Ab initio coupled-cluster theory for open-shell nuclei

G. R. Jansen, M. Hjorth-Jensen, G. Hagen, and T. Papenbrock

1Department of Physics and Center of Mathematics for Applications, University of Oslo, N–0316 Oslo, Norway
2Physics Division, Oak Ridge National Laboratory, Oak Ridge, TN 37831, USA
3Department of Physics and Astronomy, University of Tennessee, Knoxville, TN 37996, USA
4GSI Helmholtzzentrum für Schwerionenforschung GmbH, 64291 Darmstadt, Germany
5Institut für Kernphysik, Technische Universität Darmstadt, 64289 Darmstadt, Germany

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We develop a new method to describe properties of truly open-shell nuclei. This method is based on single-reference coupled-cluster theory and the equation-of-motion method with extensions to nuclei with \( A \pm 2 \) nucleons outside a closed shell. We perform proof-of-principle calculations for the ground states of the helium isotopes \(^3\text{He}, ^4\text{He}, ^5\text{He}, ^6\text{He}\) and the first excited \(^2^+\) state in \(^6\text{He}\). The comparison with exact results from matrix diagonalization in small model spaces demonstrates the accuracy of the coupled-cluster methods. Three-particle–one-hole excitations of \(^4\text{He}\) play an important role for the accurate description of \(^6\text{He}\). For the open-shell nucleus \(^6\text{He}\), the computational cost of the method is comparable with the coupled-cluster singles-and-doubles approximation while its accuracy is similar to coupled-cluster with singles, doubles and triples excitations.

I. INTRODUCTION

The nuclear shell model is the paradigm for our understanding of atomic nuclei \([1]\). Within this model, doubly magic nuclei (with fully occupied shells for protons and neutrons) are particularly important because they are stronger bound than their neighbors and can be approximated by a simple product of single-particle states. Within the nuclear shell model, doubly magic nuclei are the cornerstones for our understanding of entire regions of the nuclear chart as they can be viewed as inert cores. For the \textit{ab initio} description of doubly magic nuclei, the coupled cluster (CC) method – based on particle-hole excitations of a reference Slater determinant that obey the linked cluster theorem – is particularly well suited and arguably one of the most efficient methods \([2, 3]\). Similar remarks hold for nuclei that differ from doubly-magic nuclei by one nucleon; such nuclei still exhibit a simple structure and a single Slater determinant is a good reference state. The structure of all other nuclei is more complicated and requires the superposition of many product states and correspondingly large model spaces.

For the light \( p\)-shell nuclei, various \textit{ab initio} methods \([4–11]\) yield virtually exact results for realistic Hamiltonians. For heavier systems, one typically relies on approximations. Here, coupled-cluster theory is an ideal compromise between accuracy on the one hand and computational cost on the other. This method has been applied to various problems in nuclear structure \([2, 3, 12–17]\).

In this paper, we use equation-of-motion (EOM) techniques within the coupled-cluster method, abbreviated to EOM-CC hereafter, for the description of nuclei that differ from closed-shell references by two nucleons. This extension of the coupled-cluster method is useful for two reasons. First, it significantly enlarges the set of nuclei that can be accessed within coupled-cluster theory. For the oxygen isotopes \(^{14–26}\text{O}\), for instance, all nuclei except \(^{19}\text{O}\) differ from closed-subshell references by two neutrons or less. Similar comments apply to isotopes of helium and calcium. Questions related to the evolution of shell structure \([18, 19]\) could thus be addressed from first principles. Second, the coupled-cluster method yields a similarity transformed Hamiltonian (see Eq. \((8)\) below) for a doubly-magic nucleus. The Hamiltonians for one and two nucleons attached to this doubly magic core provide us with effective single-particle energies and an effective two-body interaction, respectively. These matrix elements could enter the construction of effective shell-model interactions \([20]\) that are the basis for large scale shell-model calculations \([21, 22]\).

Within the EOM-CC methods, the equations for excited states and one particle attached/removed are well known in quantum chemistry \([4, 23–27]\), and have also been applied to atomic nuclei \([28, 31]\). However, the corresponding equations for two particle attached/removed have seen very few applications in quantum chemistry \([32, 33]\) and are new in nuclear physics. In this article, we will thus extend the range of the EOM-CC methods to include open-shell nuclei with \( A = \pm 2 \) nucleons outside a closed shell core. We present here the necessary formalism for deriving such equations, including the pertinent diagrams and algebraic equations. To our knowledge, these details have not been presented elsewhere. The results from our EOM-CC calculations are compared with full configuration interaction (FCI) calculations for helium isotopes, demonstrating the accuracy of this approach.

This paper is organized as follows. In Sec. \(\text{II}\) we give a brief overview of the equation-of-motion method within coupled-cluster theory. The extension of this method to two valence nucleons outside a closed-shell core is presented in Sec. \(\text{II B}\). We discuss and present our results in Sec. \(\text{III}\). Section \(\text{IV}\) contains our conclusions and an outlook for future work.
II. COUPLED-CLUSTER THEORY AND EQUATIONS-OF-MOTION FOR NUCLEI

A. Single-reference coupled-cluster theory

In this section we introduce the Hamiltonian that enters our calculation, together with a brief review of single reference coupled-cluster theory. We keep the presentation limited to those details that are required for the derivation of the two particle attached/removed (2PA/PR-EOM-CC) amplitudes presented in Sec. II B. The interested reader is referred to [4] for details.

We use throughout this work the convention that the indices \( i,j,k \ldots \) denote states below the Fermi level (holes), while the indices \( a,b,c \ldots \) denote states above the Fermi level (particles). For an unspecified state, the indices \( p,q,r \ldots \) are used. The amplitudes \( \langle i_1 \ldots i_n | j_1 \ldots j_m \rangle \) will be determined by solving the coupled-cluster equations. In the singles and doubles approximation we truncate the cluster operator as

\[
\hat{T} \approx \hat{T}_{\text{CCSD}} = \hat{T}_1 + \hat{T}_2 ,
\]

which defines the coupled-cluster approach with singles and doubles excitations, the so-called CCSD approximation. The unknown amplitudes result from the solution of the CCSD equations given by

\[
\langle \Phi | \hat{H} | \Phi \rangle = 0, \\
\langle \Phi | \hat{t}_{ij}^\dagger \hat{t}_{ij} | \Phi \rangle = 0 .
\]

(7)

The term

\[
\hat{H} = \exp (-\hat{T}) \hat{H} \exp (\hat{T}) = \left( \hat{H} \exp (\hat{T}) \right)_C ,
\]

(8)

is the similarity transform of the normal-ordered Hamiltonian. The state \( | \Phi \rangle \) is a Slater determinant that differs from the reference \( | \Phi_0 \rangle \) by holes in the orbitals \( ij \ldots \) and by particles in the orbitals \( ab \ldots \). The subscript \( C \) indicates that only connected diagrams enter.

Once the \( t_{ij}^\dagger \) and \( t_{ij} \) amplitudes have been determined from Eq. (7), the correlated ground-state energy is given by

\[
E_{\text{CC}} = \langle \Phi_0 | \hat{H} | \Phi_0 \rangle + E_0 .
\]

(9)

Here, \( E_0 \) denotes the vacuum expectation value with respect to the reference state. The coupled-cluster equations (7) show that the reference state \( | \Phi_0 \rangle \) is an eigenstate of the similarity transformed Hamiltonian (8) within the space of \( 1p-1h \) and \( 2p-2h \) excitations.

B. Equation-of-motion coupled-cluster theory for open-shell nuclei

In this work, our focus is on the development of coupled-cluster theory for truly open-shell nuclei, i.e., systems where no single reference can be constructed without breaking symmetries (such as rotational invariance). One could apply the CCSD method to the deformed (symmetry breaking) Hartree-Fock ground state of an open-shell nucleus. However, the restoration of angular momentum requires more than singles and doubles cluster excitations (see for example Ref. [15]) and is computationally expensive. Here, we wish to stay within the computationally inexpensive CCSD scheme.

Open-shell systems can be computed with multi-reference methods. In such an approach, many reference wave functions are included and treated on an equal footing. However, the loss of mathematical simplicity and transparency, and problems related to intruder states make these multi-reference approaches difficult to pursue. For a detailed discussion, we refer the reader to Ref. [34]. Equation-of-motion methods (see [26, 27] for recent reviews) avoid these problems as they exhibit the transparency and computational simplicity of single-reference coupled-cluster theory.
Within the EOM-CCSD approach, the states of the $A \pm 2$ open-shell nuclei are computed from the ground state of the A-body system as
\[
|\Psi^{(A\pm 2)}\rangle = \hat{R}^{(A\pm 2)}|\Psi^{(A)}\rangle = \hat{R}^{(A\pm 2)}|\Phi_0\rangle \quad (10)
\]
Here, $\hat{R}^{(A\pm 2)}$ is a particle removal or particle addition operator that generates an ($A \pm 2$)-body state from the A-body coupled-cluster wave function. The label $\mu$ identifies the quantum numbers (energy, angular momentum, ...) of the state of interest.

The operator $\hat{R}_\mu$, and the energies $E_\mu$ of the states of interest solve the eigenvalue problem [4, 23–27]
\[
(\hat{H} \hat{R}^{(A\pm 2)}|\mu\rangle) = \omega_\mu |\hat{R}^{(A\pm 2)}|\Phi_0\rangle . \quad (11)
\]
Here, the expression $(\hat{H} \hat{R}^{(A\pm 2)}|\mu\rangle)$ denotes all terms that connect the similarity transformed Hamiltonian $\hat{H}$ with the excitation operator $\hat{R}^{(A\pm 2)}$. The energy difference $\omega_\mu \equiv E_\mu - E_0$ is the excitation energy of the state $\mu$ in the nucleus $A \pm 2$ with respect to the ground state of the reference nucleus with mass $A$.

The operators $\hat{R}_\mu$ relevant for this work are (we drop the label $\mu$ for convenience)
\[
\hat{R}^{(A+2)} = \frac{1}{2} \sum_{iab} r^{ab} a_i^\dagger a_b^\dagger + \frac{1}{6} \sum_{iabc} r^{abc} a_i^\dagger a_j^\dagger a_k^\dagger + \ldots \quad (12)
\]
\[
\hat{R}^{(A-2)} = \frac{1}{2} \sum_{ij} r_{ij} a_i a_j + \frac{1}{6} \sum_{ijk} r_{ijk} a_i^\dagger a_k a_j + \ldots \quad (13)
\]
where the unknown amplitudes $r$ (subscripts and superscripts dropped) can be grouped into a vector that solves the eigenvalue problem of Eq. (11). The operator (12) consists of a $2p$-$0h$ term, a $3p$-$1h$ term, and in general up to an $(A + 2)p$-$Ah$ term. In this work, we will truncate the operator (12) at the $3p$-$1h$ level. Clearly, this truncation will only be a good approximation for states in the $(A \pm 2)$-body system that have a relatively simple structure built on the A-body nucleus. We will introduce two different truncations for the particle attached and the particle removed method, and identify them by the number of particle-hole excitations kept in the operator (12). A truncation after the first term in Eq. (12) is referred to as 2PA-EOM-CCSD($2p$-$0h$), while the truncation after the second term is denoted as 2PA-EOM-CCSD($3p$-$1h$). Similarly for 2PR-EOM-CCSD, we will use the abbreviations 2PR-EOM-CCSD($0p$-$2h$) and 2PR-EOM-CCSD($1p$-$3h$) for truncations after the first and second term in Eq. (13), respectively. Table I shows the excitation operators used in these truncation schemes.

We construct the matrix (i.e., the connected part of $\hat{H} \hat{R}$) of the eigenvalue problem of Eq. (11) diagrammatically. As usual, lines directed upwards represent particle states, while lines directed downwards represent hole states [4]. The horizontal lines represent the operators and we use a heavy and a wiggly line too differentiate the two operators in the composite diagrams. Table II shows the diagrams corresponding to the $r$ amplitudes, while the matrix elements of the similarity-transformed Hamiltonian are represented by the diagrams shown in Table IIII. These elements are well known and computed from the corresponding contractions of the cluster operator (8) with the Hamiltonian of Eq. (11) (see, for instance, Ref. [4]). They result from the construction of the operator (8) after the CCSD equations (7) have been solved.

Table IV shows the algebraic expressions for the matrix elements of the similarity-transformed Hamiltonian $\hat{H}$. For notational efficiency, some intermediate objects (amplitudes) are defined separately in Table V. In the numerical implementation, the storage of the similarity transformed Hamiltonian requires some memory. However, this modest cost in memory yields a significant reduction
The top three diagrams represent the one-body matrix elements of the similarity-transformed Hamiltonian $\tilde{H}$, the last four diagrams result from connecting a diagram in Table III with a diagram in Table IV. Figures 1 and 2 show the diagrams of $(\tilde{H} \tilde{R})_C$ for the two truncations of the 2PA-EOM-CCSD method. Let us briefly discuss some of these diagrams.

For 2PA-EOM-CCSD(2p-0h), the relevant diagrams correspond to the matrix element $\langle \Phi | \tilde{H} | \Phi \rangle$, i.e., they have two outgoing particle lines and consist of contractions of the similarity transformed Hamiltonians with $r^{ab}$. As an example, consider the diagram of Fig. 1(b).

It results from contracting the diagram of $r^{ab}$ (cf. Table IV) in computational cycles. For a detailed analysis we refer the reader to Refs. 33, 34.

In a diagrammatic language, the left-hand-side of the eigenvalue problem $\tilde{H} | \Phi \rangle = \epsilon | \Phi \rangle$ consists of all topologically different diagrams that result from connecting a diagram from Table II with a diagram in Table III. Figures I and II show the diagrams of $(\tilde{H} \tilde{R})_C$ for the two truncations of the 2PA-EOM-CCSD method. Let us briefly discuss some of these diagrams.

For 2PA-EOM-CCSD(2p-0h), the relevant diagrams correspond to the matrix element $\langle \Phi | \tilde{H} | \Phi \rangle$, i.e., they have two outgoing particle lines and consist of contractions of the similarity transformed Hamiltonians with $r^{ab}$. As an example, consider the diagram of Fig. 1(b). It results from contracting the diagram of $r^{ab}$ (cf. Table IV) in computational cycles. For a detailed analysis we refer the reader to Refs. 33, 34.
Figure 1: Diagrams corresponding to the matrix element $\langle \Phi^{ab}|(\hat{H}\hat{R})_C|\Phi_0\rangle$ for the 2PA-EOM-CCSD(2p-0h) amplitude equation. All diagrams are constructed by contracting a diagram from Table II with a diagram from Table III. Only diagrams that satisfy the topological form of $\langle \Phi^{ab}|(\hat{H}\hat{R})_C|\Phi_0\rangle$, with two external particle lines in the upper part of the diagram and no external lines in the bottom part of the diagram, are selected.

Figure 2: Diagrams corresponding to the matrix elements $\langle \Phi^{ab}|(\hat{H}\hat{R})_C|\Phi_0\rangle$ (a-d) and $\langle \Phi^{abc}_{ijkl}|(\hat{H}\hat{R})_C|\Phi_0\rangle$ (e-k) for the 2PA-EOM-CCSD(3p-1h) amplitude equation. Let us turn to the diagrams for two-particle removal. Figures 3 and 4 show the diagrams of $(\hat{H}\hat{R})_C$ for the two truncations of the 2PR-EOM-CCSD method. Here, one needs topologies of the form $\langle \Phi_{ij}|(\hat{H}\hat{R})_C|\Phi_0\rangle$ and $\langle \Phi_{ijk}|(\hat{H}\hat{R})_C|\Phi_0\rangle$ using the $r_{ij}$ and $r_{ijk}$ diagrams in Table III.

The algebraic expressions corresponding to these diagrams are derived according to the standard rules, see for example Ref. [4], and shown in Table VI. The computational cost is $n_u^2 n_o$ for the 2PA-EOM-CCSD(3p-1h) method, and the most expensive diagram is shown in Fig. 2(h). Likewise, the most expensive diagram for the 2PR-EOM-CCSD(1p-3h) is shown in Fig. 4(i) and requires of the order of $n_u^2 n_o$ operations. Here, $n_u$ is the number of unoccupied orbitals (equal to the size of the valence space), and $n_o$ is the number of occupied orbitals in the reference state. For comparison, the computational costs of the single-reference CCSD and CCSDT meth-
The eigenvalue problem of Eq. (11) for the two-particle system thus becomes

\[ \langle \Phi_{ij} | (\hat{H} R) | \Phi_0 \rangle = -P(ij) H^m_{ij} r_{im} + \frac{1}{2} H^{mn}_{ij} r_{mn} \]

\[ \langle \Phi_{ijk} | (\hat{H} R) | \Phi_0 \rangle = -P(ijk) H^{mn}_{ijk} r_{im} + \frac{1}{2} H^a_{ijk} r_{mn} \]

Table VI: Algebraic expressions for the 2PA/2PR-EOM-CCSD(2p-0h)/0p-2h) and 2PA/2PR-EOM-CCSD(3p-1h/1p-3h) approximations. All terms are defined in Table VII.

The eigenvalue problem of Eq. (11) for the two-particle attached system thus becomes

\[ \langle \Phi_{ij} | (\hat{H} R) | \Phi_0 \rangle = \omega r_{ij} \]

\[ \langle \Phi_{ijk} | (\hat{H} R) | \Phi_0 \rangle = \omega r^a_{ijk} \]  \quad (14)

Here, the left-hand-side is a linear function of the “vector” \( R = (r^{ab}, r^{abc}) \) of amplitudes (see Table VII) and constitutes a matrix-vector product. Note that the two equations are coupled and constitute a single eigenvalue problem for the 3p-1h truncation. Likewise, we find for the two-particle removed problem

\[ \langle \Phi_{ij} | (\hat{H} R) | \Phi_0 \rangle = \omega r_{ij} \]

\[ \langle \Phi_{ijk} | (\hat{H} R) | \Phi_0 \rangle = \omega r^a_{ijk} \]  \quad (15)

We are usually only interested in the few lowest eigenvalues of Eq. (11). For this purpose, we use the Arnoldi Method for asymmetric eigenvalue problems, see for example Ref. [37] and references therein. This method is based on repeated applications of the matrix-vector product \((\hat{H} R)_C\). Specifically, our numerical implementation uses the ARPACK software [38] package. The expressions in Table VII can thus be used to solve the eigenvalue problem directly. In this paper, we employ the \(m\)-scheme basis for the vectors \( R = (r^{ab}, r^{abc}) \) of the 2PA. Within this scheme, we are limited to small model spaces. However, in this work we are mainly interested in testing the newly developed methods and in gauging the accuracy of the employed cluster truncation through comparisons with exact diagonalization. Note that the similarity-transformed Hamiltonian \( \hat{H} \) exhibits the symmetries of the underlying Hamiltonian. In the \(m\)-scheme basis, we can classify states by their projections \( J_z \) of angular momentum and \( T_z \) of isospin and their parity. Although the solutions will have a good total angular momentum \( J^2 \), we will not be able to exploit this symmetry in the \(m\)-scheme basis.

The FCI method we employ [39] is limited to relatively small model spaces. Note, however, that we are also working on an angular-momentum-coupled implementation [40] of 2PA/2PR-EOM-CCM. This will allow us to exploit rotational symmetries and give us access to much larger model spaces. These spaces are well beyond the reach of present full diagonalization methods.

### III. RESULTS

For the proof-of-principle study we consider the helium isotopes \(^3\text{He}\) to \(^6\text{He}\). Here, \(^3\text{He}\) and \(^5\text{He}\) are viewed as one neutron removed from and attached to \(^4\text{He}\), respectively. In a shell-model picture, \(^6\text{He}\) is a truly open-shell nucleus with two valence neutrons in the \(p_{1/2}\) shell. Thus, a single-reference Slater determinant may not be a good starting point for coupled-cluster calculations of this nucleus, and it seems advantageous to describe this six-nucleon system as two (halo) neutrons added to the \(^4\text{He}\) core. We do not present results for \(^2\text{He}\) using the 2PR-EOM-CCSD approach; a analysis of this method will be presented elsewhere [40].

For our calculations we use a realistic nucleon-nucleon (NN) potential derived from chiral effective field theory [41–45] at order N\(^3\)LO using interaction matrix elements from Ref. [45]. The matrix elements of the bare interaction employ a cutoff at \(\Lambda = 500\text{ MeV}\). The short-range parts of the interaction are removed via the similarity renormalization group transformation [40] with a cutoff at \(1.9\text{ fm}^{-1}\). We use the spherical harmonic oscillator with the oscillator frequency \(h\omega = 24\text{ MeV}\) as our single-particle basis. Our model space consists of five major oscillator shells, with maximum orbital angular momentum \(l_{\text{max}} = 2\). This results in a total of 76 single-particle states for neutrons and protons. We neglect three-body and four-body interactions. This leads to missing many-body physics but is not relevant for our proof-of-principle
We calculate the ground state energies of the A= 3-6 helium isotopes. For $^6$He we also compute the first 2$^+$ state and the expectation values of the total angular momentum. In addition, we also discuss the first 1$^-$ excited state, as an example where the 3p-1h truncation fails. We compare the equation-of-motion (EOM) approach to the FCI method and to three different single-reference coupled-cluster approximations. Recall that within the $m$-scheme, one can also compute open-shell nuclei directly with the coupled-cluster method without resorting to EOM techniques. While such a direct approach may suffer from the lack of a good reference state (as is the case for the ground state of $^6$He), the inclusion of more and more clusters must converge to the FCI results. Comparing the EOM-CC approach to these single-reference coupled-cluster calculations allows us to gauge the efficiency of the various coupled-cluster approximations.

The direct coupled-cluster calculations employ the CCSD approximation described above, the CCSDT approximation (that includes triples clusters) and the CCSDT-1 approach which includes some of the 3p-3h clusters of the full CCSDT approximation. For the EOM calculations, we use the CCSD wave function of $^4$He as the reference wave function and employ the intrinsic Hamiltonian of Eq. (1) with $A = 3, 5, 6$ for $^{3,5,6}$He, respectively. We compare all results to an exact calculation done with the FCI approach, using the same interaction and model space. The ground-state energies of $^{3-5}$He are shown in Table VIII. For $^3$He, CCSDT becomes an exact method and agrees with FCI. Here, the single-reference coupled-cluster calculations are superior to the EOM-CCSD approach. Evidently, the weakly bound nucleus $^3$He is not well described as a neutron removed from the tightly bound $^4$He. This means in turn that correlations beyond one-particle-two-hole excitations play a non-negligible role. For $^4$He, the EOM-CCSD approach is identical to CCSD. Here, triples corrections and full triples represent a significant improvement over CCSD, bringing the results close to the FCI ones. For $^5$He, the EOM-CCSD is superior to a single-reference CCSD calculation and competes well with the computationally more expensive triples correction CCSDT-1. Clearly, the valence neutron in $^5$He is weakly correlated with the strongly bound $^4$He core, and the PA-EOM-CCSD approach captures this state very well.

We turn to the truly open-shell nucleus $^6$He and show our results in Table VIII. Here, the 2PA-EOM-CCSD (3p-1h) approach is clearly superior to the single-reference coupled-cluster approaches, as it reproduces the energy and the spin of the ground state to a very good approximation. For the computation of the spin within the single-reference approaches, we compute the expectation value $\langle J^2 \rangle$ within the Hellmann-Feynman theorem, and define $\langle J \rangle$ from the relation $\langle J^2 \rangle = \langle J \rangle (\langle J \rangle + 1)$. A direct computation within the CCSD or CCSDT approximations can only be based on a symmetry-breaking reference state. Clearly, 2-particle-2-hole excitations (CCSD) cannot restore the symmetry, and we find that $\langle J \rangle = 0.78$ for the ground state of $^6$He. Adding 3-particle-3-hole excitations (CCSDT approximation) almost restores the rotational symmetry, but some correlation energy is still missing. In the 2PA-EOM-CCSD approach, however, the rotational symmetry is preserved throughout the calculation, and we obtain a very good approximation of the energy at a relatively low computational cost. As expected, the 2PA-EOM-CCSD (2p-0h) approach is less accurate than the 3p-1h approximation, since it is unable to account simultaneously for the correlations within the three-body system consisting of the two valence neutrons and the $^4$He core. The ground state and the first excited 2$^+$ state of $^6$He are both dominated by a configuration with two neutrons in the $p_{3/2}$ orbit. This is consistent with the shell-model picture of this nucleus. For the excited 2$^+$ state, the CCSD, CCSDT and CCSDT-1 methods result in the correct value of the angular momentum due to the choice of reference state.

Let us define the fraction $\sigma_{\text{corr}}$ of the correlation energy

| $^3$He | $^4$He | $^5$He |
|-------|-------|-------|
| CCSD  | $-6.624$ | $-27.468$ | $-22.997$ |
| CCSDT-1 | $-6.829$ | $-27.600$ | $-23.381$ |
| CCSDT  | $-6.911$ | $-27.619$ | $-23.474$ |
| EOM-CCSD | $-6.357$ | $-27.468$ | $-23.382$ |
| FCI    | $-6.911$ | $-27.640$ | $-23.640$ |

Table VII: Ground-state energies (in MeV) for $^3$He, $^4$He and $^5$He, calculated with coupled-cluster methods truncated at the 2-particle-2-hole (CCSD) level, 3-particle-3-hole (CCSDT) and a hybrid (CCSDT-1) where a small subset of the leading diagrams in CCSDT are included. For the EOM-CCSD approach, truncations have been made at the 1-particle-2-hole level, the 2-particle-2-hole level, and the 2-particle-1-hole level for $^3$He, $^4$He and $^5$He respectively. The energies are compared to the exact full configuration interaction (FCI).

| $^3$He | $^4$He | $^5$He |
|-------|-------|-------|
| CCSD  | $-22.732$ | $-20.905$ | $0.78$ | $2$ |
| CCSDT-1 | $-24.617$ | $-21.586$ | $0.25$ | $2$ |
| CCSDT  | $-24.530$ | $-21.786$ | $0.01$ | $2$ |
| 2PA-EOM-CCSD(2p-0h) | $-21.185$ | $-18.996$ | $0$ | $2$ |
| 2PA-EOM-CCSD(3p-1h) | $-24.543$ | $-21.634$ | $0$ | $2$ |
| FCI | $-24.853$ | $-21.994$ | $0$ | $2$ |

Table VIII: Energies (in MeV) for the ground state and first excited state of $^6$He and the expectation value of the total angular momentum, calculated with coupled-cluster methods truncated at the 2-particle-2-hole (CCSD) level, 3-particle-3-hole (CCSDT) and a hybrid (CCSDT-1) where the 3-particle-3-hole amplitudes are treated perturbatively. The 2PA-EOM-CCSD results are calculated with a truncation at the 2-particle-0-hole (2PA-EOM-CCSD(2p-0h)) level and at the 3-particle-1-hole (2PA-EOM-CCSD(3p-1h)) level. All energies are compared to full configuration interaction (FCI) results.
as
\[ \sigma_{\text{corr}} = \frac{E_{2\text{PA}} - E_0}{E_{\text{FCI}} - E_0}. \] (16)

Here \( E_{2\text{PA}} \) is the energy from 2PA-EOM-CCSD, \( E_{\text{FCI}} \) is the energy from the FCI, while \( E_0 \) is the energy expectation of the uncorrelated reference state \( |\Phi_0\rangle \). Both \( E_{2\text{PA}} \) and \( E_{\text{FCI}} \) are shown in Table VIII, and \( E_0 = -16.807 \text{ MeV} \). We also compute the norm
\[ \mathcal{N} = \sum_{ab} |r_{ab}|^2 + \sum_{abi} |r_{abi}|^2. \] (17)

The normalized squared weights
\[ \rho_1^2 \equiv \mathcal{N}^{-1} \sum_{ab} |r_{ab}|^2 \]
\[ \rho_2^2 \equiv \mathcal{N}^{-1} \sum_{abi} |r_{abi}|^2, \] (18)
fulfill \( \rho_1^2 + \rho_2^2 = 1 \) and measure the importance of the 2p-0h and 3p-1h amplitudes, respectively.

Table IX shows the relative correlation energy \( \sigma_{\text{corr}} \) and the relative weights \( \rho_1 \) and \( \rho_2 \) of 2PA-EOM-CCSD(3p-1h) for the three lowest states with quantum numbers \( 0^+, 1^- \) and \( 2^+ \) of \(^6\)He, respectively. We see that 2PA-EOM-CCSD accounts for more than 90\% of the correlation energy for the \( 0^+ \) and \( 2^+ \) states, and that most of the weight is carried by the 2p-0h amplitude. The \( 1^- \) state, however, is not very accurately reproduced and much of the correlation energy is lacking. For this state, the 2PA-EOM-CCSD(3p-1h) energy is \( E_{1^-} = -20.95 \text{ MeV} \) and deviates considerably from the full CI result \( E_{1^-} = -23.26 \text{ MeV} \). Consistent with this picture is the large weight carried by the 3p-1h amplitudes. For a converged computation, one would presumably also need to include 4p-2h or higher clusters.

Table IX: The relative correlation energy \( \sigma_{\text{corr}} \) defined in Eq. (16) for the lowest states with quantum numbers \( J^\pi = 0^+, 1^- \) and \( 2^+ \) in \(^6\)He, respectively. The relative weights \( \rho_1 \) and \( \rho_2 \) of the 2p-0h and 3p-1h amplitudes, respectively, are defined in Eqs. (18). The results of this section show that states in open-shell nuclei that exhibit a simple structure imposed on a correlated core can be well described by the EOM method. Moreover, the EOM wave function preserves the symmetries of the Hamiltonian and is expected to be useful in the computation of matrix elements besides the energy. Taking into account the low computational cost of the EOM-CC methods as compared to the full CCSDT approach, our EOM approach is clearly well suited for these selected states. For states where more complicated particle-hole excitations are prominent, the various truncation schemes discussed here are insufficient and we will need additional correlations in the EOM operator to reach satisfactory results. The ground state of \(^5\)He and the first excited \( 1^- \) state of \(^5\)He discussed above, are examples in case.

| \( J^\pi \) | \( \sigma_{\text{corr}} \) | \( \rho_1 \) | \( \rho_2 \) |
|---|---|---|---|
| \( 0^+ \) | 0.96 | 0.84 | 0.16 |
| \( 1^- \) | 0.64 | 0.34 | 0.66 |
| \( 2^+ \) | 0.93 | 0.81 | 0.19 |

IV. CONCLUSIONS AND FUTURE PERSPECTIVES.

We have developed and implemented the equation-of-motion coupled-cluster method and performed microscopic calculations of helium isotopes with up to two valence nucleons outside the closed-shell alpha particle. The comparison with full configuration interaction calculations shows that the equation-of-motion coupled-cluster methods yield accurate results for sufficiently simple states. The open-shell nucleus \(^6\)He, for instance, can be viewed and computed as two weakly correlated neutrons attached to the correlated core of \(^4\)He. The proof-of-principle calculations were performed in a reduced model space for a comparison with results from exact diagonalizations. We are working on implementing our formalism for the two-particle attached and removed equation-of-motion coupled-cluster methods in an angular momentum coupled basis. This will allow us to employ much larger model spaces. With this improvement, the first-principles computation of semi-magic nuclei, single-particle energies, and effective two-particle interactions for the nuclear shell-model can be addressed.

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