Psi_{\text{ref}}^{R^{(k)}}} \) and (ii) the PBE-UEG functional gives larger basis-set corrections than the PBE-OT functional. As regards the effect of the self-consistency, it is remarkable to notice that self-consistency lowers the total energy by a very small fraction of the total basis-set correction (typically less than 1%), whatever the choice of functional or local range-separation parameter. These results thus validate the previously introduced non-self-consistent approximation to the basis-set correction (see Sec. II A 3) for energy calculations.
TABLE I. Total ground-state energies (in hartree) of the Be atom calculated using the aug-cc-pCVXZ (ACVXZ) basis sets (with \( X = D, T, Q \)) with CIPSI without the frozen-core approximation and including different basis-set corrections with or without self-consistency. The energy lowering \( \Delta E_{SC} \) (in \( \mu \)hartree) from the non-self-consistent to the self-consistent version of the basis-set correction is reported in square brackets.

| Basis Set | ACVDZ       | ACVTZ       | ACVQZ       |
|-----------|-------------|-------------|-------------|
| CIPSI     | –14.651 922 5 | –14.662 397 1 | –14.665 576 7 |
| CIPSI + PBE – UEG\(_{\text{red}}\) | –14.668 361 7 | –14.668 631 4 | –14.668 102 0 |
| SC CIPSI + PBE – UEG\(_{\text{red}}\) [\( \Delta E_{SC} \)] | –14.668 387 8 [–26.1] | –14.668 363 4 [–4.0] | –14.668 102 6 [–0.6] |
| CIPSI + PBE – UEG\(_{\text{ccpsis}}\) | –14.667 703 5 | –14.668 376 2 | –14.667 964 3 |
| SC CIPSI + PBE – UEG\(_{\text{ccpsis}}\) [\( \Delta E_{SC} \)] | –14.667 739 5 [–36.0] | –14.668 387 4 [–9.2] | –14.667 968 1 [–3.8] |
| CIPSI + PBE – OT\(_{\text{unc}}\) | –14.666 337 6 | –14.667 884 6 | –14.667 978 2 |
| SC CIPSI + PBE – OT\(_{\text{unc}}\) [\( \Delta E_{SC} \)] | –14.666 374 1 [–36.5] | –14.667 895 6 [–11.0] | –14.667 802 0 [–3.8] |
| CIPSI + PBE – OT\(_{\text{ccpsis}}\) | –14.665 946 3 | –14.667 712 8 | –14.667 714 0 |
| SC CIPSI + PBE – OT\(_{\text{ccpsis}}\) [\( \Delta E_{SC} \)] | –14.665 974 8 [–28.5] | –14.667 722 3 [–9.5] | –14.667 717 1 [–3.1] |

Exact non-relativistic total energy* | –14.667 356 5

*From Ref. 41.

TABLE II. Total ground-state energies (in hartree) of the BH molecule calculated using the aug-cc-pCVXZ (ACVXZ) basis sets (with \( X = D, T, Q, 5 \)) with CIPSI without the frozen-core approximation and including different basis-set corrections with or without self-consistency. The energy lowering \( \Delta E_{SC} \) (in \( \mu \)hartree) from the non-self-consistent to the self-consistent version of the basis-set correction is reported in square brackets.

| Basis Set | ACVDZ       | ACVTZ       | ACVQZ       | ACV5Z      |
|-----------|-------------|-------------|-------------|------------|
| CIPSI     | –25.255 015 0 | –25.278 617 9 | –25.285 758 3 | –25.287 377 9 |
| CIPSI + PBE – UEG\(_{\text{red}}\) | –25.283 817 9 | –25.289 630 3 | –25.290 677 2 | –25.290 007 9 |
| SC CIPSI + PBE – UEG\(_{\text{red}}\) [\( \Delta E_{SC} \)] | –25.283 927 0 [–109.1] | –25.289 647 1 [–16.8] | –25.290 680 4 [–3.2] | … |
| CIPSI + PBE – UEG\(_{\text{ccpsis}}\) | –25.282 507 9 | –25.289 024 5 | –25.290 397 5 | … |
| SC CIPSI + PBE – UEG\(_{\text{ccpsis}}\) [\( \Delta E_{SC} \)] | –25.282 609 0 [–101.1] | –25.289 040 0 [–15.5] | –25.290 397 5 | … |
| CIPSI + PBE – OT\(_{\text{unc}}\) | –25.279 829 7 | –25.288 048 6 | –25.289 977 4 | … |
| SC CIPSI + PBE – OT\(_{\text{unc}}\) [\( \Delta E_{SC} \)] | –25.280 039 1 [–209.4] | –25.288 100 8 [–52.2] | –25.289 993 7 [–16.3] | … |
| CIPSI + PBE – OT\(_{\text{ccpsis}}\) | –25.278 973 8 | –25.287 636 3 | –25.287 788 3 | … |
| SC CIPSI + PBE – OT\(_{\text{ccpsis}}\) [\( \Delta E_{SC} \)] | –25.279 160 0 [–186.2] | –25.287 680 9 [–44.6] | … |

CIPSI total energy extrapolated to the CBS limit | –25.289 032

C. Dipole moments of the BH, FH, \( \text{H}_2\text{O} \), and \( \text{CH}_2 \) molecules

As seen from Sec. III B, the self-consistency of the basis-set correction does not lead to significant changes in the total energies. Nevertheless, one can wonder if the effective wave functions obtained with the self-consistent basis-set correction provide better properties.

We choose to investigate this aspect through the computation of the dipole moments of the BH, FH, \( \text{H}_2\text{O} \), and \( \text{CH}_2 \) molecules for several reasons: (i) The basis-set convergence of dipole moments with correlated wave function methods is known to be slow (see, e.g., Refs. 37, 40, and 42), (ii) these molecules are sufficiently small to have near-CBS reference values, and (iii) the BH and \( \text{CH}_2 \) molecules exhibit a non-trivial mixture of both strong and weak correlation due to the 2s–2p near degeneracy in the boron and carbon atoms, whereas the FH and \( \text{H}_2\text{O} \) molecules are dominated by weak correlation effects.

1. Dipole moment of the BH molecule

We start by a detailed analysis on the BH molecule. The slow convergence of the dipole moment of the BH molecule with respect to the size of the basis set has been illustrated at various correlation levels including CCSD(T) by Halkier et al.,\(^\text{43}\) and we report in Table III their CCSD(T) results at the non-frozen-core level. In order to have a reasonable estimate of the dipole moment in the CBS limit, we use the two-point \( X^3 \) extrapolation proposed in Ref. 37 using the aug-cc-pCVQZ and aug-cc-pCV5Z basis sets. At the CCSD(T) level, an error of about 0.001 \( \text{a.u.} \) with respect to the CBS limit is not even reached with the aug-cc-pCV5Z basis set, while the error is about 0.006 and 0.003 \( \text{a.u.} \) with the aug-cc-pCVTZ and aug-cc-pCVQZ basis set, respectively, showing indeed the quite slow convergence of the dipole moment of BH.

We also report in Table III the dipole moment at the CIPSI level together with the value of the PT2 energy correction \( \langle \epsilon_{\text{PT2}} \rangle \) associated with the variational wave function for which the dipole moment has
been calculated. The values of $|\mu_{\text{PT2}}|$ are all below $2 \times 10^{-5}$ hartree, which was found to be mandatory to converge the CIPSI dipole moments to a precision of about 0.0001 a.u. for the BH molecule. These represent therefore rather large calculations involving about 10^{10} Slater determinants for the largest aug-cc-pCV5Z basis set. In order to obtain an estimation of the error of the dipole moment at the CIPSI level in a given basis set with respect to FCI, we extrapolate the CIPSI dipole moment to $E_{\text{PT2}} \to 0$ using a linear extrapolation as a function of $|E_{\text{PT2}}|$ (similar to the proposal of Holmes et al.\textsuperscript{14} for the total energy) using different values of $|E_{\text{PT2}}|$. As one can see from Table III, for any basis set, the difference between the basis-set moment calculated by CIPSI with the smallest $E_{\text{PT2}}$ available and the dipole moment extrapolated with respect to $|E_{\text{PT2}}|$ is negligible. Thus, with the values of $|E_{\text{PT2}}|$ used, the CIPSI dipole moments provide reliable estimates of the FCI dipole moments. As regards basis-set errors, similar to the CCSD(T) results, an error of about 0.001 a.u. on the CIPSI dipole moment with respect to the CBS limit is not even reached using the aug-cc-pCV5Z basis set, illustrating once more the slow convergence of the dipole moment with respect to the size of the basis set.

From Table III, one can also notice that there is a small deviation between the CCSD(T) and the CIPSI dipole moments, which is of about 0.002 a.u. for the aug-cc-pCVXZ basis sets with $X = D, T, Q$ and about 0.001 a.u. for the aug-cc-pCV5Z basis set. In addition, the deviation between the CCSD(T) and CIPSI dipole moments extrapolated to the CBS limit is about 0.001 a.u., showing that CCSD(T) provides a very accurate value for the dipole moment of the BH molecule.

Coming now to the self-consistent basis-set correction calculations, we report in Table III the dipole moments obtained using the PBE – UEG and PBE – OT functionals using the local range-separation parameters $\mu_{\text{SD}}$ and $\mu_{\text{CIPSI}}$ and a graphical representation of the data for $\mu_{\text{SD}}$ is given in Fig. 1(a). Clearly, the basis-set correction strongly accelerates the convergence to the CBS limit. More specifically, it can be observed that (i) in a given basis set, all basis-set corrections significantly improve the dipole moment with respect to the CIPSI value, (ii) the PBE – OT functional gives more accurate results than the PBE – UEG functional, and (iii) an error of about 0.001 a.u. on the dipole moment is obtained already with the aug-cc-pCVTZ basis set when using the PBE – OT functional. The result

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**TABLE III.** Dipole moment (in a.u.) of the ground state of the BH molecule calculated using the aug-cc-pCVXZ (ACVXZ) basis sets (with $X = D, T, Q, 5$) by Hartree–Fock (HF), CCSD(T), and CIPSI without the frozen-core approximation and including different self-consistent basis-set corrections. For the CIPSI calculations, the PT2 energy correction $|E_{\text{PT2}}|$ (in hartree) is reported in square brackets. Extrapolations to the CBS limit are given in the last column.

| Basis Set | ACVDZ | ACVTZ | ACVQZ | ACV5Z | CBS  |
|-----------|-------|-------|-------|-------|------|
| HF        | 0.68796 | 0.68570 | 0.68489 | 0.68530 |      |
| CCSD(T)   | 0.52968 | 0.54649 | 0.54984 | 0.55125 | 0.55271 |
| CIPSI $|E_{\text{PT2}}|$ | 0.52758 [2 × 10^{-6}] | 0.54388 [4 × 10^{-6}] | 0.54975 [2 × 10^{-5}] | 0.55142 |
| CIPSI extrapolated to $E_{\text{PT2}} \to 0$ | 0.52757 | 0.54386 | 0.54970 | 0.54967 | 0.55126 |
| SC CIPSI + PBE – UEG$|E_{\text{CIPSI}}|$ | 0.53658 [2 × 10^{-6}] | 0.55092 [4 × 10^{-6}] | 0.55093 [1 × 10^{-5}] |      |
| SC CIPSI + PBE – OT$|E_{\text{CIPSI}}|$ | 0.53477 [2 × 10^{-6}] | 0.54973 [8 × 10^{-6}] |      |      |

*From Ref. 37."
The dipole moment (in a.u.) of the ground state of the BH molecule calculated using the aug-cc-pVXZ (AVXZ) basis sets (with X = D, T, Q, 5) by Hartree–Fock (HF), CCSD(T), and CIPSI with the frozen-core approximation and including different self-consistent basis-set corrections. For the CIPSI calculations, the PT2 energy correction \( |E_{PT2}| \) (in hartree) is reported in square brackets. Extrapolations to the CBS limit are given in the last column.

|          | AVDZ       | AVTZ       | AVQZ       | AVSZ       | CBS       |
|----------|------------|------------|------------|------------|-----------|
| HF       | 0.68796    | 0.68590    | 0.68494    | 0.68496    |           |
| CCSD(T)  | 0.52939    | 0.54500    | 0.54724    | 0.54843    | 0.54966   |
| CIPSI \( |E_{PT2}| \) | 0.52782 [3 \times 10^{-8}] | 0.54334 [1 \times 10^{-7}] | 0.54563 [4 \times 10^{-7}] | 0.54691 [1 \times 10^{-6}] | 0.54823   |
| SC CIPSI + PBE – UEG \( \text{end} \) \( |E_{PT2}| \) | 0.53791 [4 \times 10^{-7}] | 0.54815 [7 \times 10^{-7}] | 0.54971 [3 \times 10^{-6}] | 0.54715 [3 \times 10^{-6}] |
| SC CIPSI + PBE – OT \( \text{anh} \) \( |E_{PT2}| \) | 0.54512 [4 \times 10^{-7}] | 0.55029 [5 \times 10^{-7}] | 0.54880 [2 \times 10^{-8}] |           |

(i) shows that, although the self-consistency does not lead to significant improvement on the total energy (see Table II), it is crucial to yield effective wave functions providing better properties, illustrating the impact and the accuracy of the effective basis-set correction potentials employed. The result (ii) shows that the use of the on-top pair density of the wave function rather than that estimated from the UEG gives a better approximation of the exact on-top pair density, which can be understood as a signature of strong-correlation effects.

Finally, we also report in Table IV the dipole moments obtained with the frozen-core approximation using the aug-cc-pVXZ basis sets, and the corresponding graphical representation is given in Fig. 1(b). Again, the basis-set correction with either the PBE – UEG or PBE – OT functional yields a faster basis-set convergence of the dipole moment than that in the standard CIPSI calculations. However, in this case, the convergence toward the CBS limit is slightly better when using the PBE – UEG functional. The PBE – OT functional slightly overestimates the dipole moment by 0.002 and 0.0006 a.u. with the aug-cc-pVTZ and aug-cc-pVQZ basis sets, respectively, whereas the PBE – UEG functional yields a deviation of below 0.0005 a.u. from the aug-cc-pVTZ to the aug-cc-pV5Z basis set.

**D. Dipole moments of the FH, H\(_2\)O, and CH\(_2\) molecules**

We now pursue our analysis on the FH, H\(_2\)O, and CH\(_2\) molecules using only the frozen-core approximation. The basis-set convergence of the dipole moments of these molecules was studied in Refs. 37 and 42 at the CCSD(T) level. In Tables V–VII, we report the CCSD(T) and CIPSI results with the aug-cc-pVXZ basis sets.

The CBS estimates are obtained from a two-point \( X^{-3} \) extrapolation using the aug-cc-pVQZ and aug-cc-pV5Z basis sets at the CCSD(T) level and also at the CIPSI level in the case of CH\(_2\). One can notice that, at the CCSD(T) or CIPSI level, an error of about 0.001 a.u. with respect to the estimated CBS limit is not even reached with the aug-cc-pV5Z basis set. In addition, in the case of CH\(_2\), there is a significant discrepancy between the extrapolated CCSD(T) and CIPSI dipole moments, which might be due to some strong-correlation effects that are mistreated at the CCSD(T) level.

We also report in Tables V–VII the results using the self-consistent basis-set correction. In contrast with the BH molecule, the effective wave functions obtained with the PBE – UEG and PBE – OT functionals always yield very similar dipole moments for the FH and H\(_2\)O molecules. This can be explained by the fact that these molecules at their equilibrium geometries are weakly correlated systems for which the on-top pair density based on the UEG is a good approximation. For the FH, H\(_2\)O, and CH\(_2\) molecules, the dipole moment is overestimated with the aug-cc-pVDZ basis set using both functionals, but the results with the aug-cc-pVTZ basis set are already very close to the estimated CBS limit. From a quantitative point of view, for the FH molecule with the aug-cc-pVDZ basis set, the error with respect to the CBS dipole moment is reduced from about 0.007 a.u. at the CIPSI level to about 0.004 a.u. with the basis-set correction, whereas with the aug-cc-pV5Z basis set the error is reduced from 0.005 a.u. to below 0.0005 a.u.

For the CH\(_2\) molecule, with the aug-cc-pVDZ basis set the error with respect to the CBS extrapolated CIPSI value is about 0.008 a.u., whereas it is about 0.003 and 0.006 a.u. using the PBE – UEG and PBE – OT basis-set corrections, respectively. With the aug-cc-pVTZ basis set, the error at the CIPSI level is still about 0.005 a.u., whereas it is below 0.001 a.u. for both the PBE – UEG and PBE – OT functionals.

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**Table IV.** Dipole moment (in a.u.) of the ground state of the BH molecule calculated using the aug-cc-pVXZ (AVXZ) basis sets (with X = D, T, Q, 5) by Hartree–Fock (HF), CCSD(T), and CIPSI with the frozen-core approximation and including different self-consistent basis-set corrections. For the CIPSI calculations, the PT2 energy correction \( |E_{PT2}| \) (in hartree) is reported in square brackets. Extrapolations to the CBS limit are given in the last column.

|          | AVDZ       | AVTZ       | AVQZ       | AVSZ       | CBS       |
|----------|------------|------------|------------|------------|-----------|
| HF       | 0.75976    | 0.75750    | 0.75634    | 0.75617    |           |
| CCSD(T)  | 0.70342    | 0.70465    | 0.70707    | 0.7094     | 0.70903   |
| CIPSI \( |E_{PT2}| \) | 0.70249 [9 \times 10^{-6}] | 0.70406 [1 \times 10^{-4}] | 0.70662 [1 \times 10^{-4}] |
| CIPSI extrapolated with respect to \( E_{PT2} \) | 0.70248 | 0.70391 | 0.70646 |           |
| SC CIPSI + PBE – UEG \( \text{end} \) \( |E_{PT2}| \) | 0.71326 [3 \times 10^{-5}] | 0.70873 [1 \times 10^{-4}] |           |
| SC CIPSI + PBE – OT \( \text{anh} \) \( |E_{PT2}| \) | 0.71362 [2 \times 10^{-5}] | 0.70915 [1 \times 10^{-4}] |           |
Finally, in the case of the H$_2$O molecule, the results using the basis-set correction are actually worst than the CIPSI ones when using the aug-cc-pVDZ basis set, the error increasing from about 0.003 a.u. for CIPSI to about 0.009 a.u. with the basis-set correction. One should nevertheless keep in mind that the convergence of the dipole moment of H$_2$O is non-monotonic at the CCSD(T) level, the dipole moment obtained with the aug-cc-pVDZ basis set being closer to the CBS limit than the ones obtained using the aug-cc-pVTZ or aug-cc-pVQZ basis sets. Therefore, the seemingly good values obtained at the CCSD(T) and CIPSI levels using the aug-cc-pVDZ basis set are likely to be due to a compensation of errors. With the aug-cc-pVTZ basis set, the expected trend is recovered, with CIPSI giving an error of about 0.006 a.u. and the basis-set correction reducing this error to about 0.001 a.u.

**IV. CONCLUSION**

In the present work, we have established the fully self-consistent density-based basis-set correction scheme.\(^1\) Different from previous works where a non-self-consistent approximation was used,\(^1,13-18\) here the energy is minimized in the presence of the basis-set correction functional, which (i) guarantees to get a lower total energy with respect to the non-self-consistent approximation and (ii) allows the wave function to change under the presence of the basis-set correction. We have tested this scheme on a few atomic and molecular systems (Be, BH, FH, H$_2$O, and CH$_2$) with CIPSI wave functions and two different basis-set correction functionals PBE - UEG and PBE - OT. While PBE - UEG is a functional of the density, PBE - OT uses in addition the on-top pair density of the wave function as an independent variable.

The main results are that (i) the lowering in total energy is extremely small compared to the non-self-consistent approximations (typically less than 1%), which thus justifies this approximation for energy-only calculations, and (ii) the wave functions obtained from the self-consistent basis-set correction scheme yield dipole moments, which converge much faster with respect to the size of the basis set than standard wave function calculations, being already close to the CBS value with a triple-zeta basis set. This study thus confirms that the density-based basis-set correction scheme is not only useful for energy calculations but also for calculations of response properties.

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**DATA AVAILABILITY**

The data that support the findings of this study are available within the article and from the corresponding author upon reasonable request.

**REFERENCES**

1. E. A. Hylleraas, *Z. Phys.* 54, 347 (1929).
2. T. Kato, *Commun. Pure Appl. Math.* 10, 151 (1957).
3. T. Helgaker, W. Klopper, H. Koch, and J. Noga, *J. Chem. Phys.* 106, 9659 (1997).
4. A. Halkier, T. Helgaker, P. Jørgensen, W. Klopper, H. Koch, J. Olsen, and A. K. Wilson, *Chem. Phys. Lett.* 286, 243 (1998).
5. S. Ten-no, *Theor. Chem. Acc.* 131, 1070 (2012).
6. S. Ten-no and J. Noga, *Wiley Interdiscip. Rev.: Comput. Mol. Sci.* 2, 114 (2012).
7. C. Häfträg, W. Klopper, A. Kohn, and D. P. Tew, *Chem. Rev.* 112, 4 (2012).
8. L. Kong, F. A. Bischoff, and E. F. Valeev, *Chem. Rev.* 112, 75 (2012).
9. A. Grünneis, S. Hirata, Y.-Y. Ohnishi, and S. Ten-no, *J. Chem. Phys.* 146, 080901 (2017).

**TABLE VI.** Dipole moment (in a.u.) of the ground state of the H$_2$O molecule calculated using the aug-cc-pVXZ (AVXZ) basis sets (with X = D, T, Q, 5) by Hartree–Fock (HF), CCSD(T), and CIPSI with the frozen-core approximation and including different self-consistent basis-set corrections. For the CIPSI calculations, the PT2 energy correction $\Delta E_{\text{PT2}}$ (in hartree) is reported in square brackets. Extrapolations to the CBS limit are given in the last column.

|        | AVDZ       | AVTZ       | AVQZ       | AV5Z       | CBS       |
|--------|------------|------------|------------|------------|-----------|
| HF     | 0.786 70   | 0.780 38   | 0.779 55   | 0.779 56   | 0.729 41  |
| CCSD(T)| 0.727 03   | 0.723 64   | 0.726 95   | 0.728 15   | 0.729 41  |
| CIPSI  | 0.726 10 ($3 \times 10^{-5}$) | 0.722 94 ($2 \times 10^{-4}$) | 0.728 18 ($2 \times 10^{-5}$) | 0.728 55 ($1 \times 10^{-4}$) | 0.729 41 |
| SC CIPSI + PBE - UEG$_{\text{ex}}$ | 0.728 09 ($2 \times 10^{-5}$) | 0.728 18 ($2 \times 10^{-5}$) | 0.728 55 ($1 \times 10^{-4}$) | 0.729 41 |
| SC CIPSI + PBE - OT$_{\text{ex}}$ | 0.730 40 ($2 \times 10^{-4}$) | 0.728 55 ($1 \times 10^{-4}$) | 0.729 41 |

**TABLE VII.** Dipole moment (in a.u.) of the lowest spin-singlet state of the CH$_2$ molecule calculated using the aug-cc-pVXZ (AVXZ) basis sets (with X = D, T, Q, 5) by Hartree–Fock (HF), CCSD(T), and CIPSI with the frozen-core approximation and including different self-consistent basis-set corrections. For the CIPSI calculations, the PT2 energy correction $\Delta E_{\text{PT2}}$ (in hartree) is reported in square brackets. Extrapolations to the CBS limit are given in the last column.

|        | AVDZ       | AVTZ       | AVQZ       | AV5Z       | CBS       |
|--------|------------|------------|------------|------------|-----------|
| HF     | 0.748 78   | 0.744 78   | 0.743 55   | 0.743 53   | 0.664 16  |
| CCSD(T)| 0.656 14   | 0.660 09   | 0.662 11   | 0.663 10   | 0.664 16  |
| CIPSI  | 0.651 20 ($2 \times 10^{-5}$) | 0.654 46 ($3 \times 10^{-5}$) | 0.656 43 ($4 \times 10^{-5}$) | 0.657 80 ($1 \times 10^{-4}$) | 0.659 26 |
| SC CIPSI + PBE - UEG$_{\text{ex}}$ | 0.662 49 ($2 \times 10^{-5}$) | 0.659 58 ($3 \times 10^{-5}$) | 0.658 90 ($3 \times 10^{-5}$) | 0.659 26 |
| SC CIPSI + PBE - OT$_{\text{ex}}$ | 0.665 27 ($2 \times 10^{-4}$) | 0.660 55 ($4 \times 10^{-5}$) | 0.659 32 ($5 \times 10^{-4}$) | 0.659 26 |
10 Q. Ma and H.-J. Werner, Wiley Interdiscip. Rev.: Comput. Mol. Sci. 8, e1371 (2018).
11 D. P. Tew, W. Klopper, C. Neiss, and C. Hättig, Phys. Chem. Chem. Phys. 9, 1921 (2007).
12 G. M. J. Barca and P.-F. Loos, J. Chem. Phys. 147, 024103 (2017).
13 E. Giner, B. Pradines, A. Ferté, R. Assaraf, A. Savin, and J. Toulouse, J. Chem. Phys. 149, 194301 (2018).
14 P.-F. Loos, B. Pradines, A. Scemama, E. Giner, and J. Toulouse, J. Chem. Theory Comput. 16, 1018 (2020).
15 P.-F. Loos, B. Pradines, A. Scemama, J. Toulouse, and E. Giner, J. Phys. Chem. Lett. 10, 2931 (2019).
16 E. Giner, A. Scemama, P.-F. Loos, and J. Toulouse, J. Chem. Phys. 152, 174104 (2020).
17 Y. Yao, E. Giner, J. Li, J. Toulouse, and C. J. Umrigar, J. Chem. Phys. 153, 124117 (2020).
18 E. Giner, A. Scemama, J. Toulouse, and P.-F. Loos, J. Chem. Phys. 151, 144118 (2019).
19 M. Levy, Proc. Natl. Acad. Sci. U. S. A. 76, 6062 (1979).
20 E. H. Lieb, Int. J. Quantum Chem. 24, 243 (1983).
21 A. Seidl, A. Görling, P. Vogl, J. A. Majewski, and M. Levy, Phys. Rev. B 53, 3764 (1996).
22 J. Toulouse, in Density Functional Theory, edited by E. Cances, G. Friesecke, and L. Lin (Springer, 2021); arXiv:2103.02645.
23 J. Toulouse, P. Gori-Giorgi, and A. Savin, Theor. Chem. Acc. 114, 305 (2005).
24 A. Ferté, E. Giner, and J. Toulouse, J. Chem. Phys. 150, 084103 (2019).
25 P. Perdew, K. Burke, and M. Ernzerhof, Phys. Rev. Lett. 77, 3865 (1996).
26 P. Gori-Giorgi and A. Savin, Phys. Rev. A 73, 032506 (2006).
27 B. Huron, J. P. Malrieu, and P. Rancurel, J. Chem. Phys. 58, 5745 (1973).
28 E. Giner, A. Scemama, and M. Caffarel, Can. J. Chem. 91, 879 (2013).
29 E. Giner, A. Scemama, and M. Caffarel, J. Chem. Phys. 142, 044115 (2015).
30 Y. Garniron, A. Scemama, P.-F. Loos, and M. Caffarel, J. Chem. Phys. 147, 034101 (2017).
31 T. H. Dunning, J. Chem. Phys. 90, 1007 (1989).
32 D. E. Woon and T. H. Dunning, J. Chem. Phys. 98, 1358 (1993).
33 D. E. Woon and T. H. Dunning, J. Chem. Phys. 103, 4572 (1995).
34 K. A. Peterson and T. H. Dunning, Jr., J. Chem. Phys. 117, 10548 (2002).
35 B. P. Pritchard, D. Altarawy, B. Didier, T. D. Gibson, and T. L. Windus, J. Chem. Inf. Model. 59, 4814 (2019).
36 Y. Garniron, T. Appelencourt, K. Gaspérich, A. Benali, A. Ferté, J. Paquier, B. Pradines, R. Assaraf, P. Reinhardt, J. Toulouse, P. Barbaresco, N. Renon, G. David, J.-P. Malrieu, M. Vérl, M. Caffarel, P.-F. Loos, E. Giner, and A. Scemama, J. Chem. Theory Comput. 15, 3591 (2019).
37 A. Halkier, W. Klopper, T. Helgaker, and P. Jørgensen, J. Chem. Phys. 111, 4424 (1999).
38 K. Aidas, C. Angeli, K. L. Bak, V. Bakken, R. Bast, L. Boman, O. Christiansen, R. Cimmaruoli, S. Coriani, P. Dahle, E. K. Dalskov, U. Ekström, T. Enveneldsen, J. J. Eriksen, P. Ettenhuber, B. Fernández, L. Ferrighi, H. Flegl, L. Frediani, K. Hald, A. Halkier, C. Hättig, H. Heiberg, T. Helgaker, A. C. Hennum, H. Hettema, E. Hjertenaes, S. Höst, I.-M. Höyvik, M. F. Iozzi, B. Jansik, H. J. Aa. Jensen, D. Jonsson, P. Jørgensen, J. Kauczor, S. Kirpekar, J. Toulouse, P. Barbaresco, N. Renon, G. David, J.-P. Malrieu, M. Vérl, M. Caffarel, P.-F. Loos, E. Giner, and A. Scemama, J. Chem. Theory Comput. 15, 3591 (2019).
39 A. Halkier, W. Klopper, T. Helgaker, and P. Jørgensen, J. Chem. Phys. 111, 4424 (1999).
40 Dalton, a molecular electronic structure program, release v2020.0, 2020 (see http://daltonprogram.org).
41 D. Hait and M. Head-Gordon, J. Chem. Theory Comput. 14, 1969 (2018).
42 I. Hornyák, L. Adamowicz, and S. Bubin, Phys. Rev. A 100, 032504 (2019).
43 P. L. Bak, J. Gauss, T. Helgaker, P. Jørgensen, and J. Olsen, Chem. Phys. Lett. 319, 563 (2000).
44 A. A. Holmes, C. J. Umrigar, and S. Sharma, J. Chem. Phys. 147, 164111 (2017).