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A multiconfigurational hybrid density-functional theory

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We propose a multiconfigurational hybrid density-functional theory which rigorously combines a multiconfiguration self-consistent-field calculation with a density-functional approximation based on a linear decomposition of the electron-electron interaction. This gives a straightforward extension of the usual hybrid approximations by essentially adding a fraction λ of exact static correlation in addition to the fraction λ of exact exchange. Test calculations on the cycloaddition reactions of ozone with ethylene or acetylene and the dissociation of diatomic molecules with the Perdew-Burke-Ernzerhof (PBE) and Becke-Lee-Yang-Parr (BLYP) density functionals show that a good value of λ is 0.25, as in the usual hybrid approximations. The results suggest that the proposed multiconfigurational hybrid approximations can improve over usual density-functional calculations for situations with strong static correlation effects.

I. INTRODUCTION

Density-functional theory (DFT) [1] within the Kohn-Sham (KS) scheme [2] is the most widely used method for electronic-structure calculations in atomic, molecular and solid-state systems. With the usual approximate density functionals, such as generalized-gradient approximations (GGA) and hybrid approximations mixing in a fraction of Hartree-Fock (HF) exchange, DFT KS generally gives good results for situations in which the so-called dynamic electron correlation dominates the total correlation energy, but it can yield severe errors for systems with strong static correlation, i.e. with partially filled near-degenerate orbitals (see, e.g., Ref. 3). It has been argued that the GGA exchange density functionals actually mimic some static correlation through their self-interaction error, though in an imperfect manner (see, e.g., Refs. 4, 5). There is often indeed a partial cancellation of errors between the self-interaction error which tends to give too low an energy and the neglect of static correlation which gives too high an energy (see, e.g., Ref. 6). Hybrid approximations have a smaller selfinteraction error and are thus often worse than pure density functionals for describing systems with static correlation (see, e.g., Ref. 7).

Several approaches have been proposed to include explicit static correlation in density-functional theory. Artificially breaking (space and spin) symmetry by unrestricted KS calculations is the simplest approach to simulate static correlation (see, e.g., Ref. 8), and it often leads to reasonable potential energy surfaces but wrong spin densities. Another possible approach consists in replacing the single KS determinant by an ensemble of determinants or, equivalently, using fractional occupation numbers for the orbitals [9–22], but a successful and general method based on this idea is still lacking. Configurationinteraction schemes have also been proposed in which modified Hamiltonian matrix elements include information from DFT [23–26]. A lot of approaches consist in adding to the energy of a partially correlated wave function calculation, including near-degenerate configuration state functions coupled by the full Coulombic electronelectron interaction, an energy density functional describing the missing correlation effects [27–62]. In these last approaches, one must use a density functional which depends on the size of the multiconfigurational expansion, in order to avoid double counting of correlation between the wave function part of the calculation and the density functional. Finally, to avoiding any double counting of correlation from the beginning, it has been proposed to decompose the Coulombic electron-electron interaction into long-range and short-range components, the long-range part being treated by a method capable of describing static correlation and the short-range part being described by a density functional approximation. The methods that have been used for the long-range part are: configuration interaction [63–69], multiconfigurational self-consistent field (MCSCF) [70–72], multireference perturbation theory [73], constrained-pairing meanfield theory [74, 75], and density-matrix functional theory [76–78].

In this work, we explore the possibility to combine MC-SCF and DFT based on a simple linear decomposition of the Coulombic electron-electron interaction, in the spirit of the usual hybrid approximations, and similarly to what was recently done for constructing theoretically justified double-hybrid approximations [79]. This approach gives a straightforward multiconfigurational extension of the standard hybrid approximations, and aims at improving their description of static correlation. After deriving this multiconfigurational hybrid density-functional theory, we test this approach on situations with strong static correlation effects, namely the cycloaddition reac-

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tions of ozone with ethylene or acetylene and the dissociation of diatomic molecules, and we compare with other methods, in particular the range-separated multiconfigurational hybrid method of Ref. 70–72.

II. THEORY

Using the formalism of the multideterminant extension of the KS scheme (see, e.g., Refs. 69, 71, 79, 80), for any coupling constant λ , the *exact* energy can be expressed as the following minimization over multideterminant wave functions Ψ:

$$
E = \min_{\Psi} \left\{ \langle \Psi | \hat{T} + \hat{V}_{\text{ext}} + \lambda \hat{W}_{ee} | \Psi \rangle + \bar{E}_{\text{Hxc}}^{\lambda} [n_{\Psi}] \right\}, (1)
$$

where \hat{T} is the kinetic energy operator, \hat{V}_{ext} is a scalar external potential operator (e.g., nuclei-electron), \hat{W}_{ee} is the electron-electron interaction operator, and $\bar{E}_{\text{Hxc}}^{\lambda}[n_{\Psi}]$ is the *complement* λ -dependent Hartreeexchange-correlation density functional evaluated at the density coming from Ψ. The complement density functional, $\bar{E}_{\text{Hxc}}^{\lambda}[n] = E_{\text{Hxc}}[n] - E_{\text{Hxc}}^{\lambda}[n]$, is the difference between the usual KS density functional $E_{Hxc}[n]$ and the λ -dependent density functional $E_{\text{Hxc}}^{\lambda}[n]$ corresponding to the interaction $\lambda \hat{W}_{ee}$. The Hartree-exchange contribution is of first order in the electron-electron interaction and is thus linear in λ ,

$$
\bar{E}_{\mathrm{H}x}^{\lambda}[n] = (1 - \lambda)E_{\mathrm{H}x}[n],\tag{2}
$$

where $E_{\text{H}x}[n]$ is the usual KS Hartree-exchange density functional. The correlation contribution is obtained by uniform coordinate scaling of the density [81–84],

$$
\begin{aligned}\n\bar{E}_c^\lambda[n] &= E_c[n] - E_c^\lambda[n] \\
&= E_c[n] - \lambda^2 E_c[n_{1/\lambda}],\n\end{aligned} \tag{3}
$$

where $E_c[n]$ is the usual KS correlation functional, $E_c^{\lambda}[n]$ is the correlation functional corresponding to the interaction $\lambda \hat{W}_{ee}$, and $n_{1/\lambda}(\mathbf{r}) = (1/\lambda)^3 n(\mathbf{r}/\lambda)$ is the scaled density.

The theory is so far exact but in practice approximations must be used for the multideterminant wave function and the density functionals. In Ref. 79, by restricting the search in Eq. (1) to single-determinant wave functions Φ , we defined the *density-scaled one-parameter hy*brid (DS1H) approximation

$$
E^{\text{DS1H}} = \min_{\Phi} \left\{ \langle \Phi | \hat{T} + \hat{V}_{\text{ext}} + \lambda \hat{W}_{ee} | \Phi \rangle \right.
$$

$$
+ (1 - \lambda) E_{\text{Hx}} [n_{\Phi}] + E_c [n_{\Phi}] - \lambda^2 E_c [n_{\Phi, 1/\lambda}] \right\}, \quad (4)
$$

and, by additionally neglecting density scaling in the correlation functional, $E_c[n_{1/\lambda}] \approx E_c[n]$, we obtained the one-parameter hybrid (1H) approximation,

$$
E^{1H} = \min_{\Phi} \left\{ \langle \Phi | \hat{T} + \hat{V}_{ext} + \lambda \hat{W}_{ee} | \Phi \rangle \right.+(1 - \lambda) E_{Hx} [n_{\Phi}] + (1 - \lambda^2) E_c [n_{\Phi}] \right\}, \quad (5)
$$

which is similar to the usual one-parameter hybrid approximations [85, 86], except that the correlation functional is weighted by a factor of $(1 - \lambda^2)$. Starting from these references and applying a second-order Møller-Plesset (MP2) perturbation theory [80, 87, 88], we also defined the density-scaled one-parameter double-hybrid (DS1DH) and one-parameter double-hybrid (1DH) approximations [79], which are one-parameter versions of the original double-hybrid approximations [89]. These latter also combine HF exchange and MP2 correlation with a semilocal exchange-correlation density functional but with two empirical parameters.

Here, we follow a different route and use an MCSCF wave function in Eq. (1), expanded as a linear combination of configuration state functions Φ_I ,

$$
|\Psi\rangle = \sum_{I} c_{I} |\Phi_{I}\rangle, \tag{6}
$$

where the coefficients c_I and the orbitals are to be simultaneously optimized. With this form of wave function, we obtain a multiconfigurational density-scaled oneparameter hybrid (MCDS1H) approximation,

$$
E^{\text{MCDS1H}} = \min_{\Psi} \left\{ \langle \Psi | \hat{T} + \hat{V}_{\text{ext}} + \lambda \hat{W}_{ee} | \Psi \rangle \right.
$$

$$
+ (1 - \lambda) E_{\text{Hx}}[n_{\Psi}] + E_c[n_{\Psi}] - \lambda^2 E_c[n_{\Psi, 1/\lambda}] \right\}, (7)
$$

and, if density scaling, which is not considered in usual hybrid approximations, is neglected, we obtain a *multi*configurational one-parameter hybrid (MC1H) approximation,

$$
E^{\text{MC1H}} = \min_{\Psi} \left\{ \langle \Psi | \hat{T} + \hat{V}_{\text{ext}} + \lambda \hat{W}_{ee} | \Psi \rangle \right.\n \left. + (1 - \lambda) E_{\text{Hx}} [n_{\Psi}] + (1 - \lambda^2) E_c [n_{\Psi}] \right\}.\n \tag{8}
$$

Equation (7) is the equivalent of the range-separated multiconfigurational hybrid method of Refs. 70–72, that we will refer to as MC-srDFT, but for a linear separation of the electron-electron interaction. Notice that, if we were to use no approximations for the wave function Ψ and the exchange-correlation density functional, then Eq. (7) would give the exact energy, *independently* of λ . In practice, of course, we must use approximations, and the energy does depend on λ , which can then be considered as an empirical parameter to be optimized.

The present scheme has two advantages over the rangeseparated scheme: (a) Only one list of two-electron Coulomb integrals is needed and it is just multiplied by λ in the MCSCF part and by $(1 - \lambda)$ in the complement Hartree energy, whereas two lists of two-electron integrals are needed in the range-separated scheme for the long-range MCSCF part and for the short-range complement Hartree energy; (b) No new exchange and correlation density functionals need in principle to be developed since all the existing approximations developed for the KS scheme can be reused with a simple scaling, whereas

FIG. 1: MAEs of O3ADD6 set as functions of the parameter λ for the MC1H and MCDS1H approximations with the BLYP (left) and PBE (right) exchange-correlation density functionals. All calculations were carried out with the aug-cc-pVTZ basis set.

new short-range density-functional approximations must be developed in the range-separated scheme.

Equations (7) and (8) can be seen as straightforward multiconfigurational extensions of the usual hybrid approximations. Indeed, the expectation value of $\lambda \hat{W}_{ee}$ over the MCSCF wave function Ψ not only introduces a fraction λ of exact exchange but also a fraction of exact static correlation. Defined in the ideal limit of reference energy levels that are degenerate, the static correlation energy is linear with respect to the electron-electron interaction, and thus we can consider that the expectation value of $\lambda \hat{W}_{ee}$ introduces a linear fraction $\hat{\lambda}$ of static correlation. By contrast, the dynamic correlation energy starts at quadratic order in the electron-electron interaction, so that, for sufficiently small λ , it is justified to neglect it in the wave function expectation value. Moreover, for sufficiently small λ , the weight $(1 - \lambda^2)$ is close to 1 and thus Eq. (8) includes a nearly complete approximate correlation energy functional, that is often thought of as correctly describing dynamic correlation. Of course, if the multiconfigurational hybrid approximations of Eq. (7) or (8) are to be accurate, the fraction $(1-\lambda)$ of static correlation energy not treated by MCSCF must be accounted for by the density functional, possibly through a compensation with the self-interaction error of the scaled exchange functional $(1 - \lambda)E_x[n]$.

III. COMPUTATIONAL DETAILS

The calculations have been performed with a development version of the DALTON 2011 program [92], in which the MCDS1H and MC1H approximations have been implemented in the same way than for the MCsrDFT method [70, 71, 93], using the direct restrictedstep second-order MCSCF algorithm of Jensen and coworkers [94–99]. For $E_x[n]$ and $E_c[n]$, we use two GGA exchange-correlation density functionals, Perdew-Burke-Ernzerhof (PBE) [100] and Becke/Lee-Yang-Parr (BLYP) [101, 102], without spin-density dependence. For

implementing the density-scaled correlation functionals in the MCSCF algorithm, we need the scaling relations for the energy density, and its first- and second-order derivatives that we give in Appendix A. The computational cost of the method is essentially the same as for a standard MCSCF calculation, with a small extra cost due to the DFT contribution.

A good value for the empirical parameter λ in Eqs. (7) and (8) is determined on the O3ADD6 benchmark set [103, 104] for the 1,3-dipolar cycloaddition reactions of ozone (O_3) with ethylene (C_2H_4) or acetylene (C_2H_2) [90, 105, 106]. For these two reactions, there are three stationary points along the reaction coordinate: the van der Waals complex, the transition state, and the cycloadduct (primary ozonide), all in a closedshell spin-singlet state. The O3ADD6 set consists of the six energies of these stationary points of the two reactions, calculated relative to the energy of the separated reactants, and without zero-point vibrational energy correction. Accurate calculations of these energies are difficult and require to handle the subtle balance between static and dynamic correlation effects along the reaction coordinate. The ozone reactant, the van der Waals complex, and the transition state have a strong multiconfigurational character corresponding to the HOMO \rightarrow LUMO double excitation in ozone. In the cycloadduct, and to a less extent in the transition state, the stabilization of the ozone HOMO and the destabilization of the ozone LUMO greatly reduce this multiconfigurational character. In addition, there are small near-degeneracy correlation effects due to the π and π^* orbitals of the reactive π bond of ethylene and acetylene. For each separate reactant, a $CAS(2,2)$ wave function is chosen, the active space corresponding to the HOMO and LUMO orbitals for ozone, and to the HOMO (π) and LUMO (π^*) orbitals of the reactive π bond for ethylene and acetylene. For the van der Waals complex, the transition state, and the cycloadduct, a CAS(4,4) wave function is consistently chosen, the active space corresponding to the orbitals that connect to the ones chosen for the reactants TABLE I: Energies of the van der Waals (vdW) complex, the transition state (TS), and the cycloadduct (primary ozonide), relative to the separated reactants, and the corresponding MAEs (in kcal/mol) for the addition of ozone with acetylene or ethylene (O3ADD6 set), calculated by several methods. For the DS1DH-PBE and 1DH-BLYP double-hybrid approximations, we use the value $\lambda = 0.65$ which was previously optimized in Ref. 79. For the MCDS1H and MC1H multiconfigurational hybrid approximations, we use a value of $\lambda = 0.25$ which roughly minimizes the MAE according to Fig. 1. For the range-separated MC-srPBE multiconfigurational hybrid approximation, we use the value of the range-separation parameter $\mu = 0.40$ bohr⁻¹ which was previously determined in Ref. 71. For the multiconfigurational methods, a complete active space wave function with 2 electrons in 2 orbitals [CAS(4,4)] is chosen for the van der Waals complex, the transition state, and the cycloadduct, and a CAS(2,2) wave function for each separate reactant. All calculations were carried out with the aug-cc-pVTZ basis set. All calculations were done for M05/6-311+G(2df,2p) geometries, except for the MRMP2 results which are for CCSD(T)/cc-pVTZ geometries.

 a From Ref. 90.

 b Performed with the Gaussian09 program [91].

in the dissociation limit. We use the aug-cc-pVTZ basis set [107, 108] and the fixed geometries of Ref. 90 optimized using the hybrid meta-GGA exchange-correlation functional M05 [109] with the $6-311+G(2df,2p)$ basis set [110, 111]. The reference values for the energies are from Ref. 90 and were obtained from extensive coupledcluster calculations extrapolated to the complete basis set limit [90, 105]. We calculate the mean absolute error (MAE) over the six values as a function of the parameter λ. We compare the MCDS1H and MC1H approximations with (a) some non-hybrid methods: HF, MCSCF, MP2 [112], multireference MP2 (MRMP2) [90, 113], PBE [100], and BLYP [101, 102]; (b) some singlehybrid approximations: PBE0 [114, 115], B1LYP [116], and B3LYP [117, 118]; (c) some double-hybrid approximations: DS1DH-PBE [79], 1DH-BLYP [79], and B2- PLYP [89], all applied in a spin-restricted formalism. We also compare with the range-separated MC-srPBE multiconfigurational hybrid approximation [70–72] using the short-range PBE exchange-correlation functional of Ref. 119 and the value of the range-separation parameter $\mu = 0.40$ bohr⁻¹ which was previously determined in Ref. 71.

We also test the MCDS1H and MC1H approximations by computing the potential energy curves the five diatomic molecules H_2 , Li_2 , C_2 , N_2 , and F_2 , using in each case a full-valence CAS wave function and the cc-pVTZ basis set [107].

IV. RESULTS

A. O3ADD6 database

Figure 1 shows the MAEs for the O3ADD6 set as functions of the parameter λ for the MCDS1H and MC1H

FIG. 2: Potential energy curves of H² calculated by HF, MCSCF, and several methods based on the PBE (left) or BLYP (right) exchange-correlation density functionals. For the MCDS1H and MC1H multiconfigurational hybrid approximations, we use a value of $\lambda = 0.25$. For the range-separated MC-srPBE multiconfigurational hybrid approximation, we use a value of the range-separation parameter of $\mu = 0.40 \text{ bohr}^{-1}$. For all multiconfigurational methods, we use a full-valence CAS wave function. The basis set is cc-pVTZ basis. The accurate curve is from Ref. 28.

approximations with the BLYP and PBE exchangecorrelation density functionals. For $\lambda = 0$, the MCDS1H and MC1H approximation reduce to a standard KS calculation with the corresponding approximate density functional. For $\lambda = 1$, they reduce to a standard MCSCF calculation. Contrary to what was found for the calculation of atomization energies using double-hybrid approximations [79], here neglecting density scaling in the correlation functional makes little difference. Toward the $\lambda = 1$ end of the curves, the MCDS1H and MC1H approximations inherit the inaccuracy of MCSCF which neglects dynamic correlation. The MAE curves of the MCDS1H-BLYP and MC1H-BLYP approximations display a marked minimum at an intermediate value of λ , thus improving upon both the standard BLYP and MC-SCF calculations. The minimum is reached at $\lambda = 0.25$ for MCDS1H-BLYP and at $\lambda = 0.30$ for MC1H-BLYP. with MAEs below 2 kcal/mol. For the MCDS1H-PBE and MC1H-PBE approximations, the MAE curves have a plateau around $\lambda = 0.25$ with a MAE of about 3 kcal/mol, which is again smaller than both the standard PBE and MCSCF calculations. In view of these results, we choose the value $\lambda = 0.25$ in all MCDS1H and MC1H approximations. It is a conservative choice since it gives the same fraction of exact exchange as the one usually advocated in the usual single-hybrid approximations [120].

Table I reports the energies of the van der Waals complex, the transition state, and the cycloadduct of the two reactions of the O3ADD6 set, relative to the separated reactants, calculated with the MCDS1H and MC1H approximations at $\lambda = 0.25$. For comparison, we also report results for various non-hybrid and other hybrid methods.

During the early stages of the two reactions, a weakly bound van der Waals complex is formed which lies in a shallow minimum $(-1.90 \text{ kcal/mol}$ for C_2H_2 and -1.94 kcal/mol for C_2H_4) below the reactants. The MCDS1H and MC1H approximations give significantly underestimated well depths, which are still in improvement over standard MCSCF but not over standard KS calculations with the corresponding functionals. The range-separated MC-srPBE method does also not perform better than KS PBE for these van der Waals systems. A better description of the long-range dispersion correlation would indeed require inclusion of perturbation corrections on top of the active space [73]. As expected, the double-hybrid approximations, which include second-order perturbation corrections, tend to perform better for these van der Waals complexes. The best performance is achieved with MRMP2 which is able to correctly describe both multiconfigurational effects and dispersion correlations. However, one should keep in mind than the methods MP2, MRMP2, and the double hybrids are most likely less converged with respect to the basis size than the other methods.

The activation barriers of the transition states are underestimated (or not present at all) in KS PBE and BLYP calculations, and to a less extent with the single-hybrid and double-hybrid approximations, while they are largely overestimated in MCSCF. Note that here, contrary to the common case, MCSCF gives higher activation barriers than HF because the ozone reactant has more static correlation than the transition state. The MCDS1H and MC1H approximations give an improvement of about 5 kcal/mol over the KS calculations with the corresponding functionals. The range-separated MC-srPBE method gives activation barriers which are slightly better than the ones given by MCDS1H-PBE and MC1H-PBE, but largely worse than the ones given by MCDS1H-BLYP and MC1H-BLYP. The values obtained with MCDS1H-BLYP and MC1H-BLYP, as well as with MRMP2, are all within 1 kcal/mol of the best estimates.

The reaction energies of the formation of the cycloadducts are overestimated in MCSCF and underestimated in MRMP2 (by about 15 kcal/mol). Zhao et al. observed that even using a large (14,14) active space does not improve the MRMP2 reaction energy [90]. All

FIG. 3: Potential energy curves of Li_2 , C_2 , N_2 , and F_2 calculated with MCSCF and several methods based on the PBE (left) or BLYP (right) exchange-correlation density functionals. For the MCDS1H and MC1H multiconfigurational hybrid approximations, we use a value of $\lambda = 0.25$. For the range-separated MC-srPBE multiconfigurational hybrid approximation, we use a value of the range-separation parameter of $\mu = 0.40 \text{ bohr}^{-1}$. For all multiconfigurational methods, we use a full-valence CAS wave function. The basis set is cc-pVTZ basis. The accurate curves are from Ref. 28.

If we accept to look more closely to the MAE values in spite of the limited statistics, we see that MC1H-BLYP gives overall the smallest MAE with 1.30 kcal/mol. The multiconfigurational hybrid MCDS1H-PBE involving the PBE exchange-correlation functional gives a larger MAE of 3.11 kcal/mol, but turns out to perform better on average than the range-separated MC-srPBE method which gives a MAE of 4.27 kcal/mol. Neglecting density scaling in the correlation functional of multiconfigurational hybrids seems slightly favorable for BLYP and slightly unfavorable for PBE. This is in line with what was found for double-hybrid approximations [79], although the MP2 correlation part made it much more sensitive to the neglect of the density scaling. Even if the effect of neglecting density scaling is systematic in giving more negative complement correlation energies $\vec{E}_c^{\lambda}[n]$ for all density functionals, its effect on the MAE depends more on fortuitous compensation of errors for the approximate functional used.

B. Dissociation of diatomic molecules

We now turn to the calculation of potential energy curves of diatomic molecules. This is a harder problem since static correlation effects are dominant at dissociation. For the multiconfigurational hybrids, we report here only the curves of MCDS1H for PBE and of MC1H for BLYP according to the results of Section IV A, but the differences between the curves of MCDS1H and MC1H for both BLYP and PBE are in fact very small for these diatomic molecules.

Figure 2 shows the potential energy curve of H_2 calculated by hybrid approximations using the PBE and BLYP density functionals. Around the equilibrium internuclear distance, all DFT-based method, including MCDS1H-PBE and MC1H-BLYP, are accurate, which means that they properly describe dynamic correlation. At large distances, the single-hybrid approximations (PBE0, B1LYP, and B3LYP), which include a fraction of HF exchange energy, give less accurate potential energy curves than non-hybrid KS calculations (PBE and BLYP). By inclusion of a fraction of exact static correlation energy, the multiconfigurational hybrids (MCDS1H-PBE, MCsrPBE, and MC1H-BLYP) correct this behavior and give potential energy curves that correctly saturate beyond a distance of about 5 bohr, as the MCSCF curve does. This point is explained by a detailed analysis of the asymptotic expansion of the potential energy curves in a minimal basis in Appendix B. However, the MCDS1H-PBE, MC-srPBE, and MC1H-BLYP methods still display significant errors on the energy of the separated atoms due to the density-functional approximations. Indeed, as in restricted KS calculations, the density functionals used in the multiconfigurational hybrids depend only of the total density and do not give accurate energies in the limit of separated atoms of open-shell character. In an unrestricted KS calculation, the energy at dissociation can be improved by breaking the spin symmetry and therefore introducing a fictitious spin density which helps to describe the separated atoms. In our present implementation of the multiconfigurational hybrids, the spin symmetry is imposed on the MCSCF wave function so that there is no fictitious spin density to be used in the density functionals.

Figure 3 shows the potential energy curves of $Li₂, C₂$, N_2 , and F_2 . The results are similar than for H_2 . Around the equilibrium distance, the MCDS1H-PBE and MC1H-BLYP curves are similar to the standard hybrid or nonhybrid KS calculations. At large distances, the singlehybrid approximations give a spurious increase of the energy, whereas the MCDS1H-PBE and MC1H-BLYP curves correctly saturate. The MC1H-BLYP approximation gives good total energies, but again a significant error remains at dissociation, especially for N_2 . The DS1DH-PBE and MC-srPBE approximation gives curves of very similar shape.

V. CONCLUSIONS

We have presented a multiconfigurational hybrid density-functional theory which rigorously combines MC-SCF and DFT based on a linear decomposition of the electron-electron interaction. It is straightforward extension of the usual hybrid approximations by essentially adding a fraction λ of exact static correlation in addition to the fraction λ of exact exchange. Any existing approximate exchange-correlation density functional can be used in this scheme by using a simple scaling relation with λ . Test calculations on the cycloaddition reactions of ozone with ethylene or acetylene and the dissociation of diatomic molecules with the PBE and BLYP density functionals show that a good value of λ is 0.25, as in the usual hybrid approximations.

Interestingly, the results seem to indicate that the present approach based on a simple linear decomposition of the electron-electron interaction is at least as good as the range-separated multiconfigurational hybrid method of Ref. 70–72 for including static correlation in DFT, at least with the approximate density functionals used here. Of course, with better short-range density-functional approximations (in particular, we do not have a short-range version of the LYP correlation functional for comparison), the conclusion could be different. Also, one should note that hybrid approaches combining perturbation theory with DFT based on a linear decomposition of the interaction [79] do not have the advantages of the rangeseparated hybrid approaches for fast basis-size convergence and explicit inclusion of long-range van der Waals interactions. For MCSCF, however, basis set convergence is not so much an issue.

The present results suggest that the proposed multi-

configurational hybrid approximations can improve over usual DFT approximations for situations with strong static correlation effects. It remains however to assess the performance of this multiconfigurational hybrid method on a larger variety of systems. Future work includes adding the dependence on the spin density in the functionals to be able to properly handle open-shell systems, and possibly other additional variables such as the ontop pair density as an alternative to the spin density for improving the accuracy of closed-shell systems [121].

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Appendix A: Scaling relations for the derivatives of the density-scaled correlation functional

We give the scaling relations for the density-scaled correlation functional $E_c^{\lambda}[n] = \lambda^2 E_c[n_{1/\lambda}]$ and its derivatives in the case of generalized-gradient approximations (GGA). Starting from a standard GGA density functional written as

$$
E_{c,\text{GGA}}[n] = \int e_c(n(\mathbf{r}), |\nabla n(\mathbf{r})|) d\mathbf{r}, \quad (A1)
$$

where $|\nabla n(\mathbf{r})|$ is the norm of the density gradient, the corresponding scaled functional is

$$
E_{c,\text{GGA}}^{\lambda}[n] = \int e_c^{\lambda}(n(\mathbf{r}), |\nabla n(\mathbf{r})|) d\mathbf{r}, \quad (A2)
$$

where the energy density is obtained by scaling relation (see Ref. 79)

$$
e_c^{\lambda}(n(\mathbf{r}), |\nabla n(\mathbf{r})|) = \lambda^5 e_c\left(\frac{n(\mathbf{r})}{\lambda^3}, \frac{|\nabla n(\mathbf{r})|}{\lambda^4}\right). \tag{A3}
$$

The first-order derivatives of the energy density are

$$
\frac{\partial e_c^{\lambda}}{\partial n} (n(\mathbf{r}), |\nabla n(\mathbf{r})|) = \lambda^2 \frac{\partial e_c}{\partial n} \left(\frac{n(\mathbf{r})}{\lambda^3}, \frac{|\nabla n(\mathbf{r})|}{\lambda^4} \right), \tag{A4}
$$

and

$$
\frac{\partial e_c^{\lambda}}{\partial |\nabla n|} (n(\mathbf{r}), |\nabla n(\mathbf{r})|) = \lambda \frac{\partial e_c}{\partial |\nabla n|} \left(\frac{n(\mathbf{r})}{\lambda^3}, \frac{|\nabla n(\mathbf{r})|}{\lambda^4} \right).
$$
\n(A5)

The second-order derivatives are

$$
\frac{\partial^2 e_c^{\lambda}}{\partial n^2} (n(\mathbf{r}), |\nabla n(\mathbf{r})|) = \frac{1}{\lambda} \frac{\partial^2 e_c}{\partial n^2} \left(\frac{n(\mathbf{r})}{\lambda^3}, \frac{|\nabla n(\mathbf{r})|}{\lambda^4} \right), (A6)
$$

$$
\frac{\partial^2 e_c^{\lambda}}{\partial |\nabla n|^2} (n(\mathbf{r}), |\nabla n(\mathbf{r})|) =
$$

$$
\frac{1}{\lambda^3} \frac{\partial^2 e_c}{\partial |\nabla n|^2} \left(\frac{n(\mathbf{r})}{\lambda^3}, \frac{|\nabla n(\mathbf{r})|}{\lambda^4} \right),
$$
 (A7)

and

$$
\frac{\partial^2 e_c^{\lambda}}{\partial n \partial |\nabla n|} (n(\mathbf{r}), |\nabla n(\mathbf{r})|) =
$$

$$
\frac{1}{\lambda^2} \frac{\partial^2 e_c}{\partial n \partial |\nabla n|} \left(\frac{n(\mathbf{r})}{\lambda^3}, \frac{|\nabla n(\mathbf{r})|}{\lambda^4} \right). \tag{A8}
$$

Appendix B: Asymptotic expansion of the potential energy curve of H²

We consider the H_2 molecule in a Slater minimal basis, with a basis function a localized on the left atom and a basis function b localized on the right atom, both basis functions being identical with exponent $\zeta = 1$. In the large internuclear distance R limit, the two molecular orbitals are $1 = (a + b)/\sqrt{2}$ and $2 = (a - b)/\sqrt{2}$. The total restricted Hartree-Fock (RHF) energy writes

$$
E^{\text{RHF}} = 2h_{11} + J_{11} + \frac{1}{R},
$$
 (B1)

where $h_{11} = t_{11} + v_{11}$ is the sum of the kinetic integral $t_{11} = (1|\hat{t}|1)$ and the nuclei-electron integral $v_{11} =$ $(1|\hat{v}_{ne}|1)$, and $J_{11} = (11|11)$ is the Coulomb two-electron integral, and $1/R$ is the nuclear repulsion energy. By expanding the molecular orbital 1 into the localized functions a and b , and using the symmetry between a and b , it is easy to find the large R behavior of all these terms:

$$
t_{11} = (a|\hat{t}|a) + (a|\hat{t}|b) = \frac{1}{2} + O(e^{-R}),
$$
 (B2)

and

$$
v_{11} = (a|\hat{v}_{ne}|a) + (a|\hat{v}_{ne}|b) = -1 - \frac{1}{R} + O(e^{-R}),
$$
 (B3)

and

$$
J_{11} = \frac{(aa|aa)}{2} + \frac{(aa|bb)}{2} + 2(aa|ab) + (ab|ab)
$$

= $\frac{5}{16} + \frac{1}{2R} + O(e^{-R}),$ (B4)

where $O(e^{-R})$ stands for exponentially decaying terms in R. For the values of the integrals, see Ref. 122. Adding all the pieces together, it leads to the following asymptotic expansion of the total RHF energy

$$
E^{\rm RHF} = -\frac{11}{16} - \frac{1}{2R} + O(e^{-R}).
$$
 (B5)

At dissociation, the RHF wave function contains 50% of the incorrect ionic contribution H^+ ... H^- , which is responsible for too high an energy and for the spurious electrostatic attraction term $-1/2R$.

The full configuration interaction (FCI) correlation energy in this basis is found by diagonalizing the 2×2 Hamiltonian matrix, leading to

$$
E_c^{\text{FCI}} = \frac{1}{2} \left(E_2 - E^{\text{RHF}} - \sqrt{(E_2 - E^{\text{RHF}})^2 + 4K_{12}^2} \right),\tag{B6}
$$

where $E_2 = 2h_{22} + J_{22} + 1/R$ is the energy of the doubleexcited determinant, and $K_{12} = (12|12)$ is the exchange two-electron integral. The asymptotic behavior of E_2 is exactly the same as the one of E^{RHF} , so that $E_2 - E^{\text{RHF}}$ vanishes exponentially when $R\rightarrow\infty$ and the asymptotic behavior of $E_{\rm g}^{\rm FCI}$ is determined by K_{12} only: $E_{\rm c}^{\rm FCI}$ = $-K_{12} + O(e^{-R})$. The asymptotic behavior of K_{12} is

$$
K_{12} = \frac{(aa|aa)}{2} - \frac{(aa|bb)}{2} = \frac{5}{16} - \frac{1}{2R} + O(e^{-R}),
$$
\n(B7)

giving for the correlation energy

$$
E_c^{\text{FCI}} = -\frac{5}{16} + \frac{1}{2R} + O(e^{-R}).
$$
 (B8)

Adding the asymptotic expansions of Eqs. (B5) and (B8) gives the asymptotic expansion of the total FCI energy in this basis

$$
E^{\text{FCI}} = -1 + O(e^{-R}),\tag{B9}
$$

which implies that the FCI potential energy curve saturates quickly at large internuclear distance.

In restricted Kohn-Sham (RKS) density-functional theory, the total energy writes

$$
ERKS = 2h11 + 2J11 + Ex + Ec + \frac{1}{R},
$$
 (B10)

where E_x and E_c are the exchange and correlation energies. At large R , it behaves as

$$
ERKS = -\frac{3}{8} + E_x + E_c + O(e^{-R}).
$$
 (B11)

With local or semilocal density-functional approximations, E_x and E_c go exponentially to constants when $R \to \infty$, so that the asymptotic expansion of E^{RKS} does not contain a spurious term in 1/R.

Single-hybrid approximations introduces a fraction λ of RHF exchange which have the following asymptotic expansion

$$
E_x^{\text{RHF}} = -J_{11} = -\frac{5}{16} - \frac{1}{2R} + O(e^{-R}), \quad (B12)
$$

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and therefore introduce a wrong $-\lambda/2R$ term in the total energy,

$$
E^{\text{hybrid}} = -\frac{3}{8} - \frac{5\lambda}{16} + (1 - \lambda)E_x + E_c - \frac{\lambda}{2R} + O(e^{-R}).
$$
\n(B13)

Single-hybrid approximations thus deteriorate the large R behavior of local or semilocal density-functional approximations (see Fig. 2). The multiconfigurational hybrid approximations introduced in this work correct this behavior by adding a fraction of the FCI correlation energy which, in the limit of large R , is just λE_c^{FCI} , the linearity in λ being a signature of static correlation. For example, the MC1H approximation has the following asymptotic expansion

$$
E^{\text{MC1H}} = -\frac{3}{8} - \frac{5\lambda}{8} + (1 - \lambda)E_x + (1 - \lambda^2)E_c + O(e^{-R}),
$$
\n(B14)

with no longer any spurious $1/R$ term, and thus improves the large R behavior (see Fig. 2). It is a typical example where exact exchange and static correlation must be considered together.

For range-separated density-functional theory, the situation is similar. Range-separated single-hybrid approximations [80, 123–126] include some long-range RHF exchange and their asymptotic expansions display a wrong $-1/2R$ term, just as RHF. Their behavior for large R is in fact worse than that of usual single-hybrid approximations since the $-1/2R$ term is not weighted by λ. However, the range-separated CI-srDFT [66, 67] or MCsrDFT [70–72] methods add some exact long-range correlation energy which removes this wrong $-1/2R$ term.

Note that other forms of single-hybrid approximations which do not use RHF exchange at long range [127, 128] allow one to avoid a wrong $-1/2R$ term in the large R limit. Symmetry breaking is another way to avoid a wrong asymptotic $-1/2R$ term since the unrestricted Hartree-Fock exchange energy does not contain such a term.

The fact that local or semilocal approximations for E_x and E_c do not introduce $1/R$ terms is in agreement with the usual conviction that approximate GGA exchange functionals not only represent exchange but also static correlation, while approximate GGA correlation functionals represent dynamic correlation only [4].

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