Basis-set correction for coupled-cluster estimation of dipole moments

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ABSTRACT
The present work proposes an approach to obtain a basis-set correction based on density-functional theory (DFT) for the computation of molecular properties in wave-function theory (WFT). This approach allows one to accelerate the basis-set convergence of any energy derivative of a non-variational WFT method, generalizing previous works on the DFT-based basis-set correction where either only ground-state energies could be computed with non-variational wave functions [Loos et al., J. Phys. Chem. Lett. 10, 2931 (2019)] or properties could be computed as expectation values over variational wave functions [Giner et al., J. Chem. Phys. 155, 044109 (2021)]. This work focuses on the basis-set correction of dipole moments in coupled-cluster with single, double, and perturbative triple excitations [CCSD(T)], which is numerically tested on a set of 14 molecules with dipole moments covering two orders of magnitude. As the basis-set correction relies only on Hartree–Fock densities, its computational cost is marginal with respect to the one of the CCSD(T) calculations. Statistical analysis of the numerical results shows a clear improvement of the basis convergence of the dipole moment with respect to the usual CCSD(T) calculations.

I. INTRODUCTION
Quantum chemistry aims to provide theoretical methods to predict molecular properties starting from the many-body quantum mechanical problem. To solve this problem, a wide range of methods were developed in the last few decades mainly based on wave-function theory (WFT) and density-functional theory (DFT). The purpose of both approaches is to accurately treat correlation effects or, in other terms, the quantum effects that go beyond a mean-field description, such as Hartree–Fock (HF). In the context of WFT, there exists a wide range of methods of increasing computational cost—ranging from Møller–Plesset perturbation theory1 to coupled-cluster methods—, which, in principle, systematically converge to the full configuration interaction (FCI) limit, which is the exact solution within a given basis set. Nevertheless, the accuracy of the results of a WFT method—even at the FCI level—strongly depends on the quality of the basis set, mainly because of the slow convergence of the wave function near the electron–electron coalescence point.3,4 The combination of the slow basis-set convergence and the strong increase of the computational cost with both the size of the basis set and the number of electrons makes it very difficult to obtain well converged WFT calculations on large systems.

There are mainly two approaches to tackle the basis-set convergence problem of WFT: basis-set extrapolation techniques5,6 and explicitly correlated F12 methods.7–12 The basis-set extrapolation techniques rely on a known asymptotic behavior of the correlation energy with the size of the basis set but requires WFT calculations with basis sets of increasing sizes, which makes their application limited to small or medium system size. The F12 methods accelerate the basis-set convergence of the results, thanks to the inclusion of a correlation factor explicitly depending on electron–electron distances and restoring Kato’s electron–electron cusp condition.12 Although F12 methods improve indeed the results (typically, energy differences obtained with a F12 method using a triple-zeta basis set are as accurate as the ones obtained with the corresponding
uncorrected WFT method using a quintuple-zeta basis set\(^1\), the
F12 methods necessarily induce computational overheads due to the
large auxiliary basis sets required to resolve three- and four-electron
integrals.\(^{14}\)

An alternative path has been recently introduced by some of
the present authors in Ref. 15 where a rigorous framework was
proposed to correct for the basis-set incompleteness of WFT using
DFT. A central idea of this work is the fact that the Coulomb
electron–electron interaction projected in an incomplete basis set is
non-divergent and quite similar to the long-range interaction used
in range-separated DFT (RSDFT). A basis-set correction density
functional can then be built from RSDFT short-range correlation
functionals using a local range-separation parameter, which
automatically adapts to the basis set used. This results in a relatively
cheap way of correcting the basis-set incompleteness of WFT, which
has the desirable property of leading to an unaltered complete-basis-
set (CBS) limit. Two versions of this theory were proposed: (i) a
non-self-consistent version where the basis-set correction functional is
evaluated with any approximate accuracy of the FCI density and then
simply added to an approximation of the FCI energy in a given basis set;\(^{15,17}\) and (ii) a recently introduced self-consistent vari-
ant where the energy is minimized in the presence of the basis-set
correction functional and, therefore, allows for the wave function to be
changed by the DFT correction.\(^{15}\) The efficiency of the non-self-
consistent approach for computing the total energies and chemically
relevant energy differences of relatively large magnitudes (such as ionization potentials,\(^{18–21}\) atomization energies in Ref. 18.

In Sec. III, we provide computational details of our
strategy for the basis-set correction, which uses only densities at
its application to non-variational approaches, such as coupled-
cluster with singles, doubles, and perturbative triple excitations
\([\text{CCSD(T)}]\).

In the present work, we propose to overcome this limitation and
target the computation of first-order molecular properties as
energy derivatives of the non-self-consistent basis-set correction
approach. We apply this strategy to the computation of dipole
moments at the CCSD(T) level and propose a cheap computational
strategy for the basis-set correction, which uses only densities at the
HF level, similarly to what have been done in the context of
atomization energies in Ref. 18.

This paper is organized as follows: In Sec. II, we introduce the
theory of the basis-set correction extended to the computation of
dipole moments. In Sec. III, we provide computational details of our
study on a set of 14 molecules with dipole moments covering two
orders of magnitude. The numerical results are discussed in Sec. IV
and compared for some molecules with the fully self-consistent
formalism of Ref. 16. Detailed results, as well as the molecular
geometries used, are available in the supplementary material.

II. THEORY
A. Dipole moment from the self-consistent
basis-set correction
In this section, we generalize the framework of the basis-set cor-
rection to the presence of a static external electric field. Consider the
Hamiltonian of a \(N\)-electron system under an external electric field
\(e = cu\) of strength \(e\) along a direction \(u\),

\[ \hat{H}(\epsilon) = \hat{H}_0 - \epsilon \hat{d}, \] (1)

where \(\hat{H}_0\) is the Hamiltonian of the system without the electric field,

\[ \hat{H}_0 = \hat{T} + \hat{V}_{ne} + \hat{W}_{ee}, \] (2)

with \(\hat{T}\) being the kinetic-energy operator, \(\hat{V}_{ne}\) being the
electron–nuclei interaction operator, and \(\hat{W}_{ee}\) the electron–electron
interaction operator, and \(\hat{d} = \hat{d} \cdot u\), where \(\hat{d}\) is the total (electron + nuclear) dipole-moment operator,

\[ \hat{d} = - \sum_{i=1}^{N} r_i + \sum_{A=1}^{N_{nuclei}} Z_A R_A, \] (3)

where \(r_i\) are the electron coordinates and \(Z_A\) and \(R_A\) are the nuclear
charges and coordinates.

In the basis-set correction formalism, the ground-state energy \(E_0(\epsilon)\) of the Hamiltonian in Eq. (1) is approximated by

\[ E_0^B(\epsilon) = \min_{\Psi^B} \{ \langle \Psi^B | \hat{H}(\epsilon) | \Psi^B \rangle + E^B \{ n_{\Psi^B} \} \}, \] (4)

where the minimization is performed over the set of \(N\)-electron wave functions \(\Psi^B\) expanded on the \(N\)-electron Hilbert space
generated by the one-electron basis set \(\mathcal{B}\) and \(E^B \{ n_{\Psi^B} \}\) is the basis-set correction functional evaluated at the density \(n_{\Psi^B}\) of \(\Psi^B\). The energy functional \(E^B \{ n \}\) (introduced in Ref. 15) compensates for the restriction on the wave functions \(\Psi^B\) due to the incompleteness of the basis set \(\mathcal{B}\). The restriction coming from the basis set \(\mathcal{B}\) in Eq. (4) then applies only to densities \(n_{\Psi^B}\). Roughly speaking, since the density converges much faster than the wave function with respect to the basis set, \(E_0^B(\epsilon)\) is a much better approximation to the exact energy \(E_0(\epsilon)\) than the corresponding FCI ground-state energy \(E_{\text{FCI}}^\mathcal{B}(\epsilon)\) calculated with the same basis set \(\mathcal{B}\). Moreover, in the CBS limit, \(E^B \{ n \}\) vanishes and, thus, \(E_0^B(\epsilon)\) correctly converges to the exact energy \(E_0(\epsilon)\).

From the basis-set corrected energy \(E_0^B(\epsilon)\) in Eq. (4), one can
define the corresponding basis-set corrected dipole moment \(d^B\) as
the first-order derivative with respect to the electric field,
\[
d\beta = - \left. \frac{dE^\beta_0 (\epsilon)}{d\epsilon} \right|_{\epsilon=0} . \tag{5}
\]

It is important to stress here that \(d\beta\) is different from the FCI dipole moment \(d_{\text{FCI}}\) with the same basis set \(B\), as the former is taken as the derivative of \(E^\beta_0 (\epsilon)\), which contains the basis-set correction functional \(E^\beta [\Psi]\). Similar to the case of the energy, we expect \(d\beta\) to have a faster basis-set convergence than \(d_{\text{FCI}}\).

Since \(E^\beta_0 (\epsilon)\) is stationary with respect to \(\Psi^\beta\), the Hellmann–Feynman theorem applies and gives \(d\beta\) as a simple expectation value,

\[
d\beta = \langle \Psi^\beta_0 (\epsilon=0) | \hat{d} \Psi^\beta_0 (\epsilon=0) \rangle , \tag{6}
\]

where \(\Psi^\beta_0 (\epsilon=0)\) is the minimizing wave function of the self-consistent equation in Eq. (4) at \(\epsilon = 0\). This was the approach used in Ref. 16.

Note that the Hellmann–Feynman theorem applies because the basis set \(B\) is independent from the perturbing electric field. By contrast, if we wanted to calculate the gradient of the energy with respect to nuclear coordinates, we would have to take into account the dependence of the atomic basis set \(B\) on the nuclear coordinates. In that case, the Hellmann–Feynman theorem would not apply, and we would have to consider additional Pulay terms coming from the dependence of both the wave function \(\Psi^\beta_0\) and the basis-set correction energy \(E^\beta\) on the nuclear coordinates.

### B. Dipole moment from the non-self-consistent basis-set correction

As initially proposed in Ref. 15 for the case without the electric field, one can avoid the minimization in Eq. (4) and approximate the energy \(E^\beta_0 (\epsilon)\) by approximating the minimizing wave function \(\Psi^\beta_0 (\epsilon)\) in Eq. (4) by the FCI wave function \(\Psi^\beta_{\text{FCI}}(\epsilon)\) in a given basis set \(B\). This leads to the following estimation of the ground-state energy:

\[
E^\beta_0 (\epsilon) \approx E^\beta_{\text{FCI}} (\epsilon) + \bar{E}^\beta [n^\beta_{\text{FCI}} (\epsilon)] , \tag{7}
\]

where \(n^\beta_{\text{FCI}}(\epsilon)\) is the ground-state FCI density obtained in the presence of the electric field of strength \(\epsilon\). The corresponding non-self-consistent basis-set corrected dipole moment is thus

\[
d\beta = d_{\text{FCI}} + \bar{d}^\beta , \tag{8}
\]

where

\[
\bar{d}^\beta = - \left. \frac{dE^\beta [n^\beta_{\text{FCI}} (\epsilon)]}{d\epsilon} \right|_{\epsilon=0} . \tag{9}
\]

is the non-self-consistent basis-set correction to the dipole moment.

As obtaining both the dipole moment and the density at FCI level is often computationally prohibitive, we follow Ref. 18 and approximate the FCI energy by the CCSD(T) energy and the FCI density by the HF density (in the presence of the electric field),

\[
E^\beta_0 (\epsilon) \approx E^\beta_{\text{CCSD(T)}} (\epsilon) + \bar{E}^\beta [n^\beta_{\text{HF}} (\epsilon)] , \tag{10}
\]

Within these approximations, the basis-set corrected dipole moment in Eq. (8) becomes

\[
d^\beta = d_{\text{CCSD(T)}} + \bar{d}^\beta , \tag{11}
\]

where \(d_{\text{CCSD(T)}}\) is the dipole moment at the CCSD(T) level, and the basis-set correction \(\bar{d}^\beta\) is

\[
\bar{d}^\beta = - \left. \frac{d\bar{E}^\beta [n^\beta_{\text{HF}} (\epsilon)]}{d\epsilon} \right|_{\epsilon=0} . \tag{12}
\]

We approximate the basis-set correction functional \(\bar{E}^\beta [n]\) with the so-called (spin-dependent) PBEUEG energy functional introduced in Ref. 18 where the local range-separation parameter \(\mu^\beta (r)\) is obtained using the HF wave function in the basis set \(B\) as proposed in Refs. 15 and 18. The results obtained with Eq. (11) with the PBEUEG approximation for \(\bar{E}^\beta [n]\) evaluated at the HF density will be referred to as CCSD(T) + PBEUEG.

In practice, we calculate the CCSD(T) dipole moment and the basis-set correction to the dipole moment using a finite-difference approximation for the energy derivatives with respect to the electric field,

\[
d_{\text{CCSD(T)}}^\beta \approx - \frac{E^\beta_{\text{CCSD(T)}} (\epsilon) - E^\beta_{\text{CCSD(T)}} (-\epsilon)}{2\epsilon} \tag{13}
\]

and

\[
\bar{d}^\beta \approx - \frac{\bar{E}^\beta [n^\beta_{\text{HF}} (\epsilon)] - \bar{E}^\beta [n^\beta_{\text{HF}} (-\epsilon)]}{2\epsilon} , \tag{14}
\]

using a finite field strength of \(\epsilon = 10^{-4}\) a.u., as suggested in Ref. 23.

### III. COMPUTATIONAL DETAILS

The computation of the basis-set correction to the dipole moment \(d^\beta\) was performed using the Quantum Package program,26 and the CCSD(T) dipole moment was obtained with the Gaussian program.27 We used the augmented Dunning basis sets (Refs. 28 and 29) aug-cc-pVXZ (abbreviated as AVXZ in Figs. 1–3 and Tables I–III), where \(X\) is the cardinal number of the basis set \(X \in \{D, T, Q, S\}\). As no core-valence functions are used, the frozen-core approximation is used throughout this paper, where the 1s orbital is kept frozen for the elements from Li to F.

The tests are done on a set of \(n = 14\) molecules among which there are six open-shell molecules, for which we use restricted open-shell CCSD(T) [ROCCSD(T)] energies and restricted open-shell HF (ROHF) densities, and eight closed-shell molecules. Experimental geometries used for the computations are taken from Ref. 25 for the entire set, except in the case of BH and FH for which the geometries are taken from Ref. 23. We also report the results obtained in Ref. 16 for the BH, FH, CH\(_2\), and H\(_2\)O molecules using the self-consistent formalism [Eq. (6)] at the near-FCI level in order to compare with the present non-self-consistent formalism.
### TABLE I. HF, CCSD(T), and CCSD(T) + PBEUEG dipole moments in atomic units. For the open-shell systems, we use the spin-restricted open-shell (RO) version of these methods.

|     | AVDZ       | AVTZ       | AVQZ       | AV5Z       | CBS       |
|-----|------------|------------|------------|------------|-----------|
| CO  |            |            |            |            |           |
| HF  | 0.010199   | 0.010499   | 0.010433   | 0.010421   |           |
| CCSD(T) | 0.05550    | 0.05000    | 0.04600    | 0.04550    | 0.04485   |
| CCSD(T) + PBEUEG | 0.04398    | 0.04414    | 0.04273    | 0.04360    |           |
| BeH |            |            |            |            |           |
| ROHF | 0.111076   | 0.111076   | 0.11199    | 0.11218    |           |
| ROCCSD(T) | 0.09550    | 0.09100    | 0.09050    | 0.09050    | 0.09030   |
| ROCCSD(T) + PBEUEG | 0.08416    | 0.08746    | 0.08941    | 0.08980    |           |
| BF  |            |            |            |            |           |
| HF  | 0.34436    | 0.33390    | 0.33314    | 0.33328    |           |
| CCSD(T) | 0.34100    | 0.32700    | 0.32300    | 0.32200    | 0.32081   |
| CCSD(T) + PBEUEG | 0.33287    | 0.32351    | 0.32082    | 0.32068    |           |
| BH  |            |            |            |            |           |
| HF  | 0.68796    | 0.68649    | 0.68494    | 0.68496    |           |
| CCSD(T) | 0.52950    | 0.54500    | 0.54750    | 0.54850    | 0.54953   |
| CCSD(T) + PBEUEG | 0.54162    | 0.55002    | 0.54986    | 0.54980    |           |
| CH  |            |            |            |            |           |
| ROHF | 0.62348    | 0.62000    | 0.61871    | 0.61858    |           |
| ROCCSD(T) | 0.54150    | 0.54950    | 0.55150    | 0.55250    | 0.55368   |
| ROCCSD(T) + PBEUEG | 0.55427    | 0.55481    | 0.55405    | 0.55396    |           |
| NH  |            |            |            |            |           |
| ROHF | 0.63850    | 0.63505    | 0.63381    | 0.63384    |           |
| ROCCSD(T) | 0.59350    | 0.59950    | 0.60200    | 0.60350    | 0.60504   |
| ROCCSD(T) + PBEUEG | 0.60792    | 0.60519    | 0.60464    | 0.60506    |           |
| CH₂ (singlet) |          |            |            |            |           |
| HF  | 0.74877    | 0.74477    | 0.74355    | 0.74353    |           |
| CCSD(T) | 0.65600    | 0.66000    | 0.66200    | 0.66350    | 0.66510   |
| CCSD(T) + PBEUEG | 0.66666    | 0.66455    | 0.66420    | 0.66478    |           |
| FH  |            |            |            |            |           |
| HF  | 0.75976    | 0.75751    | 0.75634    | 0.75617    |           |
| CCSD(T) | 0.70350    | 0.70450    | 0.70700    | 0.70750    | 0.70820   |
| CCSD(T) + PBEUEG | 0.71371    | 0.70903    | 0.70946    | 0.70900    |           |
| H₂O |            |            |            |            |           |
| HF  | 0.78671    | 0.78039    | 0.77956    | 0.77956    |           |
| CCSD(T) | 0.72700    | 0.72400    | 0.72650    | 0.72800    | 0.72957   |
| CCSD(T) + PBEUEG | 0.73891    | 0.72930    | 0.72912    | 0.72920    |           |
| BN  |            |            |            |            |           |
| ROHF | 1.13451    | 1.13862    | 1.13831    | 1.13840    |           |
| ROCCSD(T) | 0.76250    | 0.77550    | 0.78400    | 0.78650    | 0.78902   |
| ROCCSD(T) + PBEUEG | 0.77517    | 0.78145    | 0.78756    | 0.78846    |           |
The accuracy of the dipole moments obtained with a given basis set and a given level of approximation is evaluated with respect to the CBS limit of the CCSD(T) dipole moments, \( d_{\text{CBS}}^{\text{CCSD(T)}} \), which are evaluated as in Ref. 23. In particular, the CBS results are computed as follows:

\[
\delta_{\text{CBS}}^{\text{CCSD(T)}} = \delta_{\text{CBS}}^{\text{HF}} + \delta_{\text{CBS}}^{\text{c}}, \tag{15}
\]

where \( \delta_{\text{CBS}}^{\text{c}} \) is the CBS limit of the correlation contribution to the CCSD(T) dipole moment that is computed using the following two-point \( X^{-3} \) extrapolation formula:

### TABLE II. Mean error (ME), mean absolute error (MAE), mean absolute relative error (MARE), maximal absolute error (MAX), and root-mean-square deviation (RMSD) (in atomic units) for the CCSD(T) and CCSD(T) + PBEUEG dipole moments of 14 molecules. See Fig. 3 for the corresponding plots of the normal distributions of errors.

|        | AVDZ | AVTZ | AVQZ | AVSZ | CBS |
|--------|------|------|------|------|-----|
| ME     |      |      |      |      |     |
| CCSD(T) | 0.01336 | 0.00579 | 0.00229 | 0.00122 |
| CCSD(T) + PBEUEG | 0.00319 | 0.00135 | 0.000004 | −0.00012 |
| MAE    |      |      |      |      |     |
| CCSD(T) | 0.01637 | 0.00579 | 0.00233 | 0.00125 |
| CCSD(T) + PBEUEG | 0.01080 | 0.00258 | 0.00086 | 0.00049 |
| MARE (in %) |      |      |      |      |     |
| CCSD(T) | 3.9 | 1.5 | 0.5 | 0.3 |
| CCSD(T) + PBEUEG | 1.6 | 0.6 | 0.4 | 0.3 |
| MAX    |      |      |      |      |     |
| CCSD(T) | 0.04518 (LiN) | 0.01418 (LiN) | 0.00502 (BN) | 0.00268 (LiN) |
| CCSD(T) + PBEUEG | 0.03504 (LiN) | 0.01004 (LiN) | 0.00254 (LiN) | 0.00136 (LiN) |
| RMSD   |      |      |      |      |     |
| CCSD(T) | 0.01484 | 0.00432 | 0.00163 | 0.00084 |
| CCSD(T) + PBEUEG | 0.01464 | 0.00376 | 0.00116 | 0.00063 |
TABLE III. Dipole moments obtained with near-FCI (CIPSI) and CCSD(T) calculations and with the self-consistent basis-set correction method (SC CIPSI + PBEUEG) of Ref. 16 and different non-self-consistent basis-set correction methods [CIPSI + PBEUEG@CIPSI, CCSD(T) + PBEUEG@CIPSI, and CCSD(T) + PBEUEG]. Estimated CBS values using Eq. (16) with $X = 5$ are reported when computations could be done with the aug-cc-pV5Z basis set.

|        | AVDZ    | AVTZ    | AVQZ    | AV5Z    | CBS    |
|--------|---------|---------|---------|---------|--------|
| **BH** |         |         |         |         |        |
| CIPSI  | 0.5278  | 0.5433  | 0.5456  | 0.5469  | 0.5482 |
| SC CIPSI + PBEUEG | 0.5379  | 0.5481  | 0.5479  | 0.5481  |        |
| CIPSI + PBEUEG@CIPSI | 0.5414  | 0.5485  | 0.5475  | 0.5482  |        |
| CCSD(T) | 0.5295  | 0.5450  | 0.5450  | 0.5485  | 0.5495 |
| CCSD(T) + PBEUEG@CIPSI | 0.5430  | 0.5502  | 0.5498  | 0.5496  |        |
| CCSD(T) + PBEUEG | 0.5416  | 0.5500  | 0.5498  | 0.5498  |        |
| **CH₂ (singlet)** |         |         |         |         |        |
| CIPSI  | 0.6512  | 0.6544  | 0.6564  | 0.6578  | 0.6592 |
| SC CIPSI + PBEUEG | 0.6624  | 0.6595  | 0.6589  |        |        |
| CIPSI + PBEUEG@CIPSI | 0.6638  | 0.6602  | 0.6595  |        |        |
| CCSD(T) | 0.6560  | 0.6600  | 0.6600  | 0.6635  | 0.6651 |
| CCSD(T) + PBEUEG@CIPSI | 0.6687  | 0.6656  | 0.6648  |        |        |
| CCSD(T) + PBEUEG | 0.6666  | 0.6645  | 0.6642  |        |        |
| **FH** |         |         |         |         |        |
| CIPSI  | 0.7024  | 0.7040  | 0.7066  |        |        |
| SC CIPSI + PBEUEG | 0.7132  | 0.7087  | 0.7112  |        |        |
| CIPSI + PBEUEG@CIPSI | 0.7139  | 0.7118  |        |        |        |
| CCSD(T) | 0.7035  | 0.7045  | 0.7045  | 0.7070  | 0.7082 |
| CCSD(T) + PBEUEG@CIPSI | 0.7142  | 0.7120  |        |        |        |
| CCSD(T) + PBEUEG | 0.7137  | 0.7093  | 0.7093  |        |        |
| **H₂O** |         |         |         |         |        |
| CIPSI  | 0.7261  | 0.7229  |        |        |        |
| SC CIPSI + PBEUEG | 0.7370  | 0.7281  |        |        |        |
| CIPSI + PBEUEG@CIPSI | 0.7365  | 0.7272  |        |        |        |
| CCSD(T) | 0.7270  | 0.7240  | 0.7265  | 0.7280  | 0.7295 |
| CCSD(T) + PBEUEG@CIPSI | 0.7373  | 0.7281  | 0.7287  |        |        |
| CCSD(T) + PBEUEG | 0.7389  | 0.7293  | 0.7291  |        |        |

$^a$From Ref. 16.

$^b$Results not available due to the computational requirement.

FIG. 1. (a) CCSD(T) and (b) CCSD(T) + PBEUEG errors on the dipole moments of 14 molecules compared to CCSD(T)/CBS reference values. The green area indicates an error of ±0.001 a.u.
FIG. 2. (a) CCSD(T) and (b) CCSD(T) + PBEUEG relative errors on the dipole moments of 14 molecules compared to CCSD(T)/CBS reference values. The green area indicates an error of ±1%.

FIG. 3. (a) CCSD(T) and (b) CCSD(T) + PBEUEG normal distribution of errors on the dipole moments of 14 molecules compared to CCSD(T)/CBS reference values. The green area indicates an error of ±0.001 a.u.

\[ \Delta_{b}^{\text{CBS}} = \frac{d_{c}^{X}X^{3} - d_{c}^{X-1}(X - 1)^{3}}{X^{3} - (X - 1)^{3}} \]  (16)

with

\[ d_{c}^{X} = d_{c}^{\text{CCSD(T)}} - d_{c}^{\text{HF}} \]  (17)

where \( d_{c}^{X} \) and \( d_{c}^{\text{HF}} \) are the correlation and HF contributions, respectively, to the CCSD(T) dipole moment using the aug-cc-pVXZ basis set. In the present calculations, we evaluate Eq. (16) at \( X = 5 \), and we estimate the CBS limit of HF dipole moment \( d_{c}^{\text{CBS}} \) in Eq. (15) simply by using the HF dipole moment in the aug-cc-pV5Z basis set. For all the systems studied here, the HF dipole moments are converged with an accuracy better than 0.001 a.u. (as measured by the difference between the aug-cc-pVQZ and aug-cc-pV5Z dipole moments).

At a given level of calculation in a basis set \( \mathcal{B} \), we report the error on the dipole moment with respect to the CBS reference \( \Delta^{\text{CBS}} = d^{\text{CBS}} - d^{\text{CCSD(T)}} \) and the relative error \( \Delta_{\text{rel}}^{\text{CBS}} = \Delta^{\text{CBS}} / d^{\text{CCSD(T)}} \). To statistically analyze the results, we also calculate the normal distribution function of the errors for a given basis set \( \mathcal{B} \),

\[ \rho(\Delta^{\text{CBS}}) = \frac{1}{\Delta_{\text{rel}}^{\text{CBS}} \sqrt{2\pi}} \exp \left[ -\frac{1}{2} \left( \frac{\Delta^{\text{CBS}} - \bar{\Delta}^{\text{CBS}}}{\Delta_{\text{rel}}^{\text{CBS}}} \right)^{2} \right] \]  (18)

where \( \bar{\Delta}^{\text{CBS}} = \left( \sum_{i=1}^{n} \Delta_{i}^{\text{CBS}} \right) / n \) is the mean error (ME) and \( \Delta_{\text{rel}}^{\text{CBS}} = \sqrt{\sum_{i=1}^{n} \left( \Delta_{i}^{\text{CBS}} - \bar{\Delta}^{\text{CBS}} \right)^{2} / (n-1) } \) is the root-mean-square deviation (RMSD).

IV. RESULTS AND DISCUSSION

In Table I, we report the dipole moments at various levels of approximations [HF, CCSD(T), and CCSD(T) + PBEUEG] with different basis sets, as well as the CCSD(T)/CBS reference values, for the set of 14 molecules. Note the wide range of magnitudes of the dipole moments (from 0.044 85 a.u. for CO to 2.787 18 a.u. for LiN). The mean error (ME), mean absolute error (MAE), mean absolute relative error (MARE), maximal absolute error (MAX), and root-mean-square deviation (RMSD) obtained with CCSD(T) and CCSD(T) + PBEUEG are reported in Table II. The graphical representations of these data are provided in Figs. 1 and 2 for the errors and relative errors and in Fig. 3 for the normal distributions of errors.
Analyzing first the results at the CCSD(T) level in Table II, one can note that, as expected, the ME and MAE systematically decrease with the size of the basis set. Moreover, as noted in previous studies, not only the average values of the errors but also the RMSD tends to decrease with the basis-set size. Nevertheless, the improvement of the results is rather slow as a MAE below 0.001 a.u. is not reached even with the aug-cc-pV5Z basis set, illustrating the slow convergence of properties with respect to the basis set at the CCSD(T) level. Regarding the relative errors in Fig. 2, not surprisingly, the largest errors with respect to the CBS reference come from the molecules with smallest dipole moments (i.e. CO and BeH). More quantitatively, an aug-cc-pVQZ basis set is needed to obtain a MARE smaller than 1%.

Going from CCSD(T) to CCSD(T) + PBEUEG, one observes a systematic decrease in the MAE, ME, MARE, and RMSD. Focusing on the MAE, an error below 0.001 a.u. is reached with the aug-cc-pVQZ basis set, whereas such an accuracy is not even reached at the CCSD(T) level with the aug-cc-pV5Z basis set. Qualitatively, for the aug-cc-pVTZ basis set and larger basis sets, the MAEs obtained with CCSD(T) + PBEUEG with a basis set of cardinal number $X$ are comparable to the MAEs obtained with CCSD(T) with a basis set of cardinal number $X + 1$. Regarding the MARE, an error below 1% is reached with CCSD(T) + PBEUEG already with the aug-cc-pVTZ basis set. One nevertheless observes that the effects of the basis-set correction on the RMSD are very weak. From the plots of Fig. 1, one notes that even if the basis-set correction systematically improves the results for the aug-cc-pVTZ basis set, its effect is less impressive when there is both a large error and a large dipole moment (i.e. for BN, BO, and LiN).

In order to further demonstrate the validity of the different approximations leading to the CCSD(T) + PBEUEG method, we conclude this study by a comparison with the self-consistent basis-set correction formalism of Ref. 16, as well as different flavors of non-self-consistent approximations. In Ref. 16, the self-consistent method referred to as SC CIPSI + PBEUEG was introduced, which can be considered as the nearly exact theory within our framework, thanks to the use of near-FCI (CIPSI; see Ref. 26 and references therein) wave functions. In all the calculations reported below, the absolute value of the second-order perturbative contribution to the energy in the CIPSI calculations is below $10^{-4}$ a.u., which implies that the CIPSI energy and density can indeed be considered as near-FCI quantities. We also consider two different levels of non-self-consistent approximations: (i) CIPSI + PBEUEG@CIPSI where the CIPSI energy is corrected with the PBEUEG functional evaluated at the CIPSI density, and (ii) CCSD(T) + PBEUEG@CIPSI where the CIPSI energy is approximated by the CCSD(T) energy but the PBEUEG functional is still evaluated at the CIPSI density. Therefore, we have a hierarchy of approximations for the basis-set correction method using the PBEUEG functional: SC CIPSI + PBEUEG as the exact self-consistent theory, then CIPSI + PBEUEG@CIPSI as the exact non-self-consistent theory, then CCSD(T) + PBEUEG@CIPSI where only the WFT energy part is approximated with respect to CIPSI + PBEUEG@CIPSI, and finally CCSD(T) + PBEUEG where both the WFT energy and the density are approximated with respect to CIPSI + PBEUEG@CIPSI.

We report in Table III the results obtained with these different levels of theory for the dipole moments of the BH, CH$_2$, FH, and H$_2$O molecules. The results obtained with the self-consistent method SC CIPSI + PBEUEG are in close agreement with that obtained with the different non-self-consistent approximations, the largest discrepancy being less than 0.006 a.u. for BH in the aug-cc-pVDZ basis set with CCSD(T) + PBEUEG@CIPSI. Comparing the two methods at the extremities of our hierarchy of approximations, one can note that the absolute deviation between CCSD(T) + PBEUEG and SC CIPSI + PBEUEG in a given basis set is never larger than 0.001 a.u. for FH and H$_2$O, and the discrepancy slightly increases up to 0.006 and 0.003 a.u. in the case of CH$_2$ and BH, respectively. Nevertheless, as originally reported in Ref. 16 and apparent from Table III, discrepancies of the same order of magnitude also appear between the uncorrected CIPSI and CCSD(T) results in the case of the CH$_2$ and BH molecules. This suggests that the main source of differences between the CCSD(T) + PBEUEG and SC CIPSI + PBEUEG methods actually comes from the parent WFT theory. Focusing now specifically on the effect of the density in the non-self-consistent basis-set correction, one can note that the use of either a HF or CIPSI density does not significantly change the results, as the largest deviation between CCSD(T) + PBEUEG@CIPSI and CCSD(T) + PBEUEG are about 0.002 a.u. in the case of CH$_2$ in the aug-cc-pVDZ basis set. These results illustrate the validity of the different approximations leading to the CCSD(T) + PBEUEG approach and are encouraging, considering that the latter has a much lower computational cost with respect to the self-consistent basis-set formalism. Indeed, CCSD(T) + PBEUEG relies only on a standard CCSD(T) calculation and HF calculations for the basis-set correction, which is of negligible computational cost with respect to CCSD(T).

V. CONCLUSION

In the present study, we have proposed an extension of the recently introduced non-self-consistent basis-set correction of CCSD(T) ground-state energies to the computation of properties as energy derivatives, focusing here on the dipole moment. The theory relies on the originally proposed DFT-based basis-set correction approach, which accelerates the basis-set convergence to the unaltered CBS limit. Numerical tests on a set of 14 molecules (including both closed and open-shell) with dipole moments spanning two orders of magnitude have been carried in order to obtain a representative study of the performance of the present approach.

Although this study aims at correcting the basis-set convergence of the CCSD(T) dipole moments, it can be formally generalized to any wave-function method and any energy derivative with respect to a static perturbation. In its present form, the basis-set correction relies only on HF calculations, which makes the basis-set correction essentially computationally free compared to the correlated wave-function calculation. This approach is an alternative to the recently proposed self-consistent basis-set correction, which allows for the computation of first-order properties through expectation values over an energy-minimized wave function. In contrast with the self-consistent formalism, the present approach does not require a variational wave function, which considerably extends the domain of application of the basis-set correction.

Regarding now the numerical results, we have shown that the present approach significantly accelerates the basis-set convergence...
of CCSD(T) dipole moments. Typically, the error obtained in a basis set of cardinal $X$ with the basis-set correction is comparable to the error of the uncorrected CCSD(T) calculation with cardinal number $X + 1$. We also compared the present non-self-consistent basis-set correction with the self-consistent formalism of Ref. 16 and showed that the two theories agree within a few milli-atomic units, illustrating the soundness of the approximations leading to the non-self-consistent approach.

Considering the generality, the global performance, and the small computational cost of the present approach, it could be an alternative to explicitly correlated approaches for calculation of molecular properties. In the near future, we will extend the method to higher-order static properties, such as static polarizabilities, and also to more general dynamic properties, leading, in particular, to the possibility of accelerating the basis-set convergence of excitation energies.

SUPPLEMENTARY MATERIAL

The supplementary material contains (i) all the geometries of the molecules studied here, (ii) a graphical representation of the convergence of the dipole moment at CCSD(T) and CCSD(T) + PBEUEG levels for each system studied, and (iii) the absolute and relative errors with respect to the estimated CBS and the CCSD(T) and CCSD(T) + PBEUEG levels.

ACKNOWLEDGMENTS

This project has received funding from the European Research Council (ERC) under the European Union’s Horizon 2020 Research and Innovation Program (Grant Agreement No. 810367) (EMC2).

AUTHOR DECLARATIONS

Conflict of Interest

The authors have no conflicts to disclose.

DATA AVAILABILITY

The data that support the findings of this study are available within the article and its supplementary material.

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