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Kyoto University
Active-space symmetry-adapted-cluster configuration-interaction and equation-of-motion coupled-cluster methods for high accuracy calculations of potential energy surfaces of radicals

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The electron-attached (EA) and ionized (IP) symmetry-adapted-cluster configuration-interaction (SAC-CI) methods and their equation-of-motion coupled-cluster (EOMCC) analogs provide an elegant framework for studying open-shell systems. As shown in this study, these schemes require the presence of higher-order excitations, such as the four-particle-three-hole (4p-3h) or four-hole–three-particle (4h–3p) terms, in the electron attaching or ionizing operator $R$ in order to produce accurate ground- and excited-state potential energy surfaces of radicals along bond breaking coordinates. The full inclusion of the 4p-3h/4h–3p excitations in the EA/IP SAC-CI and EOMCC methods leads to schemes which are far too expensive for calculations involving larger radicals and realistic basis sets. In order to reduce the large costs of such schemes without sacrificing accuracy, the active-space EA/IP EOMCC methodology [J. R. Gour et al., J. Chem. Phys. 123, 134113 (2005)] is extended to the EA/IP SAC-CI approaches with 4p-3h/4h–3p excitations. The resulting methods, which use a physically motivated set of active orbitals to pick out the most important 3p-2h/3h–2p and 4p-3h/4h–3p excitations, represent practical computational approaches for high-accuracy calculations of potential energy surfaces of radicals. To illustrate the potential offered by the active-space EA/IP SAC-CI approaches with up to 4p-3h/4h–3p excitations, the results of benchmark calculations for the potential energy surfaces of the low-lying doublet states of CH and OH are presented and compared with other SAC-CI and EOMCC methods, and full CI results.

I. INTRODUCTION

Radicals represent an important class of molecular systems due to their high reactivity and significance as reactants, products, and intermediates in elementary chemical processes. It is, therefore, essential to develop practical $ab\ initio$ approaches that would enable one to provide a highly accurate description of ground and excited electronic states of radical and other open-shell species, particularly since it is very difficult to obtain reliable information about the relevant potential energy surfaces by experimental techniques due to the short lifetimes of radical intermediates. Unfortunately, the development of accurate and, at the same time, affordable quantum chemistry methods for the ground- and excited-state potential energy surfaces of radicals poses a challenging problem for the existing $ab\ initio$ methodologies. This, in particular, applies to methods based on the coupled-cluster (CC) theory, including the equation-of-motion CC (EOMCC) approaches and their closely related symmetry-adapted-cluster (SAC)/symmetry-adapted-cluster configuration-interaction (SAC-CI) and response or time-dependent CC (Refs. 22–25) predecessors. Indeed, the low-lying electronic states of radicals often display a manifestly multideterminantal character, which cannot be captured by the basic CC singles and doubles (CCSD) and EOMCC singles and doubles (EOMCCSD) SAC singles
and doubles (SAC-SD) and SAC-CI-SD-R (SAC-CI singles and doubles), and linear response CCSD approximations, exploiting the unrestricted Hartree-Fock (UHF) or restricted open-shell Hartree-Fock (ROHF) references (see, e.g., Refs. 31–35). The basic electron-attached (EA) and ionized (IP) EOMCC methods with up to two-particle-one-hole (2p-1h) and two-hole-one-particle (2h-1p) excitations, which are referred to as the EA-EOMCCSD (Refs. 36 and 37) or EA-EOMCCSD(2p-1h) (Refs. 34 and 35) and IP-EOMCCSD (Refs. 38–41) or IP-EOMCCSD(2h-1p) (Refs. 34 and 35) schemes, and the analogous and historically older EA and IP SAC-CI approximations truncated at the 2p-1h and 2h-1p excitations,17,19,42–44 which belong to the family of the SAC-CI-SD-R approximations (cf., e.g., Ref. 44) and which are referred to here and elsewhere in this article as the SAC-CI(2p-1h) and SAC-CI(2h-1p) schemes, can be useful in calculations of electron affinities and ionization potentials, but they are, in most cases, insufficient for reliable calculations of excited states of radicals (see, e.g., Refs. 34–36 and 45–47). In these methods, one obtains the electronic states of radicals by applying the electron attaching or ionizing operator \( R \) with the 1p and 2p-1h or 1h and 2h-1p components to the ground state of the related closed-shell reference system provided by the CCSD or SAC-SD approaches, and one needs higher-than-2p-1h/2h-1p excitations in \( R \) to model the electronic structure of radical species. Breaking bonds in ground and excited states of radicals is particularly difficult to describe by the basic EOMCC and SAC-CI models.34,35

One might attempt to address the above difficulties by turning to the sophisticated multireference CC (MRCC) methodologies of either the Fock-space/valence-universal or Hilbert-space/state-universal type (cf., e.g., Refs. 51–53 for selected reviews), but none of the existing genuine MRCC methods and computer codes are ready for routine chemical applications. In fact, the use of the true multistate MRCC methods based on complete active spaces may become prohibitively expensive when the radical systems of interest become large and when one has to rely on larger multidimensional reference spaces. Moreover, all genuine MRCC methods exploiting the generalized Bloch equation and the effective Hamiltonian formalism continue to face their own nontrivial challenges, including the need to deal with unphysical multiple solutions,45–51 intruder states,54,55,58 and the existence of the so-called intruder solutions that may tremendously complicate the MRCC calculations and the subsequent analysis of the resulting wave functions and energies.54,56 This is not to say that genuine MRCC methods should not continue to be developed. On the contrary, recent years have witnessed renewed interest and significant progress in the area of valence-universal and state-universal MRCC calculations (cf., e.g., Refs. 53, 54, and 61–72). We are rather reflecting on the need for the development of relatively inexpensive, intuitive, and yet highly accurate CC or response CC, EOMCC, and SAC-CI models that would be free from the mathematical and numerical problems plaguing genuine multistate MRCC methods, that would not exceed the relatively low computer costs of the basic EOMCC and SAC-CI approaches of the CCSD type by a large factor, and that would enable one to avoid the prohibitive costs of the accurate, but very expensive conventional EOMCC and SAC-CI methods with a full treatment of triple and higher-than-triple excitations. Indeed, the complete inclusion of triple or triple and quadruple excitations in the EOMCC theory via the full EOMCC singles, doubles, and triples (EOMCCSDT) (cf., also, Ref. 75) and EOMCC singles, doubles, triples, and quadruples (EOMCCSDTQ) approaches exploiting the UHF or ROHF references greatly improves the results when the excited states of radicals of interest become significantly multideterminantal.24 However, the large computer costs of the EOMCCSDT, EOMCCSDTQ, and the analogous high-order SAC-CI schemes, referred to as the SAC-CI-general-\( R \) methods (cf., e.g., Refs. 42–45), which are characterized by the iterative steps that scale as \( n^4_o \) or \( N^8 \) in the triples case and \( n^2_o \) or \( N^{10} \) in the case of quadruple excitations, where \( n_o \) and \( n_a \) are the numbers of occupied and unoccupied orbitals in a basis set, respectively, and \( N \) is a measure of the system size, limit the applicability of the EOMCC and SAC-CI methods with a full account of triple or triple and quadruple excitations to very small molecular problems. The high cost of the open-shell EOMCCSDT calculations employing the UHF or ROHF references can be considerably reduced by exploiting the recently developed extensions of the iterative CC3 (Refs. 77–80) and noniterative CR-EOMCCSD(T) (Refs. 81–83) approaches to open shells32,33 but there may always be cases in which the CC3, CR-EOMCCSD(T), and similar approximate triples methods do not provide a sufficiently accurate description of the electronic states of radicals of interest. The situations examined in the present article, where one often has to go beyond triple excitations, are examples of such cases. Moreover, the use of the spin-orbital formalism in the existing open-shell implementations of the CC3 and CR-EOMCCSD(T) methods leads to problems such as spin contamination of the resulting electronic states. The same is true when one uses the open-shell EOMCCSDT,31 EOMCCSDTQ,31,76 and other high-order EOMCC schemes76 employing the ROHF or UHF references.

The recently implemented EA-EOMCCSDT (Ref. 84) and IP-EOMCCSDT (Refs. 85 and 86) methods, which incorporate up to 3p-2h and 3h-2p excitations in the linear excitation operator \( R \), and the analogous and historically older EA and IP SAC-CI approaches with up to 3p-2h and 3h-2p excitations in the \( R \) operator,12–44 which belong to the SAC-CI-general-\( R \) (SDT) family,42 and which are referred to here and elsewhere in this article, for consistency reasons, as the SAC-CI(3p-2h) and SAC-CI(3h-2p) approaches, lead to the orthogonally spin-adapted wave functions of radicals and should be accurate enough in at least some cases of radical excited states, but there are two problems with these kinds of methods. The first problem is the computer cost. For example, the EA-EOMCCSDT and IP-EOMCCSDT methods, as implemented in Refs. 84–86, are characterized by the expensive iterative \( n^4_o n^5 \) and \( n^2_o n^4 (N^8 \text{-like}) \) steps in the diagonalization of the similarity-transformed Hamiltonian, combined with even more expensive \( n^3_o n^5 (N^{10} \text{-like}) \) steps of the underlying ground-state CCSDT calculations. Iterative steps of these types (particularly, the \( n^2_o n^5 \) steps of EA-
EOMCCSDT and the $n^1_{p}n^5_{u}$ steps of the underlying CCSDT) are far too expensive for routine chemical applications. The original SAC-CI-general-R(SDT) [i.e., SAC-CI(3p-2h) and SAC-CI(3h-2p)] approaches\textsuperscript{42-44} and the recently developed IP-EOMCCSDT3\textsuperscript{37} IP-CC3\textsuperscript{37} IP-EOMCCSD(3h-2p),\textsuperscript{34,35,88} and EA-EOMCCSD(3p-2h)\textsuperscript{34,35,88} methods eliminate the most expensive $n^3_{p}n^5_{u}$ steps of CCSDT from the calculations, but the iterative $n^3_{p}n^4_{u}$ steps defining the diagonalization of the Hamiltonian in the IP triples case and the iterative $n^5_{p}n^5_{u}$ steps characterizing the diagonalization of the Hamiltonian in the EA triples case remain.

The second, more fundamental, problem, which is particularly relevant to the studies reported in this paper, is that even the full account of 3p-2h and 3h-2p excitations in the EA and IP EOMCC and SAC-CI calculations is, in many cases, insufficient to provide the correct description of ground- and excited-state potential energy surfaces of radical species along the relevant bond breaking coordinates. This has been pointed out in Ref. 35 and the present paper provides a clear demonstration that one has to consider at least the 4p-3h and 4h-3p terms in the electron attaching or ionizing operator $R$ defining the SAC-CI and EOMCC wave function Ansätze to obtain the high-quality potential energy surfaces of radicals, particularly when the excited states of interest gain significant 3p-2h and 3h-2p components, as is often the case when bonds are stretched or broken. As shown in this paper, the EA-EOMCCSD(3p-2h) method\textsuperscript{34,35,88} and its SAC-CI(3p-2h) counterpart\textsuperscript{42-44} which use up to 3p-2h excitations in the electron attaching operator $R$, are accurate enough for the electronic states of radicals dominated by the 1p and 2p-1h components. Similarly, the IP-EOMCCSD(3h-2p) (Refs. 34, 35, and 88) and SAC-CI(3h-2p) (Refs. 42-44) approaches provide a reasonably accurate description of the electronic states dominated by the 1h and 2h-1p components. Unfortunately, the ground- and excited-state wave functions of radicals gain significant 3p-2h and 3h-2p components relative to the corresponding closed-shell reference molecule at larger internuclear separations, which, as shown in this work, cause major problems to the EA-EOMCCSD(3p-2h)/SAC-CI(3p-2h) and IP-EOMCCSD(3h-2p)/SAC-CI(3h-2p) methods.

The fact that one needs the higher-order EA and IP SAC-CI or EOMCC schemes with at least the 4p-3h or 4h-3p components in the electron attaching or ionizing operator $R$ to study ground- and excited-state potential energy surfaces of radicals is actually not a surprise. A radical can be viewed as a system obtained by attaching an electron to or removing an electron from the related closed-shell molecule. Assuming for a moment that a reference closed-shell molecule of interest is a singly bonded species, one needs at least triple (i.e., 3p-3h) excitations to describe bond breaking in it in a quantitative manner (cf., e.g., Refs. 81, 83, and 89-92). The superposition of the 3p-3h excitations in the closed-shell reference molecule with the 1p or 1h excitations that one needs to apply in order to form a radical from it results in the 4p-3h or 4h-3p excitations that should, therefore, be present in the EA or IP SAC-CI and EOMCC calculations to obtain a quantitative description of the ground- and excited-state potential energy surfaces of radical species along the relevant bond breaking coordinates. A similar argument can also be used to explain why it is often enough to use the EA-EOMCCSD(3p-2h)/SAC-CI(3p-2h) and IP-EOMCCSD(3h-2p)/SAC-CI(3h-2p) methods, which ignore the 4p-3h or 4h-3p components, to describe ground and excited states of radicals near the equilibrium geometry (i.e., in the Franck-Condon region).\textsuperscript{34,35,88} At shorter internuclear distances, double (i.e., 2p-2h) excitations are often good enough to describe the closed-shell reference molecule in a reasonable manner. The superposition of the 2p-2h excitations in the closed-shell reference molecule with the 1p or 1h excitations that one needs to apply in order to form a radical species from it results in the need for 3p-2h or 3h-2p excitations in the EA or IP SAC-CI and EOMCC calculations, but higher-order 4p-3h or 4h-3p excitations can often be neglected.

The above discussion, supported by the ample numerical evidence provided in this paper, implies that it is essential to consider the 4p-3h and 4h-3p excitations in the electron attaching and ionizing operators defining the EA and IP SAC-CI and analogous EOMCC schemes in order to obtain the accurate ground- and excited-state potential energy surfaces of radicals. The problem is that the excitation amplitudes defining the 4p-3h and 4h-3p operators, whose numbers scale as $n^3_{p}n^6_{u}$ and $n^3_{u}n^3_{u}$ respectively, are far too numerous for routine calculations for larger radicals and larger basis sets. Moreover, the full treatment of the 4p-3h and 4h-3p excitations in the electron attaching and ionizing operators, as in the recently implemented EA- and IP-EOMCCSDTQ approaches,\textsuperscript{37} leads to schemes which use the prohibitively expensive CPU steps that scale as $n^3_{p}n^6_{u}$ and $n^3_{u}n^3_{u}$ or $N^3$ in the Hamiltonian diagonalization part, and $n^3_{p}n^6_{u}$ ($N^7$) or $n^3_{u}n^3_{u}$ ($N^{10}$) in the part that deals with the underlying ground-state CCSDT or CCSDTQ calculations for the reference closed-shell system (although this is not absolutely necessary,\textsuperscript{10} particularly when radicals are examined)\textsuperscript{34,35} the authors of Ref. 47 chose the more expensive CCSDTQ option in the underlying CC calculations for the closed-shell reference system that precede the EA- and IP-EOMCCSDTQ calculations). The complete treatment of all many-body components of the electron attaching and ionizing operators $R$ that define the SAC-CI-general-R methods with up to 4p-3h and 4h-3p excitations\textsuperscript{34,44} the EA and IP SAC-CI-general-R (SDTQ) schemes in Ref. 44 and the SAC-CI(4p-3h) and SAC-CI(4h-3p) approaches in this work] leads to similar computer costs. Clearly, methods of this kind can only be used in benchmark or small-molecule calculations. In fact, as mentioned earlier, a full treatment of the less demanding 3p-2h and 3h-2p components of the electron attaching and ionizing operators $R$ of the EA and IP SAC-CI and EOMCC methods leads to schemes that are already very expensive. It is important to develop the EA and IP SAC-CI and EOMCC methods that can reduce the large costs of handling the 3p-2h/3h-2p and 4p-3h/4h-3p terms without a substantial loss of accuracy.

The need for an approximate treatment of the 3p-2h/3h-2p and 4p-3h/4h-3p excitations has already been pointed out in the original papers on the EA and IP SAC-CI-general-R methods\textsuperscript{34,44} that precede the initial papers on the
EA and IP EOMCC approaches,\textsuperscript{36–41} which ignore these terms altogether. This has prompted the development of the highly efficient EA and IP SAC-CI methods with the perturbative selection (PS) of the 3p-2h/3h-2p, 4p-3h/4h-3p, and other higher-order excitations, which also neglect or approximate higher-order nonlinear terms in the cluster operator defining the ground electronic state of the reference closed-shell molecule\textsuperscript{42,44} (see Ref. 93 for the original idea of the SAC-CI approaches with the PS, abbreviated here and elsewhere in this article as the SAC-CI/PS schemes). There are several advantages of the SAC-CI/PS methods, the most important one being the fact that in these approaches we do not have to limit ourselves to a particular excitation order and, at least in principle, can select the 3p-2h/3h-2p, 4p-3h/4h-3p, and other appropriate higher-order excitations purely numerically, based on their significance in the perturbation theory analysis rather than on their excitation rank. As a result, the SAC-CI/PS approaches can be successfully applied to a wide range of molecular problems, including large systems such as biological molecules, without losing much accuracy. There also are disadvantages. The perturbative selection of the 3p-2h/3h-2p, 4p-3h/4h-3p, and other higher-order excitations may lead to substantial numerical noise if we try to apply the SAC-CI/PS methods to potential energy surfaces, since different sets of higher-order excitations will be selected at different geometries. This problem can be alleviated by merging the 3p-2h/3h-2p, 4p-3h/4h-3p, and other higher-order excitations selected in different regions of potential energy surfaces. In this regard, the Group SUM method,\textsuperscript{90} which gives a common set of the suitably selected excitation operators, is particularly useful, producing smooth potential energy surfaces within the SAC-CI/PS framework. However, there may be cases where it is not immediately obvious which regions of potential energy surfaces should initially be probed to obtain the adequate selection of the 3p-2h/3h-2p, 4p-3h/4h-3p, and other higher-order excitations that would work in all regions. In fact, the same remarks apply to other methods based on numerical selections of higher-order excitations, including, for example, the popular multireference CI approach with a selection of single and double excitations from a multireferential reference wave function of Refs. 95–97. It is also quite difficult to combine the PS scheme with the EA, IP, and other EOMCC methods, which traditionally rely on the many-body diagrammatic formulation and factorization of nonlinear terms that enter the EOMCC wave functions and the similarity-transformed Hamiltonian of the underlying CC theory.

The above discussion implies that it is important to consider alternative formulations of the EA and IP SAC-CI and EOMCC methods with 3p-2h/3h-2p and 4p-3h/4h-3p excitations, in which selection of the dominant 3p-2h/3h-2p and 4p-3h/4h-3p terms is done mathematically rather than via numerical thresholds. An idea, which can help us to design the relatively inexpensive EA and IP SAC-CI and EOMCC methods that could provide highly accurate results for ground- and excited-state potential energy surfaces of radicals, is that of the active-space CC or single-reference-like, state-selective MRCC approaches pioneered by Adamowicz, Piecuch, and co-workers\textsuperscript{89–92,98–106} and active-space EOMCC theories introduced and fully developed by Kowalski and Piecuch.\textsuperscript{73,74,107} In these methods and their various subsequent implementations, by Piecuch and co-workers and Adamowicz and co-workers (cf., e.g., Refs. 108–112) and others (cf., e.g., Refs. 76 and 113–118), one combines the single-reference CC/EOMCC formalism with a multireference concept of active orbitals, which are used to select a relatively small subset of the dominant triply and other higher-than-doubly excited clusters that reflect the nature of the electronic quasidegeneracy or excited states of interest, reducing the computer costs of the parent CC/EOMCC calculations with a full treatment of higher-than-double excitations by a large factor. In particular, Gour et al. have recently proposed the active-space EA and IP EOMCC methods in which one uses small sets of active orbitals to select the most important 3p-2h/3h-2p, 4p-3h/4h-3p, and other higher-order contributions to the electron attaching and ionizing operators $R$ that define the EA and IP EOMCC theories and their multiply attached and ionized variants.\textsuperscript{34,35,88} Just like the parent EA and IP EOMCC approaches with the full treatment of the 3p-2h/3h-2p, 4p-3h/4h-3p, and other higher components of $R$, the active-space EA and IP EOMCC methods and their multiply attached and ionized analogs discussed in Ref. 34 provide an orthogonally spin-adapted description of ground and excited states of radical and other open-shell species. Among the active-space EA and IP EOMCC approaches discussed in Refs. 34, 35, and 88 are the EA-EOMCCSDtq and IP-EOMCCSDtq approximations, in which the dominant 3p-2h and 3h-2p components of the electron attaching and ionizing operators $R$ defining the aforementioned EA-EOMCCSD(3p-2h) and IP-EOMCCSD(3h-2p) schemes are selected via active orbitals. The EA-EOMCCSDtq/IP-EOMCCSDtq methods based on diagonalizing the similarity-transformed Hamiltonian of CCSD in the space of all $1p/1h$ and $2p−1h/2h−1p$ excitations and a small subset of $3p-2h/3h-2p$ excitations defined through active orbitals have been tested in Refs. 34, 35, and 88, demonstrating a considerable promise in applications to excitation energies of radicals calculated in the vicinity of the equilibrium geometry. As shown in Refs. 34, 35, and 88, the EA-EOMCCSDtq/IP-EOMCCSDtq approaches give excellent vertical and adiabatic excitation energies of radical species at a small fraction of the cost associated with the parent EA-EOMCCSD(3p-2h) and IP-EOMCCSD(3h-2p) calculations, which use all 3p-2h and 3h-2p excitations. However, as indicated in Ref. 35, the EA-EOMCCSDtq/IP-EOMCCSDtq methods are not sufficiently accurate to describe potential energy surfaces along bond breaking coordinates, since ground and excited states of radicals often gain significant 3p-2h and 3h-2p components when larger internuclear separations are examined and these, as already pointed out above, cannot be well described by the EA-EOMCCSD(3p-2h) and IP-EOMCCSD(3h-2p) levels of theory. This prompts the need for the implementation of the active-space EA-EOMCCSDtq and IP-EOMCCSDtq methods, as defined in Ref. 34, and their SAC-CI counterparts, in which one uses active orbitals to select the most important 4p-3h and 4h-3p excitations, in addition to the 3p-2h and 3h-2p excitations present in the EA-EOMCCSDtq and IP-EOMCCSDtq methods.
The basic idea of EA-EOMCCSDtq, IP-EOMCCSDtq, and the analogous active-space SAC-CI methods developed and tested in this work is to reduce the enormous computer costs of the EA and IP EOMCC and SAC-CI calculations with the full treatment of $4p$-$3h$ and $4h$-$3p$ excitations by systematically selecting the dominant $4p$-$3h$ and $4h$-$3p$ components and their $3p$-$2h$ and $3h$-$2p$ counterparts via a small subset of active orbitals.

In this paper, we focus on the development, implementation, and testing of the active-space SAC-CI methods with the $3p$-$2h$/$3h$-$2p$ and $4p$-$3h$/$4h$-$3p$ terms. SAC-CI offers an advantage over the current implementations of the EOMCC theory in that it is easier to impose the active-space logic on the $3p$-$2h$/$3h$-$2p$ and $4p$-$3h$/$4h$-$3p$ components within the language of electron configurations exploited in SAC-CI, as compared with the diagrammatic many-body language used in EOMCC. We will eventually pursue the more complete active-space EA-EOMCCSDtq and IP-EOMCCSDtq methods suggested in Ref. 34, which use the CCSDt (Refs. 73, 74, 89–92, and 98–107) or CCSDt0 (Refs. 90, 99–102, and 104) wave functions to describe the ground state of the reference closed-shell system and which do not neglect any nonlinear terms in the cluster operator, but it is important to investigate first what are the potential benefits of incorporating the high-order $4p$-$3h$ and $4h$-$3p$ terms via active orbitals and what types of improvements in the full and active-space EA-EOMCCSD$(3p$-$2h$)/SAC-CI$(3p$-$2h$) and IP-EOMCCSD$(3h$-$2p$)/SAC-CI$(3h$-$2p$) results can be obtained when the ground- and excited-state potential energy surfaces of radicals along bond breaking coordinates are examined with the active-space methods truncated at the $4p$-$3h$ and $4h$-$3p$ excitations. The SAC-CI methodology is very useful in this regard.

The primary objectives of the present work are the following: (i) to show that one needs the genuine $4p$-$3h$ and $4h$-$3p$ excitations in the electron attaching and ionizing operators $R$ in order to obtain an accurate representation of the ground- and excited-state potential energy surfaces of radicals along the relevant bond breaking coordinates, demonstrating that the $3p$-$2h$ and $3h$-$2p$ components (and, of course, the lower-order $1p$ and $2p$-$1h$ or $2h$ and $2h$-$1p$ terms) are generally insufficient; (ii) to demonstrate that one can obtain accurate ground- and excited-state potential energy surfaces, which can compete with the results of SAC-CI calculations with a full treatment of $3p$-$2h$/$3h$-$2p$ and $4p$-$3h$/$4h$-$3p$ components and be close to the exact, full CI potentials, using small subsets of active orbitals to select the dominant $3p$-$2h$/$3h$-$2p$ and $4p$-$3h$/$4h$-$3p$ excitations; and (iii) to show that the active-space SAC-CI methods with the $3p$-$2h$/$3h$-$2p$ and $4p$-$3h$/$4h$-$3p$ excitations selected mathematically by redefining the relevant $R$ operators are at least as effective as the SAC-CI/PS approaches, in which the $4p$-$3h$/$4h$-$3p$ components are selected numerically based on perturbative arguments and thresholds for neglecting small contributions. We also demonstrate that it is not sufficient to use the CI methods with up to $4p$-$3h$ and $4h$-$3p$ components to obtain accurate ground- and excited-state potential energy surfaces of the CH and OH radicals. One needs to apply the electron attaching and ionizing operators $R$ to the correlated CC or SAC ground state to obtain high-quality potential energy surfaces of radical species along the relevant bond breaking coordinates. The relevant test calculations are reported for the ground- and excited-state potential energy curves representing a few low-lying doublet states of the CH (the EA case) and OH (the IP case) radicals, which are small enough to allow for the exact, full CI, calculations. The results reported in this article are encouraging enough to pursue the implementation of the active-space EA-EOMCCSDtq and IP-EOMCCSDtq methods of Ref. 34 in the future, since one can regard the EA-EOMCCSDtq and IP-EOMCCSDtq approaches as nothing else than the EA and IP EOMCC analogs of the active-space SAC-CI$(4p$-$3h$) and SAC-CI$(4h$-$3p$) methods considered here. This statement reflects on our belief that the historically older SAC-CI methodology and the more recent EOMCC formalism, being so closely related to each other, may both benefit from being developed side by side. The present study may serve as an illustration of how much one can learn by working with both methodologies at the same time. Since there have been no prior studies that use the SAC-CI and EOMCC methods together, we use this paper as an opportunity to emphasize that up to rather unimportant details, which can be dealt with if necessary, the EA and IP SAC-CI methods and their EOMCC counterparts represent essentially the same methodology.

II. THEORY

The active-space EA and IP SAC-CI and EOMCC methods are based on the idea of selecting the most important $3p$-$2h$/$3h$-$2p$, $4p$-$3h$/$4h$-$3p$, and other higher-order excitations in the corresponding full schemes. Thus, we begin this section with a review of the basic elements of the EA and IP SAC-CI and EOMCC methodologies. This enables us to make an important point that there are many similarities between the EA and IP SAC-CI methods and their EOMCC analogs, which have not always been appreciated in the literature and which may benefit the development of both methodologies, as is the case in this work.

A. An overview of the EA and IP SAC-CI and EOMCC methods

The EA and IP SAC-CI and EOMCC methods use the following wave function Ansatz to represent the electronic states of the $(N+1)$- or $(N-1)$-electron system:

$$\Psi_{(N+1)} = R^{(N+1)}_{\mu} \Psi_0,$$

where

$$\Psi_0 = e^S\Phi$$

is the SAC (in the SAC-CI case) or CC (in the EOMCC case) ground state of an $N$-electron closed-shell system, and $R^{(N+1)}_{\mu}$ and $R^{(N-1)}_{\mu}$ are the electron attaching and ionizing operators that generate the $(N+1)$- or $(N-1)$-electron states from the $N$-electron SAC/CC wave function $\Psi_0$. Here, $|\Phi\rangle$ is a closed-shell, $N$-electron reference determinant [in the applications presented in this work, the restricted Hartree-Fock (RHF) configuration] and $S$ is the cluster operator,
S = \sum_{n=1}^{M_S} S_n = \left( \frac{1}{n!} \right)^2 s_{a_{i_1}...a_{i_n}}^{\dagger} a_{a_{i_1}...a_{i_n}}^\dagger \cdots a_{a_{i_n}}^\dagger a_{i_1} ... a_{i_n}, \tag{3}

with \( s_{a_{i_1}...a_{i_n}}^{\dagger} \) representing the corresponding cluster amplitudes. We use the usual notation where \( i, j, ... \) or \( i_1, i_2, ... \) \((a, b, ... \text{ or } a_1, a_2, ...)\) are the spin-orbitals occupied (unoccupied) in the reference determinant \( |\Phi\rangle \) and \( a^\dagger(a) \) are the creation (annihilation) operators associated with the spin-orbital basis set \(|p\rangle\). Whenever appropriate, we also use the Einstein summation convention over the repeated upper and lower indices. In general, the excitation level \( M_S \) in Eq. (3) satisfies \( M_S \leq N \). In the basic SAC-SD and CCSD calculations, which provide the ground-state wave function of the reference \( N \)-electron closed-shell system for all of the EA and IP SAC-CI and EOMCC calculations reported in this work, \( M_S = 2 \).

The electron attaching and ionizing operators, \( R_{\mu}^{(N+1)} \) and \( R_{\mu}^{(N-1)} \), respectively, entering Eq. (1), are defined as

\[
R_{\mu}^{(N+1)} = \sum_{n=0}^{M_R} R_{\mu,(n+1)p-nh}
\]

and

\[
R_{\mu}^{(N-1)} = \sum_{n=0}^{M_R} R_{\mu,(n+1)h-np},
\]

where

\[
R_{\mu,(n+1)p-nh} = \frac{1}{n!(n+1)!} s_{a_{i_1}...a_{i_n}}^{\dagger} a_{a_{i_1}...a_{i_n}}^\dagger \cdots a_{a_{i_n}}^\dagger a_{i_1} ... a_{i_n},
\]

and

\[
R_{\mu,(n+1)h-np} = \frac{1}{n!(n+1)!} s_{a_{i_1}...a_{i_n}}^{\dagger} a_{a_{i_1}...a_{i_n}}^\dagger \cdots a_{a_{i_n}}^\dagger a_{i_1} ... a_{i_n},
\]

with \( M_R = N \) in the exact theory and \( M_R < N \) in the approximate approaches. In this paper, we use the following EA and IP SAC-CI and EOMCC truncation schemes:

(i) EA-EOMCCSD(2p-1h) and IP-EOMCCSD(2h-1p).

These are the basic EA and IP EOMCC approaches, in which \( M_S = 2 \) and \( M_R = 1 \). The EA-EOMCCSD(2p-1h) and IP-EOMCCSD(2h-1p) methods are also known in the literature as the EA-EOMCCSD and IP-EOMCCSD approaches.\(^{36-41}\) They use the CCSD approximation for \( S \) and the \( R_{\mu}^{(N+1)} \) and \( R_{\mu}^{(N-1)} \) operators are truncated at the 2p-1h and 2h-1p components, so that

\[
R_{\mu}^{(N+1)} = R_{\mu,1p} + R_{\mu,2p-1h} = r_{1p} a_{1} + \frac{1}{2} r_{1p} a_{1}^\dagger a_{1}^\dagger a_{1},
\]

and

\[
R_{\mu}^{(N-1)} = R_{\mu,1h} + R_{\mu,2h-1p} = r_{1h} a_{1} + \frac{1}{2} r_{1h} a_{1}^\dagger a_{1}^\dagger a_{1},
\]

respectively. The EA-EOMCCSD(2p-1h) and IP-EOMCCSD(2h-1p) methods are the EOMCC analogs of the SAC-CI(2p-1h) and SAC-CI(2h-1p) schemes, also known as the EA and IP SAC-CI-SD-R methods.\(^{42}\) We do not report the SAC-CI(2p-1h) and SAC-CI(2h-1p) numerical results in this work, since they are very similar to those obtained with the EA-EOMCCSD(2p-1h) and IP-EOMCCSD(2h-1p) approaches. Moreover, neither SAC-CI nor EOMCC schemes truncated at the 2p-1h and 2h-1p excitations provide accurate potential energy surfaces of radical species. The EA-EOMCCSD(2p-1h) and IP-EOMCCSD(2h-1p) results are sufficient to illustrate this point.

(ii) EA-EOMCCSD(3p-2h), IP-EOMCCSD(3h-2p), SAC-CI(3p-2h), and SAC-CI(3h-2p). In these schemes, \( M_S = 2 \) and \( M_R = 1 \). In other words, we continue to use the CCSD (the EOMCC case) or SACSD (the SAC-CI case) ground state of an \( N \)-electron closed-shell system as a correlated reference state \(|\Psi_0\rangle\), while truncating the \( R_{\mu}^{(N+1)} \) and \( R_{\mu}^{(N-1)} \) operators at the 3p-2h and 3h-2p components to obtain

\[
R_{\mu}^{(N+1)} = R_{\mu,1p} + R_{\mu,2p-1h} + R_{\mu,3p-2h}
\]

and

\[
R_{\mu}^{(N-1)} = R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p},
\]

respectively, where the \( R_{\mu,1p} \), \( R_{\mu,2p-1h} \), \( R_{\mu,1h} \), and \( R_{\mu,2h-1p} \) components are defined in the same manner as in Eqs. (8) and (9) and \( R_{\mu,3p-2h} \) and \( R_{\mu,3h-2p} \) are given by

\[
R_{\mu,3p-2h} = \frac{1}{12} i a_{i}^\dagger a_{j}^\dagger a_{1}^\dagger a_{1}^\dagger a_{1} a_{1},
\]

and

\[
R_{\mu,3h-2p} = \frac{1}{12} i a_{i}^\dagger a_{j}^\dagger a_{1}^\dagger a_{1}^\dagger a_{1} a_{1},
\]

respectively. We use the EA-EOMCCSD(3p-2h) and IP-EOMCCSD(3h-2p) methods and the analogous SAC-CI(3p-2h) and SAC-CI(3h-2p) schemes to show that the more complete treatment of the higher-order nonlinear terms in \( S \) in the EA-EOMCCSD(3p-2h) and IP-EOMCCSD(3h-2p) approaches is not sufficient to eliminate large errors in the SAC-CI(3p-2h) and SAC-CI(3h-2p) results at larger internuclear separations. One needs the genuine 4p-3h and 4h-3p excitations in \( R_{\mu}^{(N+1)} \) and \( R_{\mu}^{(N-1)} \), respectively, to improve the EA-EOMCCSD(3p-2h)/SAC-CI(3p-2h) and IP-EOMCCSD(3h-2p)/SAC-CI(3h-2p) results at significantly stretched geometries. The active-space variants of the EA-EOMCCSD(3p-2h)/SAC-CI(3p-2h) and IP-EOMCCSD(3h-2p)/SAC-CI(3h-2p) methods are described in Sec. II B.

(iii) SAC-CI(4p-3h) and SAC-CI(4h-3p). These are the key truncation schemes for the considerations reported in this paper. In these approximation, \( M_S = 2 \) and \( M_R = 1 \). Thus, we use the SAC-SD ground state of an \( N \)-electron closed-shell system as the correlated reference state \(|\Psi_0\rangle\), while truncating the \( R_{\mu}^{(N+1)} \) and \( R_{\mu}^{(N-1)} \) operators at the 4p-3h and 4h-3p components to obtain

\[
R_{\mu}^{(N+1)} = R_{\mu,1p} + R_{\mu,2p-1h} + R_{\mu,3p-2h} + R_{\mu,4p-3h}
\]

and

\[
R_{\mu}^{(N-1)} = R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p} + R_{\mu,4h-3p},
\]

respectively, where the \( R_{\mu,1p} \), \( R_{\mu,2p-1h} \), \( R_{\mu,1h} \), \( R_{\mu,2h-1p} \), \( R_{\mu,3p-2h} \) and \( R_{\mu,3h-2p} \) components are the same as in Eqs. (8)–(11), and \( R_{\mu,4p-3h} \) and \( R_{\mu,4h-3p} \) are defined as

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\[ R_{\mu,4p-3h} = \frac{1}{144} j_{\mu bc} d^a d^b a_i a_j \] (16)

and

\[ R_{\mu,4h-3p} = \frac{1}{144} j_{\mu bc} d^a d^b a_i a_j a_p a_q \] (17)

respectively. One could define the EA-EOMCCSD(4p-3h) and IP-EOMCCSD(4h-3p) truncation schemes in a similar manner. Although the EA-EOMCCSD(4p-3h) and IP-EOMCCSD(4h-3p) approaches would offer a complete treatment of the nonlinear terms in \( S_1 \) and \( S_2 \) resulting from the use of the CC Ansatz for the \( N \)-electron ground state \(| \Psi_0 \rangle \) [Eq. (2)], we focus on the corresponding and simpler SAC-CI(4p-3h) and SAC-CI(4h-3p) methods and their relatively inexpensive active-space variants, which are sufficient to eliminate the large errors in describing the ground- and excited-state potential energy curves of the \( \text{CH} \) and \( \text{OH} \) radicals at larger internuclear separations produced by the EA-EOMCCSD(3p-2h)/SAC-CI(3p-2h) and IP-EOMCCSD(3h-2p)/SAC-CI(3h-2p) schemes. The active-space SAC-CI(4p-3h) and SAC-CI(4h-3p) approaches are described in Sec. II B.

In addition to the above EA and IP EOMCC and SAC-CI methods, we use the CI(4p-3h) and CI(4h-3p) approaches, in which one obtains the electronic wave functions \(| \Psi_{\text{CI}(N)} \rangle \) by applying the electron attaching and ionizing operators \( R_{\mu,4p-3h}^{(N)} \) and \( R_{\mu,4h-3p}^{(N)} \) with up to 4p-3h and 4h-3p excitations, defined by Eqs. (14) and (15), respectively, directly to the \( N \)-electron reference determinant \(| \Phi \rangle \) rather than to the correlated CC or SAC ground state \(| \Psi_0 \rangle \) [Eq. (2)]. The purpose of comparing the CI(4p-3h)/CI(4h-3p) results with the corresponding results of the SAC-CI(4p-3h)/SAC-CI(4h-3p) calculations is to show that the mere inclusion of the 4p-3h and 4h-3p excitations in \( R_{\mu,4p-3h}^{(N)} \) and \( R_{\mu,4h-3p}^{(N)} \), respectively, is not sufficient to obtain the accurate potential energy curves of the \( \text{CH} \) and \( \text{OH} \) radicals. One has to apply the \( R_{\mu,4p-3h}^{(N)} \) and \( R_{\mu,4h-3p}^{(N)} \) operators with a full or active-space treatment of the 4p-3h and 4h-3p components to the correlated ground state of the \( N \)-electron reference system.

Before discussing the active-space EA and IP SAC-CI and EOMCC methods developed and/or tested in this paper, let us emphasize that although there are technical differences between the SAC-CI and EOMCC approaches, which are related to different algorithms that are used to determine the \( S \) and \( R \) operators and the additional approximations in the SAC-CI models that are not exploited in the standard implementations of the EOMCC theories, the similarities between the SAC-CI and EOMCC methodologies are so great that the main conclusions drawn from the SAC-CI calculations apply to the EOMCC calculations and vice versa. Indeed, in the EA and IP SAC-CI approaches which, historically speaking, precede the analogous EA and IP EOMCC methods by a decade or so, one determines the cluster operator \( S \) by solving the system of equations

\[ \langle \Phi | (H - E_0^{(N)}) e^S | \Phi \rangle = 0, \]

\[ \langle \Phi | e^{S_i} | e^{S_j} | \Phi \rangle = 0, \]

\[ i_1 < \cdots < i_n, \quad a_1 < \cdots < a_n, \quad n = 1, \ldots, M_S, \] (19)

where \( H \) is the electronic Hamiltonian, \( E_0^{(N)} \) is the ground-state energy of the \( N \)-electron closed-shell system, and \( | \Phi^{a_1 \cdots a_n} \rangle = e^{S_i} | a_1 \cdots a_n \rangle | \Phi \rangle \) are the \( n \)-tuple excited determinants relative to \(| \Phi \rangle \) [in practical implementations of SAC and SAC-CI, the spin- and symmetry-adapted configuration state functions (CSFs) corresponding to determinants \(| \Phi^{a_1 \cdots a_n} \rangle \)]. Equations (18) and (19) are obtained by left-projecting the electronic Schrödinger equation for the CC/ SAC wave function \(| \Psi_0 \rangle \) [Eq. (2)] on the reference determinant \(| \Phi \rangle \) and the excited determinants \(| \Phi^{a_1 \cdots a_n} \rangle \) corresponding to the excitations included in \( S \). If we do not neglect any nonlinear terms in \( S \), the SAC equations [Eqs. (18) and (19)] become equivalent to the standard CC equations of Čížek,

\[ E_0^{(N)} = \langle \Phi | H | \Phi \rangle, \]

\[ \langle \Phi^{a_1 \cdots a_n} | H | \Phi \rangle = 0, \]

\[ i_1 < \cdots < i_n, \quad a_1 < \cdots < a_n, \quad n = 1, \ldots, M_S, \] (21)

where

\[ \bar{H} = e^{-S} H e^S = (He^S)_C \] (22)

is the similarity-transformed Hamiltonian of the CC theory and subscript \( C \) refers to the connected part of a given operator expression. This is because the energy-dependent terms in Eq. (19) cancel out the disconnected terms contributing to the product of \( H \) and \( e^S \), leaving us with the energy-independent system of nonlinear CC equations defined by Eq. (21). In particular, the full SAC-SD approximation, in which all nonlinear terms in \( S_1 \) and \( S_2 \) are retained, is equivalent to the standard CCSD approximation. Although full SAC-SD calculations are, in principle, possible using the routines that form part of \textsc{gaussian 03},119 the present computational algorithm used to determine all nonlinear terms of the full SAC-SD approach is not particularly efficient; so in the conventional SAC-SD calculations, which are used in this work to generate the \( S_1 \) and \( S_2 \) clusters for the subsequent SAC-CI(2p-1h), SAC-CI(2h-1p), SAC-CI(3p-2h), SAC-CI(3h-2p), SAC-CI(4p-3h), and SAC-CI(4h-3p) calculations, all nonlinear terms in Eqs. (18) and (19) other than \( \frac{1}{2} S_2^2 \) are neglected. Thus, the SAC-SD and CCSD results differ. However, since the nonlinear terms neglected in SAC-SD calculations are usually rather unimportant, the differences between the SAC-SD and CCSD results are often small. They may become larger when the \( S_1 \) and \( S_2 \) clusters become large, as is the case at stretched nuclear geometries, but in those cases neither CCSD nor SAC-SD are accurate. This is shown in Sec. III, where we compare the SAC-SD and CCSD potential energy curves of the \( \text{CH}^+ \) and \( \text{OH}^- \) ions, which serve as reference closed-shell systems for the EA and IP SAC-CI and EOMCC calculations for the \( \text{CH} \) and \( \text{OH} \)
The fact that the cluster operator $S$ of SAC is traditionally defined via the spin- and symmetry-adapted excited CSFs and that SAC equations are programed and solved in a CI-like fashion, using the energy-dependent equations [Eqs. (18) and (19)] rather than the energy-independent equations defining all standard CC methods [Eq. (21)], which are usually derived using diagrammatic techniques, does not play any significant role in discussing the differences between the SAC-SD and CCSD results, particularly when the SAC-SD and CCSD approaches are applied to closed-shell systems and spin- and symmetry-adapted RHF reference determinants, as is the case in this work.

The above similarities between SAC/SAC-CI and CC/EOMCC methods persist when we analyze the corresponding equations defining the EA and IP formalisms. Indeed, once $S$ is determined by solving Eqs. (18) and (19), the amplitudes $r_{a}$ and $r_{a}^{(N-1)}$ defining the EA SAC-CI wave functions $|\Psi^{N-1}\rangle$ or the amplitudes $r_{a}$ and $r_{a}^{(N-1)}$ defining the IP SAC-CI wave functions $|\Psi^{N-1}\rangle (n=1, \ldots, M_R)$ are obtained by solving the eigenvalue problem

$$HR^{(N_{1})}_{\mu}|\phi\rangle = E^{(N_{1})}_{\mu}R^{(N_{1})}_{\mu}|\phi\rangle,$$

in the relevant subspace of $\mathcal{H}^{(N_{1})}$ (the EA case) or $\mathcal{H}^{(N-1)}$ (the IP case). Here and elsewhere in this paper, $\mathcal{H}^{(N_{1})}$ and $\mathcal{H}^{(N-1)}$ designate the appropriate $(N+1)$- and $(N-1)$-electron subspaces of the Fock space. Thus, the subspace of $\mathcal{H}^{(N_{1})}$ used to solve the EA SAC-CI eigenvalue problem [the $(N+1)$-electron variant of Eq. (23)] is spanned by the $|\Phi\rangle = |a_{1}\rangle|\Phi\rangle$ and $|\Phi^{a_{1}\cdots a_{n}}\rangle = |a_{1}\cdots a_{n}\rangle|\Phi\rangle (n=1, \ldots, M_R)$ determinants, while the subspace of $\mathcal{H}^{(N-1)}$ used to solve the IP SAC-CI eigenvalue problem [the $(N-1)$-electron variant of Eq. (23)] is spanned by the $|\Phi\rangle = |a_{1}\rangle|\Phi\rangle$ and $|\Phi^{a_{1}\cdots a_{n}}\rangle = |a_{1}\cdots a_{n}\rangle|\Phi\rangle (n=1, \ldots, M_R)$ determinants [in analogy to the ground-state SAC calculations for the $N$-electron closed-shell reference system, one uses the spin- and symmetry-adapted CSFs corresponding to $|\Phi\rangle$ and $|\Phi^{a_{1}\cdots a_{n}}\rangle$ in the EA case or $|\Phi\rangle$ and $|\Phi^{a_{1}\cdots a_{n}}\rangle$ in the IP case when solving Eq. (23) in the $\mathcal{H}^{(N_{1})}$ or $\mathcal{H}^{(N-1)}$ subspace]. Again, if we do not neglect any nonlinear terms in $S$, the EA/IP SAC-CI equations [Eqs. (18), (19), and (23)] are equivalent to the corresponding EA/IP EOMCC equations, which are defined by Eqs. (20) and (21), and the non-Hermitian eigenvalue problem

$$HR^{(N_{1})}_{\mu}|\phi\rangle = E^{(N_{1})}_{\mu}R^{(N_{1})}_{\mu}|\phi\rangle,$$

in which we diagonalize the similarity-transformed Hamiltonian $\tilde{H}$ [Eq. (22)] in the relevant $(N+1)$- or $(N-1)$-electron subspace, $\mathcal{H}^{(N_{1})}$ or $\mathcal{H}^{(N-1)}$, respectively, as defined above, to obtain the $r_{a}$ and $r_{a}^{(N-1)}$ amplitudes $R^{(N_{1})}_{\mu}$ in the EA case or the $r_{a}$ and $r_{a}^{(N-1)}$ amplitudes $R^{(N-1)}_{\mu}$ in the IP case, where $n=1, \ldots, M_R$.

It is only when we start neglecting nonlinear terms in $S$ in the EA and IP SAC-CI eigenvalue problem [Eq. (23)] that the mathematical equivalence between the EA/IP SAC-CI and EOMCC equations is lost. Moreover, the ad hoc truncations in the many-body expansions defining $S$ and $R^{(N_{1})}_{\mu}$ may affect the size intensity of results (discussed in detail, in the EOMCC context, in Refs. 30 and 120; cf. also Refs. 33, 116, and 121–123) by introducing the disconnected components of the operator product $HH_{\mu}^{(N_{1})}$ into Eq. (24) that do not cancel out if $M_R > M_S$. Indeed, as shown in Ref. 10, the eigenvalue problem defined by Eq. (24), understood as a diagonalization of $\tilde{H}$ [Eq. (22)] with $S$ truncated at the $M_S$-body clusters as in Eq. (3), in the $\mathcal{H}^{(N_{1})}$ subspace spanned by $|\Phi\rangle$ and $|\Phi^{a_{1}\cdots a_{n}}\rangle$ or in the $\mathcal{H}^{(N-1)}$ subspace spanned by $|\Phi\rangle$ and $|\Phi^{a_{1}\cdots a_{n}}\rangle (n=1, \ldots, M_R)$, becomes equivalent to the explicitly connected eigenvalue problem

$$\langle \Phi^{(N_{1})} \rangle |\phi\rangle = \omega^{(N_{1})}_{\mu}R^{(N_{1})}_{\mu}|\phi\rangle,$$

only when $M_R \leq M_S$ and only when the cluster operator $S$ satisfies Eq. (21), meaning that no terms nonlinear in $S$ are neglected. Here, we define

$$\omega^{(N_{1})}_{\mu} = E^{(N_{1})}_{\mu} - E^{(N)}_{0},$$

where $E^{(N)}_{0}$ is the ground-state CC energy of the $N$-electron reference system [Eq. (20)] and

$$\tilde{H}_{\text{open}} = (H_N)^{\text{C,open}} = e^{-H_N}e^{-2} - (H_N)^{\text{C,close}} = \tilde{H} - E^{(N)}_{0},$$

where $\tilde{H}$ is the similarity-transformed Hamiltonian defined by Eq. (22), with $S$ defined by Eq. (3), $H_N = H - (\langle H\rangle|\Phi\rangle \langle \Phi|\Phi\rangle)$ is the Hamiltonian in the normal-ordered form relative to the Fermi vacuum $|\Phi\rangle$, and the subscripts “open,” “closed,” and $C$ refer to open (i.e., having external lines), closed (i.e., having no external lines), and connected parts of a given operator expression. If the condition $M_R \leq M_S$ is not satisfied, we can only write

$$\tilde{H}_{\text{open}}R^{(N_{1})}_{\mu}|\phi\rangle = \omega^{(N_{1})}_{\mu}R^{(N_{1})}_{\mu}|\phi\rangle,$$

even when all relevant nonlinear terms in $S$ are included in the calculations. This introduces size-intensity errors into the resulting “excitation” energies $\omega^{(N_{1})}_{\mu}$. Eq. (26), when $M_R > M_S$. In fact, one might even argue, based on the analogy between the EA and IP EOMCC methods and the valence-universal MRCC approach in the (1,0) and (0,1) sectors of the Fock space, that it may be more appropriate to use the $M_R = M_S - 1$ condition for the truncations in $S$ and $R^{(N_{1})}_{\mu}$ to obtain the formally correct EA and IP EOMCC approximations [the simplest EA-EOMCCSD(2p-1h) = EA-EOMCCSD (Refs. 36 and 37) and IP-EOMCCSD(2h-1p) = IP-EOMCCSD (Refs. 38–41) approximations and their higher-order EA-EOMCCSDT, IP-EOMCCSDT, and EA-EOMCCSDTQ analogs in this category], but the analysis presented in Ref. 10, supported by the numerical evidence presented in Refs. 34, 35, and 88 and in Sec. III, shows that the $M_R = M_S$ condition that leads to the explicitly connected EA and IP EOMCC schemes, such as EA-EOMCCSD(3p-2h) and IP-EOMCCSD(3h-2p) (Refs. 34, 35, and 88), is perfectly acceptable in calculations of ground and excited states of radical species.

The issue that is far more important from the point of view of the accuracies in applications involving potential
energy surfaces of radicals is the inclusion of the 4p-3h/4h-3p excitations in $R^{(N+1)}_{\mu}$. Indeed, the $M_R \leq M_S$ condition, which is needed to convert Eqs. (23) and (28) into the explicitly connected Eq. (25), is satisfied by the EA-EOMCCSD(2p-1h), IP-EOMCCSD(2h-1p), EA-EOMCCSD(3p-2h), and IP-EOMCCSD(3h-2p) approaches, for which $M_S=2$ and $M_R$ equals 1, 2, and 2, respectively. All of these approaches lead to a rigorously size intensive description of the “excitation” energies $\omega^{(N+1)}_{\mu}$ defined by Eq. (26), but the corresponding results at larger intermolecular separations of radicals are generally poor. The $M_R \leq M_S$ condition is also satisfied by the SAC-CI analogs of the above four approaches, including the SAC-CI(3p-2h) and SAC-CI(3h-2p) methods and their active-space variants, although the SAC-CI(3p-2h) and SAC-CI(3h-2p) approaches tested in this work neglect the nonlinear terms in $S$ in Eq. (23), i.e., the corresponding wave functions

$$\Psi^{(N+1)}_{\mu} = (R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p} + R_{\mu,4h-3p}) e^{S_1 S_2} |\Phi\rangle \tag{29}$$

and

$$\Psi^{(N-1)}_{\mu} = (R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p}) e^{S_1 S_2} |\Phi\rangle \tag{30}$$

are approximated by

$$\Psi^{(N+1)}_{\mu} = (R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p})(1 + S_2) |\Phi\rangle \tag{31}$$

and

$$\Psi^{(N-1)}_{\mu} = (R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p})(1 + S_2) |\Phi\rangle \tag{32}$$

respectively, and this leads to a departure from a strictly size intensive behavior. This is not a problem for the analysis presented in this work, where all of the calculations are performed for small many-electron systems, for which the exact, full CI results can be generated for comparison purposes. In fact, the full treatment of nonlinear terms in $S$ offered by the corresponding EA-EOMCCSD(3p-2h) and IP-EOMCCSD(3h-2p) approaches, which rely on Eqs. (29) and (30) rather than Eqs. (31) and (32) and on the explicitly connected form of the eigenvalue problem [Eq. (25)], does not improve the relatively poor SAC-CI(3p-2h) and SAC-CI(3h-2p) results at larger intermolecular separations of the CH and OH radicals. As shown in Sec. III, one has to include the genuine 4p-3h and 4h-3p components of the $R^{(N+1)}_{\mu}$ and $R^{(N-1)}_{\mu}$ operators, as in Eqs. (14) and (15), and their active-space analogs discussed in Sec. II B to obtain an accurate description of the ground- and excited-state potential energy curves of the CH and OH. This is done in the present paper via the SAC-CI(4p-3h) and SAC-CI(4h-3p) methods and their active-space variants in which, in analogy to the aforementioned linearized forms of the SAC-CI(3p-2h) and SAC-CI(3h-2p) approaches, instead of the complete wave functions

$$\Psi^{(N+1)}_{\mu} = (R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p} + R_{\mu,4h-3p}) e^{S_1 S_2} |\Phi\rangle \tag{29}$$

and

$$\Psi^{(N-1)}_{\mu} = (R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p} + R_{\mu,4h-3p}) e^{S_1 S_2} |\Phi\rangle \tag{30}$$

we use the simplified forms

$$\Psi^{(N+1)}_{\mu} = (R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p} + R_{\mu,4h-3p}) \times (1 + S_2) |\Phi\rangle \tag{31}$$

and

$$\Psi^{(N-1)}_{\mu} = (R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p} + R_{\mu,4h-3p}) \times (1 + S_2) |\Phi\rangle \tag{32}$$

respectively, in which only the lead terms linear in $S_2$ are retained. The SAC-CI(4p-3h) and SAC-CI(4h-3p) methods are not rigorously size intensive, since they do not satisfy the condition $M_R \leq M_S (M_R=3$ and $M_S=2$ in this case) and neglect nonlinear terms in $S$ but, as shown in Sec. III, they are sufficiently accurate to provide an excellent description of the entire potential energy curves of the low-lying states of the CH and OH radicals. Clearly, it would eventually be more proper to use the more complete and size intensive EA-EOMCCSDTQ(4p-3h) and IP-EOMCCSDTQ(4h-3p) approximations, in which all nonlinear terms resulting from $e^S$ are retained and $M_R=M_S=3$, or the EA-EOMCCSDTQ(4p-3h)=EA-EOMCCSDTQ and IP-EOMCCSDTQ(4h-3p)=IP-EOMCCSDTQ approximations, as recently implemented in Ref. 47, in which $M_R=3$ and $M_S=4$, but this is not necessary to prove one of the main points of the present paper, which has not received much attention in the literature, that one needs the 4p-3h and 4h-3p components of the $R^{(N+1)}_{\mu}$ and $R^{(N-1)}_{\mu}$ operators to obtain accurate ground- and excited-state potential energy curves of radical species along the relevant bond breaking coordinates. Moreover, the development of the EA-EOMCCSDTQ(4p-3h), IP-EOMCCSDTQ(4h-3p), EA-EOMCCSDTQ(4p-3h), and IP-EOMCCSDTQ(4h-3p) codes, albeit in principle possible,47 leads to computational schemes that are prohibitively expensive anyway. Thus, for now, we will rely on the simplified SAC-CI(4p-3h) and SAC-CI(4h-3p) methods, as described by Eqs. (35) and (36), respectively, in our analyses. Even these severely truncated approaches are usually much too expensive, particularly when larger systems/larger basis sets are employed. Thus, it is essential to simplify the SAC-CI(4p-3h) and SAC-CI(4h-3p) methods even further. The idea advocated in this paper is that of the active-space SAC-CI(4p-3h) and SAC-CI(4h-3p) methods in which the large numbers of $r^{jk}_{abc}$ and $r^{kl}_{abcd}$ amplitudes defining the 3p-2h and 4p-3h components of $R^{(N+1)}_{\mu}$ and the large numbers of $r^{jk}_{abc}$ and $r^{kl}_{abcd}$ amplitudes defining the 3h-2p and 4h-3p components of $R^{(N-1)}_{\mu}$ are significantly reduced through the use of active orbitals.

### B. The active-space EA and IP SAC-CI and EOMCC approaches

As shown in this paper, the EA and IP SAC-CI approaches truncated at 4p-3h/4h-3p excitations are capable of providing a highly accurate description of the entire potential energy curves of the ground and low-lying excited states of...
radical species, but, as pointed out in the Introduction, the resulting SAC-CI(4p-3h) and SAC-CI(4h-3p) methods and their EA and IP EOMCC analogs are much too expensive to be widely applicable. In fact, even the less expensive EA and IP SAC-CI and EOMCC approaches truncated at 3p-2h/3h-2p excitations, which work well in the Frank-Condon region,\textsuperscript{34,35,43,88} are not very practical, particularly when the number of electrons in a radical system of interest is larger or a larger basis set is employed. This is a consequence of the fact that the numbers of $r_{ijk}^{ab}/r_{ij}^{ab}$ and $r_{abcd}^{ijkl}/r_{abcd}^{ijkl}$ amplitudes defining the relevant $K^{(N\alpha)}$ operators, particularly the numbers of the 4p-3h and 4h-3p amplitudes $r_{ijkl}^{hijk}$ and $r_{ijkl}^{ijkl}$, which are $\sim n_{p}^{3}$ and $\sim n_{p}^{3}$ respectively, are far too large for the majority of applications. The SAC-CI(4p-3h)/PS and SAC-CI(4h-3p)/PS methods mentioned in the Introduction, in which the small $r_{ijkl}^{hijk}$ and $r_{ijkl}^{ijkl}$ amplitudes that do not significantly perturb the suitably chosen zero-order wave functions, obtained with the CI(3p-2h) and CI(3h-2p) approaches, are eliminated from the calculations with the help of numerical thresholds, following the recipes described in Refs. 42, 44, and 93, substantially reduce the costs of the SAC-CI(4p-3h) and SAC-CI(4h-3p) calculations, but there are good reasons, also discussed in the Introduction, for seeking alternative ways of reducing these costs. The active-space SAC-CI(4p-3h) and SAC-CI(4h-3p) methods developed in this work and their recently proposed EA and IP EOMCC analogs, such as the EA-EOMCCSDt, EA-EOMCCSDtq, IP-EOMCCSDt, and IP-EOMCCSDtq approaches defined in Ref. 34, represent one of the most promising solutions to this problem.

The active-space EA and IP SAC-CI methods examined in this paper and their EOMCC counterparts discussed in Refs. 34, 35, and 88 are based on the observation that not all 3p-2h/3h-2p and 4p-3h/4h-3p amplitudes $r_{ijkl}^{hijk}$ and $r_{ijkl}^{ijkl}$ are equally important. In many cases, one can a priori select the dominant 3p-2h/3h-2p and 4p-3h/4h-3p patterns based on the model in which a radical is interpreted as a species obtained by either attaching an electron to one of the lowest-energy unoccupied orbitals or removing an electron from one of the highest-energy occupied orbitals of the related closed-shell system. For example, the CH and OH radicals examined in Sec. III are obtained, at least at the zero-order level, by attaching an electron to the lowest-energy unoccupied 1$\pi$ orbitals of the CH$^+$ closed-shell ion (the CH case) or by removing an electron from the highest-energy occupied 1$\sigma$ orbitals of the OH$^-$ closed-shell ion (the OH case), respectively. Thus, in analogy to the active-space CC and EOMCC methods, pioneered by Adamowicz, Piecuch, and co-workers,\textsuperscript{73,74,90,98-100,107} and multireference theories, one can treat the 1$\pi$ shells of CH$^+$ and OH$^-$ as active orbitals for selecting the most important 3p-2h/3h-2p and 4p-3h/4h-3p amplitudes in the EA and IP SAC-CI and EOMCC calculations for CH and OH. The choice of 1$\sigma$ shells of CH$^+$ and OH$^-$ as active orbitals may not be sufficient for the calculations at larger C-H and O-H distances that interest us in this work, since the 1$\sigma$ orbitals of the CH$^+$ and OH$^-$ systems become asymptotically degenerate with the corresponding 3$\sigma$ and 4$\sigma$ orbitals, but this can be easily addressed by selecting all valence orbitals of CH$^+$ and OH$^-$ that correlate with the 2$p$ shells of C and O and 1$s$ shell of H (the 3$\sigma$, 1$\pi$, 1$\pi$, and 4$\sigma$ orbitals of CH$^+$ and OH$^-$) as active orbitals for the EA and IP SAC-CI and EOMCC calculations with 3p-2h/3h-2p and 4p-3h/4h-3p excitations (see Fig. 1). This is equivalent to stating that at larger C-H distances, the low-lying electronic states of the CH radical are formed by the process of adding an electron to the 1$\pi$ or 4$\sigma$ unoccupied valence orbitals of CH$^+$, which can be combined with the electronic excitations from the highest occupied 3$\sigma$ orbital that belongs to the same valence shell as the 1$\pi$ or 4$\sigma$ orbitals. In the case of OH, we might say that at larger O-H separations, the low-lying electronic states of the OH radical are formed by the process of removing an electron from the 1$\sigma$ shell of OH$^-$, which can be coupled with the electronic excitations from the 3$\sigma$ and 1$\pi$ occupied orbitals to the 4$\sigma$ orbital that again belongs to the same valence shell as the 3$\sigma$ and 1$\pi$ orbitals.

The above situation, where the process of forming radical species from the related closed-shell species is defined by a relatively small subset of orbitals which are involved in the relevant electron attachment or electron removal process, occurs in a large number of molecular systems. It is, therefore, reasonable, in a manner similar to that of multireference approaches, to use these orbitals as active orbitals in the EA-EOMCC and IP-EOMCC calculations. Formally, in order to define the active-space EA and IP EOMCC and SAC-CI methods, we first divide the available spin-orbitals of a closed-shell $N$-electron system into four disjoint groups of core or inactive occupied spin-orbitals (labeled by lowercase bold letters $i,j,\ldots$), active spin-orbitals occupied in the closed-shell reference determinant $|\Phi\rangle$ (labeled by uppercase bold letters $I,J,\ldots$), active spin-orbitals unoccupied in $|\Phi\rangle$ (labeled by uppercase bold letters $A,B,\ldots$), and virtual or inactive unoccupied spin-orbitals (labeled by lowercase bold letters $a,b,\ldots$). We continue to designate the occupied and unoccupied spin-orbitals in $|\Phi\rangle$ by the italic characters $i,j,\ldots$ and $a,b,\ldots$, respectively, if the active/inactive character of the spin-orbitals is not specified. Once the above orbital classification scheme is established, we use it to define the electron attaching and electron removing operators $K_{\mu}^{(N\alpha)}$ of the active-space EA and IP EOMCC and SAC-CI methods, following the general strategy described in Ref. 34.
For example, the active-space EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) methods are obtained by replacing the 3p-2h component $R^{(N+1)}_{\mu}$ of the electron attaching operator $R^{(N+1)}_{\mu}$ in Eq. (10), by

$$r_{\mu,3p-2h} = \sum_{j>k,\lambda} r^{jk}_{\lambda\mu} a_\lambda^j a_\mu^k,$$  \hspace{1cm} (37)

Thus, the active-space EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) methods reduce the relatively large number of all 3p-2h amplitudes $r^{jk}_{\lambda\mu}$ that enter $R^{(N+1)}_{\mu}$, which is (if we ignore symmetries) $\sim n^2_{\mu} n^2_{\mu}$ to $\sim N_n n^2_{\mu}$, where $N_n$ is the number of active orbitals occupied in the reference determinant $|\Phi\rangle$. Assuming that $N_2 < n_\mu$, this is a major reduction in the computational effort compared to full EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) calculations. The active-space EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) methods approach uses $N_n$ active orbitals are designated by EA-EOMCCSD(3p-2h)$|N_n\rangle$ and SAC-CI(3p-2h)$|N_n\rangle$, respectively, clearly, the EA-EOMCCSD(3p-2h)$|N_n\rangle$ and SAC-CI(3p-2h)$|N_n\rangle$ methods, in which all orbitals are chosen to be active, are equivalent to the full EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) approaches. The relatively small set of the unknown amplitudes $r^{jk}_{\lambda\mu}$, defining $r_{\mu,3p-2h}$ [Eq. (37)], in which at least one of the three unoccupied spin-orbital indices is active, and the remaining 1p and 2p-1h amplitudes $r_a$ and $r_{ab}$ that enter the $(N+1)$-electron wave functions of the active-space EA-EOMCCSD(3p-2h)$|N_n\rangle$ approach,

$$\Psi^{(N+1)}_{\mu} = (R_{\mu,1p} + R_{\mu,2p-1h} + r_{\mu,3p-2h})e^{S_1S_2}|\Phi\rangle,$$  \hspace{1cm} (38)

where $S_1$ and $S_2$ are the singly and doubly excited clusters obtained in the preceding CCSD calculations for the closed-shell, $N$-electron, reference system, are obtained by diagonalizing the similarity-transformed Hamiltonian of CCSD,

$$H^{\text{CCSD}}_{\text{open}} = (H_N e^{S_1S_2})_{\text{open}},$$  \hspace{1cm} (39)

in the subspace of $H^{(N+1)}$ spanned by the $|\Phi^o\rangle$, $|\Phi^{ab}\rangle$, and $|\Phi^{jk}\rangle$ determinants [cf. Eq. (25)]. In the case of the active-space SAC-CI(3p-2h)$|N_n\rangle$ approach, as implemented in this paper, where the wave functions $|\Psi^{(N+1)}_{\mu}\rangle$, [Eq. (38)], are simplified to [see Eq. (31)]

$$\Psi^{(N+1)}_{\mu} = (R_{\mu,1p} + R_{\mu,2p-1h} + r_{\mu,3p-2h})(1 + S_2)|\Phi\rangle,$$  \hspace{1cm} (40)

where $S_2$ is obtained in the ground-state SAC-SD calculations for the $N$-electron reference system, the relevant $r_a$, $r_{ab}$, and $r^{jk}_{\lambda\mu}$ amplitudes defining $R^{(N+1)}_{\mu}$, $R^{(N+1)}_{\mu}$, and $R^{(N+1)}_{\mu}$ are obtained by solving the system of equations obtained by projecting the eigenvalue problem given by Eq. (23), in which $e^{S}$ is replaced by $(1+S_2)$ and $R^{(N+1)}_{\mu}$ are replaced by $(1+S_2)$ and $R^{(N+1)}_{\mu}$, respectively, onto the CSFs corresponding to the $|\Phi^o\rangle$, $|\Phi^{ab}\rangle$, and $|\Phi^{jk}\rangle$ determinants.

The savings in the computer effort offered by the active-space EA and IP SAC-CI and EOMCC approaches truncated at the 3p-2h and 3h-2p excitations can be quite significant compared to the parent EA-EOMCCSD(3p-2h)/SAC-CI(3p-2h) and IP-EOMCCSD(3h-2p)/SAC-CI(3h-2p) approximations, particularly in the EA case where the $N_n/n_\mu$ ratio which defines these savings is usually a small number. These savings become even larger when the active-space schemes with 4p-3h and 4h-3p excitations are considered, since in that case the active-space restrictions on the indices defining the relevant $r^{jk}_{\lambda\mu}$ and $r^{jk}_{\lambda\mu}$ amplitudes apply to both occupied and unoccupied spin-orbitals. For example, following the general prescription described in Sec. II C of Ref. 34 and using Eqs. (35) and (36), we define the wave functions $|\Psi^{(N+1)}_{\mu}\rangle$ and $|\Psi^{(N+1)}_{\mu}\rangle$ exploited in the active-space SAC-CI(4p-3h) and SAC-CI(4h-3p) methods employing $N_n$ active occupied and $N_n$ active unoccupied spin-orbitals,
designated as the SAC-CI(4p-3h)\{N_o,N_a\} and SAC-CI(4h-3p)\{N_o,N_a\} schemes, respectively, in the following manner:

\[
|\Psi^{(N+1)}_\mu\rangle = (R_{\mu,1p} + R_{\mu,2p-1h} + R_{\mu,3p-2h} + R_{\mu,4p-3h}) \\
\times (1 + S_2)|\Phi\rangle
\] (44)

and

\[
|\Psi^{(N-1)}_\mu\rangle = (R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p} + R_{\mu,4h-3p}) \\
\times (1 + S_2)|\Phi\rangle,
\] (45)

where \(S_2\) is a doubly excited cluster operator obtained in the ground-state SAC-SD calculations for the N-electron reference system. The \(r_{\mu,3p-2h}\) and \(r_{\mu,3h-2p}\) operators entering Eqs. (44) and (45) are exactly the same as those used in the active-space EA-EOMCCSD(3p-2h)/SAC-Cl(3p-2h) and IP-EOMCCSD(3h-2p)/SAC-Cl(3h-2p) calculations [see Eqs. (37) and (41)], whereas the \(r_{\mu,4p-3h}\) and \(r_{\mu,4h-3p}\) components are given by

\[
r_{\mu,4p-3h} = \sum_{J>k-1,A<B<cd} r_{\text{ABcd}} a_{\text{A}}^\dagger a_{\text{B}}^\dagger a_{\text{a}} a_{\text{d}},
\] (46)

and

\[
r_{\mu,4h-3p} = \sum_{J>k-1,A<B<cd} r_{\text{ABcd}} a_{\text{a}} a_{\text{d}} a_{\text{A}}^\dagger a_{\text{B}}^\dagger,
\] (47)

respectively. The explicit equations for the unknown \(r_{\text{AB}}\), \(r_{\text{ABC}}^j\), \(r_{\text{ABcd}}\), and \(r_{\text{ABCD}}^{jk}\) amplitudes, which define the \(R_{\mu,1p}\), \(R_{\mu,2p-1h}\), \(R_{\mu,3p-2h}\), and \(R_{\mu,4p-3h}\) operators entering the wave functions \(|\Psi^{(N+1)}_\mu\rangle\) of the SAC-CI(4p-3h)\{N_o,N_a\} approach [Eq. (44)], are obtained by projecting the SAC-CI eigenvalue problem given by Eq. (23), in which \(e^\dagger\) is replaced by \((1 + S_2)\) and \(R^{(N+1)}\) by \((R_{\mu,1p} + R_{\mu,2p-1h} + R_{\mu,3p-2h} + R_{\mu,4p-3h})\), onto the \(\mu\) CSFs corresponding to the \(|\Phi_j\rangle\), \(|\Phi_{jk}\rangle\), \(|\Phi_{jkl}\rangle\), and \(|\Phi_{jklm}\rangle\) determinants. Similarly, the explicit equations for the \(r_{\text{AB}}^{i}\), \(r_{\text{ABC}}^{ij}\), and \(r_{\text{ABCD}}^{jkl}\) amplitudes, which define the \(R_{\mu,1h}\), \(R_{\mu,2h-1p}\), \(R_{\mu,3h-2p}\), and \(R_{\mu,4h-3p}\) operators entering the wave functions \(|\Psi^{(N-1)}_\mu\rangle\) of the SAC-CI(4h-3p)\{N_o,N_a\} approach [Eq. (45)], are obtained by projecting the SAC-CI eigenvalue problem [Eq. (23)], in which \(e^\dagger\) is replaced by \((1 + S_2)\) and \(R^{(N-1)}\) by \((R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p} + R_{\mu,4h-3p})\), onto the CSFs corresponding to determinants \(|\Phi_j\rangle\), \(|\Phi_{jk}\rangle\), \(|\Phi_{jkl}\rangle\), and \(|\Phi_{jklm}\rangle\). Clearly, we could easily write the analogous expressions for the active-space EA-EOMCCSD(4p-3h)\{N_o,N_a\} and IP-EOMCCSD(4h-3p)\{N_o,N_a\} methods. In the case of EA-EOMCCSD(4p-3h)\{N_o,N_a\}, the explicit equations for the relevant \(r_{\text{AB}}\), \(r_{\text{ABC}}^j\), \(r_{\text{ABcd}}\), and \(r_{\text{ABCD}}^{jk}\) amplitudes, which define the EA-EOMCCSD(4p-3h)\{N_o,N_a\} wave functions \(|\Psi^{(N+1)}_\mu\rangle\) via the formula

\[
|\Psi^{(N+1)}_\mu\rangle = (R_{\mu,1p} + R_{\mu,2p-1h} + R_{\mu,3p-2h} + R_{\mu,4p-3h}) e^{S_1+S_2} |\Phi\rangle,
\] (48)

where \(S_1\) and \(S_2\) are determined by solving the CCSD equations for the N-electron reference system, would be obtained by projecting the eigenvalue problem involving the similarity-transformed Hamiltonian of CCSD [Eq. (39)], as given by Eq. (28), on \(|\Phi_{i}^{0}\rangle\), \(|\Phi_{i}^{1}\rangle\), \(|\Phi_{i}^{2}\rangle\), and \(|\Phi_{i}^{3}\rangle\). Similarly, the explicit equations for the \(r_{\text{AB}}\), \(r_{\text{ABC}}^j\), \(r_{\text{ABcd}}\), and \(r_{\text{ABCD}}^{jk}\) amplitudes, which define the IP-EOMCCSD(4h-3p)\{N_o,N_a\} wave functions \(|\Psi^{(N-1)}_\mu\rangle\) via

\[
|\Psi^{(N-1)}_\mu\rangle = (R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p} + R_{\mu,4h-3p}) e^{S_1+S_2} |\Phi\rangle,
\] (49)

would be obtained by projecting Eq. (28), with the similarity-transformed Hamiltonian \(\hat{H}_{\text{N,open}}\) replaced by \(\hat{H}_{\text{N,open}}^{\text{CSF}}\) [Eq. (39)], and \(R^{(N-1)}\) approximated by \((R_{\mu,1h} + R_{\mu,2h-1p} + R_{\mu,3h-2p} + R_{\mu,4h-3p})\), on \(|\Phi_j\rangle\), \(|\Phi_{jk}\rangle\), \(|\Phi_{jkl}\rangle\), and \(|\Phi_{jklm}\rangle\) determinants. Based on the discussion in Sec II A, ultimately one would like to pursue the active-space variants of the more complete and rigorously size intensive EA-EOMCCSDT(4p-3h) and IP-EOMCCSDT(4h-3p) approximations or their recently implemented EA-EOMCCSDTQ(4p-3h) = EA-EOMCCSDTQ and IP-EOMCCSDTQ(4p-3h) = IP-EOMCCSDTQ analogs, designated in Ref. 34 as the EA-EOMCCSDtq and IP-EOMCCSDtq approaches, in which the electron attaching and electron removing operators, \(R^{(N+1)}\) and \(R^{(N-1)}\), respectively, are defined in exactly the same manner as in the active-space SAC-CI(4p-3h)\{N_o,N_a\} and SAC-CI(4h-3p)\{N_o,N_a\} schemes and in which the ground state of the N-electron reference system is obtained in the active-space CCSDT (Ref. 73, 74, 89–92, and 98–107) or CCSDTq (90, 99–102, and 104) calculations. We plan to work on such approaches in the future, following the efficient computer coding strategy described in Ref. 88. The primary objective of this study is to show that one can accurately and efficiently handle the most expensive 4p-3h and 4h-3p components of the \(R^{(N+1)}\) and \(R^{(N-1)}\) operators, which are crucial for obtaining high quality potential energy curves of radical systems, via small active orbital spaces. The active-space SAC-CI(4p-3h)\{N_o,N_a\} and SAC-CI(4h-3p)\{N_o,N_a\} approaches are sufficient to prove this point.

Clearly, the SAC-CI(4p-3h)\{N_o,N_a\} and SAC-CI(4h-3p)\{N_o,N_a\} approaches, in which all orbitals are active, are equivalent to the parent SAC-CI(4p-3h) and SAC-CI(4h-3p) theories. This simple relationship between the active-space and parent methods is one of the advantages of all active-space approaches, since we always know that by increasing the active orbital space we can systematically approach the parent approximations. Of course, the main advantage of the active-space SAC-CI(4p-3h)\{N_o,N_a\} and SAC-CI(4h-3p)\{N_o,N_a\} approaches and their EA and IP EOMCC analogs with an active-space treatment of the 3p-2h/4p-3h and 3h-2p/4h-3p excitations, which we plan to develop in the future, is that they offer substantial savings in the computer effort compared to the parent SAC-CI(4p-3h) and SAC-CI(4h-3p) theories, and similar EOMCC approximations. In particular, the SAC-CI(4p-3h)\{N_o,N_a\} method reduces the large number of 4p-3h amplitudes used in the regular, all-orbital SAC-CI(4p-3h) approach, which can be estimated at \(-n_o^4n_a^2\) to \(-n_o^3n_a^2n_o^2\). Since \(n_o < n_a\) and, in the vast majority of cases, \(n_a < n_o\), the number of 4p-3h amplitudes used in the
active-space SAC-CI(4p-3h)\{N_o,N_u\} calculations is a tiny fraction of all amplitudes \( r_{ijkl}^{abed} \) and is true even for smaller radicals and smaller basis sets, such as the CH radical described by the \[5s3p1d/3s1p] basis set of Ref. 125, tested in this work. As shown in Table I, the number of the spin- and symmetry-adapted CSFs corresponding to the largest block of 4p-3h excitations entering the SAC-CI(4p-3h)\{1,3\} calculations for the CH/\[5s3p1d/3s1p] system, using the 3\( \sigma \), 1\( \pi_\sigma \), 1\( \pi_c \), and 4\( \pi \) orbitals of the CH* reference ion as active orbitals, is approximately 10\% of all 4p-3h CSFs needed to define the full SAC-CI(4p-3h) eigenvalue problem. There is also a significant reduction in the number of 3p-2h excitations (by a factor of \( \sim 2.6-2.7 \)), which entering the SAC-CI(3p-2h) and SAC-CI(4p-3h) calculations, when we use the above active orbitals. All of this translates into a reduction of the CPU time by almost two orders of magnitude, when we compare the active-space SAC-CI(4p-3h)\{1,3\} and regular SAC-CI(4p-3h) calculations for the CH/\[5s3p1d/3s1p] system. As shown in Table II, the savings offered by the SAC-CI(4h-3p)\{3,1\} approach, when applied to the OH radical, as described by the 6-31G(d,p) basis set, are less impressive, partly due to the fact that in this case \( N_o (\sim 3) \) is almost identical to \( n_o (\sim 4) \) and partly due to the small dimension of the employed basis set, but they are still substantial. In fact, in this case the number of spin- and symmetry-adapted 3h-2p amplitudes \( r_{bc}^{jk} \) is the same as the number of all 3h-2p amplitudes \( r_{ab}^{jk} \), since there is only one correlated occupied orbital outside the active space. Thus, all of the observed savings in the computer effort are due to the active-space treatment of 4h-3p excitations. Clearly, the savings offered by the SAC-CI(4h-3p)\{N_o,N_u\} would be greater if larger systems/larger basis sets were considered. In general, the SAC-CI(4h-3p)\{N_o,N_u\} method and its IP-EOMCC analogs, including the IP-EOMCCSD(4h-3p)\{N_o,N_u\} and IP-EOMCCSDtq approximations discussed above and in Ref. 34, reduce the large number of all 4h-3p amplitudes used, for example, in the regular SAC-CI(4h-3p) approach, which can be estimated at \( \sim n_o^3 n_u^2 \) to \( \sim N_o^7 N_u^2 \). Since \( N_o < n_o \) and for larger basis sets \( N_u < n_u \), the number of 4h-3p amplitudes used in the active-space SAC-CI(4h-3p)\{N_o,N_u\} calculations is much smaller than a number of all amplitudes \( r_{ijkl}^{abcd} \).

Tables I and II also illustrate the point that one can achieve significant reductions in the numbers of 4p-3h and 4h-3p amplitudes by exploiting the SAC-CI/PS approaches. The main difference between the active-space EA and IP SAC-CI and EOMCC methods pursued in this work and the corresponding SAC-CI/PS approaches, including the SAC-CI(4p-3h)/PS and SAC-CI(4h-3p)/PS methods used in our calculations for CH and OH, in which the 4p-3h/4h-3p components are selected numerically based on the suitable perturbative analysis exploiting the CI(3p-2h)/CI(3h-2p) reference states, is that in the active-space methods we use the a priori selected active orbitals which reflect on the electron attachment or ionization process when going from the N-electron reference system to the (N\( \pm 1 \))-electron radical of interest, whereas in the SAC-CI/PS methods we let the numerical thresholds decide which 4p-3h/4h-3p amplitudes are kept in the calculations and which are removed.

The active-space EA and IP EOMCC and SAC-CI methods, including the active-space SAC-CI(4p-3h) and SAC-CI(4h-3p) methods developed in this work, offer considerable savings in the computer effort when compared to the corresponding parent approximations. The key question is if the significant cost reduction offered by the active-space methods is not done at the expense of sacrificing the accuracy of the parent EA and IP EOMCC and SAC-CI calculations. This and related questions are addressed in the next section.

### III. NUMERICAL EXAMPLES AND DISCUSSION: POTENTIAL ENERGY CURVES OF THE LOW-LYING ELECTRONIC STATES OF CH AND OH

In order to examine the significance of the 4p-3h/4h-3p excitations in the EA/IP SAC-CI and, potentially, EOMCC calculations for ground- and excited-state potential energy surfaces of radical species along bond breaking coordinates and to demonstrate the effectiveness of the active-space EA/IP SAC-CI and EOMCC methods in recovering the dominant 3p-2h/3h-2p and 4p-3h/4h-3p contributions that
are needed to achieve high accuracies, we present the results of various SAC-CI and EOMCC calculations for the low-lying doublet states of the CH radical, as obtained with the [5s3p1d/3s1p] basis set of Ref. 125, and the low-lying doublet states of the OH radical, as computed with the 6–31G(d,p) basis set.\cite{126,127} In each case, we considered a wide range of nuclear geometries, including larger internuclear separations from the stretched-bond and asymptotic regions and several geometries from the region of the minimum on the corresponding ground-state potential energy curve.

The ground-state RHF orbitals of the closed-shell reference systems (CH\(^+\) for CH and OH\(^-\) for OH) were employed throughout and the spherical components of the \(d\) functions were used in the case of the 6–31G(d,p) basis set of the OH/OH\(^-\) system. In all correlated (SAC, CC, SAC-CI, EOMCC, and CI) calculations, the lowest-energy orbital of CH\(^+\) (the CH case) and the lowest-energy orbital of OH\(^-\) (the OH case) were kept frozen. The [5s3p1d/3s1p] basis set of Ref. 125 used in the calculations for CH and the 6–31G(d,p) basis set used in the calculations for OH are small enough to enable the exact, full CI calculations, which we performed using the GAMESS package.\cite{128} The availability of the full CI results, which are provided in the supplementary material,\cite{129} enables us to assess the accuracy of various EA and IP SAC-CI and EOMCC approaches. By having access to full CI wave functions of the CH and OH radicals, we can also analyze the relationship between the performance of various EA/IP SAC-CI and EOMCC methods and the significance of the 1\(p\)1\(h\), 2\(p\)-1\(h\)/2\(h\)-1\(p\), 3\(p\)-2\(h\)/3\(h\)-2\(p\), etc., contributions to these wave functions. All calculations for CH and OH reported in this work were based on exploiting the \(C_2\) symmetry.  

All of the EA and IP SAC-CI calculations, including the SAC-CI(3\(p\)-2\(h\)), SAC-CI(3\(h\)-2\(p\)), SAC-CI(4\(p\)-3\(h\)), and SAC-CI(4\(h\)-3\(p\)) calculations based on the wave function Ansätze defined by Eqs. (31), (32), (35), and (36), respectively, the corresponding SAC-CI(4\(p\)-3\(h\))/PS and SAC-CI(4\(h\)-3\(p\))/PS calculations, in which the smallest 4\(p\)-3\(h\) and 4\(h\)-3\(p\) contributions in the \(R^{(N+1)}\) and \(R^{(N-1)}\) operators that do not significantly perturb the zero-order states obtained in the Cl(3\(p\)-2\(h\)) and Cl(3\(h\)-2\(p\)) calculations are ignored, and the underlying SAC-SD calculations for the closed-shell reference systems (CH\(^+\) for CH and OH\(^-\) for OH) were performed using the routines developed at Kyoto University that form part of GAUSSIAN 03.\cite{130,131} The threshold \(\lambda_\mu\) for selection of the 4\(p\)-3\(h\) and 4\(h\)-3\(p\) components of the \(R^{(N+1)}\) and \(R^{(N-1)}\) operators in the SAC-CI(4\(p\)-3\(h\))/PS and SAC-CI(4\(h\)-3\(p\))/PS calculations was set at 10\(^{-7}\) hartree. The active-space SAC-CI(3\(p\)-2\(h\)), SAC-CI(3\(h\)-2\(p\)), SAC-CI(4\(p\)-3\(h\)), and SAC-CI(4\(h\)-3\(p\)) calculations were carried out using the new computer programs developed in this work, which were obtained by properly imposing the selection schemes discussed in detail in Sec. II B on the lists of spin- and symmetry-adapted CSFs corresponding to the \(|\Phi^{ab}_{ij}|\) and \(|\Phi^{cd}_{ijkl}\rangle\) (the EA case) and \(|\Phi^{bc}_{ijk}\rangle\) and \(|\Phi^{cd}_{ijk}\rangle\) (the IP case) determinants that define the corresponding Hamiltonian eigenvalue problems. These programs are very efficient and fully utilize the advantages of the active-space approaches, primarily by reducing the dimensions of the Hamiltonian matrices to be evaluated in the EA and IP SAC-CI calculations, as discussed in detail in Sec. II B (cf. Tables I and II). Once the final lists of spin- and symmetry-adapted CSFs that correspond to the \(|\Phi^{ab}_{ij}|\), \(|\Phi^{cd}_{ijkl}\rangle\), or \(|\Phi^{bc}_{ijk}\rangle\), \(|\Phi^{cd}_{ijk}\rangle\) or \(|\Phi^{ab}_{ij}|\), \(|\Phi^{cd}_{ijkl}\rangle\), \(|\Phi^{ab}_{ijk}\rangle\), \(|\Phi^{cd}_{ijkl}\rangle\), or \(|\Phi^{ab}_{ij}|\), \(|\Phi^{cd}_{ijkl}\rangle\), \(|\Phi^{ab}_{ijk}\rangle\), \(|\Phi^{cd}_{ijkl}\rangle\), \(\Phi^{bc}_{ijk}\rangle\), and \(\Phi^{cd}_{ijk}\rangle\) determinants are constructed, our programs proceed to the calculation of the nonzero Hamiltonian matrix elements that enter the eigenvalue problem defining the active-space SAC-CI(3\(p\)-2\(h\))/\(N\), SAC-CI(3\(h\)-2\(p\))/\(N\), SAC-CI(4\(p\)-3\(h\))/\(N\), and SAC-CI(4\(h\)-3\(p\))/\(N\), approach of interest. The full and active-space EA-EOMCCSD(3\(h\)-2\(p\)) and IP-EOMCCSD(3\(h\)-2\(p\)) calculations and the corresponding EA-EOMCCSD(2\(p\)-1\(h\)) and IP-EOMCCSD(2\(h\)-1\(p\)) calculations were performed using the highly efficient, vectorized computer codes employing the suitably defined recursively generated intermediates and fast matrix multiplication routines, which were described elsewhere.\cite{132,133,134} In particular, for the details of the EA-EOMCCSD(3\(p\)-2\(h\))/\(N\) and IP-EOMCCSD(3\(h\)-2\(p\))/\(N\) computer programs used in this work and the examination of their efficiency versus parent EA-EOMCCSD(3\(p\)-2\(h\)) and IP-EOMCCSD(3\(h\)-2\(p\)) calculations, we refer the reader to Ref. \(88\). The EA and IP EOMCC codes and the CCSD program used in this work have been interfaced with the RHF and integral routines in GAMESS.\cite{128}

A. The CH radical

Based on the model of the CH radical discussed in Sec. II B (cf. Fig. 1), in which the low-lying electronic states of CH are formed by the process of adding an electron to the 1\(\pi\) or 4\(\sigma\) unoccupied valence orbitals of CH\(^+\), which can be combined with the electronic excitations from the highest occupied 3\(\sigma\) orbital that belongs to the same valence shell as the 1\(\pi\) or 4\(\sigma\) orbitals, we have chosen the 3\(\sigma\), 1\(\pi\), 1\(\pi\), and 4\(\sigma\) valence orbitals of CH\(^+\) that correlate with the 2\(p\) shell of C and 1\(s\) shell of H as active orbitals for the SAC-CI(3\(p\)-2\(h\))/\(N\), EA-EOMCCSD(3\(p\)-2\(h\))/\(N\), and SAC-CI(4\(p\)-3\(h\))/\(N\) calculations discussed in this subsection. Thus, our choice of \(N\) for the active-space EA SAC-CI and EOMCC calculations reported in this work is \(N=1\) and \(N=3\). Before discussing the results of the active-space SAC-CI(3\(p\)-2\(h\))/\(N\), EA-EOMCCSD(3\(p\)-2\(h\))/\(N\), and SAC-CI(4\(p\)-3\(h\))/\(N\) calculations discussed in this subsection, we have used the EA SAC-CI and EOMCC calculations reported in this work for the 1\(\pi\), 2\(p\)-1\(h\), and 3\(p\)-2\(h\) contributions to the full CI wave functions representing the low-lying doublet states of CH (shown in Table IV) and the performance of the regular EA SAC-CI and EOMCC schemes in which all orbitals are active. We focus on the \(X^2\Pi\) ground state and the \(A^2\Delta\), \(B^2\Sigma^+\), and \(C^2\Sigma^+\) excited states. The \(X^2\Pi\) and \(B^2\Sigma^+\) states correlate with the lowest-energy \(C(3\,P)\)+\(H(2\,S)\) asymptote, whereas the \(A^2\Delta\) and

C $^2\Sigma^+$ states correlate with the C(1$D$)+H(2$S$) asymptote [see Fig. 2(a), which shows the full CI curves]. One of the interesting features of the low-lying states of CH is the crossing between the potential energy curves representing the $A^2\Delta$ and $B^2\Sigma^-$ states in the Franck-Condon region. It is interesting to examine what are the minimum levels of the EA SAC-CI and EOMCC theories that can properly describe various features of the full CI curves shown in Fig. 2(a), including, among others, the crossing of the $A^2\Delta$ and $B^2\Sigma^-$ curves in the Franck-Condon region and the asymptotic degeneracies of the $X^2\Pi$ and $B^2\Sigma^-$ as well as $A^2\Delta$ and $C^2\Sigma^+$ states.

The results in Table III indicate that the basic level of the EA EOMCC or SAC-CI theory, in which the electron attaching operator $K^{(N+1)}_\mu$ is truncated at the 2p-1h excitations, is practically useless. In the case of excited states, the differences between the EA-EOMCCSD(2p-1h) and full CI energies in the $R_{\text{CI}}$=0.75–4.0 Å region ($R_{\text{CI}}$ is the C–H separation and $R_{\text{CI}}$=1.119 786 Å is the experimental equilibrium bond length in CH taken from Ref. 130) are 57.722–278.327 mhartree for the $A^2\Delta$ state, 82.720–323.029 mhartree for the $B^2\Sigma^-$ state, and 18.885–89.126 mhartree for the $C^2\Sigma^+$ state. The analogous SAC-CI(2p-1h) results (not shown) are equally poor. As shown in Fig. 2(b), the EA-EOMCCSD(2p-1h) curves for the $A^2\Delta$, $B^2\Sigma^-$, and $C^2\Sigma^+$ states do not resemble the corre-

<table>
<thead>
<tr>
<th>State</th>
<th>Method</th>
<th>0.75</th>
<th>1.119 786</th>
<th>1.30</th>
<th>1.50</th>
<th>1.75</th>
<th>2.00</th>
<th>2.50</th>
<th>3.00</th>
<th>4.00</th>
<th>MUE</th>
<th>NPE</th>
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<tbody>
<tr>
<td>$X^2\Pi$</td>
<td>EA-EOMCCSD(2p-1h)</td>
<td></td>
<td>0.561</td>
<td>1.094</td>
<td>1.905</td>
<td>3.576</td>
<td>7.160</td>
<td>12.539</td>
<td>25.450</td>
<td>32.192</td>
<td>27.295</td>
<td>32.192</td>
</tr>
<tr>
<td></td>
<td>SAC-CI(3p-2h)</td>
<td></td>
<td>-0.236</td>
<td>-0.221</td>
<td>-0.212</td>
<td>-0.198</td>
<td>-0.152</td>
<td>0.009</td>
<td>0.785</td>
<td>1.526</td>
<td>2.227</td>
<td>2.227</td>
</tr>
<tr>
<td></td>
<td>SAC-CI(3p-2h)(3)$^b$</td>
<td></td>
<td>0.679</td>
<td>1.123</td>
<td>1.199</td>
<td>0.848</td>
<td>0.567</td>
<td>0.552</td>
<td>1.123</td>
<td>1.787</td>
<td>2.464</td>
<td>2.464</td>
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<tr>
<td></td>
<td>SAC-CI(4p-3h)</td>
<td></td>
<td>-0.103</td>
<td>-0.043</td>
<td>-0.176</td>
<td>-0.502</td>
<td>-1.996</td>
<td>-2.615</td>
<td>-3.834</td>
<td>-4.462</td>
<td>-4.946</td>
<td>4.946</td>
</tr>
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<td></td>
<td>SAC-CI(4p-3h)(1,3)$^c$</td>
<td></td>
<td>0.011</td>
<td>0.045</td>
<td>0.241</td>
<td>0.201</td>
<td>-1.012</td>
<td>-1.768</td>
<td>-2.896</td>
<td>-3.381</td>
<td>-3.815</td>
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<td></td>
<td>SAC-CI(4p-3h)/PS</td>
<td></td>
<td>0.170</td>
<td>0.408</td>
<td>0.222</td>
<td>0.023</td>
<td>-0.068</td>
<td>-0.913</td>
<td>-1.891</td>
<td>-2.251</td>
<td>-1.975</td>
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<tr>
<td></td>
<td>CI(4p-3h)</td>
<td></td>
<td>1.093</td>
<td>1.237</td>
<td>1.373</td>
<td>1.630</td>
<td>2.175</td>
<td>2.977</td>
<td>4.749</td>
<td>5.778</td>
<td>6.474</td>
<td>6.474</td>
</tr>
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</table>

$^a$Equilibrium geometry taken from Ref. 130.
$^b$The active space consisted of the 1$p_\pi$, 1$p_\pi$, 1$p_\sigma$, and 4$s$ orbitals of CH$^+$.  
$^c$The active space consisted of the 3$s$, 1$p_\pi$, 1$p_\pi$, and 4$s$ orbitals of CH$^+$.  

$C^2\Sigma^+$ states correlate with the C(1$D$)+H(2$S$) asymptote [see Fig. 2(a), which shows the full CI curves]. One of the interesting features of the low-lying states of CH is the crossing between the potential energy curves representing the $A^2\Delta$ and $B^2\Sigma^-$ states in the Franck-Condon region. It is interesting to examine what are the minimum levels of the EA SAC-CI and EOMCC theories that can properly describe various features of the full CI curves shown in Fig. 2(a), including, among others, the crossing of the $A^2\Delta$ and $B^2\Sigma^-$ curves in the Franck-Condon region and the asymptotic degeneracies of the $X^2\Pi$ and $B^2\Sigma^-$ as well as $A^2\Delta$ and $C^2\Sigma^+$ states.

The results in Table III indicate that the basic level of the EA EOMCC or SAC-CI theory, in which the electron attaching operator $K^{(N+1)}_\mu$ is truncated at the 2p-1h excitations, is practically useless. In the case of excited states, the differences between the EA-EOMCCSD(2p-1h) and full CI energies in the $R_{\text{CI}}$=0.75–4.0 Å region ($R_{\text{CI}}$ is the C–H separation and $R_{\text{CI}}$=1.119 786 Å is the experimental equilibrium bond length in CH taken from Ref. 130) are 57.722–278.327 mhartree for the $A^2\Delta$ state, 82.720–323.029 mhartree for the $B^2\Sigma^-$ state, and 18.885–89.126 mhartree for the $C^2\Sigma^+$ state. The analogous SAC-CI(2p-1h) results (not shown) are equally poor. As shown in Fig. 2(b), the EA-EOMCCSD(2p-1h) curves for the $A^2\Delta$, $B^2\Sigma^-$, and $C^2\Sigma^+$ states do not resemble the corre-
Energies are in hartree and the C–H distance $R_{\text{C-H}}$ is in Å. (a) The full CI results, (b) the EA-EOMCCSD(2p-1h) results, (c) the EA-EOMCCSD(3p-2h) results, (d) the SAC-CI(4p-3h) results, (e) the SAC-CI(4p-3h)[1,3] results, and (f) the SAC-CI(4p-3h)/PS results.

The two different coefficients shown are for the two doublet configuration state functions, each corresponding to a different intermediate spin state, that result from coupling the spins of the three unpaired electrons in this orbital occupation scheme.

This poor agreement in the calculations of excited states of CH can be understood if we realize that the $A^2\Delta$, $B^2\Sigma^-$, and $C^2\Sigma^+$ states have significant 2p-1h components at all C–H distances (see Table IV). They also gain substantially more at C=H distances and cannot describe the crossing of the $A^2\Delta$ and $B^2\Sigma^-$ states at $R_{\text{C-H}}=1.3$ Å. The situation only worsens as $R_{\text{C-H}}$ approaches the asymptotic region.

TABLE IV. An analysis of the major full CI configurations (all configurations with a coefficient $\geq 0.15$ for at least one of the selected values of $R_{\text{C-H}}$ are included) for the low-lying doublet states of the CH radical, as described by the [5s3p1d/3s1p] basis set of Ref. 125.

<table>
<thead>
<tr>
<th>State</th>
<th>Configuration orbital occupancy</th>
<th>Coefficients for various values of $R_{\text{C-H}}$</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>$X^2\Pi$</td>
<td>$(1\sigma^2)(2\sigma^2)(3\sigma^2)(1\pi_1^2)$</td>
<td>0.942</td>
<td>0.913</td>
<td>0.828</td>
<td>0.573</td>
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<td></td>
<td>$(1\sigma^2)(2\sigma^2)(3\sigma^2)(1\pi_1^2)(4\sigma^2)$</td>
<td>$&lt;0.01$</td>
<td>0.029</td>
<td>$-0.042, 0.212$</td>
<td>$-0.171, 0.370$</td>
</tr>
<tr>
<td></td>
<td>$(1\sigma^2)(2\sigma^2)(1\pi_1^2)(4\sigma^2)$</td>
<td>$&lt;0.01$</td>
<td>$-0.088$</td>
<td>$-0.245$</td>
<td>$-0.408$</td>
</tr>
<tr>
<td>$A^2\Delta$</td>
<td>$(1\sigma^2)(2\sigma^2)(3\sigma^2)(1\pi_1^2-1\pi_2^2)$</td>
<td>$-0.681$</td>
<td>$-0.648$</td>
<td>$-0.534$</td>
<td>0.362</td>
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<tr>
<td></td>
<td>$(1\sigma^2)(2\sigma^2)(1\pi_1^2-1\pi_2^2)(4\sigma^2)$</td>
<td>0.014</td>
<td>0.163</td>
<td>0.415</td>
<td>$-0.569$</td>
</tr>
<tr>
<td>$B^2\Sigma^-$</td>
<td>$(1\sigma^2)(2\sigma^2)(3\sigma^2)(1\pi_1^2)$</td>
<td>0.472, 0.818</td>
<td>0.440, 0.763</td>
<td>0.357, 0.619</td>
<td>$-0.250, -0.432$</td>
</tr>
<tr>
<td></td>
<td>$(1\sigma^2)(2\sigma^2)(1\pi_1^2)(4\sigma^2)$</td>
<td>$&lt;0.01, -0.013$</td>
<td>0.144, 0.249</td>
<td>0.310, -0.536</td>
<td>$-0.401, 0.694$</td>
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<tr>
<td>$C^2\Sigma^+$</td>
<td>$(1\sigma^2)(2\sigma^2)(3\sigma^2)(4\sigma^2)$</td>
<td>$-0.033$</td>
<td>0.166</td>
<td>0.710</td>
<td>0.730</td>
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<tr>
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<td>$(1\sigma^2)(2\sigma^2)(3\sigma^2)(4\sigma^2)$</td>
<td>0.015</td>
<td>0.053</td>
<td>0.161</td>
<td>0.295</td>
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<tr>
<td></td>
<td>$(1\sigma^2)(2\sigma^2)(3\sigma^2)(1\pi_1^2+1\pi_2^2)$</td>
<td>0.675</td>
<td>0.628</td>
<td>0.334</td>
<td>0.209</td>
</tr>
<tr>
<td></td>
<td>$(1\sigma^2)(2\sigma^2)(1\pi_1^2+1\pi_2^2)(4\sigma^2)$</td>
<td>$-0.011$</td>
<td>$-0.174$</td>
<td>$-0.288$</td>
<td>$-0.334$</td>
</tr>
</tbody>
</table>

$^a$ Equilibrium bond length taken from Ref. 130.
$^b$ Relative to the ground-state reference configuration of CH$^*$. $|1\sigma^22\sigma^23\sigma^2|^2$.
$^c$ The two different coefficients shown are for the two doublet configuration state functions, each corresponding to a different intermediate spin state, that result from coupling the spins of the three unpaired electrons in this orbital occupation scheme.
tial 3p-2h components, when $R_{C-H} > 1.5$ Å. This is particularly true for the $A^2 \Delta$ and $B^2 \Sigma^-$ states (for the $C^2 \Sigma^+$ state, the increase of the 3p-2h components is less pronounced). The EA-EOMCCSD(2p-1h) and SAC-CI(2p-1h) approaches can provide reasonable results only when the electronic states of interest are of an almost pure 1p character. As shown in Table IV, the $A^2 \Delta$, $B^2 \Sigma^-$, and $C^2 \Sigma^+$ states do not satisfy this requirement and this leads to huge errors in the EA-EOMCCSD(2p-1h) results for these states. The only state for which the EA-EOMCCSD(2p-1h) approach can find some limited use is the ground state. The ground state of CH is dominated by the 1p excitations in the $R_{C-H} > 2.0$ Å region and this correlates quite well with the relatively small errors in the EA-EOMCCSD(2p-1h) results for the $X^2 \Pi$ state in this region.

As shown in Table III and Fig. 2(c), the explicit inclusion of the 3p-2h excitations in the electron attaching operator $R_0^{(2p+1)}$ leads to substantial improvements in the poor EA-EOMCCSD(2p-1h) and SAC-CI(2p-1h) results. These improvements are particularly impressive when $R_{C-H} \leq 1.5$ Å. Indeed, the 57.722–98.442, 82.720–143.887, and 46.283–89.126 mhartree errors in the EA-EOMCCSD(2p-1h) results for the $A^2 \Delta$, $B^2 \Sigma^-$, and $C^2 \Sigma^+$ states, respectively, in the $R_{C-H} = 1.5$ Å region reduce to 1.260–5.061, 2.679–10.506, and 2.489–5.185 mhartree, respectively, when the EA-EOMCCSD(3p-2h) approach is employed, and 1.292–5.596, 2.740–11.141, and 2.489–5.488 mhartree, respectively, when the SAC-CI(3p-2h) method is used. The EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) results for the ground state are virtually perfect in the $R_{C-H} \leq 1.5$ Å region as well. For example, there is practically no difference between the EA-EOMCCSD(3p-2h) and full CI potential energy curves in the $R_{C-H} \leq 1.5$ Å region and the EA-EOMCCSD(3p-2h) approach restores the crossing of the $A^2 \Delta$ and $B^2 \Sigma^-$ states at $R_{C-H} \approx 1.3$ Å, which the EA-EOMCCSD(2p-1h) approach could not describe [cf. Figs. 2(a) and 2(c)].

The fact that the EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) approaches work so well in the $R_{C-H} \leq 1.5$ Å region is a consequence of the absence of the significant higher-than-2p-1h contributions in the $X^2 \Pi$, $A^2 \Delta$, $B^2 \Sigma^-$, and $C^2 \Sigma^+$ states in this region. This is particularly true in the vicinity of the minimum on the ground-state potential energy curve. For the $X^2 \Pi$, $A^2 \Delta$, and $C^2 \Sigma^+$ states, the 3p-2h and higher-than-3p-2h contributions remain relatively small in the entire $R_{C-H} \leq 1.5$ Å region. For the $B^2 \Sigma^-$ state, they begin to grow as $R_{C-H}$ approaches 1.5 Å and this results in the increase of the errors characterizing the EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) results for this state at $R_{C-H} = 1.5$ Å.

The EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) approaches provide considerable improvements in the poor EA-EOMCCSD(2p-1h) and SAC-CI(2p-1h) results in the $R_{C-H} > 1.5$ Å region as well. For example, the EA-EOMCCSD(3p-2h) approach reduces the $131.490–278.327$ and $185.074–323.029$ mhartree errors in the EA-EOMCCSD(2p-1h) results for the $A^2 \Delta$ and $B^2 \Sigma^-$ states in the $R_{C-H} = 1.75–4.0$ Å region down to 10.780–29.483 and 17.021–33.358 mhartree, respectively. SAC-CI(3p-2h) provides very similar improvements. In fact, since the 3p-2h contributions to the $X^2 \Pi$ and $C^2 \Sigma^+$ states do not grow as fast as in the case of the $A^2 \Delta$ and $B^2 \Sigma^-$ states and since the $X^2 \Pi$ and $C^2 \Sigma^+$ states remain largely dominated by the 1p and 2p-1h excitations for almost all values of $R_{C-H}$, the EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) approaches provide a very good description of the entire $X^2 \Pi$ and $C^2 \Sigma^+$ potential energy curves, with the MUE and NPE values ranging between 2 and 7 mhartree in the entire 0.75 Å $\leq R_{C-H} \leq 4.0$ Å region. This can also be seen by comparing the $X^2 \Pi$ and $C^2 \Sigma^+$ potentials obtained with EA-EOMCCSD(3p-2h) [shown in Fig. 2(c)] with the corresponding full CI curves [shown in Fig. 2(a)].

In spite of the considerable improvements in the results offered by the EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) approaches, one cannot rely on these approaches in quantitative calculations of the entire potential energy curves. The 17.315–29.483 and 18.367–33.800 mhartree errors resulting from the EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) calculations for the $A^2 \Delta$ state in the $R_{C-H} \geq 2.0$ Å region and the 22.849–33.358 and 23.928–36.910 mhartree errors obtained with the EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) methods for the $B^2 \Sigma^-$ state in the same region are too large to be acceptable in high accuracy calculations. The relatively good description of the entire $X^2 \Pi$ and $C^2 \Sigma^+$ curves and the relatively poor description of the $A^2 \Delta$ and $B^2 \Sigma^-$ curves at larger C–H distances by the EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) approaches result in a significant breakdown of the asymptotic degeneracies of the electronic states of CH shown in Fig. 2(a). For example, the $X^2 \Pi$ and $B^2 \Sigma^-$ states should become degenerate as $R_{C-H} \rightarrow \infty$, but they are far from being degenerate at larger C–H distances when the EA-EOMCCSD(3p-2h) method is employed, since the $B^2 \Sigma^-$ state dissociates incorrectly in the EA-EOMCCSD(3p-2h) calculations [see Fig. 2(c)]. A similar remark is true for the $A^2 \Delta$ and $C^2 \Sigma^+$ states, which should become degenerate as $R_{C-H} \rightarrow \infty$, but remain separated by a rather large energy gap at larger C–H distances in the EA-EOMCCSD(3p-2h) calculations, since only one of the two states ($C^2 \Sigma^+$) is reasonably well described by the EA-EOMCCSD(3p-2h) approach.

The above analysis demonstrates that one has to go beyond the EA-EOMCCSD(3p-2h) and SAC-CI(3p-2h) approximations to obtain a uniformly accurate description of the entire potential energy curves representing the low-lying states of CH. This is a consequence of the presence of the relatively large 3p-2h contributions to the electronic states of CH (particularly, the $A^2 \Delta$ and $B^2 \Sigma^-$ states) at larger internuclear separations. If we did not have access to the EA-EOMCCSD(3p-2h) data and had only to rely on the results obtained with the quasilinearized forms of the SAC-CI(3p-2h) wave functions used in this paper, we might speculate that larger errors in the results for the $A^2 \Delta$ and $B^2 \Sigma^-$ states at larger C–H distances are a consequence of ignoring the nonlinear terms in the cluster operator $S$ in Eq. (31) defining the SAC-CI(3p-2h) states rather than due to the neglect of higher-than-3p-2h components in the electron attaching operator $R_0^{(3p+1)}$. We might even speculate that perhaps the nonlinear terms in $S$ neglected in the SAC-SD cal-
citations for the CH⁺ reference system are responsible for the inaccuracies observed in the SAC-CI(3p-2h) calculations at larger C–H distances. Fortunately, we have access to the full CCSD and EA-EOMCCSD(3p-2h) results, in addition to the SAC-SD and SAC-CI(3p-2h) data, so that we do not have to speculate about these issues. As shown in Table III, there is virtually no difference between the results obtained with the quasilinearized form of SAC-CI(3p-2h) used in this paper and the results of the EA-EOMCCSD(3p-2h) calculations, which do not neglect any nonlinear terms in S that enter the EA-EOMCCSD(3p-2h)/SAC-CI(3p-2h) eigenvalue problem. Moreover, as shown in Table V, there is almost no difference in the results of the SAC-SD calculations for the ground-state potential energy curve of CH⁺, in which all nonlinear terms other than 1/2S₂ are neglected, and the analogous results obtained with the full CCSD approach, in which all nonlinear terms in S₁ and S₂ are kept. The SAC-SD and CCSD curves begin to deviate only in the R_{C-H} ≈ 2.5 Å region. In any case, since we use the full CCSD approach to produce the correlated reference states for the EA-EOMCCSD(3p-2h) calculations and since we do not neglect any relevant nonlinear terms in S in the EA-EOMCCSD(3p-2h) approach, and yet the EA-EOMCCSD(3p-2h) results for the Λ²Δ and Λ²Σ⁻ states at larger C–H separations are rather poor, we believe that the primary reason for the observed failures of the SAC-CI(3p-2h) and EA-EOMCCSD(3p-2h) approximations in the region of larger R_{C-H} values is the absence of the 4p-3h excitations in the electron attaching operator R^{(N+1)}_{µ} defining the EA SAC-CI and EOMCC theories.

The importance of the 4p-3h components of R^{(N+1)}_{µ} at larger C–H distances can be seen if we analyze the CI(4p-3h) data in Table III. The description of the X²Π, Λ²Δ, Λ²Σ⁻, and Λ²Σ⁺ potential energy curves of CH by the CI(4p-3h) approach, in which the electron attaching operator R^{(N+1)}_{µ} truncated at 4p-3h excitations is directly applied to the reference determinant |Φ⟩, is not perfect and the maximum errors in the CI(4p-3h) results range from 6.474 to 9.497 mhartree, but we already observe substantial improvements in the results for the Λ²Δ and Λ²Σ⁻ states, compared to the SAC-CI(3p-2h) and EA-EOMCCSD(3p-2h) calculations (MUE reduction by a factor of 4), when the CI(4p-3h) method is employed.

The most accurate results are obtained when we apply the electron attaching operator R^{(N+1)}_{µ} truncated at 4p-3h excitations to the properly correlated ground state of CH⁺ rather than to the uncorrelated, single-determinantal state |Φ⟩. This is clearly seen when we look at the SAC-CI(4p-3h) results in Table III and when we compare the potential energy curves resulting from the SAC-CI(4p-3h) calculations shown in Fig. 2(d) with the corresponding full CI curves shown in Fig. 2(a). The 29.483, 33.800, and 8.234 mhartree maximum errors in the EA-EOMCCSD(3p-2h), SAC-CI(3p-2h), and CI(4p-3h) results for the Λ²Δ state, the maximum errors of 33.358, 36.910, and 7.683 mhartree characterizing the EA-EOMCCSD(3p-2h), SAC-CI(3p-2h), and CI(4p-3h) calculations for the Λ²Σ⁻ state, and the 5.691, 5.488, and 9.497 mhartree maximum errors characterizing the EA-EOMCCSD(3p-2h), SAC-CI(3p-2h), and CI(4p-3h) results for the Λ²Σ⁺ state reduce to 1.381, 1.286, and 0.866 mhartree, respectively, when the SAC-CI(4p-3h) approach exploiting the quasilinearized form of the wave function [Eq. (35)] is employed. Based on the above discussion of the relative performance of EA SAC-CI, EOMCC, and CI(4p-3h) methods and based on the remarks made in Sec. II A, one of the best solutions that would, most likely, enable us to obtain the virtually exact description of the entire potential energy curves of the ground and excited states of CH should originate from the EA-EOMCCSDT(4p-3h) calculations, which would employ the CCSDT ground state of CH⁺ and the R^{(N+1)}_{µ} operator truncated at the 4p-3h excitations. Unfortunately, the EA-EOMCCSDT(4p-3h) approach has not been implemented yet [the automated implementation of the related EA-EOMCCSDTQ(4p-3h) = EA-EOMCCSDTQ approximation that uses the CCSDTQ rather than the CCSDT wave function as a reference ground state in the EA EOMCC calculations truncated at 4p-3h excitations has recently been reported,47 but we have no access to this implementation]. As pointed out in Sec. II A, the EA-EOMCCSDT(4p-3h) method would provide a rigorously size intensive description of the electronic excitations in CH. Moreover, as implied by the excellent full CCSDT results for CH⁺ and the results obtained by the recently formulated CR-CC(2,3) approach,31,132 which offers an approximate and yet highly accurate treatment of S₁ clusters, both shown in Table V, the explicit inclusion of S₁ clusters in the underlying calculations for CH⁺ would lead to the virtually exact description of the ground state of CH⁺, which serves as a reference for the EA EOMCC and SAC-CI calculations. These observations, combined with the great im-
provenments in the description of the ground and excited states of CH offered by the presence of the 4p-3h excitations in the electron attaching operator $R_{\mu}^{(N+1)}$, make us believe that the use of the EA-EOMCCSDT(4p-3h) approach would result in the virtually exact description of the entire potential energy curves of CH. On the other hand, it is encouraging to observe that the SAC-CI(4p-3h) method, in which most of the nonlinear terms in $S$ and the $S_3$ components are ignored and which is less expensive than EA-EOMCCSDT(4p-3h), provides a highly accurate description of the entire ground- and excited-state potential energy curves of CH. We might wonder if one could obtain similarly good results with the EA-EOMCCSDT(3p-2h) approach, in which $S$ is truncated at $S_3$ and the 4p-3h excitations in $R_{\mu}^{(N+1)}$ are ignored, since CASSCF provides the perfect description of CH; we will examine this issue in the future, but right now we believe that the SAC-CI(3p-2h) provides the perfect description of CH+; we will see if one could obtain similarly good results with the SAC-CI(3p-2h) approach and similar to the 2.251, 2.481, 3.713, and 3.601 mhartree maximum errors characterizing the corresponding SAC-CI(4p-3h)/PS curves shown in Figs. (e) and (f), respectively. There is also almost no difference between the SAC-CI(3p-2h)[1,3] curves shown in Table III, and the full CI curves shown in Fig. (a). All essential features of the full CI curves, including the crossing of the $A^2\Delta$ and $B^2\Sigma^+$ curves in the Franck-Condon region and the asymptotic degeneracies of the X$^2\Pi$ and $B^2\Sigma^+$ as well as $A^2\Delta$ and $C^2\Sigma^+$ states of CH are accurately reproduced by the SAC-CI(4p-3h)[1,3] calculations.

As one might expect, the SAC-CI(3p-2h)[3] and EA-EOMCCSD(3p-2h)[3] schemes are less accurate than the SAC-CI(4p-3h)[1,3] method, particularly in the region of larger C–H separations, but it is encouraging to observe that the SAC-CI(3p-2h)[3] and EA-EOMCCSD(3p-2h)[3] approaches, which reduce the dimensionality of the SAC-CI(3p-2h) and EA-EOMCCSD(3p-2h) eigenvalue problems by a factor of 2.6–2.7, faithfully reproduce the results of the parent SAC-CI(3p-2h) and EA-EOMCCSD(3p-2h) calculations at all C–H separations. The relatively small 0.679–1.200, 1.764–6.185, 3.337–11.738, and 3.207–6.496 mhartree errors in the SAC-CI(3p-2h)[3] results for the $X^2\Pi$, $A^2\Delta$, $B^2\Sigma^+$, and $C^2\Sigma^+$ states, respectively, in the $R_{CH}=1$ Å region and the very similar errors of 1.769–2.641, 1.733–5.646, 3.284–11.097, and 3.186–6.189 mhartree characterizing the EA-EOMCCSD(3p-2h)[3] results for the $X^2\Pi$, $A^2\Delta$, $B^2\Sigma^+$, and $C^2\Sigma^+$ states in the same region are not much different than the 0.198–0.236, 1.922–5.596, 2.740–11.141, and 2.489–5.488 mhartree errors in the corresponding SAC-CI(3p-2h) results and the 0.818–1.520, 1.260–5.061, 2.679–10.506, and 2.489–5.488 mhartree errors in the corresponding SAC-CI(3p-2h)/PS calculations (see Table III and Ref. 129). As in the case of the parent SAC-CI(3p-2h) and EA-EOMCCSD(3p-2h) approximations, the active-space SAC-CI(3p-2h)[3] and EA-EOMCCSD(3p-2h)[3] approaches fail to provide an accurate description of the asymptotic region, but the differences between the SAC-CI(3p-2h)[3]/EA-EOMCCSD(3p-2h)[3] energies and their SAC-CI(3p-2h)/EA-EOMCCSD(3p-2h) counterparts remain small even when $R_{CH}>1$ Å. For example, the relatively large errors...
in the EA-EOMCCSD(3p-2h){3} results for the $A^2\Delta$ and $B^2\Sigma^-$ states in the $R_{\text{OH}}=1.75-4.0$ Å region, where these states gain significant $3p-2h$ components, of 11.213-29.772 and 17.433-33.821 mhartree, respectively, are similar to the 10.780-29.483 and 17.021-33.358 mhartree errors obtained with the all-orbital EA-EOMCCSD(3p-2h) approach (cf. Table III). It is virtually impossible to distinguish between the ground- and excited-state potential energy curves of the low-lying states of CH obtained in the active-space SAC-CI(3p-2h){3} and EA-EOMCCSD(3p-2h){3} calculations and the analogous potential energy curves resulting from the parent SAC-CI(3p-2h) and EA-EOMCCSD(3p-2h) calculations.

We can conclude this subsection by stating that the active-space SAC-CI(3p-2h){3}, EA-EOMCCSD(3p-2h){3}, and SAC-CI(4p-3h){3} methods are as accurate as their considerably more expensive SAC-CI(3p-2h), EA-EOMCCSD(3p-2h), and SAC-CI(4p-3h) counterparts. In particular, the active-space SAC-CI(4p-3h){1,3} approach, in which one uses only four active orbitals to select the most numerous $3p-2h$ and $4p-3h$ excitations, provides an excellent description of the entire potential energy curves of the ground and low-lying excited states of the OH radical, including the Franck-Condon and asymptotic regions, at a small fraction of the computer effort involved in the SAC-CI(4p-3h) calculations.

B. The OH radical

The zero-order description of the OH radical discussed in Sec. II B (see Fig. 1), in which the low-lying electronic states of OH are formed by removing an electron from the occupied $1\pi$ shell of OH, which can be further coupled with the electronic excitations from the $3\sigma$ and $1\pi$ occupied orbitals to the $4\sigma$ unoccupied orbital that belongs to the same valence shell as the $3\sigma$ and $1\pi$ orbitals, suggests that the most natural choice of active orbitals for the SAC-CI(3h-2p){3}, IP-EOMCCSD(3h-2p){3}, and SAC-CI(4h-3p){3} calculations for OH is represented by the $3\sigma$, $1\pi_a$, $1\pi_b$, and $4\sigma$ valence orbitals of OH. These orbitals correlate with the $2p$ shell of O and $1s$ shell of H. This is the choice made in the active-space SAC-CI(3h-2p){3}, IP-EOMCCSD(3h-2p){3}, and SAC-CI(4h-3p){3} calculations for OH discussed in this subsection. In other words, in the active-space IP SAC-CI and EOMCC calculations performed in this study, the $N_\sigma$ and $N_\pi$ values are set at 3 and 1, respectively. As explained in Sec. II B, with this particular choice of the active space, the number of spin- symmetry-adapted $3h-2p$ amplitudes $f_{ij}^{\mu}$ is the same as the number of all $3h-2p$ amplitudes $f_{ij}^{\mu}$, since there is only one correlated occupied orbital outside the active space. This immediately implies that the SAC-CI(3h-2p){3} and IP-EOMCCSD(3h-2p){3} calculations for OH are equivalent to the regular, all-orbital SAC-CI(3h-2p) and IP-EOMCCSD(3h-2p) calculations, respectively. Thus, in Table VI and Fig. 3, and in the more detailed additional tables in the supplementary material, where the results of the IP SAC-CI and EOMCC calculations for OH are collected, we focus on the energies obtained in the active-space SAC-CI(4h-3p){3,1} calculations, which we compare with the corresponding full CI, IP-EOMCCSD(2h-1p), IP-EOMCCSD(3h-2p), SAC-CI(3h-2p), SAC-CI(4h-3p), and SAC-CI(4h-3p)/PS data.

In analogy to the CH radical, we first analyze the relationship between the importance of the $1h$, $2h-1p$, and $3h-2p$ contributions to the full CI wave functions representing the low-lying doublet states of OH (shown in Table VII) and the performance of the normal IP SAC-CI and EOMCC schemes in which all orbitals are active. We focus on the $X^2\Pi$ ground state and the $1^2\Delta$, $2^2\Sigma^+$, $1^2\Sigma^-$, and $B^2\Sigma^+$ excited states. As shown in Fig. 3(a), which displays the exact potential energy curves obtained in full CI calculations, the $X^2\Pi$ and $1^2\Sigma^-$ states correlate with the lowest-energy $O(3\Pi)+H(2\Sigma)$ asymptote, the $1^2\Delta$, $2^2\Pi$, and $A^2\Sigma^+$ states correlate with the next $O(3\Delta)+H(2\Sigma)$ asymptote, and the $B^2\Sigma^+$ state dissociates into $O(1\Sigma)+H(2\Sigma)$. We begin our analysis by examining the effectiveness of various IP SAC-CI and EOMCC approximations in describing the most important features of the full CI curves shown in Fig. 3(a), including, for example, the state ordering, the two interesting crossings of the $A^2\Sigma^+$ and $1^2\Sigma^-$ states at the O-H distance $R_{\text{O-H}}=1.5$ Å and of the $2^2\Pi$ and $B^2\Sigma^+$ states at $R_{\text{O-H}}=1.3$ Å, and the asymptotic degeneracies of the $X^2\Pi$ and $1^2\Sigma^-$ states and the $1^2\Delta$, $2^2\Delta$, and $A^2\Sigma^+$ states.

The performance of the basic IP EOMCC and SAC-CI approximations, in which the electron removing operator $K^{(3-1)}$ is truncated at the $2h-1p$ excitations, is generally very poor. In the following, we only discuss the results of the IP-EOMCCSD(2h-1p) calculations, since the analogous SAC-CI(2h-1p) results are essentially identical. As shown in Table VI, the only two states that are reasonably well described by the IP-EOMCCSD(2h-1p) method are the $X^2\Pi$ and $A^2\Sigma^+$ states, and even in this case the applicability of the IP-EOMCCSD(2h-1p) and SAC-CI(2h-1p) approaches is limited to the region of the minimum on the $X^2\Pi$ curve and to the region of the relatively small stretches of the O-H bond which do not exceed $R_{\text{O-H}}=1.5$ Å ($R_{\text{O-H}}$ is the O-H separation and $R_{\text{O-H}}=0.96966$ Å is the experimental equilibrium bond length in OH taken from Ref. 133). This can be understood by analyzing the leading contributions to the full CI wave functions of the $X^2\Pi$ and $A^2\Sigma^+$ states shown in Table VII. In the $R_{\text{O-H}}=1.5$ Å region, the $X^2\Pi$ and $A^2\Sigma^+$ states are dominated by $1h$ excitations. The IP EOMCC and SAC-CI approximations truncated at $2h-1p$ excitations work well for such states. Unfortunately, once we enter the $R_{\text{O-H}}=1.5$ Å region, the $X^2\Pi$ and $A^2\Sigma^+$ states gain significant $2h-1p$ components and, in the $R_{\text{O-H}}=3.0$ Å region, the relatively large $3h-2p$ contributions. The presence of the relatively large higher-than-$1h$ contributions in the wave functions representing the $X^2\Pi$ and $A^2\Sigma^+$ states in the $R_{\text{O-H}}=1.5$ Å region results in a rapid deterioration of the quality of the IP-EOMCCSD(2h-1p) results. Indeed, the relatively small errors in the IP-EOMCCSD(2h-1p) results relative to full CI in the $R_{\text{O-H}}=1.5$ Å region, which do not exceed 6.231 mhartree for the $X^2\Pi$ state and 8.921 mhartree

[129]
for the \( A^{2\Sigma^+} \) state, increase to 16.711–73.473 and 15.291–31.552 mhartree, respectively, when the 1.75 Å equilibrium bond length taken from Ref. 133.

The situation for the remaining states of OH considered in this work, which have large \( 2h\text{-}1p \) and, in some cases, large \( 3h\text{-}2p \) components independent of the value of \( R_{O-H} \) (cf. Table VII), is even more dramatic. Indeed, the errors in the IP-EOMCCSD(2h-1p) results for the \( 1^2\Delta, 2^2\Pi, 1^2\Sigma^- \), and \( B^{2\Sigma^+} \) states are 225.279–378.026, 67.703–256.555, 262.378–450.090, and 188.632–335.385 mhartree, respectively, when the entire 0.77 Å \( \leq R_{O-H} \leq 3.0 \) Å region is considered (see Table VI and Ref. 129). In consequence, the IP-EOMCCSD(2h-2p) potential energy curves for the \( 1^2\Delta, 2^2\Pi, 1^2\Sigma^- \), and \( B^{2\Sigma^+} \) states are completely pathological.
[cf. Figs. 3(a) and 3(b)]. The IP-EOMCCSD(2h-1p) curves are characterized by the very large MUE and NPE values. The energy gaps between electronic states and their relative ordering resulting from the IP-EOMCCSD(2h-1p) calculations are incorrect, even in the vicinity of the minimum on the ground-state potential energy curve. The $A^2\Sigma^+$ and $1^2\Sigma^-$ curves, which should cross at $R_{O-H}=1.5$ Å, do not cross in the IP-EOMCCSD(2h-1p) calculations, since the $1^2\Sigma^-$ state is significantly shifted up in energy. The $2^2\Pi$ and $B^3\Sigma^+$ curves obtained in the IP-EOMCCSD(2h-1p) calculations cross, but not at the right geometry. The asymptotic degeneracies of the $X^2\Pi$ and $1^2\Sigma^-$ states and the $1^2\Delta$, $2^2\Pi$, and $A^2\Sigma^+$ states are completely broken by the IP-EOMCCSD(2h-1p) method.

As shown in Table VI and Fig. 3(c), the inclusion of the 3h-2p excitations in the $R_0^{\text{CI}(2h-1p)}$ operator leads to large improvements in the poor IP-EOMCCSD(2h-1p) [and SAC-CI(2h-1p)] results. These improvements are particularly substantial in the region of the minimum on the ground-state potential energy curve and for the $1^2\Delta$, $2^2\Pi$, $1^2\Sigma^-$, and $B^3\Sigma^+$ states, which have significant 2h-1p components. For example, the 225.597, 171.011, 262.749, and 188.632 mhartree errors in the IP-EOMCCSD(2h-1p) results for the $1^2\Delta$, $2^2\Pi$, $1^2\Sigma^-$, and $B^3\Sigma^+$ states at $R_{O-H}=0.969$ 66 Å (the experimental equilibrium geometry of the ground-state OH radical) reduce to 7.191, 4.281, 12.392, and 4.203 mhartree, respectively, when the IP-EOMCCSD(3h-2p) approach is used, and 6.879, 4.259, 11.957, and 3.851 mhartree, respectively, when the SAC-CI(3h-2p) method is employed (see Table VI). For the remaining $X^2\Pi$ and $A^2\Sigma^+$ states, which in the Franck-Condon region are dominated by the $1h$ excitations and which are adequately described by the IP-EOMCCSD(2h-1p) approach when $R_{O-H}=0.969$ 66 Å, we observe further error reduction in the already good IP-EOMCCSD(2h-1p) results (the $X^2\Pi$ case) or no essential changes in the quality of the IP-EOMCCSD(2h-1p) energies (the $A^2\Sigma^+$ case). If we limit ourselves to the Franck-Condon region, there is almost no difference between the IP-EOMCCSD(3h-2p) potential energy curves representing the $X^2\Pi$, $1^2\Delta$, $2^2\Pi$, $A^2\Sigma^+$, $1^2\Sigma^-$, and $B^3\Sigma^+$ states of OH [shown in Fig. 3(c)] and the corresponding full CI curves [shown in Fig. 3(a)]. In particular, the IP-EOMCCSD(3h-2p) method restores the correct state ordering in the region of the minimum on the ground-state curve of OH, destroyed by the IP-EOMCCSD(2h-1p) model. It also provides a reasonable description of the crossing of the $A^2\Sigma^+$ and $1^2\Sigma^-$ states, which in the full CI calculations appears at $R_{O-H}=1.5$ Å and in the IP-EOMCCSD(3h-2p) calculations at $R_{O-H}=1.6$ Å, and of the crossing of the $2^2\Pi$ and $B^3\Sigma^+$ states, which both in the full CI and in the IP-EOMCCSD(3h-2p) calculations appear at $R_{O-H}=1.3$ Å. The IP-EOMCCSD(3h-2p) method provides an overall very good description of the ground and excited states of OH in the Franck-Condon region. The same is true for SAC-CI(3h-2p).

The IP-EOMCCSD(3h-2p) and SAC-CI(3h-2p) approaches are also quite effective in describing the entire potential energy curves of the $X^2\Pi$, $2^2\Pi$, and $A^2\Sigma^+$ states, which (with an exception of the $X^2\Pi$ state in the $R_{O-H}=3.0$ Å region) are dominated by the $1h$ and 2h-1p excita-
tions, with the errors relative to full CI that do not exceed a few millihartrees, but they fail to accurately describe the $1^{2}\Delta$, $1^{2}\Sigma^{-}$, and $B^{2}\Sigma^{+}$ states at larger O–H separations (see Table VI and Ref. 127). Indeed, although the IP-EOMCCSD(3h-2p) and SAC-CI(3h-2p) methods improve the pathological description of the $1^{2}\Delta$, $1^{2}\Sigma^{-}$, and $B^{2}\Sigma^{+}$ states by the IP-EOMCCSD(2h-1p) and SAC-CI(2h-1p) approaches, reducing the 225.279–378.026, 262.378–450.090, and 188.632–335.385 mhartree errors in the IP-EOMCCSD(2h-1p) results for these three states in the entire 0.77 Å $\leq R_{O-H} \leq$ 3.0 Å region to 3.033–42.747, 7.410–49.731, and 0.765–26.164 mhartree, respectively, in the IP-EOMCCSD(3h-2p) case and 2.801–61.313, 7.043–67.298, and 0.515–39.324 mhartree, respectively, in the SAC-CI(3h-2p) case, the errors in the IP-EOMCCSD(3h-2p) and SAC-CI(3h-2p) results for the $1^{2}\Delta$, $1^{2}\Sigma^{-}$, and $B^{2}\Sigma^{+}$ states at larger O–H distances are too large for high accuracy calculations. The IP-EOMCCSD(3h-2p) and SAC-CI(3h-2p) approaches break the asymptotic degeneracies of the $X^{2}\Pi$ and $1^{2}\Sigma^{-}$ states, which should dissociate into $O^{3}(P)+H^{(2)S}$, and of the $1^{2}\Delta$, $2^{2}\Pi$, and $A^{2}\Sigma^{+}$ states, which should dissociate into $O^{1}(D)+H^{(2)S}$, since they fail to correctly describe the $1^{2}\Sigma^{-}$ and $1^{2}\Delta$ states at larger O–H separations. For example, the IP-EOMCCSD(3h-2p) method restores the asymptotic degeneracy of the $2^{2}\Pi$ and $A^{2}\Sigma^{+}$ states, broken by IP-EOMCCSD(2h-1p), but this is not sufficient to obtain a correct description of the asymptotic region, since the $2^{2}\Pi$ and $A^{2}\Sigma^{+}$ states should also become degenerate with the $1^{2}\Delta$ state as $R_{O-H} \rightarrow \infty$ [see Fig. 3(c)].

The IP-EOMCCSD(3h-2p) and SAC-CI(3h-2p) approximations cannot provide an accurate description of the asymptotic region, since the $1^{2}\Delta$, $1^{2}\Sigma^{-}$, and $B^{2}\Sigma^{+}$ states gain large 3h-2p contributions at stretched O–H distances and, as argued in the Introduction, one has to include the 4h-3p excitations in the IP EOMCC and SAC-CI calculations in order to obtain an accurate description of the electronic states characterized by larger 3h-2p contributions. The significance of the 4h-3p components in the electron removing operator $R_{\mu}^{(N-1)}$ at larger O–H separations, particularly when the $1^{2}\Delta$, $1^{2}\Sigma^{-}$, and $B^{2}\Sigma^{+}$ states are examined, can be seen by comparing the IP-EOMCCSD(3h-2p) and SAC-CI(3h-2p) results, which ignore the 4h-3p contributions altogether, with the results of the CI(4h-3p) calculations, in which the $R_{\mu}^{(N-1)}$ operator truncated at 4h-3p excitations is directly applied to the reference determinant $|\Phi\rangle$. Although, as shown in Table VI, the description of the potential energy curves representing the $1^{2}\Delta$, $1^{2}\Sigma^{-}$, and $B^{2}\Sigma^{+}$ states of OH by the CI(4h-3p) approach is not fully quantitative and the maximum errors in the CI(4h-3p) results are still as large as 28.181 mhartree for the $1^{2}\Delta$ state, 25.486 mhartree for the $1^{2}\Sigma^{-}$ state, and 26.759 mhartree for the $B^{2}\Sigma^{+}$ state, we already observe major improvements in
the results, compared to the SAC-Cl(3h-2p) and IP-EOMCCSD(3h-2p) calculations, both in the maximum errors and in the corresponding NPE values, when the CI(4h-3p) method is employed. This implies that the inclusion of the genuine 4h-3p components in the $R^{(N-1)}_\mu$ operator is the key step toward improving the overall description of the potential energy curves of the low-lying states of OH. Other factors, such as the use of the quasilinearized form of the SAC-Cl(3h-2p) wave functions [Eq. (32)] and the neglect of several nonlinear terms in the cluster operator $S$ in the SAC-SD calculations for the OH$^-$ reference system that precede the SAC-Cl(3h-2p) calculations for OH are of lesser importance, since the IP-EOMCCSD(3h-2p) approach, which uses a more complete form of the wave function compared to SAC-Cl(3h-2p), provides results which are essentially as inaccurate in the region of larger O–H distances as those obtained with SAC-Cl(3h-2p). As shown in Table VI, there are relatively small differences between the results obtained with the quasilinearized form of SAC-Cl(3h-2p) used in this paper and the results of the IP-EOMCCSD(3h-2p) calculations, which do not neglect any nonlinear terms in $S$ that enter the full IP-EOMCCSD(3h-2p)/SAC-Cl(3h-2p) eigenvalue problem. Moreover, as shown in Table VIII, with an exception of the $R_{O-H} \approx 2.5$ Å region, the differences between the results of the SAC-SD calculations for the ground-state potential energy curve of OH$^-$, in which all nonlinear terms other than $\frac{1}{2}S_2$ are neglected, and the analogous results obtained with the full CCSD approach, in which all nonlinear terms in $S_1$ and $S_2$ are retained, are on the order of a fraction of a millihartree or a few millihartrees, at most. The SAC-SD and CCSD curves for OH$^-$ begin to differ more substantially in the $R_{O-H} \gtrsim 2.5$ Å region, but this is not too important in the context of the present discussion, since neither SAC-SD nor CCSD works well in this region (see Table VIII). A similar remark applies to the SAC-Cl(3h-2p) versus IP-EOMCCSD(3h-2p) results for the most difficult 1$^2\Delta$, 1$^2\Sigma^-$, and 2$^2\Sigma^+$ states, which begin to deviate more substantially only in the $R_{O-H} \approx 2.5$ Å region where neither SAC-Cl(3h-2p) nor IP-EOMCCSD(3h-2p) works well. We can conclude that since we do not neglect any relevant nonlinear terms in $S$ in the IP-EOMCCSD(3h-2p) calculations and yet the IP-EOMCCSD(3h-2p) results for the 1$^2\Delta$, 1$^2\Sigma^-$, and 2$^2\Sigma^+$ states of OH at larger O–H distances remain quite poor, the failures of the SAC-Cl(3h-2p) and IP-EOMCCSD(3h-2p) approximations in the region of larger $R_{O-H}$ values are primarily due to the absence of the 4h-3p excitations in the electron removing operators $R^{(N-1)}_\mu$ defining the SAC-Cl(3h-2p) and IP-EOMCCSD(3h-2p) schemes.

The CI(4h-3p) approach improves the poor description of the 1$^2\Delta$, 1$^2\Sigma^-$, and 2$^2\Sigma^+$ states at larger O–H separations by the IP-EOMCCSD(3h-2p), SAC-Cl(3h-2p), and other lower-order IP EOMCC and SAC-Cl methods, but, as shown in Table VI, the most accurate results for the electronic states of OH discussed in this work are obtained when we apply the electron removing operator $R^{(N-1)}_\mu$ truncated at 4p-3h excitations to the correlated ground state of OH$^-$, as is done in the SAC-Cl(4h-3p) calculations, rather than to the uncorrelated, single-determinant state $|\Phi\rangle$ used in the CI(4h-3p) model. This becomes clear when we examine the SAC-Cl(4h-3p) results in Table VI and when we compare the SAC-Cl(4h-3p) potential energy curves shown in Fig. 3(d) with the corresponding full CI curves shown in Fig. 3(a). The maximum errors of 42,747, 61,313, and 28,181 mhartree characterizing the IP-EOMCCSD(3h-2p), SAC-Cl(3h-2p), and CI(4h-3p) calculations for the 1$^2\Delta$ state, the 49,731, 67,298, and 25,486 mhartree maximum errors characterizing the IP-EOMCCSD(3h-2p), SAC-Cl(3h-2p), and CI(4h-3p) results for the 1$^2\Sigma^-$ state, and the maximum errors of 26,164, 39,324, and 26,759 mhartree characterizing the IP-EOMCCSD(3h-2p), SAC-Cl(3h-2p), and CI(4h-3p) calculations for the 2$^2\Sigma^+$ state reduce to 0.781, 0.698, and 1.464 mhartree, respectively, when the SAC-Cl(4h-3p) approach is employed (see Table VI). For the remaining states of OH listed in Table VI, the results of the SAC-Cl(4h-3p) calculations are equally good. We might be able to obtain additional small improvements if we used the rigorously size extensive IP-EOMCCSDT(4h-3p) approach, in which the $R^{(N-1)}_\mu$ operator truncated at the 4h-3p excitations is applied to the full CCSDT ground state of OH$^-$, but the IP-EOMCCSDT(4h-3p) method has not been implemented yet [again, the related IP-EOMCCSDT(4h-3p) = IP-EOMCCSDTQ(4h-3p) approach that uses the CCSDTQ rather than the CCSDT reference ground state in the IP EOMCC calculations truncated at 4h-3p excitations has recently been implemented, but this program is not available to us]. As implied by the excellent performance of the full CCSDT approach in the calculations for OH$^-$ and the analogous results obtained with the CR-CC(2,3) approach, which offers

<table>
<thead>
<tr>
<th>Method</th>
<th>0.77</th>
<th>0.969 666</th>
<th>1.07</th>
<th>1.27</th>
<th>1.50</th>
<th>1.75</th>
<th>2.00</th>
<th>2.50</th>
<th>3.00</th>
<th>MUE</th>
<th>NPE</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCSDT</td>
<td>0.464</td>
<td>0.594</td>
<td>0.652</td>
<td>0.735</td>
<td>0.779</td>
<td>0.826</td>
<td>0.927</td>
<td>1.341</td>
<td>1.840</td>
<td>1.840</td>
<td>1.376</td>
</tr>
<tr>
<td>CR-CC(2,3)</td>
<td>0.338</td>
<td>0.446</td>
<td>0.479</td>
<td>0.501</td>
<td>0.458</td>
<td>0.435</td>
<td>0.618</td>
<td>1.179</td>
<td>0.701</td>
<td>1.179</td>
<td>0.841</td>
</tr>
</tbody>
</table>

*The equilibrium bond length of OH taken from Ref. 133.*
an approximate and yet highly accurate treatment of $S_1$ clusters in all cases involving single bond breaking.131,132 both shown in Table VIII, the explicit inclusion of $S_1$ clusters would lead to the virtually exact description of the ground state of OH$^+$. which serves as a reference for the IP EOMCC and SAC-CI calculations for OH. This observation, combined with the great improvements in accuracy offered by the presence of the 4h-3p excitations in the electron removing operator $R^{\mu(N-1)}$, makes us believe that the IP-EOMCCSDT(4p-3h) approach would give the virtually exact description of the entire potential energy curves of OH. Having said all this, it is most encouraging to observe that the presence of the 4h-3p excitations in the asymptotic region. In addition to the small maximum errors and NPE values characterizing the SAC-CI(4h-3p) results for OH, the SAC-CI(4h-3p) approach restores the asymptotic degeneracies of the $X^2\Pi$ and $1^2\Sigma^-$ states, which dissociate into $\text{O}(^3P) + \text{H}(^2S)$, and of the $1^3\Delta$, $2^3\Pi$, and $A^2\Sigma^+$ states, which dissociate into $\text{O}(^1D) + \text{H}(^2S)$, which are not properly described by the lower-order IP SAC-CI and EOMCC models [cf. Figs. 3(d) and 3(a)]. Moreover, as shown in Fig. 3(d), the SAC-CI(4h-3p) approach provides the virtually perfect description of the crossings of the $A^2\Sigma^+$ and $1^2\Sigma^-$ states at $R_{\text{O-H}} \approx 1.5$ Å and of the $2^3\Pi$ and $B^2\Sigma^+$ states at $R_{\text{O-H}} \approx 1.3$ Å, further improving the IP-EOMCCSD(3h-2p) and SAC-CI(3h-2p) results in this regard. In analogy to the CH case, we might wonder if one could obtain similarly accurate results with the IP-EOMCCSD(3h-2p) method, in which $S$ is truncated at $S_3$ and the 4h-3p components of $R^{\mu(N-1)}$ are ignored, since CCSDT works perfectly for OH$^+$; we plan to study this issue in the future, but at this time we believe that one needs the 4h-3p excitations in $R^{\mu(N-1)}$, since the CI(4h-3p) calculations greatly improve the IP-EOMCCSD(3h-2p) results for the most difficult $1^3\Delta$, $2^3\Pi$, $1^2\Sigma^-$, and $B^2\Sigma^+$ states.

Thus far, we have demonstrated that the SAC-CI(4h-3p) approach provides high-quality potential energy curves for the ground and excited states of OH. The final question remains if one can retain the high accuracy of the SAC-CI(4h-3p) calculations without a full inclusion of all 4h-3p amplitudes $r^{\mu(N-1)}_{hi,ij}$. In analogy to the CH case, one can reduce large costs of the SAC-CI(4h-3p) calculations by using the SAC-CI(4h-3p)/PS approach, in which the 4h-3p components of the $R^{\mu(N-1)}_{\mu}$ operators that do not significantly perturb the CI(3h-2p) reference wave function and whose energy contributions are smaller than the threshold $\lambda_\mu$ (set in our calculations at 10$^{-1}$ hartree) are neglected and eliminated from the SAC-CI(4h-3p) diagonalization. As shown in Table VI, the SAC-CI(4h-3p)/PS results are essentially as good as those obtained with the complete SAC-CI(4h-3p) approach, in spite of the fact that the total number of spin- and symmetry-adapted amplitudes defining the $R^{\mu(N-1)}_{\mu}$ operator used in the SAC-CI(4h-3p)/PS calculations represents 12%–46% of all $r$ amplitudes used in the SAC-CI(4h-3p) calculations (see Table II). There is, however, a problem, mentioned in the Introduction, of the numerical noise which the SAC-CI(4h-3p)/PS calculations may produce due to the fact that different sets of the 4h-3p contributions to the $R^{\mu(N-1)}_{\mu}$ operators are selected at different nuclear geometries. The active-space SAC-CI(4h-3p) approach developed in this work uses the same set of $r$ amplitudes defining the $R^{\mu(N-1)}_{\mu}$ operator at all nuclear geometries, producing smooth potential energy curves. At the same time, the active-space SAC-CI(4h-3p) method offers a substantial reduction in the dimensionality of the corresponding eigenvalue problem through the selection of the dominant 3h-2p and 4h-3p amplitudes via active orbitals, as discussed in Sec. II B (see Table II). As shown in Table VI, the active-space SAC-CI(4h-3p)$[3,1]$ approach employing only three occupied and one unoccupied active orbitals, and using approximately 29% of all amplitudes $r$ defining the $R^{\mu(N-1)}_{\mu}$ operator truncated at 4h-3p excitations, is as effective in producing highly accurate potential energy curves of OH as the SAC-CI(4h-3p)/PS and complete SAC-CI(4h-3p) methods. The small maximum errors relative to full CI of 3.381, 2.577, 2.270, 2.505, 3.107, and 3.452 mhartree characterizing the SAC-CI(4p-3h)$[3,1]$ results for the entire potential energy curves of the $X^2\Pi$, $1^3\Delta$, $2^3\Pi$, $A^2\Sigma^+$, $1^2\Sigma^-$, and $B^2\Sigma^+$ states of OH are similar to or only slightly larger than the 4.776, 0.781, 1.531, 3.651, 0.698, and 1.464 mhartree maximum errors obtained with the SAC-CI(4h-3p) approach and the 3.773, 0.803, 1.591, 3.193, 1.942, and 2.118 mhartree maximum errors resulting from the SAC-CI(4h-3p)/PS calculations. Similar remarks apply to the NPE values characterizing the potential energy curves of OH obtained in the SAC-CI(4h-3p)$[3,1]$ calculations. In consequence, the SAC-CI(4h-3p)$[3,1]$ potential energy curves shown in Fig. 3(e) can hardly be distinguished from the highly accurate SAC-CI(4h-3p) and SAC-CI(4h-3p)/PS curves shown in Figs. 3(d) and 3(f), respectively. There is practically no difference between the ground- and excited-state potential energy curves obtained in the relatively inexpensive active-space SAC-CI(4h-3p)$[3,1]$ calculations shown in Fig. 3(e) and the full CI curves shown in Fig. 3(a). All essential features of the full CI curves, including the asymptotic degeneracies of the $X^2\Pi$ and $1^2\Sigma^-$ states, which dissociate into $\text{O}(^3P) + \text{H}(^2S)$, and of the $1^3\Delta$, $2^3\Pi$, and $A^2\Sigma^+$ states, which dissociate into $\text{O}(^1D) + \text{H}(^2S)$ and the aforementioned crossings of the $A^2\Sigma^+$ and $1^2\Sigma^-$ states at $R_{\text{O-H}} \approx 1.5$ Å and of the $2^3\Pi$ and $B^2\Sigma^+$ states at $R_{\text{O-H}} \approx 1.3$ Å, are accurately described by the active-space SAC-CI(4h-3p)$[3,1]$ approach.

We conclude by stating that the active-space SAC-CI(4h-3p)$[N_{\mu}, N_{\nu}]$ approach exploiting small numbers of occupied and unoccupied active orbitals is practically as accurate as its considerably more expensive SAC-CI(4h-3p) counterpart. The active-space SAC-CI(4h-3p)$[3,1]$ approach employing four active orbitals to select the dominant 3h-2p and 4h-3p excitations provides an excellent description of the potential energy curves of the ground and excited states of the OH radical, including the Franck-Condon and asymptotic regions, at the relatively small fraction of the effort involved in the parent SAC-CI(4h-3p) calculations.
IV. SUMMARY AND CONCLUDING REMARKS

In the present paper, we have focused on one of the major challenges of modern electronic structure theory, which is the development of practical methods that can accurately describe ground- and excited-state potential energy surfaces of radical species along bond breaking coordinates. Specifically, we have shown that one can develop relatively inexpensive and easy-to-use \textit{ab initio} schemes for high accuracy calculations of ground- and excited-state potential energy surfaces of radicals by combining the EA/IP SAC-CI and EOMCC methods with 3p-2h/3h-2p and 4p-3h/4h-3p excitations with the idea of using the physically motivated sets of active orbitals to select the most important 3p-2h/3h-2p and 4p-3h/4h-3p excitations. The active-space variants of the SAC-CI(4p-3h) and SAC-CI(4h-3p) approaches turned out to be particularly successful, enabling us to obtain highly accurate potential energy curves of the ground and excited states of the CH and OH radicals along the relevant bond breaking coordinates, including the Franck-Condon and asymptotic regions, at a small fraction of the computer cost associated with the regular SAC-CI(4p-3h) and SAC-CI(4h-3p) calculations. We have demonstrated that it is sufficient to use small numbers of active orbitals that correlate with the valence shells of the radical species of interest and small fractions of all 3p-2h/3h-2p and 4p-3h/4h-3p excitations selected via active orbitals to obtain excellent results of the full SAC-CI(4p-3h)/SAC-CI(4h-3p) quality. We have also shown that in general the active-space EA and IP EOMCC and SAC-CI methods with 3p-2h/3h-2p or 3p-2h/3h-2p and 4p-3h/4h-3p excitations faithfully reproduce the results obtained with the parent EA and IP EOMCC and SAC-CI approaches including these high-order excitations.

By comparing the potential energy curves of the ground and excited states of the CH and OH systems obtained in the full and active-space SAC-CI(4p-3h) and SAC-CI(4h-3p) calculations with the corresponding full CI curves, full and active-space EA-EOMCCSD(3p-2h)/SAC-CI(3p-2h) and IP-EOMCCSD(3h-2p)/SAC-CI(3h-2p) results, and potential energy curves obtained with the basic EA-EOMCCSD(2p-1h) and IP-EOMCCSD(2h-1p) approximations, we have shown that one needs the 4p-3h and 4h-3p excitations in the electron attaching and ionizing operators $R_{\mu}^{(N\pm 1)}$ in order to obtain an accurate representation of the entire ground- and excited-state potential energy surfaces of radicals along bond breaking coordinates. The full and active-space EA-EOMCCSD(3p-2h)/SAC-CI(3p-2h) and IP-EOMCCSD(3h-2p)/SAC-CI(3h-2p) methods, in which the 4p-3h and 4h-3p components of $R_{\mu}^{(N\pm 1)}$ are ignored, provide accurate excitation energies in the Franck-Condon region, but they fail at larger internuclear separations, where electronic states of radicals often gain significant 3p-2h and 3h-2p contributions. A rationale for this behavior of the EA/IP SAC-CI and EOMCC methods has been provided in the Introduction. The EA/IP SAC-CI and EOMCC methods with 4p-3h/4h-3p excitations are expected to work well in the Franck-Condon and asymptotic regions of many radical species, since radicals can be viewed as systems obtained by attaching an electron to or removing an electron from the related closed-shell molecule and it is well known that the triply excited (i.e., 3p-3h) clusters are usually sufficient to obtain a quantitative description of single bond breaking in closed-shell systems. Our numerical results for the CH and OH radicals obtained with the full and active-space SAC-CI(4p-3h) and SAC-CI(4h-3p) methods confirm this expectation.

As shown in this study, the active-space variants of the SAC-CI(4p-3h) and SAC-CI(4h-3p) methods are at least as effective in describing ground and excited states of radicals as the SAC-CI(4p-3h)/PS and SAC-CI(4h-3p)/PS approaches, in which one selects the 4p-3h and 4h-3p excitations numerically, based on their significance in perturbing the CI(3p-2h) and CI(3h-2p) wave functions. The active-space SAC-CI(4p-3h) and SAC-CI(4h-3p) methods and their parent SAC-CI(4p-3h) and SAC-CI(4h-3p) approximations, in which all orbitals are active, have also been shown to be significantly more accurate than the analogous CI methods with up to 4p-3h and 4h-3p excitations. One has to apply the electron attaching and ionizing operators $R_{\mu}^{(N\pm 1)}$ truncated at the 4p-3h and 4h-3p components to the correlated CC or SAC ground state of the $N$-electron reference system to obtain high-quality potential energy surfaces of the $(N\pm 1)$-electron radical species along bond breaking coordinates.

Based on the numerical results and theoretical analysis presented in this paper, we expect that the active-space variants of the EA-EOMCCSD(4p-3h) and IP-EOMCCSD(4h-3p) approaches, which would represent the EOMCC analogs of the active-space SAC-CI(4p-3h) and SAC-CI(4h-3p) methods examined in this work, and, particularly, the active-space variants of the rigorously size intensive EA-EOMCCSDT(4p-3h) and IP-EOMCCSDT(4h-3p) methods, referred to in Ref. 34 as the EA-EOMCCSDtq and IP-EOMCCSDtq schemes, should be at least as effective as the active-space SAC-CI(4p-3h) and SAC-CI(4h-3p) approximations discussed here. The efficient computer implementation of the EA-EOMCCSDtq and IP-EOMCCSDtq methods proposed in Ref. 34, which would use active orbitals to select the most important $S_\mu$ components in the cluster operator $S$ defining the underlying CCSDT/SAC-SDT calculations for the $N$-electron reference systems and the dominant 3p-2h/3h-2p and 4p-3h/4h-3p excitations in the corresponding operators $R_{\mu}^{(N\pm 1)}$, would be a useful next step toward the development of high accuracy methods for radicals and other open-shell systems. On the other hand, it is encouraging to observe that the active-space SAC-CI(4p-3h) and SAC-CI(4h-3p) methods, which are less expensive than the EA- and IP-EOMCCSDtq schemes, provide highly accurate potential energy surfaces of radical species.

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