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# PAPER



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# Accelerated basis-set convergence of coupled-cluster excitation energies using the density-based basis-set correction method

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We present the first application to real molecular systems of the recently proposed linearresponse theory for the density-based basis-set correction method [J. Chem. Phys., 158, 234107 (2023)]. We apply this approach to accelerate the basis-set convergence of excitation energies in the equation-of-motion coupled-cluster singles and doubles (EOM-CCSD) method. We use an approximate linear-response framework that neglects the second-order derivative of the basis-set correction density functional and consists in simply adding to the usual Hamiltonian the one-electron potential generated by the first-order derivative of the functional. This additional basis-set correction potential is evaluated at the Hartree–Fock density, leading to a very computationally cheap basisset correction. We tested this approach over a set of about 30 excitation energies computed for five small molecular systems and found that the excitation energies from the ground state to Rydberg states are the main source of basis-set error. These excitation energies systematically increase when the size of the basis set is increased, suggesting a biased description in favour of the excited state. Despite the simplicity of the present approach, the results obtained with the basis-set-corrected EOM-CCSD method are encouraging as they yield a mean absolute deviation of 0.02 eV for the aug-cc-pVTZ basis set, while it is 0.04 eV using the standard EOM-CCSD method. This might open the path to an alternative to explicitly correlated approaches to accelerate the basis-set convergence of excitation energies.

# 1. Introduction

One of the main bottlenecks of computational electronic-structure wave-function methods is the slow convergence of the energy and properties with the size of the one-electron basis set used to expand the wave function. At the origin of this limitation lies the divergence of the Coulomb electron–electron interaction as the

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interelectronic distance goes to zero, creating derivative discontinuities in the wave function<sup>1</sup> that cannot be represented with a finite one-electron basis set.<sup>2</sup> One can drastically improve the basis-set convergence by using geminals explicitly depending on the interelectronic distance.<sup>3,4</sup> Among the various types of such explicitly correlated approaches, the so-called F12 method<sup>5-9</sup> has proven to be a remarkably efficient tool to describe ground-state properties. Application of the F12 or R12 methodology to excited states<sup>10-18</sup> is nevertheless not straightforward and the initial attempt led to relatively disappointing results.<sup>10</sup> The main issue came from a strong bias toward the ground state: in the usual formulation the geminals are applied only on the Hartree-Fock (HF) Slater determinant, which dominates the ground-state wave function but has only a small contribution to the excited states. Further developments, in which the geminals were also applied on singly excited configurations, led to a strong reduction of this bias.<sup>11</sup> While the latter developments used the so-called linear R12 geminals,<sup>10,11</sup> the modern types of explicitly correlated methods use Slater-type geminals (F12), which, unlike the R12 geminals, decay at large interelectronic distances, and the so-called SP ansatz introduced by Ten-No,19 which avoids the optimization of the geminal amplitudes by using first- and second-order cusp conditions. In this spirit, Köhn proposed the so-called XSP ansatz,13 which is an extended version of the SP ansatz adapted to response theory by adding the single hole-particle excitation channels in the SP geminals. The initial application of the XSP ansatz to the equation-of-motion coupled-cluster singles and doubles (EOM-CCSD) method (see, e.g., ref. 20), in addition to the use of the complementary auxiliary basis-set one-electron correction, not only suppressed the ground-state bias of the standard SP approach but also improved the basis-set convergence of the excitation energies. Further extensions of the XSP ansatz to the second-order coupled-cluster (CC2) method were reported with applications to organic molecules.18 The XSP ansatz can therefore be considered as the state-of-the-art method for treating excited states within the F12 framework.

Despite the undeniable successes of F12 theories, we might point out some limitations. First, they are usually formulated for single-reference methods and are not often generally available for multi-reference methods. Second, they rely on a relatively involved formulation, which makes their incorporation in computer software a relatively complex task. Third, the quality of the results for excitation energies particularly depends on the Slater geminal parameter  $\gamma$ . Last but not least, F12 theories need three- and four-electron integrals, which have to be approximated through resolution-of-identity techniques.

An alternative to F12 approaches was recently proposed by some of the present authors,<sup>21</sup> with the so-called density-based basis-set correction (DBBSC) method. This method uses range-separated density-functional theory (RSDFT) (see ref. 22 and references therein) in order to capture the short-range correlation energy missing from the description of wave-function approaches with an incomplete one-electron basis set  $\mathcal{B}$ . The DBBSC method relies on the determination of an effective local range-separation parameter  $\mu^{\mathcal{B}}(\mathbf{r})$ , depending on the spatial position  $\mathbf{r}$ , which provides a local measure of the incompleteness of a given basis set  $\mathcal{B}$ . One can then simply use a short-range correlation density functional designed for multideterminant RSDFT<sup>23,24</sup> with this local range–separation parameter  $\mu^{\mathcal{B}}(\mathbf{r})$  to obtain an estimation of the correlation energy missing in the basis set  $\mathcal{B}$ . The DBBSC method has been tested and validated for the calculation of ground-state atomization

energies,<sup>25-28</sup> including for light and transition-metal elements and strongly correlated systems, ionization potentials,<sup>21,29</sup> and dipole moments.<sup>30,31</sup> Efficient implementations using density-fitting technologies were recently reported.<sup>32,33</sup>

Being based on the Levy-Lieb formulation of density-functional theory (DFT), the DBBSC method is a ground-state theory and an attempt to apply it to excited states was proposed<sup>34</sup> in a state-specific way by simply evaluating the basis-set correction energy functional at the density (and, in some cases, the pair density) of each excited state. While the results obtained were encouraging,34 this approach is nevertheless not rigorous, as it consists in the unjustified application of a ground-state theory to excited states. Recently, the present authors derived the general equations of linear-response theory for the DBBSC method<sup>35</sup> and applied them to the calculation of excitation energies in a one-dimensional model system. This linear-response formalism relies on a variational self-consistent version of the DBBSC method that was implemented and tested in ref. 30. In the latter framework, the wave function is changed through the addition of a basis-set correction potential that is self-consistently determined, and numerical tests showed that, while self-consistency makes very little changes to the energy, the modified wave function leads to a significant change in the density that greatly accelerates the basis-set convergence of dipole moments. Nevertheless, the self-consistent approach is difficult to realise with a non-variational wave-function ansatz, which therefore restricts its domain of applicability. Recently, the present authors<sup>31</sup> proposed calculating the basis-set correction to coupled-cluster dipole moments through a numerical energy derivative of the non-self-consistent basis-set correction functional evaluated at the HF density. The results showed that the accuracy is similar to that obtained with the selfconsistent approach. Such a finding supports the idea that using the selfconsistently optimised density in the presence of the basis-set correction is often not mandatory and that one can simply use the HF density in the basis-set correction functional in many cases.

In the present work, for calculating excitation energies, we propose using a simple linear-response variant of the DBBSC method, which consists in several approximations: (i) neglecting the second-order derivative of the basis-set correction functional, (ii) approximating the density at the HF level, (iii) using the simplest (semi)local basis-set correction functionals, (iv) approximating fullconfiguration-interaction (FCI) excitation energies at the EOM-CCSD level. The paper is organised as follows. In Section 2 we present the theory: a summary of the main equations of the ground-state DBBSC method is provided in Section 2.1, then the approximate linear-response theory is presented in Section 2.2, and eventually its application to the EOM-CCSD method is sketched out in Section 2.3. Numerical results are presented and discussed in Section 3 for a set of 30 excitation energies in  $NH_3$ ,  $H_2O$ , CO,  $N_2$  and  $N_2CH_2$  molecules comprising Rydberg and valence excited states with singlet and triplet symmetry. Finally, Section 4 contains our conclusions.

# 2. Theory

## 2.1. Ground-state DBBSC method

As introduced in ref. 21, in the DBBSC method, one defines an approximation to the ground-state energy  $E_0^{\mathcal{B}}$  for a given basis set  $\mathcal{B}$  by restricting the DFT ground-

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state energy minimization to  $\mathcal{B}$ -representable densities  $n^{\mathcal{B}}$  (*i.e.*, densities that can be obtained from a wave function  $\Psi^{\mathcal{B}}$  in the many-electron Hilbert space  $\mathcal{H}^{\mathcal{B}}$  generated by the basis set  $\mathcal{B}$ ):

$$E_0^{\mathcal{B}} = \min_{n^{\mathcal{B}}} \left( F[n^{\mathcal{B}}] + \int d\mathbf{r} \ v_{\rm ne}(\mathbf{r}) n(\mathbf{r}) \right), \tag{1}$$

where  $F[n] = \min_{\Psi \to n} \langle \Psi | \hat{T} + \hat{W}_{ee} | \Psi \rangle$  is the usual Levy-Lieb universal density functional. Here,  $\hat{T}$  and  $\hat{W}_{ee}$  are the kinetic-energy and Coulomb electron–electron operators, and  $\nu_{ne}$  is the nuclei–electron potential. As the basis-set restriction in eqn (1) is only on the density and not on the wave function, the energy  $E_0^{\mathcal{B}}$  is a much better approximation to the exact ground-state energy  $E_0$  than the groundstate FCI energy  $E_{FCI}^{\mathcal{B}}$  in the same basis set  $\mathcal{B}$ . One can in fact rewrite eqn (1) in terms of a minimization over wave functions  $\Psi^{\mathcal{B}}$  restricted to the Hilbert space  $\mathcal{H}^{\mathcal{B}}$ :

$$E_0^{\mathcal{B}} = \min_{\Psi^{\mathcal{B}}} \left( \langle \Psi^{\mathcal{B}} | \hat{H} | \Psi^{\mathcal{B}} \rangle + \overline{E}^{\mathcal{B}} [n_{\Psi^{\mathcal{B}}}] \right), \tag{2}$$

where  $\hat{H} = \hat{T} + \hat{W}_{ee} + \hat{V}_{ne}$  is the total Hamiltonian, including the nuclei-electron operator  $\hat{V}_{ne} = \int d\mathbf{r} v_{ne}(\mathbf{r}) \hat{n}(\mathbf{r})$  expressed with the density operator  $\hat{n}(\mathbf{r})$ , and  $\overline{E}^{\mathcal{B}}[n_{\Psi^{\mathcal{B}}}]$ is the basis-set correction density functional evaluated at the density of  $\Psi^{\mathcal{B}}$ . For a  $\mathcal{B}$ -representable density  $n^{\mathcal{B}}$ , this functional is defined as

$$\overline{E}^{\mathcal{B}}[n^{\mathcal{B}}] = \min_{\Psi \to n^{\mathcal{B}}} \langle \Psi | \hat{T} + \hat{W}_{ee} | \Psi \rangle - \min_{\Psi^{\mathcal{B}} \to n^{\mathcal{B}}} \langle \Psi^{\mathcal{B}} | \hat{T} + \hat{W}_{ee} | \Psi^{\mathcal{B}} \rangle,$$
(3)

and corrects for the error due to the basis-set restriction on the wave functions  $\Psi^{\mathcal{B}}$ . A minimizing wave function  $\Psi^{\mathcal{B}}_0$  in eqn (2) satisfies the following self-consistent Schrödinger-like equation:

$$\hat{P}^{\mathcal{B}}\widehat{\overline{H}}^{\mathcal{B}}\left[n_{\Psi_{0}^{\mathcal{B}}}\right]\left|\Psi_{0}^{\mathcal{B}}\right\rangle = \mathcal{E}_{0}^{\mathcal{B}}\left|\Psi_{0}^{\mathcal{B}}\right\rangle,\tag{4}$$

where  $\mathcal{E}_0^{\mathcal{B}}$  is the Lagrange multiplier imposing the normalization condition of the wave function,  $\hat{P}^{\mathcal{B}}$  is the projector on the Hilbert space  $\mathcal{H}^{\mathcal{B}}$ , and  $\hat{\overline{H}}^{\mathcal{B}}[n]$  is an effective Hamiltonian,

$$\widehat{\overline{H}}^{\mathcal{B}}[n] = \widehat{H} + \widehat{\overline{V}}^{\mathcal{B}}[n],$$
(5)

where  $\widehat{\overline{V}}^{\mathcal{B}}[n]$  is the basis-set correction potential operator generated by the derivative of the basis-set correction functional:

$$\widehat{\overline{V}}^{\mathcal{B}}[n] = \int \mathrm{d}\mathbf{r} \frac{\delta \overline{E}^{\mathcal{B}}[n]}{\delta n(\mathbf{r})} \hat{n}(\mathbf{r}), \tag{6}$$

which corresponds to a one-electron potential.

Instead of performing the minimization in eqn (2), one can use a non-selfconsistent approximation:<sup>21,25</sup>

$$E_0^{\mathcal{B}} \approx E_X^{\mathcal{B}} + \overline{E}^{\mathcal{B}} [n_Y^{\mathcal{B}}], \tag{7}$$

where  $E_X^B$  is an approximation of the FCI energy in the basis set  $\mathcal{B}$  calculated with a method X and  $n_Y^B$  is the density calculated with a method Y and the same basis set  $\mathcal{B}$ . This non-self-consistent variant was successfully applied in ref. 25 and 31 using coupled-cluster singles, doubles and perturbative triples [CCSD(T)] for

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method X and HF for method Y to compute ground-state atomization energies and dipole moments. A detailed study was carried out in ref. 30, where it was shown that the non-self-consistent approximation was good enough to calculate the energy.

Unlike the Levy–Lieb density functional, the basis-set correction functional  $\overline{E}^{\mathcal{B}}[n]$  is no longer universal as it depends explicitly on the basis set  $\mathcal{B}$ , which is in practice system-dependent. Nevertheless, as most of the basis-set incompleteness consists in missing correlation effects occurring at short interelectronic distances (*i.e.*, in the vicinity of the universal electron–electron cusp), one can expect to find generic approximations for  $\overline{E}^{\mathcal{B}}[n]$ . As originally proposed in ref. 21, the basis-set correction functional  $\overline{E}^{\mathcal{B}}[n]$  can be approximated by the short-range (sr) multi-determinant (md) correlation functional from RSDFT,<sup>23,24</sup> evaluated with a basis-set dependent and local range-separation parameter  $\mu^{\mathcal{B}}(\mathbf{r})$ . A semilocal version of it that is appropriate for weakly correlated systems is:<sup>25</sup>

$$\overline{E}^{\mathcal{B}}[n] \approx \int \mathrm{d}\mathbf{r} \ e_{\mathrm{c,md}}^{\mathrm{sr}}(n(\mathbf{r}), \nabla n(\mathbf{r}), \mu^{\mathcal{B}}(\mathbf{r})), \tag{8}$$

where  $\nabla n$  is the density gradient and  $e_{c,md}^{sr}(n, \nabla n, \mu)$  is the following correlation energy density:

$$e_{\rm c,md}^{\rm sr}(n,\nabla n,\mu) = \frac{e_{\rm c}(n,\nabla n)}{1+\beta(n,\nabla n)\mu^3},\tag{9}$$

with

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$$\beta(n, \nabla n) = \frac{3}{2\sqrt{\pi}(1-\sqrt{2})} \frac{e_{\rm c}(n, \nabla n)}{n_2^{\rm UEG}(n)},\tag{10}$$

where  $e_c(n, \nabla n)$  can be any approximate Kohn–Sham (semi)local correlation energy density, and  $n_2^{\text{UEG}}(n) = n^2 g_0(n)$  is the on-top pair density of the uniform electron gas (UEG),<sup>36</sup> written in terms of the UEG on-top pair-distribution function  $g_0(n)$  as parametrised in eqn (46) of ref. 37. The function  $e_{c,md}^{\text{sr}}(n, \nabla n, \mu)$  is designed such that it interpolates between the exact large- $\mu$  behavior<sup>22,37,38</sup> in  $1/\mu^3$  and the Kohn–Sham correlation energy density  $e_c(n, \nabla n)$  at  $\mu = 0$ . In the present work, we use two different approximations for the correlation functional  $e_c(n, \nabla n)$ : the local-density approximation (LDA)<sup>39</sup> and the Perdew–Burke–Ernzerhof (PBE) approximation.<sup>40</sup>

Although in its most general form  $\mu^{\mathcal{B}}(\mathbf{r})$  depends on a correlated wave function  $\Psi_{\text{loc}}^{\mathcal{B}}$  used to localise the Coulomb two-electron interaction projected in the basis set,<sup>21</sup> it was shown in a series of studies<sup>21,25,29-33</sup> that simply using the HF wave function (*i.e.*,  $\Psi_{\text{loc}}^{\mathcal{B}} = \Phi_{\text{HF}}^{\mathcal{B}}$ ) is enough to obtain reliable results for weakly correlated systems. Also, as in most cases wave-function calculations are performed with the frozen-core (FC) approximation, a corresponding FC version for the calculation of  $\mu^{\mathcal{B}}(\mathbf{r})$  was introduced in ref. 25. The use of the FC version with the HF wave function leads to the following expression for  $\mu^{\mathcal{B}}(\mathbf{r})$ :

$$\mu^{\mathcal{B}}(\mathbf{r}) = \frac{\sqrt{\pi}}{2} \frac{f_{\rm HF}^{\mathcal{B}}(\mathbf{r})}{n_{2,\rm HF}^{\mathcal{B}}(\mathbf{r})},\tag{11}$$

with the function

$$f_{\rm HF}^{\mathcal{B}}(\mathbf{r}) = 2 \sum_{p,q\in \, \text{all}^{i,j\in \, \text{act}}} \sum_{\phi_p(\mathbf{r})\phi_q(\mathbf{r})\phi_i(\mathbf{r})\phi_j(\mathbf{r}) V_{pq}^{ij},\tag{12}$$

and the HF on-top pair density

$$n_{2,\mathrm{HF}}^{\mathcal{B}}(\mathbf{r}) = 2 \sum_{i,j\in\,\mathrm{act}} \left(\phi_i(\mathbf{r})\phi_j(\mathbf{r})\right)^2,\tag{13}$$

where  $\{\phi_p\}$  are the (real-valued) spatial HF orbitals,  $V_{pq}^{ij} = \langle pq | ij \rangle$  are the usual twoelectron Coulomb integrals, and p, q run over all (occupied + virtual) HF spatial orbitals in the basis set  $\mathcal{B}$  and i, j run over only the active (*i.e.*, non-core occupied) spatial HF orbitals. Correspondingly, with the FC approximation, the basis-set correction functional is evaluated at the active HF density (removing the contribution from the core orbitals):

$$n_{\rm HF}^{\mathcal{B}}(\mathbf{r}) = 2 \sum_{i \in \rm act} \phi_i(\mathbf{r})^2.$$
(14)

As shown in ref. 21, this local range-separation parameter  $\mu^{\mathcal{B}}(\mathbf{r})$  automatically adapts to the basis set and tends to infinity in the complete-basis-set (CBS) limit. This makes the basis-set correction functional  $\overline{E}^{\mathcal{B}}[n]$  in eqn (8) correctly vanish in the CBS limit.

### 2.2. Approximate linear-response DBBSC method

The extension of the DBBSC method to linear-response theory was recently proposed by the present authors in ref. 35 when using a FCI wave function.

In this case, the (normalised) ground-state wave function  $\Psi_0^{\mathcal{B}}$  satisfying eqn (4) is expanded in terms of *N* orthonormal Slater determinants  $\{\Phi_i\}_{1 \le i \le N}$  spanning the Hilbert space  $\mathcal{H}^{\mathcal{B}}$ :

$$\left|\Psi_{0}^{\mathcal{B}}
ight
angle =\sum_{i=1}^{N}c_{0,i}|\Phi_{i}
angle,$$
 (15)

and the (real-valued) coefficients  $\{c_{0,i}\}$  satisfy the stationary equation, for  $1 \le i \le N$ ,

$$\left\langle \overline{\Psi}_{i} \middle| \widehat{H}^{\mathcal{B}} \Big[ n_{\Psi_{0}^{\mathcal{B}}} \Big] \middle| \Psi_{0}^{\mathcal{B}} \right\rangle = 0, \tag{16}$$

where  $\bar{\psi}_i$  are the wave-function derivatives

$$\left|\overline{\Psi}_{i}\right\rangle = \left|\Phi_{i}\right\rangle - c_{0,i}\left|\Psi_{0}^{\mathcal{B}}\right\rangle,\tag{17}$$

which are orthogonal to  $\Psi_0^{\mathcal{B}}$ , *i.e.*,  $\langle \overline{\Psi}_i | \Psi_0^{\mathcal{B}} \rangle = 0$ .

In the linear-response equations of ref. 35, if we neglect the kernel contribution coming from the second-order derivative of the basis-set correction functional  $\overline{E}^{\mathcal{B}}[n]$ , we obtain the following approximate linear-response equations:

$$\overline{\mathbf{A}} \ \mathbf{X}_n = \omega_n^{\mathcal{B}} \ \overline{\mathbf{S}} \ \mathbf{X}_n, \tag{18}$$

where  $\mathbf{X}_n$  are eigenvectors,  $\omega_n^{\mathcal{B}}$  are the eigenvalues corresponding to excitation energies ( $1 \le n \le N - 1$ ), and the matrix elements of  $\mathbf{\bar{A}}$  and  $\mathbf{\bar{S}}$  are, for  $2 \le i, j \le N$ ,

$$\overline{A}_{ij} = \left\langle \overline{\Psi}_i \middle| \widehat{\overline{H}}^{\mathcal{B}} \Big[ n_{\Psi_0^{\mathcal{B}}} \Big] - \mathcal{E}_0^{\mathcal{B}} \middle| \overline{\Psi}_j \right\rangle, \tag{19}$$

and

$$\bar{S}_{ij} = \langle \bar{\psi}_i | \bar{\psi}_j \rangle = \delta_{ij} - c_{0,i} c_{0,j}.$$
(20)

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To avoid the parameter redundancy due to the normalization constraint on the wave function, the first wave-function derivative  $|\overline{\Psi}_1\rangle = |\Phi_1\rangle - c_{0,1}|\Psi_0^B\rangle$ , involving the HF Slater determinant  $\Phi_1 \equiv \Phi_{\text{HF}}^B$ , has been dropped in the linear-response equations. Thus, only N - 1 equations remain. These approximate linear-response equations are in fact completely equivalent to the FCI equations for the effective Hamiltonian  $\widehat{H}^B[n_{\Psi_0^B}]$ . To see this, we begin by rewriting eqn (18) as

$$\overline{\mathbf{H}}\mathbf{X}_n = \mathcal{E}_n^{\mathcal{B}} \overline{\mathbf{S}} \mathbf{X}_n, \tag{21}$$

where  $\overline{H}_{ij} = \langle \overline{\Psi}_i | \widehat{\overline{H}}^{\mathcal{B}}[n_{\Psi_0^{\mathcal{B}}}] | \overline{\Psi}_j \rangle$  and  $\mathcal{E}_n^{\mathcal{B}} = \mathcal{E}_0^{\mathcal{B}} + \omega_n^{\mathcal{B}}$  are the excited-state total energies. Clearly, eqn (21) is the eigenvalue equation for the Hamiltonian  $\widehat{\overline{H}}^{\mathcal{B}}[n_{\Psi_0^{\mathcal{B}}}]$  in the non-orthogonal basis  $\{\overline{\psi}_i\}_{2 \leq i \leq N}$ . We then add the ground-state FCI wave function  $\Psi_0^{\mathcal{B}}$  to this basis, which, using the stationary equation of eqn (16), leads to the following  $N \times N$  eigenvalue equation, for  $0 \leq n \leq N - 1$ :

$$\begin{pmatrix} \mathcal{E}_{0}^{\mathcal{B}} & 0\\ 0 & \overline{\mathbf{H}} \end{pmatrix} \begin{pmatrix} \delta_{0n} \\ \mathbf{X}_{n} \end{pmatrix} = \mathcal{E}_{n}^{\mathcal{B}} \begin{pmatrix} 1 & 0\\ 0 & \overline{\mathbf{S}} \end{pmatrix} \begin{pmatrix} \delta_{0n} \\ \mathbf{X}_{n} \end{pmatrix},$$
(22)

and the FCI ground state is recovered for n = 0 with  $\mathbf{X}_0 = \mathbf{0}$ . Eqn (22) is thus the eigenvalue equation for the Hamiltonian  $\widehat{\overline{H}}^{\mathcal{B}}[n_{\Psi_0^{\mathcal{B}}}]$  in the non-orthogonal basis  $\{\Psi_0^{\mathcal{B}}\} \cup \{\overline{\Psi}_i\}_{2 \le i \le N}$ . If we rewrite this eigenvalue equation in the orthonormal basis of the *N* Slater determinants  $\{\Phi_i\}_{1 \le i \le N}$ , we recover a standard FCI eigenvalue equation for the effective Hamiltonian  $\widehat{\overline{H}}^{\mathcal{B}}[n_{\Psi_0^{\mathcal{B}}}]$ :

$$\mathbf{H}\mathbf{c}_n = \mathcal{E}_n^{\mathcal{B}}\mathbf{c}_n,\tag{23}$$

where  $H_{ij} = \langle \Phi_i | \widehat{\overline{H}}^B[n_{\Psi_0^B}] | \Phi_j \rangle$  and  $\mathbf{c}_n$  are the eigenvectors. Equivalently, this eigenvalue equation can be written as

$$\hat{P}^{\mathcal{B}}\widehat{\overline{H}}^{\mathcal{B}}\left[n_{\Psi_{0}^{\mathcal{B}}}\right]\left|\Psi_{n}^{\mathcal{B}}\right\rangle = \mathcal{E}_{n}^{\mathcal{B}}\left|\Psi_{n}^{\mathcal{B}}\right\rangle,\tag{24}$$

and the excited-state wave functions are

$$\left|\Psi_{n}^{\mathcal{B}}\right\rangle = \sum_{i=1}^{N} c_{n,i} |\Phi_{i}\rangle.$$
(25)

Therefore, in the linear-response DBBSC method, when neglecting the kernel coming from the second-order derivative of the basis-set correction functional, the excitation energies  $\omega_n^{\mathcal{B}}$  can be obtained directly by solving the FCI eigenvalue equation with the effective Hamiltonian  $\widehat{H}^{\mathcal{B}}[n_{\Psi_0^{\mathcal{B}}}] = \hat{H} + \widehat{\nabla}^{\mathcal{B}}[n_{\Psi_0^{\mathcal{B}}}]$  containing the basis-set correction potential operator  $\widehat{\nabla}^{\mathcal{B}}[n_{\Psi_0^{\mathcal{B}}}]$ .

## 2.3. Application to the EOM-CCSD method

Although eqn (24) can *a priori* be solved using any wave-function method targeting excited states, the fact that the basis-set correction potential must be evaluated at the density of the ground-state wave function  $\Psi_0^B$  is not convenient. Indeed, it requires performing a self-consistent ground-state calculation for obtaining  $\Psi_0^{\mathcal{B}}$ . Nevertheless, as shown in previous works,<sup>21,25,29,31</sup> good results can be obtained when the density  $n_{\Psi_0^{\mathcal{B}}}$  is approximated by the HF density  $n_{\text{HF}}^{\mathcal{B}}$ . Therefore, we use here the following approximation:

$$\widehat{\overline{H}}^{\mathcal{B}}\left[n_{\Psi_{0}^{\mathcal{B}}}\right] \approx \widehat{\overline{H}}^{\mathcal{B}}\left[n_{\mathrm{HF}}^{\mathcal{B}}\right] \equiv \widehat{\overline{H}}^{\mathcal{B}},\tag{26}$$

where we have dropped the explicit dependence on  $n_{\rm HF}^{\mathcal{B}}$  in the Hamiltonian for the sake of simplicity.

We then approximately solve eqn (24), with the approximation of eqn (26), using the EOM-CCSD method (see, *e.g.*, ref. 20) as follows. First, the ground-state wave function is approximated as a coupled-cluster singles and doubles (CCSD) ansatz:

$$\left|\Psi_{0}^{\mathcal{B}}\right\rangle = \mathbf{e}^{\hat{T}} \left|\Phi_{\mathrm{HF}}^{\mathcal{B}}\right\rangle,\tag{27}$$

with  $\hat{T} = \hat{T}_1 + \hat{T}_2$ , where  $\hat{T}_1$  and  $\hat{T}_2$  are the usual single- and double-excitation operators in the basis set  $\mathcal{B}$ . The single- and double-excitation amplitudes are determined from the ground-state CCSD amplitude equations using the Hamiltonian  $\hat{\overline{H}}^{\mathcal{B}}$ ,

$$\left\langle \Phi_{\mu} \middle| e^{-\hat{T}} \widehat{\overline{H}}^{\mathcal{B}} e^{\hat{T}} \middle| \Phi^{\mathcal{B}}_{\rm HF} \right\rangle = 0, \tag{28}$$

for all singly and doubly excited Slater determinants  $\Phi_{\mu}$  with respect to  $\Phi_{\text{HF}}^{\mathcal{B}}$  (which we will denote with  $\mu \in \text{SD}$ ). Note of course that the optimal single- and doubleexcitation amplitudes are not the same as in standard CCSD, since we use the effective Hamiltonian  $\widehat{\overline{H}}^{\mathcal{B}}$ . The corresponding ground-state CCSD energy eigenvalue is

$$\mathcal{E}_{0}^{\mathcal{B}} = \left\langle \Phi_{\rm HF}^{\mathcal{B}} \middle| e^{-\hat{T}} \widehat{\overline{H}}^{\mathcal{B}} e^{\hat{T}} \middle| \Phi_{\rm HF}^{\mathcal{B}} \right\rangle.$$
<sup>(29)</sup>

Then, we solve the EOM-CCSD equations with the fixed similarity-transformed Hamiltonian,  $e^{-\hat{T}} \hat{\overline{H}}^{\mathcal{B}} e^{\hat{T}}$ :

$$\left\langle \Phi_{\mu} \middle| \mathbf{e}^{-\hat{T}} \left( \widehat{\overline{H}}^{\mathscr{B}} - \mathcal{E}_{0}^{\mathscr{B}} \right) \mathbf{e}^{\hat{T}} \middle| \Psi_{n}^{\mathscr{B}} \right\rangle = \omega_{n}^{\mathscr{B}} \left\langle \Phi_{\mu} \middle| \Psi_{n}^{\mathscr{B}} \right\rangle, \tag{30}$$

where  $\omega_n^{\mathcal{B}}$  are the excitation energies  $(n \ge 1)$  and the excited-state wave functions are expanded on all single and double excitations:

$$|\Psi_{n}^{\mathcal{B}}\rangle = \sum_{\mu \in \mathrm{SD}} c_{n,\mu} |\Phi_{\mu}\rangle.$$
 (31)

In practice, the only change to make in the standard EOM-CCSD algorithm is thus to replace the usual one-electron integrals  $h_{pq} = \langle \phi_p | \hat{T} + \hat{V}_v | \phi_q \rangle$  with

$$h_{pq} \to h_{pq} + \overline{\nu}_{pq}^{\mathcal{B}},\tag{32}$$

where  $\overline{v}_{pq}^{\mathcal{B}} = \langle \phi_p | \widehat{\overline{v}}^{\mathcal{B}} [n_{\text{HF}}^{\mathcal{B}}] | \phi_q \rangle$  are the integrals of the basis-set correction potential. Consistently with the FC approximation, the one-electron integral  $\overline{v}_{pq}^{\mathcal{B}}$  is set to zero if *p* or *q* refers to a core orbital.

# 3. Numerical results

# 3.1. Computational details

We computed systematically the first three excited states of both singlet and triplet spin symmetry for  $NH_3$ ,  $H_2O$ , CO,  $N_2$ , and  $N_2CH_2$  molecules, whose geometries have been taken from ref. 41. This constitutes a set of 30 excited states, among which 14 have a Rydberg character and 16 have a valence character (according to the classification reported in previous works<sup>20,41</sup>).

Standard EOM-CCSD calculations (i.e., using the standard Hamiltonian) have been performed using the Gaussian-16 software<sup>42</sup> with the aug-cc-pVXZ (X = D, T, O, 5) basis sets,<sup>43</sup> abbreviated as AVXZ, except for the N<sub>2</sub>CH<sub>2</sub> molecule with the AV5Z basis set, for which the PySCF software<sup>44</sup> was used. The EOM-CCSD calculations using the effective Hamiltonian  $\widehat{\overline{H}}^{\mathcal{B}}$  have been performed using the PySCF software<sup>44</sup> by reading the one- and two-electron integrals defining the Hamiltonian  $\widehat{\overline{H}}^{\mathcal{B}}$  from a FCIDUMP format for the AVDZ and AVTZ basis sets. Limitations of the FCIDUMP format prevented us from performing calculations in larger basis sets, but we believe that the results presented here are sufficient to discuss the main trends. The one-electron integrals, including the basis-set correction potential integrals  $\bar{v}_{pq}^{\mathcal{B}}$  [eqn (32)], and the two-electron integrals have been computed with the Quantum Package software.<sup>45</sup> The integrals  $\overline{v}_{pq}^{\mathcal{B}}$  have been computed using a standard Becke-type⁴6 spatial grid with 75 radial points and 302 Lebedev angular points. All calculations have been performed within the FC approximation, both for the EOM-CCSD part and the computation of all quantities related to the basis-set correction [see eqn (11)-(14)]. The EOM-CCSD calculations using either the LDA or PBE versions of the basis-set correction functional [see eqn (9) and (10)] will be referred to as EOM-CCSD-LDA and EOM-CCSD-PBE, respectively.

# 3.2. Results and discussion

We report the EOM-CCSD, EOM-CCSD-LDA, and EOM-CCSD-PBE excitation energies for the NH<sub>3</sub>, H<sub>2</sub>O, CO, N<sub>2</sub>, and N<sub>2</sub>CH<sub>2</sub> molecules in Tables 1–5, respectively. We can notice that, except for the <sup>1</sup>A<sub>1</sub> and <sup>3</sup>A<sub>1</sub> states of the NH<sub>3</sub> molecule, all standard EOM-CCSD excitation energies computed with the AV5Z basis set can be considered as converged with respect to the basis set to within less than 0.02 eV. For each system and each excited state, we will therefore use the EOM-CCSD excitation energies computed with the AV5Z basis set for the CBS limit. For each system we also report the mean absolute deviation (MAD) with respect to the reference AV5Z basis set calculation, and when possible we also report the MAD computed with only Rydberg excited states [MAD(R)] or only valence excited states [MAD(V)] in order to differentiate these two types of excitation energies. We also report in Table 6 the MAD computed over the whole set of 28 converged excitation energies, together with MAD(R) and MAD(V) obtained with the 12 and 16 converged Rydberg and valence excitation energies, respectively.

A detailed look at all the tables reveals two interesting general trends: (i) except for the  ${}^{3}\Sigma$  valence state of the CO molecule and for the two unconverged  ${}^{1}A_{1}$  and  ${}^{3}A_{1}$  Rydberg states of the NH<sub>3</sub> molecule, all the excitation energies corresponding to Rydberg states increase when the size of the basis set is increased, while the excitation energies corresponding valence excited states tend to be stable or

Table 1 NH<sub>3</sub> molecule: standard EOM-CCSD and basis-set-corrected EOM-CCSD-LDA and EOM-CCSD-PBE excitation energies (eV) with the AVXZ basis sets (with X = D, T, Q, 5). The letter "R" indicates the Rydberg nature of the excited states. The MAD reported was calculated with all the excitation energies, except for the <sup>1</sup>A<sub>1</sub> and <sup>3</sup>A<sub>1</sub> excited states, for which convergence is not yet reached with the AVSZ basis set

	EOM-CO	CSD		EOM-CCSD- LDA		EOM-CCSD- PBE		
	AVDZ	AVTZ	AVQZ	AV5Z	AVDZ	AVTZ	AVDZ	AVTZ
${}^{1}A_{2}(R)$	6.45	6.60	6.65	6.67	6.59	6.65	6.58	6.65
$^{1}E(R)$	8.02	8.15	8.19	8.20	8.16	8.20	8.15	8.20
${}^{1}A_{1}(R)$	9.65	9.33	9.13	8.93	9.77	9.39	9.77	9.39
${}^{3}A_{2}(R)$	6.15	6.30	6.35	6.37	6.28	6.35	6.28	6.35
${}^{3}E(R)$	7.89	8.02	8.07	8.08	8.02	8.08	8.02	8.08
${}^{3}A_{1}(R)$	8.45	8.70	8.60	8.48	8.98	8.76	8.98	8.76
MAD(R)	0.20	0.06	0.01	—	0.07	0.01	0.07	0.01

**Table 2** H<sub>2</sub>O molecule: standard EOM-CCSD and basis-set-corrected EOM-CCSD-LDA and EOM-CCSD-PBE excitation energies (eV) with the AVXZ basis sets (with X = D, T, Q, 5). The letter "R" indicates the Rydberg nature of the excited states. The MAD for all six excitation energies is reported

	EOM-CO	CSD			EOM-CCSD- LDA		EOM-CCSD- PBE	
	AVDZ	AVTZ	AVQZ	AV5Z	AVDZ	AVTZ	AVDZ	AVTZ
${}^{1}B_{1}(R)$	7.45	7.60	7.66	7.68	7.63	7.67	7.62	7.67
${}^{1}A_{2}(R)$	9.21	9.36	9.42	9.44	9.40	9.44	9.39	9.44
${}^{1}A_{1}(R)$	9.86	9.96	10.00	10.01	10.02	10.02	10.02	10.02
${}^{3}B_{1}(R)$	7.04	7.20	7.28	7.30	7.22	7.28	7.21	7.27
${}^{3}A_{2}(R)$	9.05	9.20	9.26	9.28	9.23	9.27	9.21	9.27
${}^{3}A_{1}(R)$	9.39	9.49	9.54	9.56	9.55	9.55	9.54	9.55
MAD(R)	0.21	0.08	0.02	—	0.04	0.01	0.05	0.01

decrease with the basis set, and (ii) the basis-set error in the excitation energies of the Rydberg excited states is much larger than that of the valence excited states. Quantitatively, for the Rydberg excited states, the overall MADs are 0.21 eV, 0.07 eV, and 0.02 eV for standard EOM-CCSD with the AVDZ, AVTZ, and AVQZ basis sets, while for the valence excited states they are 0.07 eV, 0.01 eV, and 0.01 eV with the same basis sets. A qualitative explanation of this observation could be that, in a Rydberg excited state, one electron is in a diffuse orbital relatively far from the bulk of the electronic density of the molecule, and therefore the correlation effects of this electron are much smaller than in the ground state, leading, in a small basis set, to a description biased toward the excited state and therefore a too-small excitation energy. By contrast, in a valence excited state, the excited electron remains in a valence orbital and is much closer to the bulk of the electronic density, and therefore the correlation effects are much more comparable to those of the ground state, leading to a much smaller basis-set error. We therefore conclude from this part of the study that the main source of basis-set error for the description of a set of excited states comes from the Rydberg excited states.

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**Table 3** CO molecule: Standard EOM-CCSD and basis-set-corrected EOM-CCSD-LDA and EOM-CCSD-PBE excitation energies (eV) with the AVXZ basis sets (with X = D, T, Q, 5). The letter "V" indicates the valence nature of the excited states. The MAD for all six excitation energies is reported

	EOM-CO	CSD			EOM-CCSD- LDA		EOM-CCSD- PBE	
	AVDZ	AVTZ	AVQZ	AV5Z	AVDZ	AVTZ	AVDZ	AVTZ
${}^{1}\Pi$ (V)	8.67	8.59	8.57	8.57	8.76	8.63	8.74	8.62
${}^{1}\Sigma$ (V)	10.10	9.99	9.99	10.00	10.15	10.01	10.15	10.01
$^{1}\Delta$ (V)	10.21	10.12	10.13	10.13	10.26	10.15	10.26	10.14
$^{3}\Pi$ (V)	6.38	6.36	6.36	6.36	6.43	6.38	6.41	6.38
$^{3}\Sigma$ (V)	8.34	8.34	8.37	8.39	8.39	8.37	8.39	8.37
$^{3}\Delta$ (V)	9.29	9.24	9.24	9.25	9.35	9.25	9.35	9.26
MAD(V)	0.06	0.02	0.01		0.11	0.02	0.10	0.02

**Table 4** N<sub>2</sub> molecule: standard EOM-CCSD and basis-set-corrected EOM-CCSD-LDA and EOM-CCSD-PBE excitation energies (eV) with the AVXZ basis sets (with X = D, T, Q, 5). The letter "V" indicates the valence nature of the excited states. The MAD for all six excitation energies is reported

	EOM-CO	CSD			EOM-CCSD- LDA		EOM-CCSD- PBE	
	AVDZ	AVTZ	AVQZ	AV5Z	AVDZ	AVTZ	AVDZ	AVTZ
${}^{1}\Pi_{g}(V)$	9.50	9.41	9.40	9.40	9.60	9.45	9.57	9.45
${}^{1}\Sigma_{u}(V)$	10.20	10.00	9.98	9.98	10.24	10.01	10.24	10.01
$^{1}\Delta_{u}(V)$	10.61	10.44	10.42	10.42	10.66	10.46	10.66	10.46
$^{3}\Sigma_{u}(V)$	7.69	7.66	7.69	7.69	7.71	7.67	7.71	7.67
${}^{3}\Pi_{g}(V)$	8.12	8.09	8.10	8.10	8.20	8.13	8.18	8.13
$^{3}\Delta_{u}(V)$	9.04	8.91	8.91	8.91	9.07	8.92	9.07	8.92
MAD(V)	0.11	0.01	0.00	—	0.16	0.03	0.15	0.03

Moving now to the our EOM-CCSD-LDA and EOM-CCSD-PBE calculations, we see that, with respect to the standard EOM-CCSD calculations in a given basis set, the effect of the basis-set correction potential is always to increase the excitation energies. In the Appendix, we provide a simplified model that rationalizes this systematic increase in the excitation energies. As the valence excitation energies tend to decrease with the basis set, our approach cannot improve these excitation energies, while it will improve the description of the Rydberg excitation energies, which tend to be underestimated in a finite basis set. The present test set consists of 12 Rydberg excited states and 16 excited valence states, and represents a relatively balanced selection between excitation energies that the basis-set correction method with the current approximations can improve and excitation energies that it will tend to deteriorate.

From a quantitative point of view, the EOM-CCSD-LDA and EOM-CCSD-PBE approximations give very similar results to the AVDZ basis set (the larger difference of the MAD is 0.01 eV) and essentially indistinguishable results from the AVTZ basis set. We also notice that the MADs for the Rydberg excitation energies are drastically reduced by the basis-set correction potential. With the AVDZ basis set,

Table 5 N<sub>2</sub>CH<sub>2</sub> molecule: standard EOM-CCSD and basis-set-corrected EOM-CCSD-LDA and EOM-CCSD-PBE excitation energies (eV) with the AVXZ basis sets (with X = D, T, Q, 5). The letter "R" or "V" indicates the Rydberg or valence nature of the excited states. The MAD for all six excitation energies is reported, together with the MAD calculated with the two Rydberg excited states and the MAD calculated with the four valence excited states

	EOM-CO	CSD			EOM-CCSD- LDA		EOM-CCSD- PBE	
	AVDZ	AVTZ	AVQZ	AV5Z	AVDZ	AVTZ	AVDZ	AVTZ
${}^{1}A_{2}(V)$	3.23	3.19	3.19	3.20	3.26	3.20	3.25	3.20
${}^{1}B_{1}(R)$	5.43	5.57	5.62	5.65	5.58	5.63	5.58	5.63
${}^{1}A_{1}(V)$	5.90	5.94	5.96	5.97	5.99	5.98	5.99	5.98
${}^{3}A_{2}(V)$	2.90	2.88	2.88	2.90	2.94	2.89	2.93	2.89
${}^{3}B_{1}(V)$	3.99	3.95	3.95	3.95	3.99	3.95	3.98	3.94
${}^{3}A_{1}(R)$	5.26	5.42	5.46	5.50	5.41	5.48	5.41	5.48
MAD	0.10	0.04	0.02		0.05	0.01	0.05	0.01
MAD(R)	0.23	0.08	0.04		0.08	0.02	0.08	0.02
MAD(V)	0.03	0.01	0.01	—	0.04	0.01	0.03	0.01

Table 6 Total MAD calculated over the set of 28 excitation energies, together with the MADs calculated over the set of Rydberg excited states [MAD(R)] and the set of valence excited states [MAD(V)]

	MAD	MAD(R)	MAD(V)
	EOM-CCSD		
AVDZ	0.14	0.21	0.07
AVTZ	0.04	0.07	0.01
AVQZ	0.01	0.02	0.01
,	EOM-CCSD-LDA	L	
AVDZ	0.09	0.06	0.10
AVTZ	0.02	0.01	0.02
	EOM-CCSD-PBE		
AVDZ	0.08	0.07	0.09
AVTZ	0.02	0.01	0.02

the MADs obtained with EOM-CCSD-PBE are 0.07 eV, 0.05 eV, and 0.08 eV for the NH<sub>3</sub>, H<sub>2</sub>O, and N<sub>2</sub>CH<sub>2</sub> molecules, respectively, smaller by at least a factor of two with respect to the MADs obtained with the standard EOM-CCSD method, and thus reaching an accuracy similar to standard EOM-CCSD with the AVTZ basis set. With the AVTZ basis set, the MADs obtained with the basis-set-corrected EOM-CCSD method for the Rydberg excitation energies are 0.01 eV for both the NH<sub>3</sub> and H<sub>2</sub>O molecules, and 0.02 eV for the N<sub>2</sub>CH<sub>2</sub> molecule, which is as accurate as standard EOM-CCSD with the AVQZ basis set for the NH<sub>3</sub> molecule, and even more accurate in the case of the H<sub>2</sub>O and N<sub>2</sub>CH<sub>2</sub> molecules. We therefore conclude that the addition of the basis-set correction potential drastically improves the basis-set convergence of the excitation energies for the Rydberg excited states at virtually no cost with respect to standard EOM-CCSD calculations.

Turning now to the set of valence excited states, as anticipated above, the basis-set correction potential overall deteriorates the accuracy of the excitation energies, but this deterioration becomes smaller as the basis-set size increases.

More quantitatively, the MADs obtained with EOM-CCSD-PBE with the AVDZ basis set are 0.10 eV, 0.15 eV, and 0.03 eV for the CO,  $N_2$ , and  $N_2CH_2$  molecules, respectively, which are larger than the standard EOM-CCSD values by 0.04 eV for the CO and  $N_2$  molecules, but identical for the  $N_2CH_2$  molecule. With the AVTZ basis set, the MADs obtained with EOM-CCSD-PBE decrease to 0.02 eV, 0.03 eV, and 0.01 eV, representing an accuracy comparable to that of standard EOM-CCSD with the same basis set.

# 4. Conclusion

In the present work we proposed and tested a novel scheme based on the DBBSC method<sup>21</sup> to improve the basis-set convergence of the excitation energies in wavefunction calculations. This is based on the recently introduced linear-response theory<sup>35</sup> for the DBBSC method, which was only tested on a one-dimensional model system. In order to treat real molecular systems, we use basis-set correction density functionals previously developed in the DBBSC method for ground-state calculations, and we introduce a further approximation in the response equations, which consists in neglecting the kernel contribution coming from the second-order derivatives of the basis-set correction functional. The advantage of the approximation is that the response equations can be rewritten as a usual FCI eigenvalue equation with an additional one-electron potential coming from the first-order derivative of the basis-set correction functional. Therefore, by simply modifying the one-electron integrals in the Hamiltonian, this scheme can be applied to any wave-function methods targeting excited states.

Applying this scheme to the EOM-CCSD method, we presented numerical tests performed on a set of 30 excitation energies for the NH<sub>3</sub>, H<sub>2</sub>O, CO, N<sub>2</sub>, and N<sub>2</sub>CH<sub>2</sub> molecules. The results were analysed based on a partition of the excitation energies: the ones corresponding to Rydberg excited states and the ones corresponding to valence excited states. We found that the global basis-set error is dominated by the Rydberg excited states, whose excitation energies tend to increase when the size of the basis set is increased, while the valence excitation energies tend to be much less sensitive to the basis set and overall tend to decrease with the basis set. The increase in the excitation energy with the basis set for a Rydberg excited state can be qualitatively understood by the fact that the excited electron is relatively far from the molecule and is therefore much less correlated than in the ground state, which leads to a biased description in favour of the excited state in small basis sets. By contrast, in a valence excited state, the excited electron remains in the valence region and its correlation effects are much more comparable to that of the ground state, leading to a much smaller sensitivity to finite basis-set effects. We found that the present basis-set-corrected EOM-CCSD method always increases the excitation energies, and therefore it tends to improve Rydberg excitation energies, while it tends to deteriorate the valence excitation energies. Indeed, with the AVTZ basis set, the DBBSC scheme reduces the MAD on the Rydberg excitation energies obtained with standard EOM-CCSD from 0.07 eV to 0.01 eV, which is a large improvement. With the same AVTZ basis set, the DBBSC scheme increases the MAD on the valence excitation energies from 0.01 eV to 0.02 eV, which still represents a reasonable accuracy.

We therefore conclude that the present basis-set-corrected EOM-CCSD method allows one to overall reduce the basis-set error for the computation of excitation

energies, at virtually no additional cost with respect to standard EOM-CCSD calculations. In forthcoming works, we will investigate the impact of taking into account the basis-set correction kernel, together with the dependency on the density used.

# 5. Rationalization of the effect of the basis-set correction potential on a simplified model

Here, we present a simplified model that rationalizes the observed systematic increase in the excitation energies when the basis-set correction potential  $\hat{\overline{v}}^{\mathcal{B}}$  is added to the standard Hamiltonian.

We consider the case of a spin-singlet ground state described by the HF determinant  $|\Phi_{\rm HF}\rangle$ , and a spin-singlet excited state described by a single spin-adapted excitation from an occupied orbital *i* to a virtual orbital *a*:

$$|\Phi_i^{a}
angle = \frac{1}{\sqrt{2}} \sum_{\sigma \in \{\uparrow,\downarrow\}} \hat{a}_{a,\sigma}^{\dagger} \hat{a}_{i,\sigma} |\Phi_{\mathrm{HF}}
angle.$$
 (33)

This is of course a very simplified model, but most of the low-lying excited states treated in the present study are dominated by singly excited Slater determinants. Because of Brillouin's theorem, the standard Hamiltonian  $\hat{H}$  is diagonal in the basis  $\{|\Phi_{\rm HF}\rangle, |\Phi_i^a\rangle\}$  and therefore the corresponding excitation energy  $\Delta_{ia}$  is obtained simply as

$$\Delta_{ia} = \langle \Phi_i^a | \hat{H} | \Phi_i^a \rangle - \langle \Phi_{\rm HF} | \hat{H} | \Phi_{\rm HF} \rangle > 0.$$
(34)

We also assume here that the orbitals *i* and *a* have different spatial symmetries, such that any totally symmetrical one-electron operator, such as the basis-set correction potential  $\hat{\overline{V}}^{\mathcal{B}}$ , has vanishing matrix elements between orbitals *i* and *a*. Thus, the effective Hamiltonian  $\hat{\overline{H}}^{\mathcal{B}} = \hat{H} + \hat{\overline{V}}^{\mathcal{B}}$  is also diagonal in the basis  $\{|\Phi_{\rm HF}\rangle, |\Phi_i^a\rangle\}$ , and thus the basis-set-corrected excitation energy  $\bar{\Delta}_{ia}$  is simply

$$\overline{\varDelta}_{ia} = \langle \Phi_i^{\ a} | \widehat{\overline{H}}^{\ B} | \Phi_i^{\ a} \rangle - \langle \Phi_{\rm HF} | \widehat{\overline{H}}^{\ B} | \Phi_{\rm HF} \rangle = \varDelta_{ia} + \overline{v}_{aa}^{\ B} - \overline{v}_{ii}^{\ B}, \qquad (35)$$

where  $\overline{v}_{pp}^{\mathcal{B}} = \langle \phi_p | \widehat{\overline{V}}^{\mathcal{B}} | \phi_p \rangle$ . Therefore, the effect of  $\widehat{\overline{V}}^{\mathcal{B}}$  is determined in this case by the sign of  $\overline{v}_{aa}^{\mathcal{B}} - \overline{v}_{ii}^{\mathcal{B}}$ . In the LDA-UEG approximation, which in practice yields essentially the same results as the PBE-UEG approximation, the integrals  $\overline{v}_{pp}^{\mathcal{B}}$  are of the form

$$\overline{\nu}_{pp}^{\mathcal{B}} = \int d\mathbf{r} \ \overline{\nu}^{\mathcal{B}} \left( n_{\rm HF}^{\mathcal{B}}(\mathbf{r}), \mu^{\mathcal{B}}(\mathbf{r}) \right) \phi_p(\mathbf{r})^2, \tag{36}$$

where the basis-set correction potential  $\overline{v}^{\mathcal{B}}(n,\mu)$  has the property of being negative for all densities and range-separation, parameters

$$\overline{v}^{\mathcal{B}}(n,\mu) < 0, \quad \forall \ n,\mu, \tag{37}$$

and we can therefore conclude that

$$\overline{v}_{pp}^{\mathcal{B}} < 0, \quad \forall \ \phi_p.$$
(38)

Hence, the integral  $\overline{\nu}_{pp}^{\mathcal{B}}$  is the overlap between the negative potential  $\overline{\nu}^{\mathcal{B}}(n_{\mathrm{HF}}^{\mathcal{B}}(\mathbf{r}), \mu^{\mathcal{B}}(\mathbf{r}))$  and the charge density  $\phi_p(\mathbf{r})^2$ . Also, for a given  $\mu$ ,  $|\overline{\nu}^{\mathcal{B}}(n, \mu)|$  is an increasing function of the density n, and thus the maximum of  $|\overline{\nu}^{\mathcal{B}}(n_{\mathrm{HF}}^{\mathcal{B}}(\mathbf{r}), \mu^{\mathcal{B}}(\mathbf{r}))|$  is located at the maximum of the density  $n_{\mathrm{HF}}^{\mathcal{B}}(\mathbf{r})$ . Therefore, as the virtual orbitals overlap less with  $\overline{\nu}^{\mathcal{B}}(n_{\mathrm{HF}}^{\mathcal{B}}(\mathbf{r}), \mu^{\mathcal{B}}(\mathbf{r}))$  than the occupied orbitals, one can expect that

$$\overline{v}_{aa}^{\mathcal{B}} - \overline{v}_{ii}^{\mathcal{B}} > 0, \tag{39}$$

so that

$$\Delta_{ia} + \overline{v}^{\mathcal{B}}_{aa} - \overline{v}^{\mathcal{B}}_{ii} > \Delta_{ia}, \tag{40}$$

explaining why the basis-set-corrected excitation energy  $\bar{\Delta}_{ia}$  is always larger than the uncorrected excitation energy  $\Delta_{ia}$ .

# Conflicts of interest

There are no conflicts to declare.

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