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Calculations of non-adiabatic couplings within equation-of-motion coupled-cluster framework: Theory, implementation, and validation against multi-reference methods

Shirin Faraji,1,2,a) Spiridoula Matsika,3 and Anna I. Krylov1
1Department of Chemistry, University of Southern California, Los Angeles, California 90089, USA
2Zernike Institute for Advanced Materials, Groningen, The Netherlands
3Department of Chemistry, Temple University, Philadelphia, Pennsylvania 19122, USA

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We report an implementation of non-adiabatic coupling (NAC) forces within the equation-of-motion coupled-cluster with single and double excitations (EOM-CCSD) framework via the summed-state approach. Using illustrative examples, we compare NAC forces computed with EOM-CCSD and multi-reference (MR) wave functions (for selected cases, we also consider configuration interaction singles). In addition to the magnitude of the NAC vectors, we analyze their direction, which is important for the calculations of the rate of non-adiabatic transitions. Our benchmark set comprises three doublet radical-cations (hexatriene, cyclohexadiene, and uracil), neutral uracil, and sodium-doped ammonia clusters. When the characters of the states agree among different methods, we observe good agreement between the respective NAC vectors, both in the Franck-Condon region and away. In the cases of large discrepancies between the methods, the disagreement can be attributed to the difference in the states’ character, which, in some cases, is very sensitive to electron correlation, both within single-reference and multi-reference frameworks. The numeric results confirm that the accuracy of NAC vectors depends critically on the quality of the underlying wave functions. Within their domain of applicability, EOM-CC methods provide a viable alternative to MR approaches. Published by AIP Publishing. https://doi.org/10.1063/1.5009433

I. INTRODUCTION

Transitions between electronic states play a key role in excited-state processes. They are essential in photochemistry, photobiology, and the operation of photovoltaic materials. These transitions are responsible for radiationless relaxation, a process in which electronic energy is converted into nuclear motions. Radiationless relaxation protects living organisms from the damaging effects of UV radiation by converting energy imparted by solar photons into heat and efficiently quenching unwanted excited-state chemistry.1 Radiationless relaxation is a key step in vision—it allows photoexcited rhodopsin’s chromophore to return to the initial state, upon completing the isomerization process, so the cycle can be repeated over and over again. In other instances, radiationless relaxation can be viewed as a nuisance. For example, in imaging applications that use fluorescent tags, radiationless relaxations compete with fluorescence (radiative relaxation), leading to reduced optical output.2 In solar energy harvesting, radiationless relaxation is also often a parasitic process as it competes with charge separation or reactions producing solar fuels.3 However, non-adiabatic transitions can be exploited to increase the efficiency of solar cells via multi-exciton generation. In organic photovoltaics, this process is called singlet fission;4,5 it involves a non-adiabatic transition between the initially excited bright state and a dark state of multi-exciton character.6

Within the Born-Oppenheimer separation of nuclear (R) and electronic (r) coordinates, the exact molecular wave function is represented by the following ansatz:

$$\Psi_K(r, R) = \sum I \Phi_I(r; R) \xi^K_I(R),$$

where $\Phi_I(r; R)$ and $\xi^K_I(R)$ denote electronic and nuclear wave functions, respectively. The electronic wave functions are defined as the solutions of the electronic Schrödinger equation,

$$H^{el} \Phi_I(r; R) = E_I(R) \Phi_I(r; R),$$

where the Hamiltonian, which includes all terms (electron kinetic energy $T_e$ and Coulomb interactions between all charged particles) except nuclear kinetic energy $T_n$, depends on nuclear positions via $V_{en}$. Equation (2) is solved at each nuclear configuration, giving rise to eigenstates and eigenenergies depending parametrically on $R$; at each $R$, the electronic wave functions $\Phi_I$ form an orthonormal basis. Ansatz (1) gives rise to the following equations for nuclear wave functions:

$$\langle T_n + E_I(R) - E_K \rangle \xi^K_J = \sum A \left( \sum J \langle \Phi_I | V^2_{R_J} \Phi_I \rangle \xi^K_J + 2 \sum J \langle \Phi_I | V_{R_J} \Phi_I \rangle \nabla_{R_J} \xi^K_J \right),$$

a)Author to whom correspondence should be addressed: shirin.faraji@gmail.com

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where $A$ denotes a particular nucleus and $E_K$ is the total energy of wave function (1). Within the Born-Oppenheimer (or adiabatic) approximation, the terms on the right-hand side of Eq. (4) are neglected giving rise to separable wave function, $\Phi_I e^{E_K}$. In this case, the nuclear motions are confined to a single potential energy surface, $E_K(R)$, and nuclear wave functions corresponding to each electronic state are independent (or uncoupled) from nuclear wave functions corresponding to other electronic states. Physically, this means that electrons adjust to moving nuclei’s positions instantly so that the electronic state of the system never changes. Electronic transitions are only possible beyond adiabatic approximation. They arise due to the terms on the right-hand side of Eq. (4), which couple nuclear motions on different potential energy surfaces (PESs). These terms originate in the parametric dependence of the electronic wave functions on the nuclear coordinates and are called derivative couplings. They become large when the changes in electronic wave functions due to nuclear motions are large, which is quantified by the derivative with respect to nuclear coordinates. These terms lead to electronic inertia so that electrons are no longer following nuclei adiabatically; rather, upon some nuclear motions, electrons can remain frozen at their configuration leading to the change of electronic state. Of the two derivative terms, only the first term contributes to the diagonal Born-Oppenheimer energy correction, which is computed when high-accuracy thermochemical data are desired. This term may cause discontinuous behavior in calculations of nonadiabatic dynamics in adiabatic representation and is omitted in the original surface-hopping model. Recently, Meek and Levine have shown that this term can be accurately accounted for within multiple-spawning calculations on adiabatic surfaces by using locally diabatized Gaussian basis functions. Interestingly, numeric simulations of non-adiabatic relaxation in ethylene have shown that the results are nearly identical whether this term is fully accounted for or completely omitted.

In this paper, we focus on the term called derivative coupling or non-adiabatic coupling (NAC),

$$d_{IJ} \equiv \langle \Phi_I | \nabla_R \Phi_J \rangle. \quad (5)$$

As clearly seen from Eq. (5), NAC is a 3N vector, where $N$ is the number of nuclei. $d_{IJ}$ couples nuclear motions in electronic states $I$ and $J$ (for real-valued wave functions, the diagonal NAC elements vanish, $d_{II} = 0$, by virtue of the normalization condition). As one can see from Eq. (4), the total magnitude of the coupling is given by the scalar product of $d_{IJ}$ and the nuclear momentum, $\nabla_R$. Thus, the probability of electronic transitions depends not only on the electronic coupling $d_{IJ}$ but also on how fast the nuclei are moving as well as the direction of their motion. Obviously, for infinitesimally slow nuclei, the coupling vanishes and the adiabatic approximation is restored. The symmetry of $d_{IJ}$ is determined by the symmetries of $\Phi_I$ and $\Phi_J$ and is equal to the product of the two irreps. Consequently, the two states are only coupled by the nuclear motions of that symmetry (otherwise, the scalar product vanishes). For example, in a $C_2v$ molecule, electronic states from the $A_2$ and $B_1$ irreps are coupled by $B_2$ vibrations (and the symmetry of $d_{IJ}$ vector is $B_2$). An important aspect of non-adiabatic dynamics in polyatomic systems is that, in contrast to the one-dimensional case, one can distinguish between molecular motions inducing the transition (those along $d_{IJ}$) and motions that modulate the magnitude of $d_{IJ}$ (those along which the wave-function composition changes significantly); for concrete examples, see Refs. 6 and 13.

For the exact wave functions, NAC vector $d_{IJ}$ can be expressed as a matrix element of the derivative of the electronic Hamiltonian, $H^{el}$,

$$d_{IJ} = \frac{\hbar}{E_J - E_I}, \quad (6)$$

$$h_{IJ} = \langle \Phi_J | \nabla_R H^{el} | \Phi_I \rangle. \quad (7)$$

Thus, one can expect large values of NAC when (i) the electronic Hamiltonian depends strongly on nuclear positions (large $h_{IJ}$) and (ii) the energy gap between the two states is small (as in the regions of PESs’ near-degeneracy). The two vectors, $h_{IJ}$ and $d_{IJ}$, are parallel. Vector $h_{IJ}$, called the non-adiabatic force matrix element, can be described as an interstate generalization of the nuclear gradient,

$$G_I \equiv \nabla_R (\langle \Psi_I | H^{el} | \Psi_I \rangle) = \langle \Psi_I | \nabla_R H^{el} | \Psi_I \rangle. \quad (8)$$

The second equality holds only when the Hellman-Feynman condition is satisfied.

This connection to the nuclear gradient can be exploited for practical calculations of NACs based on the following observation. Consider two electronic states, $\Phi_I$ and $\Phi_J$, and a fictitious summed state $\Phi_{(I+J)} \equiv \Phi_I + \Phi_J$. The energy (defined as an expectation value) of $\Phi_{(I+J)}$ equals $E_I + E_J$, but $\Phi_{(I+J)}$ is not an eigenstate—thus, the Hellman-Feynman theorem does not hold for it. One can define (and compute) Hellman-Feynman gradients for all three states, which are $G_K \equiv \langle \Phi_K | \nabla_R H | \Phi_K \rangle$, $K = I, J, I + J$. At the same time,

$$G_{I+J} \equiv \langle \Phi_{(I+J)} | \nabla_R H | \Phi_{(I+J)} \rangle = (\langle \Phi_I | \nabla_R H | \Phi_J \rangle + \langle \Phi_J | \nabla_R H | \Phi_I \rangle) + \langle \Phi_I | \nabla_R H | \Phi_I \rangle + \langle \Phi_J | \nabla_R H | \Phi_J \rangle = G_I + G_J + \langle \Phi_I | \nabla_R H | \Phi_J \rangle + \langle \Phi_J | \nabla_R H | \Phi_I \rangle. \quad (9)$$

Thus

$$h_{IJ} \equiv \langle \Phi_I | \nabla_R H | \Phi_J \rangle = \frac{1}{2} \left( G_{I+J} - G_I - G_J \right) \quad (10)$$

and, consequently,

$$d_{IJ} \equiv \frac{G_{I+J} - G_I - G_J}{2(E_J - E_I)}. \quad (11)$$

Equation (10) provides an insight into the nature of nonadiabatic force matrix element. It is the non-Hellman-Feynman part of the gradient of the summed state, i.e., the difference between the expectation value of $\nabla_R$ and the sum of the gradients of the two states. When the two states belong to different irreps, this is the part of the summed-state gradient which is not fully symmetric but has symmetry of the product of the two irreps.

Because the magnitude of NACs becomes large when the energy gap between the two electronic states is small, the $F - 2$ dimensional seams along which the surfaces cross are of special significance in non-adiabatic processes ($F$ denotes
the number of internal degrees of freedom; for a non-linear molecule, \( F = 3N - 6 \). The minimum-energy crossing points\(^{15-17}\) along these seams (called MECPs) play a role akin to transition states in transition-state theory (TST)—their energetic accessibility is one of the factors controlling the rate of non-adiabatic transitions. In the non-adiabatic extension of TST, MECP structures are used to compute the density of states, electronic couplings, and Arrhenius factors that together define the rate of non-adiabatic reaction flow. The degeneracy seams are also called conical intersections, as the topology of the two PESs around such seams has a conical shape. The conical shape arises because the degeneracy between the two states, \( I \) and \( J \), is lifted along the following two coordinates,\(^{17,18}\) \( h_{IJ} \) [defined by Eq. (29)] and \( g_{IJ} \equiv G_I - G_J \). These two orthogonal vectors give rise to the \( g - h \) plane defining the conical intersection (Fig. 1). Thus, \( h \) is needed for characterizing the conical intersection and for computing the rate of non-adiabatic transitions (we note that efficient algorithms\(^{19,20}\) for locating MECPs also use \( h \)).

What is needed for computing MECPs and NACs by electronic structure methods? First, the method should be capable of describing multiple electronic states on the same footing, so the relative state energies and, consequently, the degeneracy seams are correctly described. Second, the method should be able to handle multi-configurational wave functions in a sufficiently flexible way, correctly reproducing the changes in the adiabatic wave functions upon changes in nuclear geometries. Third, the expressions for the exact states should be reformulated for approximate wave functions using techniques similar to those used in analytic gradient and response theory. Pioneering studies of conical intersections and non-adiabatic phenomena employed multi-reference configuration interaction (MRCI) method.\(^{21-25}\) The formalism and computer implementations for the calculation of NACs using complete active space self-consistent field (CASSCF) wave functions\(^{26,27}\) and within multi-state (CASSCF corrected by second-order perturbation theory) framework\(^{28,29}\) have also been reported. Calculations of MECPs and NACs by using more economical methods, configuration interaction singles (CISs) and time-dependent density functional theory (TDDFT), have been presented.\(^{30-34}\)

These single-reference methods use linear parameterization of target states and are, therefore, capable of describing conical intersections between excited states in the vicinity of the Franck-Condon region, where the reference state retains its single-configurational character. However, the intersections between the ground and excited states cannot be treated by these approaches, owing to single-determinantal reference state.\(^{35}\) This issue is addressed by the spin-flip (SF) approach in which a high-spin reference is used, and the target states (both the ground state and excited states) are described as spin-flipping excitations.\(^{36-39}\) Recently, calculations of NACs and MECPs using SF-CIS and SF-DFT have been reported\(^{34,40,41}\) and used in \textit{ab initio} surface-hopping dynamics.\(^{42}\)

Equation-of-motion coupled-cluster (EOM-CC) methods\(^{43-45}\) share many common features with CI. The target states are described by the CI-like excitation operators \( R \) from the CC reference state,

\[
\Phi = R^T |0\rangle,
\]

where \( |0\rangle \) is the reference determinant and amplitudes of \( T \) satisfy CC equations for the reference state. Amplitudes \( R \) are found by solving a non-Hermitian eigenproblem,

\[
\bar{H} R^K = E_K R^K, \\
\bar{H} \equiv e^{-T} H e^T.
\]

Depending on the specific choice of the reference and excitation operators, different types of target states can be accessed, as illustrated in Fig. 2. Most commonly used variant is EOM for excitation energies in which the spin- and particle-conserving excitation operators are used, and the reference corresponds to the ground state of the system.\(^{36-38}\) EOM operators that change the number of electrons (or their spins) open access to various open-shell wave functions.\(^{36,49-53}\) EOM-CC is capable of describing many types of multi-configurational wave functions within single-reference formalism.\(^{43,54}\) Because of

\[\text{FIG. 1.} \quad \text{Two PESs can intersect in the } 3N - 8 \text{ space, which is the dimensionality of conical intersection. The degeneracy is lifted along the } g \text{ and } h \text{ vectors. Vector } g \text{ points in the direction of maximal energy splitting, whereas } h \text{ points in the direction of the maximal non-adiabatic interaction.}\]

\[\text{FIG. 2.} \quad \text{Different EOM models are defined by choosing the reference } (\Phi_0) \text{ and the form of the operator } R. \text{ EOM-EF allows access to electronically excited states of closed-shell molecules, EOM-IP and EOM-EA can describe doublet target states, and EOM-SF describes multi-configurational wave functions encountered in bond-breaking and transition states.}\]
its multi-state nature, EOM-CC can treat iterations and crossings between multiple interacting states. Thus, EOM-CC is an attractive platform for computing MECPs and NACs.

Formulation of NACs within EOM-CC theory have been described for EE (excitation energy) and IP (ionization potential) variants. Here we extend the theory by including the EOM-EA (EOM for electron affinity) method. We also extend the list of examples for EOM-EE and EOM-IP methods and provide detailed comparisons with CASSCF and MRCI. Our focus is on NACs rather than MECPs, as the former has not received as much attention as the latter. The outline of this paper is as follows. Section II presents the formalism for computing NAC forces for inexact states and within non-Hermitian framework, as needed for EOM-CC wave functions. Section III summarizes computational details. The benchmark results and analysis are presented in Sec. IV.

II. THEORY

A. NACs for inexact states

For inexact states, calculations of properties, such as nuclear gradients, require extra steps, relative to simply computing an expectation value of the appropriate derivative of the Hamiltonian using approximate wave functions. Additional terms (often referred to as “Pulay terms”), which include derivatives of the wave function parameters, arise because the Hellman-Feynman theorem is not satisfied. The efficient evaluation of these terms can be derived using the Lagrangian approach.

For example, in CC/EOM-CC theory, the analytic gradient can be formulated as a contraction of derivatives of the Hamiltonian with one- and two-particle density matrices which include an unrelaxed part (the equivalent of the expectation-value calculation) and amplitude and orbital-response parts. The latter two contributions are computed by solving auxiliary equation responses. This formalism has been extended by Stanton and co-workers for calculating NAC forces, which they refer to as quasiadiabatic coupling strength; see Eq. (57) in Ref. 55.

\[
\lambda_{IJ}^x = \langle 0 | L_I e^{-T} \frac{\partial H}{\partial x} e^{-T} R_J | 0 \rangle + \langle 0 | L_I e^{-T} H e^{-T} \frac{\partial T}{\partial x} R_J | 0 \rangle \\
+ \langle 0 | L_I e^{-T} \frac{\partial T}{\partial x} H e^{-T} R_J | 0 \rangle,
\]

(15)

where \( x \) denotes a particular Cartesian coordinate with respect to which the derivative is taken. Here the first term corresponds to the full derivative of the Hamiltonian (i.e., including the derivatives of the molecular orbitals), and the second two terms include EOM-CC amplitude response contributions. The full derivation of the EOM-CC NAC force and the comparison with quasiadiabatic couplings \( \lambda \) defined in Ref. 55 is given in Appendix A.

In the CI formalism, it is convenient to cast the gradient and NAC expressions in the generalized Hellman-Feynman form, where \( \mathcal{H} \) denotes the projected Hamiltonian (e.g., in the case of CIS, it is the Hamiltonian projected into the space of singly excited determinants) and superscript \( x \) denotes the full derivative (which includes the derivative of the Hamiltonian as well as the derivative of the projector operator). Given the CI-like form of the EOM-CC equations, the expression for the NAC force in EOM-CC theory can be cast in the same form,

\[
h_{IJ}^x = \langle \Phi_I | \mathcal{H}^x | \Phi_J \rangle,
\]

(16)

where \( \mathcal{H} \) is the similarity transformed Hamiltonian projected onto the space of the reference, singly, and doubly excited determinants, and \( h_{IJ}^x \) denotes the \( x \) component of \( h_{IJ} \). By taking the derivative, one arrives to formally exact \( h_{IJ} \), which are identical to a calculation based on the non-Hermitian generalization of Eq. (10).

For approximate wave functions, the connection between the NAC vector (\( d_{IJ} \)) and the NAC force (\( h_{IJ} \)) is not as simple

\[
d_{IJ} = d_{IJ}^c + d_{IJ}^\phi,
\]

(18)

\[
d_{IJ} = \frac{h_{IJ}}{E_I - E_J}.
\]

(19)

The orbital-derivative part, \( d_{IJ}^\phi \), can be evaluated, giving rise to the “proper” \( d_{IJ} \), which has been done within MRCI and EOM-CC frameworks. The expression for \( d_{IJ}^c \) is given in Appendix B. Interestingly, full \( d_{IJ} \) computed for approximate wave functions is not translationally invariant. That is, one can induce non-adiabatic transitions by simply moving the entire molecule around, which is unphysical and problematic for dynamics simulations. Subotnik and co-workers have developed a correction restoring translational invariance of \( d_{IJ} \) within the CIS framework; the correction was called ETFs (electronic translation factors). Later, Zhang and Herbert have shown that the correction is equal to the orbital contribution part and cancels it out exactly. By virtue of Eq. (10), it is clear that \( h_{IJ} \) and, consequently, \( d_{IJ}^c \) are translationally invariant, as they are expressed in terms of expectation values and gradients. Thus, the unphysical violation of the translational invariance can only come from \( d_{IJ}^\phi \), and by omitting this unphysical term, one naturally arrives to the ETF-corrected NACs. Therefore, in the present study, we focus on the translationally invariant NAC,

\[
d_{IJ}^{ETF} = d_{IJ}^c = \frac{h_{IJ}}{E_I - E_J}.
\]

(20)

Another rational for omitting \( d_{IJ}^\phi \) as has been done in the definition of quasi-diabatic couplings is that all important couplings leading to rapid variation of the wave function come from the diagonalization, i.e., the EOM or CI part.

B. NACs within EOM-CC

The theory of calculations of NAC within the EOM-CC framework has been described in detail in Refs. 14, 55, and 68. Here we employ the approach of Tajji and Szalay who generalized Eq. (10) to the EOM-CC theory. Since EOM-CC theory is non-Hermitian, the left and right EOM states are not
the conjugate of each other but form a biorthogonal set,
\[ \tilde{H} | \psi_I \rangle = E_I | \psi_I \rangle, \]  
(21)
\[ \langle \psi_I | L_J^\dagger \tilde{H} = \langle \psi_I | L_J^\dagger E_J, \]  
(22)
\[ \langle \psi_I | L_J^\dagger R_I | \psi_I \rangle = \delta_{IJ}. \]  
(23)
The norms of left or right EOM states are arbitrary, which leads to an ambiguity in calculations of the interstate matrix elements. The first (minor) issue is that the magnitudes of these elements are not equal to $| |$ elements. The first (minor) issue is that $| |$ leads to an ambiguity in calculations of the interstate matrix element. The norms of left or right EOM states are arbitrary, which implies that the computed NAC forces: for all considered cases, the effect on the NAC was less than $10^{-6}$. Thus, for the sake of simplicity we use $| |$ unit weights.

The second issue is that the magnitudes of these elements depend on the arbitrary normalization. One possible solution is to consider geometric average,\(^{48}\)
\[ A_{IJ} \equiv \left( \frac{W_I A_{IJ} + W_J A_{JI}}{2} \right)^{1/2}. \]  
(25)
While the signs of $A_{IJ}$ and $A_{JI}$ are arbitrary, they are not independent; biorthornormality condition requires that if the sign of right EOM state $I$ is flipped, then the sign of the left EOM state $J$ must be flipped as well; consequently, both $A_{IJ}$ and $A_{JI}$ change signs. Thus, one can use the sign of either $A_{IJ}$ or $A_{JI}$ to assign a sign to $A_{IJ}$.

An equally justified approach of defining $A_{IJ}$ would be to consider an arithmetic average,\(^{14}\)
\[ A_{IJ} \equiv \frac{W_I A_{IJ} + W_J A_{JI}}{2}, \]  
(26)
where weights $W_K$ depend on the norms of the left and right EOM states,\(^{14}\) $N^I_J$ and $N^J_I$,
\[ W_I = N^I_J N^J_I, \]  
(27)
\[ W_J = N^J_I N^I_J. \]  
(28)
The definition and the expressions of the norms are given in Appendix C. Tajti and Szalay\(^{14}\) pointed out that in the limit of exact solution, the EOM values of interstate matrix elements defined by Eq. (26) should agree with full configuration interaction (FCI) ones.

We note that the sign of the NAC vector (or any other interstate matrix element) is not uniquely defined because the phases of adiabatic states are not defined by the electronic Schrödinger equation, i.e., flipping the sign of eigenstate $\Phi_I(r; R)$ does not affect Eq. (2). The sign issue is related to a well-known geometric phase effect,\(^{69}\) which requires proper handling of the phase of the electronic wave function when solving the nuclear problem for the overall wave function defined by Eq. (1).

Tajti and Szalay have shown that the EOM NAC force defined by using Eq. (26) can be computed from the gradients of the two states and a fictitious summed state,
\[ \mathbf{h}_{IJ} = \frac{2 \mathbf{G}_{I(J+J)} - W_I \mathbf{G}_J - W_J \mathbf{G}_I}{2}, \]  
(29)
where left and right amplitudes for the summed state are
\[ L_{I+J} = \frac{1}{\sqrt{2}} (W_I L_I + W_J L_J), \]  
(30)
\[ R_{I+J} = \frac{1}{\sqrt{2}} (R_I + R_J). \]  
(31)
Here we introduced normalization factors $\frac{1}{\sqrt{3}}$ for convenience so that the separable part of the respective density matrices (reference contribution to $\mathbf{G}_K$’s) is handled correctly by the regular EOM-CC code (we note that one can introduce weights $W_K$ when computing the right summed state and use simple summation of the left amplitudes; the results are numerically identical). This approach, which allows for a straightforward evaluation of the EOM-CC NACs by trivial modifications of the analytic gradient code,\(^{67}\) is exploited in this work.\(^{70}\) Our benchmark calculations showed that the computed $W_K$ are always nearly equal to one and have a minute effect on the computed NAC forces: for all considered cases, the effect on the NAC was less than $10^{-6}$. Thus, for the sake of simplicity we use $W_I = W_J = 1$. All reported values use these unit weights.

In the context of conical intersections, the non-Hermiticity of EOM-CC theory might be a concern.\(^{71}\) As pointed out by Köhn and Tajti, a straightforward extension of Wigner-Neumann derivation\(^{72}\) to non-symmetric model Hamiltonians suggests that the dimensionality of a conical intersection between the states of the same symmetry becomes $F-3$ instead of $F-2$. That means that the apex of the cone in Fig. 1 becomes a cylinder. They illustrated this troublesome behavior by the EOM-EE calculations of excited states in formaldehyde, where in a small region around conical intersection, EOM-EE-CCSD roots become complex (with real parts of the energies degenerate). Yet, the exact EOM-CC theory, which is equivalent to FCI, describes the intersections correctly. This paradox has been recently explained by Koch and co-workers;\(^{73}\) these authors also suggested a possible solution restoring the correct topolgy of conical intersections, as in Fig. 1, in inexact EOM-CC theories.\(^{74}\)

### III. COMPUTATIONAL DETAILS

MRCl calculations were performed by Columbus.\(^{75}\) EOM-CC calculations were performed with Q-Chem.\(^{76,77}\) The C4or calculations were performed using the modified version, as described in Appendix A.\(^{102}\) An EOM-CC wave function analysis was performed using the libw module\(^{78,79}\) of Q-Chem.

CASSCF and MRCl calculations in Columbus report both $h_{IJ}$ and $d_{IJ}$, but within MRCl, $h_{IJ}$ depends on the choice of orbital resolution, while the total coupling $d_{IJ}$ does not.\(^{24}\) In order to avoid this dependence, we use $d_{IJ}$ and extract $h_{IJ}$ from it. To do so, we project out the part violating translational invariance and then multiply the so corrected $d_{IJ}$ by the respective energy gap,
\[ \tilde{d}_{IJ}^{\alpha} = d_{IJ}^{\alpha} - \frac{1}{N} \sum_{n=1}^{N} d_{IJ}^{\alpha n}, \]  
(32)
\[ h_{IJ} = \tilde{d}_{IJ}(E_J - E_I), \]  
(33)
where $\alpha = x, y, z$ and index $n$ runs over all atoms. Although the NAC values corrected by ETF and by projecting out the translationally non-invariant part are not identical, they are reasonably close.\(^{31}\)
The Cartesian geometries and the values of NAC vectors are given in the supplementary material. The details of EOM-CC and MR calculations are given below.

**Cis-1,3,5-hexatriene, 1,3-cyclohexadiene, and uracil cations.** The cc-pVDZ basis was used in all calculations. In EOM-IP-CCSD calculations, core electrons were frozen. The complete details of the MR calculations can be found in Ref. 80. For the uracil and hexatriene cations, the state-averaged (SA) CASSCF method was used. For hexatriene, the active space included six \( \pi \)-orbitals and the corresponding five electrons [denoted (5,6)] and state-averaging was performed over three states. For uracil, the active space was (13,10), comprising all eight \( \pi \)-orbitals and two lone pairs on the oxygen atoms; the averaging was performed over four states. For cyclohexadiene, a restricted active space SCF (RASSCF) was used in order to obtain the same ordering of states as in EOM-IP-CCSD. The details of validation of these expansions and correct state ordering are given in Ref. 80.

**Excited states of uracil.** The cc-pVDZ basis was used in all calculations. In the EOM-EE-CCSD calculations, core electrons were frozen. An active space of 12 electrons in 9 orbitals (12,9) was used in all cases, comprising all \( \pi \) orbitals and one lone pair on oxygen, leading to 2520 reference CSFs (configuration state functions). The CASSCF calculations were performed using three-states averaging (SA3-CASSCF) in most cases. For additional tests along mode 25, SA4-CASSCF was performed. Single excitations from the active space were used in the MRCI expansion (denoted MR-CIS) using SA-CASSCF orbitals. In an additional MRCI calculation denoted MRCI1 (for mode 25), a larger expansion that included also \( \sigma \) and 3 lone pair orbitals on oxygen, leading to 2520 reference CSFs (configuration state functions). The CASSCF calculations were previously benchmarked and used in many previous studies of uracil.81–83

\( \text{Na(NH}_3)_n \), \( n = 1 \ldots 3 \). A mixed double-zeta basis set (aug-cc-pVDZ on heavy atoms and cc-pVDZ on hydrogens) was used for sodium-doped ammonia clusters. All electrons were active in the EOM-EA-CCSD calculations. In MRCI calculations, orbitals obtained from CASSCF averaged over four states (SA4-CASSCF) were used. The active space in CASSCF and MRCI consisted of 1-electron-in-4-orbitals (3\( \sigma \) and 3\( \pi \) of Na). The MRCI expansion included all single and double excitations from all orbitals except the core, giving rise to 893 703, 5 304 376, and 18 072 665 CSFs for the complexes with 1, 2, and 3 \( \text{NH}_3 \) respectively.

### IV. RESULTS AND DISCUSSION

#### A. Validation

We validated our implementation by comparing EOM-IP-CCSD and EOM-EE-CCSD NAC forces against a modified Cfour implementation of quasidiabatic couplings.55 (see Appendix A) for the selected examples. In all cases, the NAC forces computed by the two programs agreed within 6 decimal points. EOM-EA-CCSD NAC forces were validated against EOM-EE using the diffuse orbital trick.84 The Cartesian geometries, details of the benchmark calculations, and the computed values of the NAC forces are given in the supplementary material. We note that since the values of the NAC force depend on the molecular orientation, for proper comparison between the methods, the same Cartesian geometries must be used.

#### B. Examples

For reliable treatment of non-adiabatic dynamics, an electronic structure method should be capable of describing accurately the shape and relative energies of the PESs to ensure the correct description of the MECP as well as the magnitude and the direction of the NAC force. Many previous studies focused on the MECPs.34,35,60–62,85–88 In this paper, we focus on the NAC force vector and compare its magnitude and direction between \( \mathbf{h} \) computed by EOM-CC and multireference methods, e.g., CASSCF, RASSCF, and MRCI. To put the differences between these two high-level methods into a context, for selected examples, we also compare EOM-CC with CIS.

We consider the norm of the difference and the angle between the NAC forces computed by different methods,

\[
\Delta^{AB} \equiv || \mathbf{h}_I^A - \mathbf{h}_I^B ||,
\]

\[
\cos \theta^{AB} \equiv \frac{\mathbf{h}_I^A \cdot \mathbf{h}_I^B}{|| \mathbf{h}_I^A || \times || \mathbf{h}_I^B ||}.
\]

Here \( A \) and \( B \) denote two different methods. For selected examples, we also report the following quantity (computed in the Franck-Condon region) related to the rate of non-adiabatic transition:80

\[
r_{IJ} = || \mathbf{d}_{IJ} \mathbf{G}_J ||.
\]

This is the dot product between the NAC force and the nuclear gradient on the upper (i.e., initially excited) PES. The correlation between \( r_{IJ} \) and the rate on non-adiabatic relaxation, suggested by the last term in Eq. (4), has been illustrated by full-dimensional simulations of non-adiabatic transitions.
dynamics in several molecules using the surface-hopping approach.\textsuperscript{80}

1. EOM-IP NAC force in selected radical cations

A recent study has investigated ultrafast non-adiabatic dynamics in three radical cations: \textit{cis}-1,3,5-hexatriene, 1,3-cyclohexadiene, and uracil cations.\textsuperscript{80} We use these three cations as a benchmark for comparing EOM-IP NAC force against MR values. In all three cases, we compute electronic properties at the equilibrium geometry of the neutral ground state. The dynamical simulations\textsuperscript{80} have shown that non-adiabatic transitions in these systems occur on a femtosecond time scale and mostly in the Franck-Condon region and that the computed rate of non-adiabatic relaxation correlates well with $r_{1/2}$ of Eq. (36).

Tables I–III summarize the results for uracil, hexatriene, and cyclohexadiene cations. The state energies, characters, and relevant molecular orbitals are given in the supplementary material (Figs. S1–S3 and Tables S1–S3). As discussed in Ref. 80, for these systems, EOM-IP-CCSD energies and state characters agree well with the multi-reference values. We begin with comparing NAC forces computed by EOM-IP-CCSD and CASSCF/RASSCF. We note that since $d_{ij}$ depends on the energy gap between the states, this quantity is more sensitive to the variations of the state energies computed by different methods than the NAC force. As one can see from Tables I–III, the magnitudes of $h$, as quantified by its norm, are in good agreement between EOM-IP and CASSCF. For most states, the norms of $\mathbf{h}$ are within 10% from each other. The only significant difference is observed for the NAC force between states $D_0-D_2$ and $D_1-D_3$ in uracil, where the two values differ by $\sim30\%$. Considering $\Delta$ and $\cos \theta$, Eqs. (34) and (35) afford a more detailed comparison. While the norms of $\Delta$ vary from 0.01 to 0.03, the values of $\cos \theta$ are consistently close to 1 (the smallest value is 0.91) confirming that the NAC forces computed by the two methods are nearly parallel.

The differences between the values of $r$, Eq. (36), computed by EOM-IP-CCSD and MR methods are somewhat larger than the differences in $||\mathbf{h}||$ because these quantities also depend on the energy gaps and gradients. Tables I and III show reasonable correlation, that is, MRCI and EOM-CCSD agree in which cases $r$ are small and in which cases $r$ is large. In the latter set of cases, the discrepancies vary from perfect agreement to a factor of 10. In the case of hexatriene (Table II), we observe good agreement for $r$ involving states $D_0-D_2$ and huge discrepancies for all cases involving $D_1$. We traced this problem to different state character: apparently, state 3 in CASSCF calculations has different character compared to EOM-CCSD. CASSCF calculations with 5 states revealed that the state which has the same character as $D_3$ in EOM-CCSD appears as state number 4. The values for CASSCF state $D_4$ agree very well with EOM-CC values for $D_3$ (see the supplementary material, Table S4). This case highlights the utility of using several metrics: the changes in the state’s character were revealed by $r$ more prominently than by slightly increased difference in $\Delta$.

2. EOM-EE non-adiabatic coupling force in uracil

As a benchmark system for EOM-EE-CCSD NACs, we consider $\pi\pi^*$ and $\pi\pi^*$ states of uracil. Figure 3 shows natural transition orbitals (NTOs) and the respective singular values for the $S_0-S_1$, $S_0-S_2$, and $S_1-S_2$ transitions. The $S_1-S_2$ transition is well described by one NTO pair (participation ratio is 1.02) and has substantial one-electron character ($l||y|| = 0.6$). The radiationless relaxation in this system was investigated

### Table II. Energy gaps ($\Delta E$), norms of the NAC force ($h$), and derivative coupling vector ($d$) between the three lowest electronic states of the hexatriene cation. Energy gaps are in eV; all other quantities are in a.u. Superscripts A and B denote the EOM-IP-CCSD/cc-pVDZ and CASSCF/cc-pVDZ values.

| States | $\Delta E^A$ | $||h^A||$ | $||d^A||$ | $\Delta E^B$ | $||h^B||$ | $||d^B||$ | $\Delta^{AB}$ | $\cos \theta^{AB}$ | $\rho^A$ | $\rho^B$ |
|--------|-----------|----------|----------|-----------|----------|----------|-------------|----------------|--------|--------|
| 0-1    | 1.97      | 0.08     | 1.130    | 2.05      | 0.09     | 1.088    | 0.01         | 0.99           | 4.2 $\times 10^{-9}$ | 1.7 $\times 10^{-7}$ |
| 0-2    | 3.01      | 0.08     | 0.684    | 3.00      | 0.08     | 0.816    | 0.02         | 0.96           | 0.006             | 0.005             |
| 0-3    | 3.61      | 0.05     | 0.407    | 3.85      | 0.04     | 0.272    | 0.07         | 0.97           | 0.012             | 1.7 $\times 10^{-7}$ |
| 1-2    | 1.04      | 0.06     | 1.617    | 0.95      | 0.08     | 2.176    | 0.03         | 0.93           | 1.04 $\times 10^{-7}$ | 4.6 $\times 10^{-7}$ |
| 1-3    | 1.65      | 0.07     | 1.118    | 1.80      | 0.11     | 1.632    | 0.13         | 0.95           | 2.04 $\times 10^{-7}$ | 0.43               |
| 2-3    | 0.60      | 0.07     | 3.343    | 0.85      | 0.06     | 1.904    | 0.09         | 0.96           | 0.31              | 3.1 $\times 10^{-8}$ |

### Table III. Energy gaps ($\Delta E$), norms of the NAC force ($h$), and derivative coupling vector ($d$) between the three lowest electronic states of the cyclohexadiene cation. Energy gaps are in eV; all other quantities are in a.u. Superscripts A and B denote the EOM-IP-CCSD/cc-pVDZ and RASSCF/cc-pVDZ values.

| States | $\Delta E^A$ | $||h^A||$ | $||d^A||$ | $\Delta E^B$ | $||h^B||$ | $||d^B||$ | $\Delta^{AB}$ | $\cos \theta^{AB}$ | $\rho^A$ | $\rho^B$ |
|--------|-----------|----------|----------|-----------|----------|----------|-------------|----------------|--------|--------|
| 0-1    | 2.73      | 0.09     | 0.884    | 2.65      | 0.10     | 1.088    | 0.03         | 0.97           | 5.3 $\times 10^{-20}$ | 5.8 $\times 10^{-7}$ |
| 0-2    | 2.94      | 0.06     | 0.574    | 3.15      | 0.07     | 0.544    | 0.01         | 0.98           | 0.025             | 0.025             |
| 0-3    | 3.41      | 0.05     | 0.397    | 3.69      | 0.06     | 0.544    | 0.01         | 0.98           | 6.5 $\times 10^{-8}$ | 9.5 $\times 10^{-8}$ |
| 1-2    | 0.21      | 0.04     | 5.819    | 0.50      | 0.05     | 2.448    | 0.01         | 0.94           | 4 $\times 10^{-7}$   | 1.1 $\times 10^{-6}$ |
| 1-3    | 0.68      | 0.06     | 2.231    | 1.05      | 0.06     | 1.360    | 0.02         | 0.95           | 0.1               | 0.01              |
| 2-3    | 0.47      | 0.15     | 8.707    | 1.05      | 0.18     | 8.979    | 0.03         | 0.99           | 2.5 $\times 10^{-7}$ | 6.9 $\times 10^{-6}$ |
by several groups with a variety of approaches,81,89–94 most recently by Zhang and Herbert95 using SF-TDDFT.38 This study identified two major pathways for excited-state deactivation: direct $1\pi\pi^* \rightarrow S_0$ and indirect $1\pi\pi^* \rightarrow n\pi^* \rightarrow S_0$ relaxation. The authors characterized two relevant MECPs: one between $1\pi\pi^*$ and $1n\pi^*$ and one between $1\pi\pi^*$ and $S_0$. Here we denote the two MECPs as CI($S_2S_1$) and CI($S_2S_0$).

In addition to MECPs, Zhang and Herbert reported the optimized excited-state structure of the $1\pi\pi^*$ state: MIN($S_1$). We use these structures to investigate the variations in NAC forces outside the Franck-Condon region, along geometric distortions most relevant to the radiationless relaxation process. Specifically, we consider the displacements along normal modes connecting the Franck-Condon structure ($S_0$ equilibrium geometry) with the CI($S_2S_1$), CI($S_2S_0$), and MIN($S_1$) structures. We use ground-state normal modes for this calculation, as often done in parallel-mode double-harmonic calculations of the Franck-Condon factors. The ground-state structure and normal modes are computed by $\omega$B97X-D/cc-pVDZ. We identified normal modes that have the largest displacements between the MIN($S_0$) and CI($S_2S_1$), CI($S_2S_0$), and MIN($S_1$) structures from Ref. 95. Table IV lists the normal modes selected for these calculations; the displacements corresponding to these normal modes are visualized in Fig. S4.
in the supplementary material. Table S5 in the supplementary material lists the displacements along all normal modes.

Figure 4 shows potential energy profiles along the selected normal modes. The PES computed by different methods looks qualitatively similar. Along modes 3 and 23, the energies of all three states are nearly flat. Along modes 10 and 25, all energies go up steeply. The energy gap between $S_1$ and $S_2$ does not change much; however, the gap between $S_1$ and $S_0$ shrinks. Along mode 10, there is a kink in the CASSCF and MRCI curves at points 9 and 10 due to an abrupt change in the state character. Along mode 25, both the variations in PESs and the differences in the PES shapes computed by different methods appear most prominent. This is not surprising because from the 4 modes considered here, mode 25 is the stiffest and the displacement along this mode is large.

Figure 5 shows the absolute values of the NAC force between $S_1$ and $S_2$ states along the selected modes for CASSCF, MRCI, CIS, and EOM-CCSD wave functions. The raw data are collected in Tables S6–S9 in the supplementary material. Tables S10–S13 in the supplementary material show differences ($\Delta$ and $\cos \theta$) between the NAC forces computed by different methods against MRCI.

In the Franck-Condon region (point 0 in Tables S6–S13 of the supplementary material and in Fig. 5 and Fig. S5 of the supplementary material), the magnitude of the difference between EOM and MRCI vectors is $\Delta = 0.012$, which is consistent with the values of the respective $\|	extbf{h}\|$; 0.0217 (EOM) and 0.0078 (MRCI). CASSCF shows the smallest difference (0.004). By looking at $\cos \theta$, we see that it is not only the magnitude of $\textbf{h}$ but also the direction of the vector which is different: the value of $\cos \theta_{\text{EOM}}$ is 0.75, whereas for CASSCF and (surprisingly) CIS, these values are larger (0.87 and 0.83, respectively).

As one can see from Fig. 5, the magnitude of $\|	extbf{h}\|$ does not vary much along modes 3 and 23 (except for an increasing difference between CASSCF and MRCI at points 8–10 along mode 3). To better illustrate the trends, Fig. S5 in the supplementary material shows the values of $\|	extbf{h}\|$ normalized to their respective Franck-Condon values. Along mode 23, all methods show a small decline (less than 5%). Along mode 3, EOM shows a slight increase (1%), whereas MRCI and CIS values drop by about 10%. Table S12 of the supplementary material confirms that along mode 23, the differences between the EOM-CC NAC force vectors and reference MRCI values do not change much, for example, $\Delta_{\text{EOM}}$ is constant and $\cos \theta_{\text{EOM}}$ varies between 0.745 and 0.743. Mode 3 shows more complex behavior: at large displacements, the alignment between EOM and MRCI vectors decreases, dropping below 0.5 at points 8–10 (Table S10 of the supplementary material). Interestingly, $\Delta_{\text{EOM}}$ shows only a relatively moderate increase at these displacements. We also note that $\cos \theta_{\text{CASSCF}}$ decreases consistently.

---

**FIG. 5.** Magnitude of the NAC force ($\|	extbf{h}\|$) along selected modes for CASSCF, MRCI, CIS, and EOM-EE-CCSD wave functions.
with the trends in the respective \(|llh|ll\). These increasing differences between MRCI and other methods at large displacements are suggestive of changing the wave function composition. For example, at large displacements, there is a noticeable difference even between CASSCF and MRCI. The reason is that at the CASSCF level, states \(S_2\) and \(S_3\) are very close and interact strongly, changing the wave function and, consequently, the coupling.

For mode 10, all \(|llh|ll\) slightly increase and the trends among all methods are consistent (except points 9 and 10, where CASSCF and MRCI potential energy curves show kinks). Table S11 of the supplementary material confirms this—the changes in \(A\) and \(\cos \theta\) are small (we note that \(\cos \theta^{EOM}\) increases along this mode, reaching 0.94 at point 7). This is a case where the states change character leading to sudden changes in the coupling. The left panel of Fig. S6 in the supplementary material shows the four electronic states of uracil at the SA4-CASSCF level. At point 9, states \(S_2\) and \(S_3\) approach each other and possibly cross. NAC highlights this crossing. As can be seen in the right panel at point 9, the values of \(|llh|ll\) suddenly switch for both \(S_2\) and \(S_3\) states. This leads to a sudden increase in the \(S_1-S_2\) coupling that we see in Fig. 5.

In contrast to modes 3, 10, and 23, mode 25 shows very large variations between the NAC forces computed by different methods. Regardless of the metric used to quantify the coupling vector (norms, normalized norms, \(A\), and \(\cos \theta\)), we clearly see that the discrepancies between the methods along this coordinate are large. Apparently, the changes in the wave functions’ characters are very sensitive to the level of correlation treatment. To further investigate this issue, we performed additional MRCI calculation (denoted MRCI1, see Sec. III) using a larger expansion. The results of the two MRCI calculations are shown in Fig. S7 in the supplementary material. As one can see, \(|llh|ll\) from the two calculations are quite different: in CASSCF and MRCI (which uses a rather compact expansion), the magnitude of \(|llh|ll\) first increases by 40% (reaching maximum at point 2) and then slightly decreases, whereas in MRCI1, the trend is reversed: \(|llh|ll\) first drops by 25% (reaching minimum at point 1), and then increases by 40%. This illustrates the sensitivity of NAC along this mode to dynamical correlation. To understand the changes in states’ characters, we analyzed EOM-CCSD wave functions along mode 25 using NTOs of the \(S_2-S_1\) transition. We found that the changes in the NTOs are subtle, and the transition retains its \(nn^*\) character at all displacements. Important quantities, such as participation ratio and \(lyll\), do not change much. A close inspection of NTOs reveals that the particle NTO for the \(S_2-S_1\) transition does not change much, retaining its out-of-plane \(n(O_2)\) character (see Fig. 5). However, the character of the hole orbital evolves (this is shown in Fig. S8 in the supplementary material) from delocalized \(\pi\) orbital to localized out-of-plane \(n(O_2)\). Figure S8 of the supplementary material also shows a continuous decrease in electron density in \(n(O_1)\) and an increase in \(n(O_2)\). The sharp increase in NAC at point 3 occurs when the amplitude of the hole NTO on \(O_1\) disappears. To further understand these changes, we consider natural orbitals of the \(S_1\) and \(S_2\) states. Figure S9 in the supplementary material compares frontier natural orbitals (NOs) computed for the EOM-CC and MRCI wave functions. As one can see, most of the changes occur in the \(S_2\) state. We note that these changes in \(S_2\) (the flow of density from \(n(O_1)\) to \(n(O_2)\)) have been observed before in a computational study of radiationless relaxation of uracil and related molecules.  The NOs computed by MRCI and EOM-CC appear to be very similar. The only noticeable difference is that EOM-CC wave function at point 4 shows more open-shell character than MRCI (natural occupations of 1.3 and 0.7 versus 1.5 and 0.6). We conclude that the magnitude of NAC is very sensitive to these small variations in the wave function character, which, in turn, are very sensitive to the correlation treatment. Thus, even when qualitatively wave functions are similar, one can observe quantitative discrepancies among different methods.

Figure 6 shows the MRCI NAC force at the Franck-Condon geometry (point 0) and at displaced geometries along modes 23 and 25. As expected from the symmetries and orbital characters (see Fig. 3) of \(S_1\) (\(A''\), \(n\pi^*\)) and \(S_2\) (\(A', \pi\pi^*\)), the NAC vector has \(a''\) symmetry (out-of-plane). Thus, only out-of-plane motions are expected to be effective in promoting non-adiabatic transition between these states. Among the four modes considered here, only mode 10 corresponds to out-of-plane distortion. In terms of Eq. (36), only the out-of-plane component of the gradient will contribute to the rate. The NAC force vector along mode 23 looks qualitatively similar to that at the Franck-Condon structure, whereas displacement along mode 25 results in noticeable changes.

3. EOM-EA NAC force in sodium-doped ammonia clusters

We use sodium-doped clusters as a model system for benchmarking EOM-EA. The low-lying electronic states in these clusters, recently studied theoretically and experimentally, can be described as \(s\)- and \(p\)-like states of

![Fig. 6. MRCI S1/S2 NAC force in uracil at the Franck-Condon geometry (a), at point 1 along mode 23 (b), and at point 1 along mode 25 (c).](image-url)
TABLE V. Energy gaps ($\Delta E$, eV), the NAC force ($||h||$), and derivative coupling vector ($||d||$) between the three lowest electronic states of the sodium-doped clusters Na(NH$_3$)$_n$. All quantities are in a.u. Superscripts A and B denote the EOM-IP-CCSD/aug-cc-pVDZ and MRCI/aug-cc-pVDZ values.

| States | $\Delta E^A$ | $||h^A||$ | $||d^A||$ | $\Delta E^B$ | $||h^B||$ | $||d^B||$ | $\Delta^A_{\theta}$ | $\cos \theta^A$ | $r^A$ | $r^B$ |
|--------|--------------|-----------|-----------|--------------|-----------|-----------|--------------------|--------------|------|------|
| NaNH$_3$ | 1-2 | 1.501 | 0.009 | 0.164 | 1.503 | 0.015 | 0.281 | 0.012 | 0.000 | 0.000 |
|         | 1-3 | 1.501 | 0.009 | 0.164 | 1.503 | 0.015 | 0.280 | 0.012 | 0.599 | 0.000 |
|         | 1-4 | 1.523 | 0.007 | 0.120 | 1.568 | 0.011 | 0.199 | 0.008 | 0.689 | 0.003 |
| Na(NH$_3$)$_2$ | 1-2 | 1.107 | 0.011 | 0.266 | 1.124 | 0.016 | 0.389 | 0.019 | 0.035 | 0.006 |
|         | 1-3 | 1.141 | 0.010 | 0.233 | 1.127 | 0.014 | 0.338 | 0.016 | 0.169 | 0.000 |
|         | 1-4 | 1.147 | 0.011 | 0.260 | 1.164 | 0.013 | 0.308 | 0.013 | 0.473 | 0.000 |
| Na(NH$_3$)$_3$ | 1-2 | 0.646 | 0.012 | 0.389 | 0.803 | 0.013 | 0.456 | 0.016 | 0.220 | 0.000 |
|         | 1-3 | 0.650 | 0.012 | 0.386 | 0.806 | 0.014 | 0.457 | 0.014 | 0.423 | 0.001 |
|         | 1-4 | 0.684 | 0.012 | 0.369 | 0.819 | 0.015 | 0.496 | 0.011 | 0.670 | 0.006 |

FIG. 7. NAC force in doped ammonia clusters computed using MRCI (a) and EOM-CCSD (b) wave functions.

surface-bound electrons, stabilized by a positively charged Na$_{n_{\delta+}}$(NH$_3$)$_n$ core. Dyson orbitals of the four lowest states are shown in Fig. S10 in the supplementary material (and in Fig. 1 in Ref. 96). The target doublet states are best described by EOM-EA using the closed-shell cation as a reference. Non-adiabatic relaxation of $p$-like states has been studied in related systems, Ba(Ar)$_n$ clusters.

Table V shows energy gaps and the NACs between the three lowest electronic states of the sodium-doped clusters, Na(NH$_3$)$_n$, with $n = 1-3$. The NAC forces for Na(NH$_3$) and Na(NH$_3$)$_2$ are shown in Fig. 7. As one can see, the NAC vectors are dominated by the relative Na-ammonia coordinate. Interestingly, while differences between the methods in the magnitude of $||h||$ are consistent with $\Delta$, the value of $\cos \theta$ appears rather low. The reason is that in these three clusters, the three $p$-like states are nearly degenerate, which means that small differences in the methods can mix them. Since the mixing of the $p$-orbitals only affects their direction, such mixing only affects the direction of the NAC forces but not their magnitude. In support of this explanation, we note that the largest $\cos \theta$ values correspond to the NAC forces between the $s$-state with the highest $p$-state (states 1-4) in NaNH$_3$ and Na(NH$_3$)$_3$ in which the gap between state 4 and states 2-3 is the largest.

V. CONCLUSION

We presented the theory and implementation of the NAC forces within the EOM-CCSD framework. The main focus of this paper is on the comparison between the EOM-CCSD and MRCI NAC forces. Using selected examples, we analyzed NACs computed using EOM-IP-CCSD, EOM-EE-CCSD, and EOM-EA-CCSD wave functions. In contrast to previous studies, which focused primarily on the structures of conical intersections, here we analyzed the direction of the NAC vector. We note that the direction of NAC vector, not only its magnitude, is very important for the calculations of the rate of non-adiabatic transitions. Our examples illustrate the utility of using different metrics of comparison. The numeric results confirm that the accuracy of NAC forces depends critically on the quality of the underlying wave functions, which are often very sensitive to correlation treatment. Within their domain of applicability, EOM-CC methods provide a viable alternative to MR approaches.
SUPPLEMENTARY MATERIAL

See supplementary material for computational details, additional validation and benchmark calculations, electronically excited states of uracil and sodium-doped ammonia clusters, and relevant Cartesian coordinates.

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APPENDIX A: DERIVATION OF EOM-CC NAC FORCE AND CONNECTION TO QUASIDIABATIC COUPLINGS

To derive expressions for the EOM-CC NACs, we begin by differentiating an interstate matrix element of $\hat{H}$, which, obviously, is zero,

$$\hat{H}_{IJ} \equiv \langle 0|L_I|\hat{H}|R_J\rangle = 0. \quad \text{(A1)}$$

By defining $\hat{H}^* \equiv e^{-T} \hat{H}^* e^T$ and using the usual notation $f^* \equiv \frac{\partial f}{\partial \hat{H}}$, we obtain

$$0 = \langle 0|L_I\hat{H}^* R_J\rangle = \langle 0|L_I\hat{H}^* R_J\rangle + \langle 0|L_I\hat{H} R_J^*\rangle$$

$$+ \langle 0|L_I\hat{H}^* T\rangle - \langle 0|L_I^T \hat{H} R_J^*\rangle$$

$$= \langle 0|L_I\hat{H}^* R_J\rangle + E_I\langle 0|L_I^T \hat{H} R_J^*\rangle$$

$$+ \langle 0|L_I\hat{H}^* q|q R_J^* T\rangle - \langle 0|L_I^T \hat{H}^* q|q R_J^* T\rangle$$

$$- \langle 0|L_I^T \hat{H}^* q|q R_J^*\rangle,$$  \hspace{1cm} \text{(A2)}

where $p$ denotes the EOM-CCSD manifold of Slater determinants: $p = 0 + S + D$ (and the complimentary space of higher excitations is denoted by $q = T + Q + \cdots$).

By using the amplitude-response operator $\Xi$,

$$\langle 0|\Xi|p\rangle \equiv \langle 0|L_I\hat{H}|q]\langle q| R_p\rangle,$$  \hspace{1cm} \text{(A3)}

we obtain

$$0 = \langle 0|L_I\hat{H}^* R_J\rangle + \langle 0|\Xi|p\rangle \langle p| T\rangle - \langle 0|L_I\hat{H}^* q|q R_J^* T\rangle$$

$$+ \langle 0|L_I^T \hat{H}^* q|q R_J^* T\rangle - \langle 0|L_I^T \hat{H}^* q|q R_J^*\rangle$$

$$+ \langle 0|L_I^T \hat{H}^* q|q R_J^* T\rangle = \langle 0|L_I^T \hat{H}^* q|q R_J^* T\rangle$$

$$+ \langle 0|L_I^T \hat{H}^* q|q R_J^*\rangle - \langle 0|L_I^T \hat{H}^* q|q R_J^* T\rangle.$$  \hspace{1cm} \text{(A4)}

Because $\langle 0|L_I^T \hat{H}^* q|q R_J^*\rangle$ is a constant,

$$\langle 0|L_I^T \hat{H}^* q|q R_J^* T\rangle = -\langle 0|L_I^T R_J^* T\rangle.$$  \hspace{1cm} \text{(A5)}

Thus, we obtain

$$0 = \langle 0|L_I\hat{H}^* R_J\rangle + \langle 0|\Xi|p\rangle \langle p| T\rangle$$

$$+ (E_I - E_J)\langle 0|L_I^T R_J^|p| T\rangle - \langle 0|L_I^T R_J^|p| T\rangle.$$  \hspace{1cm} \text{(A6)}

We now define EOM-EE derivative coupling as

$$d_{IJ}^\text{c} = \langle 0|L_I^T R_J^|p| T\rangle.$$  \hspace{1cm} \text{(A7)}

In this definition, we neglect orbital response terms except for those that arise from the differentiation of $\hat{H}$.

By combining Eqs. (A6) and (A7), we obtain the following expression for $d_{IJ}^\text{c}$:

$$d_{IJ}^\text{c} = \frac{\langle 0|L_I\hat{H}^* R_J\rangle + \langle 0|\Xi|p\rangle \langle p| T\rangle}{E_I - E_J}.$$  \hspace{1cm} \text{(A8)}

The numerator in this equation is the EOM NAC force from Eq. (15). It is equal to quasiadiabatic couplings from Ref. 55 minus the following term:

$$(E_I - E_J)\langle 0|\Xi|p\rangle \langle p| T\rangle = (E_I - E_J)\langle 0|L_I R_J^|p| T\rangle.$$  \hspace{1cm} \text{(A9)}

As our benchmarks illustrate, the magnitude of this term is rather small. Thus, the EOM-CC $d_{IJ}^\text{c}$ is nearly identical to the quasiadiabatic couplings $\lambda_{IJ}$ from Ref. 55.

APPENDIX B: ORBITAL-RESPONSE PART OF NAC

The orbital-response part of the full derivative coupling is given by the following expression:

$$d_{IJ}^\text{o} = \sum_{pq} \langle \phi_p | \Phi_q^\text{o} | D_{pq}^\text{c} \rangle,$$  \hspace{1cm} \text{(B1)}

where $D_{pq}^\text{c}$ is the antisymmetric one-particle transition density matrix,

$$D_{pq}^\text{c} = \frac{1}{2} \langle 0|(W_I L_I - W_J L_J)^{T}\langle q R_I + R_J\rangle|\rangle$$

$$- \frac{1}{2} \left(W_I^\gamma J^\gamma - W_J^\gamma J^\gamma + W_I^\gamma J^\gamma - W_J^\gamma J^\gamma \right).$$  \hspace{1cm} \text{(B2)}

and $|\Phi_q^\text{o}\rangle$ is

$$\langle \phi_p | \phi_q^\text{o} \rangle = \langle \phi_p | \Phi_q^\text{o} \rangle + U_{pq}^\text{o},$$  \hspace{1cm} \text{(B3)}

where the first term is the derivative of the overlap matrix and the second part is the vector of coupled-perturbed Hartree-Fock coefficients describing the response of the MO coefficients to the moving of the $\xi$ atomic basis functions. The calculation of this term has been described by Gauss et al. 8 The $d_{IJ}^\text{o}$ term violates the translational invariance of the NAC and is omitted in our implementation. It is also omitted in the implementation of quasidiabatic couplings of Stanton and co-workers. 55

APPENDIX C: NORMALIZATION FACTORS FOR EOM-CCSD STATES

The normalization factors for left EOM states are defined as

$$(N_{L}^\text{e})^{-2} \equiv \langle \Phi_0 | L_{e^{-T} L_{a}} | \Phi_0 \rangle.$$  \hspace{1cm} \text{(C1)}

This equation can be tackled by inserting the resolution of the identity, $\sum_{x} \langle \Phi_0 | \Phi_x | \rangle (x$ denotes the excitation level).

For EOM-EE-CCSD, only the following three terms survive,
giving rise to the following programmable expression:

\[
(N_d)_{\mu \nu} = \langle \Phi_0 | L_d e^{-T} | \Phi_0 \rangle + \sum_{ia} \langle \Phi_0 | L_d e^{-T} | \Phi_i \rangle^2 \\
+ \frac{1}{4} \sum_{ijab} \langle \Phi_0 | L_d e^{-T} | \Phi_{ijab} \rangle^2 + \ldots \\
= \left( \sum_{ia} \langle \Phi_0 | L_d e^{-T} | \Phi_i \rangle^2 + \frac{1}{4} \sum_{ijab} \langle \Phi_0 | L_d e^{-T} | \Phi_{ijab} \rangle^2 + \ldots \right)^2 \\
+ \sum_{ia} X_i^a X_i^a + \frac{1}{4} \sum_{ij} \langle \Phi_0 | L_d e^{-T} | \Phi_{ij} \rangle_0^2, \\
X_i^a = l_i - \sum_{ja} \langle \Phi_0 | L_d e^{-T} | \Phi_{ja} \rangle_0^2, \quad (C2)
\]

The \( N_R \) is approximated as

\[
N_R = \frac{1}{N_d} \quad (C3)
\]

This is exact in the case of FCI.\(^{14}\)

We note that in EOM-SF version, the first term in the above expression is zero. For EOM-IP and EOM-EA, the expressions for the left state norms are

\[
(N_d)_{\mu \nu} = \sum_{ia} \langle \Phi_0 | L_d e^{-T} | \Phi_i \rangle^2 \\
X_i^a = l_i - \sum_{ja} \langle \Phi_0 | L_d e^{-T} | \Phi_{ja} \rangle_0^2, \quad (C4)
\]

and

\[
(N_d)_{\mu \nu} = \sum_{ia} \langle \Phi_0 | L_d e^{-T} | \Phi_i \rangle^2 \\
X_i^a = l_a - \sum_{jb} \langle \Phi_0 | L_d e^{-T} | \Phi_{jb} \rangle_0^2, \quad (C5)
\]

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An important modification of the EOM-EE-CCSD programmable expressions from Ref. 67 is that for summed states, the \( \omega_{ij} \) terms in density matrices should be replaced by \( \sum_{\omega} \omega_{ij} r_{\omega}^2 + \frac{1}{2} \sum_{\omega} \omega_{ij} r_{\omega}^2 (|\langle \omega | l | \omega \rangle|)^2 \).


In Ref. 14, there is no factor² [i.e., (N_{Lx}^2)^{-1} = ...] which we believe is a typo. The expression is clearly the square of the norm since it is the dot product of the left vector with itself. Using square root or not does not affect the results all that much though since they mostly depend on W_{ij} and not on the individual N_{Lx}².