DENSITY CUMULANT THEORY
FOR GROUND AND EXCITED ELECTRONIC STATES

by

Andreas Victor Copan

(Under the Direction of Henry F. Schaefer III)

Abstract

Density cumulant theory (DCT) describes the state of an electronic system by variationally optimizing a parametrization of the two-particle density cumulant. The cumulant is a statistical descriptor of the electronic density distribution which naturally decouples independent subsystems, leading to desirable properties like size-extensivity which have given the coupled-cluster methods pride of place in electronic structure theory. We present benchmarks calculations demonstrating the superior performance of density cumulant theory relative to coupled-cluster theory for the description of ground-state properties. Next, we extend this method for the description of excited states using linear response theory. Finally, we develop algorithms for the new excited state model which enable us to study larger systems such as hexatriene with this new model.

INDEX WORDS: Electronic structure theory, cumulants, density matrices, linear response theory, excited states, coupled-cluster theory

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# DENSITY CUMULANT THEORY FOR GROUND AND EXCITED ELECTRONIC STATES

by

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In	memory	of	Valery	And reiye vich	and	Valtraut	Kirsch	Copan

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### Chapter 1

## $Introduction^*$

The goal of electronic structure theory is to solve the time-independent Born-Oppenheimer Schrödinger equation with an optimal balance of accuracy and efficiency for the problem of interest.

$$\hat{H}\Psi_k = E_k \Psi_k \tag{1.1}$$

$$\hat{H} = V_{\text{nuc}} + \hat{H}_e = \sum_{a < b}^{\text{nuc.}} \frac{Z_a Z_b}{|\mathbf{R}_a - \mathbf{R}_b|} - \frac{1}{2} \sum_{i}^{\text{elec.}} \nabla_i^2 - \sum_{a}^{\text{nuc. elec.}} \frac{Z_a}{|\mathbf{R}_a - \mathbf{r}_i|} + \sum_{i < j}^{\text{elec.}} \frac{1}{|\mathbf{r}_i - \mathbf{r}_j|}$$
(1.2)

The most accurate solution possible for a given one-particle basis set of spinorbitals  $\{\psi_p\}$  results from expanding the wavefunction as a linear combination of all antisymmetrized products of these one-particle states, which are known as Slater determinants  $\{\Phi_{\mu}\}$ .

$$\Psi_k = \sum_{\mu} \Phi_{\mu} c_{\mu k} \tag{1.3}$$

<sup>\*</sup>Adapted from my lecture notes at https://github.com/CCQC/chem-8950.

The expansion coefficients  $(\mathbf{c})_k = c_{\mu k}$  are eigenvectors of the matrix  $(\mathbf{H})_{\mu\nu} = \langle \Phi_{\mu} | \hat{H} | \Phi_{\nu} \rangle$ , which is the matrix representation of the Hamiltonian in the determinant basis. This is called the *full configuration-interaction* (FCI) solution.

Any one-electron basis spans the same "function space" as the AO basis set itself, and the full n-electron basis  $\{\Phi_{\mu}\}$  spans the same space of n-electron functions regardless of how one forms spin orbitals from the AO basis set. As a result, one obtains the same FCI solution for any choice of spin-orbitals. In general, however, FCI solutions are completely unfeasible for basis sets of sufficient size to approach the complete basis set limit because the number of determinants grows exponentially with the dimension of the one-particle basis. The upshot is that we usually have to omit some Slater determinants in order to get an answer in a reasonable amount of time. As soon as we truncate our determinant expansion (1.3), our choice of spin MOs makes a significant difference in the quality of our results. In particular, we need to choose our set of one-electron functions to minimize the number of Slater determinants it takes to "get close to" the exact wavefunction. One approach is to determine the best single-determinant approximation to the wavefunction by optimizing its energy expectation value with respect to variations in the orbitals

$$\frac{\delta\langle\Phi|\hat{H}_e|\Phi\rangle}{\delta\psi_i} \stackrel{!}{=} 0 \qquad \Phi(1,\dots,n) = \frac{1}{\sqrt{n!}} \begin{vmatrix} \psi_1(1) & \psi_2(1) & \cdots & \psi_n(1) \\ \psi_1(2) & \psi_2(2) & \cdots & \psi_n(2) \\ \vdots & \vdots & \ddots & \vdots \\ \psi_1(n) & \psi_2(n) & \cdots & \psi_n(n) \end{vmatrix}$$
(1.4)

where the arguments  $1, \ldots, n$  are short hand for the spin,  $s_1, \ldots, s_n$ , and spatial,  $\mathbf{r}_1, \ldots, \mathbf{r}_n$ , degrees of freedom of individual electrons. This variational condition can be written as an eigenvalue equation of a one-body "Fock" operator which represents an effective Hamiltonian for the interaction of an electron with the mean field of the other electrons.

$$\hat{f}\psi_i = \epsilon_i \psi_i \tag{1.5}$$

Once we have solved for these variationally optimal "Hartree-Fock" orbitals, the expectation value  $\langle \Phi | \hat{H}_e | \Phi \rangle$  is itself a good first approximation to the electronic energy. More importantly, however, when we use this new set of Hartree-Fock spin-orbitals,  $\{\psi_p\}$ , the FCI expansion tends to converge much more quickly to the true wavefunction. Specifically, when we rewrite equation 1.3 in terms of single  $\{\Phi_{ij}^a\}$ , double  $\{\Phi_{ijk}^{abc}\}$ , etc. replacements of the orbitals in the Hartree-Fock determinant  $\Phi$  with the remaining orbitals in the basis

$$\Psi = \Phi + \sum_{\substack{a \\ i}} \Phi_i^a c_a^i + \sum_{\substack{a < b \\ i < j}} \Phi_{ij}^{ab} c_{ab}^{ij} + \sum_{\substack{a < b < c \\ i < j < k}} \Phi_{ijk}^{abc} c_{abc}^{ijk} + \dots$$
 (1.6)

the coefficients tend to be very small, and are often virtually negligible for higher than quadruple replacements.

Our starting point in density cumulant theory (DCT) is to express the electronic energy as a trace of the one- and antisymmetrized two-electron integrals ( $h_p^q$  and  $\overline{g}_{pq}^{rs}$ ) with the reduced one- and two-body density matrices ( $\gamma_q^p$  and  $\gamma_{rs}^{pq}$ ):

$$E = h_p^q \gamma_q^p + \frac{1}{4} \overline{g}_{pq}^{rs} \gamma_{rs}^{pq} \tag{1.7}$$

where summation over the repeated indices is implied. In DCT, the two-body density matrix  $\gamma_{rs}^{pq}$  is expanded in terms of its connected part, the two-body density cumulant  $(\lambda_{rs}^{pq})$ , and its disconnected part, which is given by an antisymmetrized product of one-body density matrices:<sup>1</sup>

$$\gamma_{rs}^{pq} = \langle \Psi | a_{rs}^{pq} | \Psi \rangle = \lambda_{rs}^{pq} + P_{(r/s)} \gamma_r^p \gamma_s^q \tag{1.8}$$

where  $P_{(r/s)}v_{rs} = v_{rs} - v_{sr}$  denotes antisymmetrization and  $a_{rs}^{pq} = a_p^{\dagger} a_q^{\dagger} a_s a_r$  is the two-body operator in second quantization. The one-body density matrix  $\gamma_q^p$  is determined from its non-linear relationship to the cumulant's partial trace:<sup>2</sup>

$$\gamma_q^p = \gamma_r^p \gamma_q^r - \lambda_{qr}^{pr} \tag{1.9}$$

This reduces eq. (1.7) to a functional of the two-body cumulant and the basis of spin-orbitals, thereby defining the DCT energy functional. The density cumulant is parametrized by choosing a specific Ansatz for the wavefunction  $|\Psi\rangle$  such that<sup>3</sup>

$$\lambda_{rs}^{pq} = \langle \Psi | a_{rs}^{pq} | \Psi \rangle_{c} \tag{1.10}$$

where c indicates that only fully connected terms are included in the parametrization. Equation (1.10) can be considered as a set of n-representability conditions that ensure that the resulting one- and two-electron density matrices represent a physical n-electron wavefunction. To compute the DCT energy, the functional (1.7) is made stationary with respect to all of its parameters. Importantly, due to the connected nature of eq. (1.10), DCT is both size-consistent and size-extensive for any parametrization of  $|\Psi\rangle$ , and is exact in the limit of a complete parametrization.<sup>3</sup>

In this work, we consider the ODC-12 method,<sup>2;4</sup> which parametrizes the cumulant through a unitary treatment of single excitations and a linear expansion of double excitations.

$$|\Psi\rangle = e^{T_1 - T_1^{\dagger}} (1 + T_2) |\Phi\rangle \tag{1.11}$$

$$T_1 = \mathbf{t}_1 \cdot \mathbf{a}_1 = t_a^i a_i^a \tag{1.12}$$

$$T_2 = \mathbf{t}_2 \cdot \mathbf{a}_2 = \frac{1}{4} t_{ab}^{ij} a_{ij}^{ab} \tag{1.13}$$

The exponential singles operator  $e^{T_1-T_1^{\dagger}}$  has the effect of a unitary transformation of the spin-orbital basis and is incorportated in our implementation of the ODC-12 method by optimizing the orbitals.<sup>4</sup> The  $\mathbf{t}_1$  and  $\mathbf{t}_2$  parameters are obtained from the stationarity conditions

$$\frac{\partial E}{\partial \mathbf{t}_{1}^{\dagger}} \stackrel{!}{=} 0 , \qquad \frac{\partial E}{\partial \mathbf{t}_{2}^{\dagger}} \stackrel{!}{=} 0 \qquad (1.14)$$

and are used to compute the ODC-12 energy. Explicit equations for the stationarity conditions are given in Refs. 2 and 4. Although in ODC-12 the wavefunction parametrization is linear with respect to double excitations (eq. (1.11)), the

ODC-12 energy stationarity conditions are non-linear in  $\mathbf{t}_2$  due to the non-linear relationship between the one-particle density matrix and the density cumulant (eq. (1.9)). Neglecting the non-linear  $\mathbf{t}_2$  terms in eq. (1.14) results in the equations that define the linearized orbital-optimized coupled cluster doubles method (OLCCD). This method is equivalent to the orbital-optimized coupled electron pair approximation zero (OCEPA<sub>0</sub>).

## Chapter 2

Ground-State Density Cumulant Theory:

Thermochemical and Kinetic Benchmark Calculations\*

#### 2.1 Abstract

We present an extensive benchmark study of density cumulant functional theory (DCFT) for thermochemistry and kinetics of closed- and open-shell molecules. The performance of DCFT methods (DC-06, DC-12, ODC-06, and ODC-12) is compared to that of coupled-electron pair methods (CEPA<sub>0</sub> and OCEPA<sub>0</sub>) and coupled-cluster theory (CCSD and CCSD(T)) for the description of noncovalent interactions (A24 database), barrier heights of hydrogen-transfer reactions (HTBH38), radical stabilization energies (RSE30), adiabatic ionization energies (AIE), and covalent bond stretching in diatomic molecules. Our results indicate that out of four DCFT methods the ODC-12 method is the most reliable and accurate DCFT formulation to date. Compared to CCSD, ODC-12 shows superior results for all benchmark tests employed in our study. With respect to coupled-pair

<sup>\*</sup>A. V. Copan, A. Yu. Sokolov, and H. F. Schaefer., J. Chem. Theory Comput. **10**, 2389 (2014). Adapted with permission of the American Chemical Society.

theories, ODC-12 outperforms CEPA<sub>0</sub>, and shows similar accuracy to the orbital-optimized CEPA<sub>0</sub> variant (OCEPA<sub>0</sub>) for systems at equilibrium geometries. For covalent bond stretching, ODC-12 is found to be more reliable than OCEPA<sub>0</sub>. For the RSE30 and AIE datasets, ODC-12 shows competitive performance with CCSD(T). In addition to benchmark results, we report new reference values for the RSE30 dataset computed using coupled cluster theory with up to perturbative quadruple excitations.

#### 2.2 Introduction

Recent developments in *ab initio* quantum chemistry have resulted in a variety of computational models for studying molecules. Apart from concerns about efficiency and accuracy, several concepts have evolved as criteria for judging the merits of a particular method. Energy-based criteria typically define an "ideal" approximation as one yielding correlation energies that are size-consistent, extensive<sup>6</sup>, well-defined (giving continuous, unique potential surfaces), and variational. While it has been argued that the practical benefits of variationality are rather limited, the efficiency of gradient computations, at least, is improved by formulating a theory in terms of a Hermitian and stationary energy functional. With respect to scope and stability, methods that show consistent performance for open-shell systems, strongly correlated states, and non-equilibrium geometries are particularly valuable.

The incorrect scaling of truncated configuration interaction (CI) energies with system size has inspired the development of size-extensive alternatives. Among the earliest formulations, the coupled electron pair approximations (CEPAs)<sup>10-14</sup> attracted much attention in 1970s, 15-19 offering rigorous extensivity and sizeconsistency while retaining much of the linearity <sup>20</sup> of CI in their equations. CEPA methods, however, have been shown to rapidly deteriorate as the molecular geometry deviates from equilibrium<sup>20</sup> and yield energies that vary under the rotation of the occupied orbitals. 13 Partly in light of such defects, CEPA has been largely displaced by coupled-cluster (CC) theory. 8:21-27 In addition to size-extensivity, CC offers orbital invariance and improved stability for non-equilibrium structures<sup>20</sup>, but has a non-Hermitian energy functional and non-linear equations which are not readily amenable to parallel implementation. Although neither class of methods is strictly variational, VCEPA (variational CEPA) has been shown to be effectively equivalent to its non-variational counterpart. 28 Various other modifications to resolve the deficiencies of traditional CEPA have been explored, including self-consistent size-consistent CI,  $^{29;30}$  orbital-invariant CEPA,  $^{31;32}$  and orbital-invariant CEPA, optimized CEPA formulations. 5;33-35 Recently, the CEPA methods have been revived by Neese and co-workers <sup>28;36;37</sup> who developed the local pair-natural-orbital CEPA (LPNO-CEPA) methods and have implemented them for massively parallel computer architectures.

It has recently been demonstrated <sup>1;38–40</sup> that CEPA methods naturally arise in the context of theories that obtain the molecular energies from density cumulants, the connected and extensive components of the reduced density matrices (RDMs). <sup>41–46</sup> The advantage of cumulant-based theories is that, unlike their RDM-based counterparts, <sup>47–49</sup> they are naturally size-extensive and size-consistent. <sup>44;50</sup>

We have recently achieved the first implementation  $^{51;52}$  of density cumulant functional theory (DCFT), proposed by Kutzelnigg in 2006. In DCFT, the molecular energy is obtained in terms of a mean-field one-particle RDM and the two-particle density cumulant, constrained to be at least approximately N-representable (*i.e.* to correspond to a physical N-electron wavefunction). Like traditional CC theory, DCFT is size-extensive and orbital-invariant, but it has the additional advantage of a stationary and Hermitian energy functional, which simplifies the computation of molecular properties. In the original DCFT formulation (DC-06)  $^{1;51;52}$  N-representability conditions derived from second-order Møller-Plesset perturbation theory (MPPT) were used,  $^{53}$  yielding equations similar to those of the simplest CEPA model (CEPA<sub>0</sub>),  $^{12;14}$  but including higher-order terms in the description of one-particle correlation effects. Using the same set of conditions, we have developed new formulations of DCFT that take advantage of an improved description of the one-particle density matrix (DC-12) and full orbital optimization (ODC-06 and ODC-12 methods).

Our previous studies<sup>2;4;51;52</sup> demonstrated for a limited set of systems that the DC-06, DC-12, ODC-06 and ODC-12 methods generally yield molecular energies and properties competitive with those obtained by CCSD and CCSD(T), but may exhibit unstable performance due to imbalances in the description of electron correlation. Herein, we present an extensive benchmark of the DCFT methods with respect to thermochemical and kinetic molecular properties, including noncovalent interactions, barrier heights in hydrogen-transfer reactions, radical stabilization energies, and adiabatic ionization energies for challenging electron-dense systems.

We conclude our benchmark study by testing the performance of DCFT for covalent bond stretching in diatomic molecules.

#### 2.3 Overview of DCFT

In this section a short overview of DCFT is presented. For details on the theory the reader is referred to our earlier publications.  $^{2;4;51}$  In the RDM methods  $^{54}$  the exact molecular energy is expressed as a functional of the one- and two-particle reduced density matrices,  $\gamma_1$  and  $\gamma_2$  (1-RDM and 2-RDM):

$$E = h_p^q \gamma_q^p + \frac{1}{2} g_{pq}^{rs} \gamma_{rs}^{pq} , \qquad [\boldsymbol{\gamma}_1]_q^p \equiv \gamma_q^p , \qquad [\boldsymbol{\gamma}_2]_{rs}^{pq} \equiv \gamma_{rs}^{pq} . \tag{2.1}$$

In eq. (2.1),  $h_p^q$  and  $g_{pq}^{rs}$  are the usual one- and two-electron integrals in the orthonormal spin-orbital basis  $\{\psi_p\}$  and summation over the repeated indices is implied. Expressing  $\gamma_1$  through  $\gamma_2$  via the partial trace relation  $\sum_r \gamma_{qr}^{pr} = (N-1)\gamma_q^p$ , the energy functional (2.1) can be minimized by varying  $\gamma_2$  subject to N-representability constraints. This is the essence of the variational 2-RDM approach. <sup>54</sup>

In DCFT, some of the challenges of the 2-RDM approach are circumvented by expanding  $\gamma_2$  in terms of its irreducible components – the 1-RDM and the two-particle cumulant (denoted by  $\lambda_2$ ):

$$\gamma_{rs}^{pq} = \gamma_r^p \gamma_s^q - \gamma_r^q \gamma_s^p + \lambda_{rs}^{pq}. \tag{2.2}$$

In eq. (2.2),  $\lambda_2$  describes the correlated part of  $\gamma_2$  that cannot be expressed via  $\gamma_1$ . The cumulant also determines the correlation contribution to  $\gamma_1$ , allowing

the 1-RDM to be decomposed as the sum of an idempotent 1-RDM ( $\kappa$ ) and a correlation correction ( $\tau$ ):

$$\gamma_1 = \kappa + \tau \,. \tag{2.3}$$

The correlation component  $\tau$  is fully specified by  $\lambda_2$ , whereas  $\kappa$  is independent of  $\lambda_2$ . Equations (2.2) and (2.3) allow us to write an equivalent energy expression with  $\kappa$  and  $\lambda_2$  as independent functional parameters:

$$E[\kappa, \lambda_2] = \frac{1}{2} (h_p^q + f_p^q) (\kappa_q^p + \tau_q^p) + \frac{1}{4} \overline{g}_{pq}^{rs} \lambda_{pq}^{rs}, f_p^q = h_p^q + \overline{g}_{pr}^{qs} (\kappa_s^r + \tau_s^r), \quad \overline{g}_{rs}^{pq} = g_{rs}^{pq} - g_{rs}^{qp}.$$
(2.4)

Here, the generalized Fock operator f differs from that of Hartree-Fock theory by the presence of an external potential  $\bar{g}_{pr}^{qs}\tau_s^r$  due to electron correlation.<sup>1</sup>

To date, all DCFT formulations make the energy (2.4) stationary with respect to variations of  $\lambda_2$ , subject to cumulant N-representability constraints derived from second-order Møller-Plesset perturbation theory (MPPT). <sup>53</sup> To account for orbital relaxation effects, the two earliest DCFT methods, DC-06<sup>1;51;52</sup> and DC-12<sup>2</sup>, determined the orbitals by diagonalizing the generalized Fock operator f defined in eq. (2.4). These two methods differ in their description of 1-RDM N-representability. Whereas DC-06 employs an approximate expression for  $\tau$  in terms of  $\lambda_2$ , DC-12 uses the exact relationship. Recently, we proposed orbital-optimized variants of DC-06 and DC-12 (ODC-06 and ODC-12), <sup>4</sup> which fully account for orbital relaxation effects.

#### 2.4 Computational Details

All computations were performed using the Psi4 package. <sup>55</sup> The results were benchmarked against coupled cluster theory with single and double excitations  $(CCSD)^{25-27}$ , CCSD with perturbative triple excitations [CCSD(T)], <sup>56;57</sup> coupled electron pair approximation zero  $(CEPA_0)$ , <sup>12;14</sup> and the orbital-optimized variant of CEPA<sub>0</sub>  $(OCEPA_0)^5$ . All electrons were correlated in all computations. The cc-pCVXZ<sup>58;59</sup> and aug-cc-pVXZ<sup>60</sup> basis sets (X = T, Q) were used (see text for details). Noncovalent interaction energies, hydrogen-transfer barrier heights, and radical stabilization energies were computed using geometries from the A24<sup>61</sup>, HTBH38<sup>62</sup>, and RSE30<sup>63</sup> benchmark databases, respectively, available in Psi4. Adiabatic ionization energies were computed from neutral and cation geometries optimized at each level of theory, with added harmonic zero-point vibrational energy corrections. Harmonic frequencies were computed by numerical differentiation of analytic energy gradients. Single-point energies were converged to  $10^{-8} E_h/a_0$  for geometry optimizations.

#### 2.5 Results

#### 2.5.1 Noncovalent Interactions

We begin by testing the accuracy of DCFT methods for the description of noncovalent interactions in 24 closed-shell molecular dimers, which are listed in table 2.1. These molecular complexes comprise the A24 dataset <sup>61</sup> developed by Řezáč and

14

Table 2.1: Errors in interaction energies (kcal mol<sup>-1</sup>) for 24 noncovalently bound molecular dimers comprising the A24 database<sup>61</sup> computed using seven methods with the aug-cc-pVTZ basis set. The errors are relative to CCSD(T) reference values (kcal mol<sup>-1</sup>) shown in the rightmost column. For each method the mean absolute deviations from CCSD(T) ( $\Delta_{MAE}$ , kcal mol<sup>-1</sup>) and the standard deviations from the mean signed error ( $\Delta_{SD}$ , kcal mol<sup>-1</sup>) are also shown.

Complex (Sym.)	$\Delta \text{CEPA}_0$	$\Delta DC-06$	$\Delta$ DC-12	$\Delta \text{CCSD}$	$\Delta  ext{OCEPA}_0$	$\Delta \text{ODC-06}$	$\Delta \text{ODC-12}$	CCSD(T)
$H_2O\cdots NH_3$ (C <sub>s</sub> )	0.26	0.24	0.22	0.36	0.19	0.20	0.18	-7.18
$\mathrm{H_2O\cdots H_2O}$ (C <sub>s</sub> )	0.19	0.18	0.16	0.25	0.13	0.14	0.12	-5.71
$HCN\cdots HCN (C_s)$	0.21	0.27	0.16	0.15	0.18	0.26	0.14	-7.12
$\mathrm{HF}$ ··· $\mathrm{HF}$ ( $\mathrm{C_s}$ )	0.14	0.13	0.11	0.16	0.08	0.09	0.07	-5.20
$NH_3 \cdots NH_3 (C_{2h})$	0.15	0.13	0.14	0.26	0.12	0.12	0.12	-3.43
$\mathrm{HF}\cdots\mathrm{CH}_{4}\ (\mathrm{C}_{3\mathrm{v}})$	0.17	0.16	0.20	0.23	0.12	0.12	0.16	-2.30
$NH_3\cdots CH_4 (C_{3v})$	0.07	0.05	0.05	0.13	0.05	0.05	0.04	-1.08
$\mathrm{H_2O}$ ···· $\mathrm{CH_4}$ ( $\mathrm{C_s}$ )	0.06	0.05	0.04	0.11	0.05	0.05	0.04	-1.03
$CH_2O\cdots CH_2O$ ( $C_s$ )	0.89	0.99	0.65	0.46	0.62	0.87	0.46	-5.23
$H_2O\cdots C_2H_4$ (C <sub>s</sub> )	0.15	0.16	0.15	0.31	0.20	0.26	0.21	-3.33
$\mathrm{CH_2O\cdots C_2H_4}$ (C <sub>s</sub> )	0.21	0.18	0.14	0.27	0.19	0.24	0.16	-2.24
$\frac{\text{HCCH} \cdot \cdot \cdot \text{HCCH (C}_{2v})}{\text{HCCH} \cdot \cdot \cdot \text{HCCH (C}_{2v})}$	) 0.07	0.05	0.05	0.20	0.10	0.12	0.10	-2.57

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Complex (Sym.)	$\Delta \text{CEPA}_0$	$\Delta DC-06$	$\Delta DC-12$	$\Delta \text{CCSD}$	$\Delta  ext{OCEPA}_0$	$\Delta \text{ODC-06}$	$\Delta \text{ODC-12}$	CCSD(T)
$\overline{\mathrm{NH_3\cdots C_2H_4\ (C_s)}}$	0.09	0.06	0.08	0.24	0.12	0.15	0.13	-2.07
$C_2H_4\cdots C_2H_4$ ( $C_{2v}$ )	0.10	0.02	0.07	0.33	0.14	0.13	0.15	-1.81
$\mathrm{CH_4\cdots C_2H_4}$ (C <sub>s</sub> )	0.02	-0.02	0.01	0.14	0.05	0.04	0.06	-0.92
$\mathrm{BH_3\cdots CH_4}\ (\mathrm{C_s})$	0.23	0.18	0.24	0.37	0.18	0.16	0.22	-2.52
$\mathrm{CH_4\cdots C_2H_4}$ (C <sub>s</sub> )	0.13	0.09	0.13	0.23	0.10	0.09	0.09	-1.37
$\mathrm{CH_4\cdots C_2H_6}$ (C <sub>s</sub> )	0.09	0.06	0.09	0.17	0.07	0.06	0.09	-1.14
$CH_4\cdots CH_4 (D_{3d})$	0.08	0.06	0.08	0.14	0.06	0.05	0.08	-0.93
$Ar \cdots CH_4 (C_{3v})$	0.07	0.05	0.07	0.10	0.05	0.05	0.06	-0.78
$Ar \cdots C_2 H_4 (C_{2v})$	0.03	-0.01	0.02	0.11	0.05	0.03	0.05	-0.63
$C_2H_4\cdots HCCH (C_{2v})$	-0.02	-0.19	-0.01	0.38	0.07	-0.06	0.11	0.43
$C_2H_4\cdots C_2H_4$ (D <sub>2h</sub> )	-0.05	-0.30	-0.03	0.43	0.04	-0.16	0.11	0.41
$HCCH\cdots HCCH$ ( $D_{2h}$	0.01	-0.09	0.02	0.34	0.10	0.02	0.12	0.91
$\Delta_{ ext{MAE}}$ :	0.14	0.16	0.12	0.25	0.13	0.15	0.13	
$\Delta_{\mathrm{SD}}$ :	0.18	0.23	0.13	0.11	0.12	0.18	0.09	

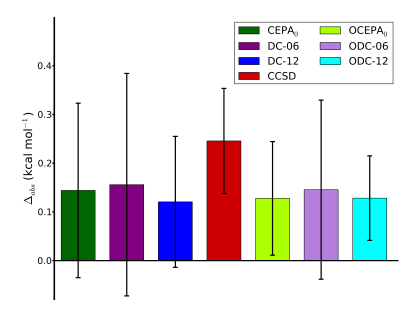


Figure 2.1: Mean absolute deviations ( $\Delta_{MAE}$ , kcal mol<sup>-1</sup>) and the standard deviations from the mean signed error ( $\Delta_{SD}$ , kcal mol<sup>-1</sup>) of the interaction energies for 24 noncovalently bound molecular dimers (A24 database) computed using seven methods with the aug-cc-pVTZ basis set. The errors are relative to CCSD(T)/aug-cc-pVTZ reference values. The  $\Delta_{MAE}$  value is represented as a height of each colored box, while the  $\Delta_{SD}$  value is depicted as a radius of the black vertical bar. See table 2.1 for data on individual database members.

Hobza to include a variety of noncovalent interactions, including hydrogen bonding and  $\pi$ - $\pi$  stacking. Although Řezáč and Hobza reported the interaction energies at the CCSD(T) complete basis set (CBS) limit, we use CCSD(T)/aug-cc-pVTZ energies as reference values in order to effectively exclude basis-set incompleteness error from the comparison.

Figure 2.1 depicts mean absolute error ( $\Delta_{\text{MAE}}$ ) relative to CCSD(T) in the binding energies of CEPA<sub>0</sub>, OCEPA<sub>0</sub>, CCSD, and the four DCFT methods (DC-

06, DC-12, ODC-06, and ODC-12), as well as the root mean square deviation from the average signed error ( $\Delta_{\rm SD}$ ). All methods but CCSD give similar  $\Delta_{\rm MAE}$  values (0.14±0.02 kcal mol<sup>-1</sup>), and a comparison between CEPA<sub>0</sub>, DC-06, and DC-12 and their orbital-optimized variants (OCEPA<sub>0</sub>, ODC-06, and ODC-12) shows negligible 0.01 kcal mol<sup>-1</sup> differences in each case. CCSD gives a significantly larger  $\Delta_{\rm MAE}$  (0.25 kcal mol<sup>-1</sup>) than the other methods, exceeding the DC-12  $\Delta_{\rm MAE}$  by a factor of two (0.12 kcal mol<sup>-1</sup>). The  $\Delta_{\rm SD}$  values are much more sensitive to the choice of method than the  $\Delta_{\rm MAE}$  values, and are noticeably affected by orbital optimization. ODC-12 gives the smallest standard deviation ( $\Delta_{\rm SD}=0.09$  kcal mol<sup>-1</sup>), while the largest  $\Delta_{\rm SD}$  value was found for DC-06 (0.23 kcal mol<sup>-1</sup>). The OCEPA<sub>0</sub>, ODC-06, and ODC-12 methods ( $\Delta_{\rm SD}=0.12, 0.18, {\rm and 0.09 \, kcal \, mol^{-1}}, {\rm respectively})$  exhibit much more consistent performance than their non-orbital-optimized analogues, with  $\Delta_{\rm SD}$  smaller by 0.05 ± 0.01 kcal mol<sup>-1</sup> in each case. CCSD also exhibits a relatively small  $\Delta_{\rm SD}$  value (0.11 kcal mol<sup>-1</sup>), possibly due to its inclusion of single excitations which partly account for orbital relaxation.

Errors in interaction energy and CCSD(T) reference values for each molecular complex are shown in table 2.1. The largest deviations from CCSD(T) were obtained for the formaldehyde dimer (CH<sub>2</sub>O···CH<sub>2</sub>O, complex 9 in table 2.1), for which DC-06, CEPA<sub>0</sub>, and OCEPA<sub>0</sub> yield errors of 0.99, 0.89, and 0.87 kcal mol<sup>-1</sup>, respectively. For this system, the best performance is shown by CCSD and ODC-12, both of which give an error of 0.46 kcal mol<sup>-1</sup>. For systems with  $\pi$ -stacking interactions (complexes 22-24 in table 2.1), CCSD shows large errors (0.38, 0.43, 0.34 kcal mol<sup>-1</sup>) relative to the magnitude of the interaction energy (0.43, 0.41,

0.91 kcal mol<sup>-1</sup>, respectively). Here CEPA<sub>0</sub>, DC-12, and their orbital-optimized variants offer much better agreement with CCSD(T), with errors ranging from 0.01 to 0.15 kcal mol<sup>-1</sup>.

#### 2.5.2 Hydrogen-Transfer Reaction Barrier Heights

We continue by assessing the performance of DCFT methods in predicting barrier heights for 18 hydrogen-transfer reactions from the HTBH38 database: <sup>62</sup>

$$R_1 + R_2 H \longrightarrow [R_1 R_2 H]^* \longrightarrow R_1 H + R_2$$
 (2.5)

These reactions<sup>†</sup> involve molecules ( $R_1$  and  $R_2$ ) and transition states ( $[R_1R_2H]^*$ ) with open-shell character, making their properties more sensitive to electron correlation effects. We employ barrier heights computed at the CCSD(T)/aug-cc-pVTZ level of theory as our reference rather than the values provided by Lynch<sup>64</sup> in order to effectively exclude basis-set incompleteness effects. We also omit the DC-06 and ODC-06 methods, which encounter frequent convergence problems due to the poor description of N-representability (see Supporting Information for incomplete DC-06 results).

Mean absolute deviations ( $\Delta_{MAE}$ ) and standard deviations ( $\Delta_{SD}$ ) for the hydrogentransfer barrier heights are presented in table 2.2 and plotted in fig. 2.2. The largest  $\Delta_{MAE}$  values come from CEPA<sub>0</sub> and DC-12 (2.77 and 2.49 kcal mol<sup>-1</sup>, respectively). Orbital optimization greatly improves the accuracy of these methods,

<sup>&</sup>lt;sup>†</sup>Reaction 19 in HTBH38, the cis-trans isomerization of piperylene, is omitted in the present study.

Table 2.2: Errors in barrier heights (kcal mol<sup>-1</sup>) for 18 hydrogen-transfer reactions ( $R_1 + R_2H \rightarrow R_1H + R_2$ ) comprising the HTBH38 database<sup>64</sup> computed using five methods with the aug-cc-pVTZ basis set. The errors are relative to CCSD(T) reference values (kcal mol<sup>-1</sup>) shown in the rightmost column. Each reaction includes barrier heights in the forward ( $R_1 + R_2H \rightarrow [R_1R_2H]^*$ ) and reverse ( $[R_1R_2H]^* \leftarrow R_1H + R_2$ ) directions, respectively, except in the case of  $R_1 = R_2 = H$  where they are the same. The mean absolute ( $\Delta_{MAE}$ , kcal mol<sup>-1</sup>) and the mean percent ( $\Delta_{rel}$ , %) errors with respect to CCSD(T), as well as the standard deviations from the mean signed error ( $\Delta_{SD}$ , kcal mol<sup>-1</sup>) are also shown.

	Reaction Barrier	$\Delta \text{CEPA}_0$	$\Delta$ DC-12	$\Delta \text{CCSD}$	$\Delta \text{OCEPA}_0$	$\Delta \text{ODC-12}$	$\overline{\mathrm{CCSD}(\mathrm{T})}$
1	$H + HCl \rightarrow [HHCl]^*$	0.74	0.49	0.09	-0.41	-0.28	5.22
2	$OH + H_2 \rightarrow [OHH_2]^*$	3.77	3.38	1.82	0.88	1.24	4.99
3	$CH_3 + H_2 \rightarrow [CH_3H_2]^*$	1.60	1.46	1.37	0.46	0.70	11.29
4	$OH + CH_4 \rightarrow [OHCH_4]^*$	4.26	3.85	2.61	1.22	1.65	5.64
5	$H + H_2 \rightarrow [HH_2]^*$	0.80	0.69	0.30	-0.27	-0.05	9.77
6	$OH + NH_3 \rightarrow [OHNH_3]^*$	6.02	5.25	3.54	1.18	1.82	3.17
7	$HCl + CH_3 \rightarrow [HClCH_3]^*$	1.93	1.78	1.79	0.68	0.92	0.10
8	$OH + C_2H_6 \rightarrow [OHC_2H_6]^*$	4.66	4.21	2.69	1.28	1.72	2.69
9	$F + H_2 \rightarrow [FH_2]^*$	3.40	3.14	1.20	0.52	0.78	1.13
10	$O + CH_4 \rightarrow [OHCH_3]^*$	3.40	3.12	2.37	0.70	1.20	13.62
11	$H + PH_3 \rightarrow [HPH_3]^*$	0.93	0.86	0.59	-0.16	0.10	2.29
12	$H + HO \rightarrow [OHH]^*$	2.03	1.59	0.44	-0.61	-0.26	10.25
13	$H + H_2S \rightarrow [HH_2S]^*$	1.01	0.92	0.65	-0.11	0.14	3.17
14	$O + HCl \rightarrow [OHCl]^*$	6.33	6.01	3.58	0.79	1.51	9.74
15	$NH_2 + CH_3 \rightarrow [CH_3NH_2]^*$	2.48	2.22	1.99	0.49	0.86	7.66
16	$NH_2 + C_2H_5 \rightarrow [NH_2C_2H_5]^*$	2.48	2.22	2.09	0.55	0.92	8.21
17	$C_2H_6 + NH_2 \rightarrow [C_2H_6NH_2]^*$	3.30	3.00	2.73	1.23	1.62	10.39
18	$\mathrm{NH_2} + \mathrm{CH_4} {\rightarrow} [\mathrm{NH_2CH_4}]^*$	2.98	2.72	2.55	1.11	1.48	13.23

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	⊃

	Reaction Barrier	$\Delta \text{CEPA}_0$	$\Delta$ DC-12	$\Delta \text{CCSD}$	$\Delta \text{OCEPA}_0$	$\Delta$ ODC-12	CCSD(T)
1	$[HHCl]^* \leftarrow H_2 + Cl$	1.44	1.31	1.61	0.53	0.77	7.39
2	$[OHH_2]^* \leftarrow H + H_2O$	2.09	1.66	0.09	-0.91	-0.58	21.07
3	$[CH_3H_2]^* \leftarrow H + CH_4$	0.95	0.80	0.38	-0.38	-0.11	14.91
4	$[OHCH_4]^* \leftarrow CH_3 + H_2O$	3.23	2.80	1.87	0.27	0.65	18.09
6	$[OHNH_3]^* \leftarrow H_2O + NH_2$	5.46	4.62	3.14	0.79	1.33	13.17
7	$[HClCH_3]^* \leftarrow Cl + CH_4$	1.97	1.94	2.31	0.78	1.16	5.89
8	$[OHC_2H_6]^* \leftarrow H_2O + C_2H_5$	3.34	2.89	1.85	0.28	0.64	18.49
9	$[FH_2]^* \leftarrow HF + H$	1.27	0.88	-0.78	-1.47	-1.33	32.95
10	$[OHCH_3]^* \leftarrow OH + CH_3$	2.62	2.29	1.82	0.32	0.68	7.43
11	$[HPH_3]^* \leftarrow PH_2 + H_2$	1.14	1.11	1.37	0.39	0.63	23.21
12	$[OHH]^* \leftarrow H_2 + O$	3.47	3.08	1.99	0.62	1.07	12.81
13	$[HH_2S]^* \leftarrow H_2 + HS$	1.51	1.51	1.88	0.65	0.97	16.41
14	$[OHCl]^* \leftarrow OH + Cl$	5.59	5.35	3.55	0.51	1.24	9.35
15	$[CH_3NH_2]^* \leftarrow CH_4 + NH$	2.77	2.49	2.26	0.73	1.12	21.32
16	$[\mathrm{NH_2C_2H_5}]^* \leftarrow \mathrm{C_2H_6} + \mathrm{NH}$	3.06	2.75	2.46	0.85	1.26	18.52
17	$[C_2H_6NH_2]^* \leftarrow NH_3 + C_2H_5$	2.54	2.30	2.30	0.63	1.02	16.20
18	$[\mathrm{NH_2CH_4}]^* \leftarrow \mathrm{CH_3} + \mathrm{NH_3}$	2.51	2.29	2.21	0.56	0.96	15.69
	$\Delta_{ ext{MAE}}$ :	2.77	2.49	1.84	0.67	0.94	
	$\Delta_{ ext{SD}}$ :	1.51	1.39	1.06	0.62	0.71	
	$\Delta_{ m rel}$ %:	99	90	77	29	40	

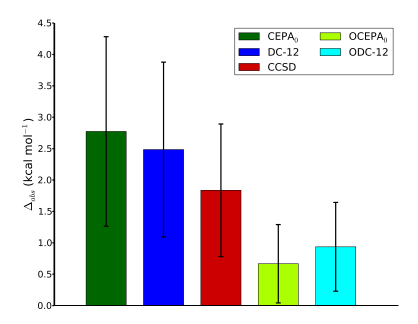


Figure 2.2: Mean absolute deviations ( $\Delta_{MAE}$ , kcal mol<sup>-1</sup>) and the standard deviations from the mean signed error ( $\Delta_{SD}$ , kcal mol<sup>-1</sup>) of barrier heights for 18 hydrogen-transfer reactions ( $R_1 + R_2H \rightarrow R_1H + R_2$ , HTBH38 database) computed using five methods with the aug-cc-pVTZ basis set. The errors are relative to CCSD(T)/aug-cc-pVTZ reference values. The  $\Delta_{MAE}$  value is represented as a height of each colored box, while the  $\Delta_{SD}$  value is depicted as a radius of the black vertical bar. See table 2.2 for data on individual database members.

resulting in  $\Delta_{\text{MAE}}$  values of just 0.67 and 0.94 kcal mol<sup>-1</sup> for OCEPA<sub>0</sub> and ODC-12, respectively. The CCSD method shows intermediate performance with  $\Delta_{\text{MAE}}$  = 1.84 kcal mol<sup>-1</sup>. A similar trend is observed for the  $\Delta_{\text{SD}}$  values, with OCEPA<sub>0</sub> (0.62 kcal mol<sup>-1</sup>) and ODC-12 (0.71 kcal mol<sup>-1</sup>) significantly improving upon CEPA<sub>0</sub> (1.51 kcal mol<sup>-1</sup>), DC-12 (1.39 kcal mol<sup>-1</sup>), and CCSD (1.06 kcal mol<sup>-1</sup>). In addition to  $\Delta_{\text{MAE}}$  and  $\Delta_{\text{SD}}$ , table 2.2 includes mean percent error ( $\Delta_{\text{rel}}$ ) values,

which are commonly used to benchmark performance for reaction kinetics. The smallest  $\Delta_{\rm rel}$  values are 29% and 40% for OCEPA<sub>0</sub> and ODC-12, respectively.

Turning to barrier heights for individual hydrogen-transfer reactions (table 2.2), the largest errors are observed for reactions 6 and 14, both involving the OH radical, for which CEPA<sub>0</sub> and DC-12 give errors of  $\sim 5$ -6 kcal mol<sup>-1</sup>. The best results for these reactions are obtained from OCEPA<sub>0</sub>, with errors ranging from 0.51 to 1.18 kcal mol<sup>-1</sup>. The ODC-12 method tends to predict larger barrier heights than OCEPA<sub>0</sub>, yielding smaller errors only when OCEPA<sub>0</sub> underestimates the barrier heights.

#### 2.5.3 Radical Stabilization Energies

In this section we study the performance of DCFT methods for predicting radical stabilization energies (RSEs). An R-group's RSE is defined as the enthalpy of a homodesmotic reaction

$$RH + CH_3 \longrightarrow R + CH_4$$
 (2.6)

where exothermic (negative) values indicate that the radical 'R is more thermodynamically stable than 'CH<sub>3</sub>. <sup>65</sup> For our benchmark we use the RSE30 dataset <sup>63</sup>, which provides a diverse variety of 'R species (listed in table 2.3). Since the performance of CCSD(T) is known to deteriorate for strongly spin-contaminated UHF references, <sup>66–70</sup> we augment CCSD(T) energies with a quadruples correction ( $\Delta Q = E_{CCSDT(Q)} - E_{CCSD(T)}$ ) and use these as our benchmark. CBS-extrapolated CCSD(T) reference values have been published for this dataset, <sup>34</sup> but we use

Table 2.3: Errors in radical stabilization energies (RSEs, kcal mol<sup>-1</sup>) for 30 open-shell doublet species ('R) comprising the RSE30 database<sup>63</sup> computed using six methods with the cc-pCVTZ basis set. The errors are relative to CCSD(T) with an added quadruples correction ( $\delta Q = E_{\text{CCSDT(Q)}} - E_{\text{CCSD(T)}}$ ) shown in the rightmost column in kcal mol<sup>-1</sup>. The  $\delta Q$  correction was computed using the cc-pCVDZ basis set. RSE is defined as the reaction enthalpy for the homodesmotic reaction 'CH<sub>3</sub>+RH $\rightarrow$ CH<sub>4</sub>+'R. To indicate the degree of spin-contamination in the UHF reference, the spin expectation values ( $\langle \hat{S}^2 \rangle_{\text{SCF}}$ ) are also shown in units of  $\hbar^2$ . For each method the mean absolute deviations from CCSD(T)+ $\delta Q$  ( $\Delta_{\text{MAE}}$ , kcal mol<sup>-1</sup>) and the standard deviations from the mean signed error ( $\Delta_{\text{SD}}$ , kcal mol<sup>-1</sup>) are also presented.

·R	$\langle \hat{S}^2 \rangle_{\mathrm{SCF}}$	$\Delta \text{CEPA}_0$	$\Delta DC-12$	$\Delta \text{CCSD}$	$\Delta  ext{OCEPA}_0$	$\Delta \text{ODC-12}$	$\Delta \text{CCSD}(T)$	$CCSD(T) + \delta Q$
$^{\cdot}\mathrm{CH_{2}NO_{2}}$	0.78	1.24	0.95	0.66	0.16	0.27	0.32	-3.50
$^{\cdot}\mathrm{CH_{2}OCHO}$	0.76	1.16	1.12	0.63	0.40	0.48	0.10	-4.84
$^{\cdot}\mathrm{CH}_{2}\mathrm{SCH}_{3}$	0.76	1.89	1.70	0.81	0.63	0.72	0.15	-11.01
$^{\cdot}CF=CH_2$	0.94	6.12	3.71	0.96	0.42	0.64	0.46	6.26
$^{\cdot}\mathrm{CH_{2}CH_{2}F}$	0.76	0.30	0.27	0.13	0.08	0.10	0.04	-1.53
$^{\cdot}\mathrm{CH_{2}CHO}$	0.93	5.01	2.86	0.32	-0.16	0.02	0.46	-10.11
$^{\cdot}\mathrm{CH_{2}CN}$	0.94	6.36	3.52	0.65	-0.02	0.21	0.46	-8.66
$^{\cdot}\mathrm{CH_{2}F}$	0.76	1.03	1.00	0.52	0.55	0.57	0.06	-4.22
$^{\cdot}\mathrm{CH_{2}NH_{2}}$	0.76	1.28	1.18	0.59	0.50	0.52	0.06	-12.06
${}^{\cdot}\mathrm{CH_2NH_3}^+$	0.76	0.16	0.10	0.08	0.06	0.03	0.02	4.58
$^{\cdot}\mathrm{CH_{2}NHOH}$	0.77	1.76	1.57	0.78	0.58	0.64	0.15	-8.81
$^{\cdot}\mathrm{CH_{2}OH}$	0.76	1.29	1.23	0.62	0.57	0.60	0.07	-9.27
$^{\cdot}\mathrm{CH_{2}PH_{3}}^{+}$	0.76	0.21	0.14	0.01	0.01	-0.02	0.05	0.49
$^{\cdot}\mathrm{CH_{2}SH_{2}^{-+}}$	0.77	0.41	0.30	0.12	0.11	0.08	0.06	2.29
$^{\cdot}\mathrm{CH_{2}SH}^{-}$	0.76	1.60	1.43	0.68	0.57	0.63	0.12	-9.68

·R	$\langle \hat{S}^2 \rangle_{\rm SCF}$	$\Delta \text{CEPA}_0$	$\Delta DC$ -12	$\Delta \text{CCSD}$	$\Delta  ext{OCEPA}_0$	$\Delta \text{ODC-12}$	$\Delta \text{CCSD}(T)$	$CCSD(T) + \delta Q$
·CH <sub>2</sub> C≡CH	1.00	6.23	3.47	0.82	-0.03	0.23	0.52	-13.17
$^{\cdot}\mathrm{CH}_{2}\mathrm{CH}_{3}$	0.76	0.30	0.26	0.11	0.08	0.10	0.03	-3.36
$^{\cdot}\mathrm{CH_{2}Cl}$	0.77	1.13	1.02	0.50	0.48	0.51	0.09	-5.67
$^{\cdot}\mathrm{CH_{2}BH_{2}}$	0.76	0.17	0.17	0.05	0.03	0.04	0.05	-11.66
.CHO	0.77	2.26	2.24	1.48	1.55	1.56	0.20	-17.61
$^{\cdot}\mathrm{CH_{2}PH_{2}}$	0.76	1.17	1.02	0.39	0.36	0.39	0.12	-6.50
·CHClF	0.76	1.61	1.52	0.76	0.78	0.81	0.13	-6.61
$\cdot$ CHFCH $_3$	0.76	1.07	1.01	0.50	0.51	0.53	0.08	-5.87
$^{\cdot}\mathrm{CH}(\mathrm{OH})_2$	0.76	1.30	1.22	0.60	0.60	0.61	0.08	-6.67
$^{\cdot}\mathrm{CHCl}_{2}$	0.77	1.78	1.57	0.72	0.72	0.75	0.15	-9.56
$^{\cdot}\mathrm{CHF}_{2}$	0.76	1.50	1.48	0.78	0.83	0.85	0.10	-4.07
$\mathrm{CH_2}{=}\mathrm{C'}{-}\mathrm{CN}$	1.39	19.10	11.50	2.36	-0.31	0.29	1.80	1.98
.С≡СН	1.15	11.20	6.51	0.77	-0.78	-0.07	0.82	26.25
$^{\cdot}CH=CH_2$	0.94	5.42	3.01	0.58	0.11	0.31	0.40	5.49
$^{\cdot}\mathrm{CH}_{2}\mathrm{-CH}\mathrm{=CH}_{2}$	0.97	4.98	3.17	0.51	0.11	0.31	0.48	-17.53
	·	$\Delta_{ ext{MAE}}$ :	2.97	2.01	0.62	0.40	0.43	0.25
		$\Delta_{\mathrm{SD}}$ :	3.97	2.27	0.45	0.43	0.35	0.35

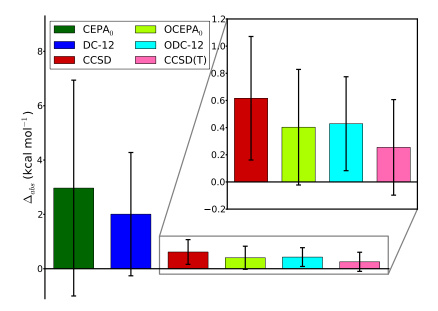


Figure 2.3: Mean absolute deviations ( $\Delta_{\text{MAE}}$ , kcal  $\text{mol}^{-1}$ ) and the standard deviations from the mean signed error ( $\Delta_{\text{SD}}$ , kcal  $\text{mol}^{-1}$ ) of the radical stabilization energies (RSEs) for 30 open-shell doublet species (RSE30 database) computed using six methods with the cc-pCVTZ basis set. The errors are relative to CCSD(T) with an added quadruples correction ( $\delta Q = E_{\text{CCSDT(Q)}} - E_{\text{CCSD(T)}}$ ). The  $\delta Q$  correction was computed using the cc-pCVDZ basis set. RSE is defined as the reaction enthalpy for the homodesmotic reaction 'CH<sub>3</sub> + RH $\rightarrow$ CH<sub>4</sub> + 'R. The  $\Delta_{\text{MAE}}$  value is represented as a height of each colored box, while the  $\Delta_{\text{SD}}$  value is depicted as a radius of the black vertical bar. See table 2.3 for data on individual database members.

CCSD(T) values computed with the cc-pCVTZ basis set to avoid basis-set incompleteness effects. The  $\delta Q$  correction was evaluated using the cc-pCVDZ basis set. As in the previous Section, DC-06 and ODC-06 computations cannot be converged for all database members and are omitted in the analysis below (see Supporting Information for incomplete DC-06 and ODC-06 data).

The relative performance of the DCFT, CEPA, and CC methods for the RSE30 dataset is shown in fig. 2.3. The effect of orbital-optimization on accuracy is now even more pronounced, reducing the large  $\Delta_{\text{MAE}}$  errors of CEPA<sub>0</sub> (2.97 kcal mol<sup>-1</sup>) and DC-12 (2.01 kcal mol<sup>-1</sup>) to 0.40 and 0.43 kcal mol<sup>-1</sup> for OCEPA<sub>0</sub> and ODC-12, respectively. CCSD has a slightly larger  $\Delta_{\text{MAE}}$  value (0.62 kcal mol<sup>-1</sup>), while CCSD(T) has the smallest overall  $\Delta_{\text{MAE}}$  (0.25 kcal mol<sup>-1</sup>). Both CEPA<sub>0</sub> and DC-12 show large standard deviations again (3.97 and 2.27 kcal mol<sup>-1</sup>, respectively). For OCEPA<sub>0</sub>, the standard deviation (0.43 kcal mol<sup>-1</sup>) is similar to that of CCSD (0.45 kcal mol<sup>-1</sup>). ODC-12 and CCSD(T) exhibit the most consistent performance with the same  $\Delta_{\text{SD}}$  value of 0.35 kcal mol<sup>-1</sup>.

Deviations from CCSD(T)+ $\delta$ Q for individual RSEs predicted by each method are tabulated in table 2.3. In addition, table 2.3 includes expectation values of the square-norm spin operator computed for the UHF wavefunction of 'R ( $\langle \hat{S}^2 \rangle_{SCF}$ ). The largest errors in computed RSEs were obtained for 'R species with  $\langle \hat{S}^2 \rangle_{SCF} > 0.9 \,\hbar^2$  (radicals 4, 6, 7, 16, and 27-30 in table 2.3). For these systems, the average CEPA<sub>0</sub> and DC-12 errors are 8.05 and 4.72 kcal mol<sup>-1</sup>, and the average CCSD(T) error is 0.68 kcal mol<sup>-1</sup>. OCEPA<sub>0</sub> and ODC-12 offer remarkably better performance for this subset, with average errors of 0.24 kcal mol<sup>-1</sup> and 0.26 kcal mol<sup>-1</sup>.

#### 2.5.4 Adiabatic Ionization Energies in Electron-Dense Molecules

We conclude the assessment of DCFT methods for the description of thermodynamic properties by computing adiabatic ionization energies (AIEs) for a set of 10

Table 2.4: Errors in adiabatic ionization energies (AIEs, eV) for 10 di- and triatomic molecules computed using six methods with the cc-pCVQZ basis set. The errors are relative to experimental values (IE<sub>ref</sub>, eV) from Ref. 71, unless noted otherwise. For all AIEs the harmonic zero-point vibrational energy corrections were included. For each method the mean absolute deviations from IE<sub>ref</sub> ( $\Delta_{MAE}$ , eV) and the standard deviations from the mean signed error ( $\Delta_{SD}$ , eV) are also shown.

Molecule	Transition	$\Delta \text{CEPA}_0$	$\Delta DC-12$	$\Delta \text{CCSD}$	$\Delta  ext{OCEPA}_0$	$\Delta \text{ODC-}12$	$\Delta \text{CCSD}(T)$	$ ext{IE}_{ ext{ref}}$
$ m N_2$	$^{1}\Sigma_{\rm g}^{+} \rightarrow ^{2}\Sigma_{\rm g}^{+}$	0.08	0.17	0.12	-0.05	0.07	-0.03	$15.581 \pm 0.008$ a
$\mathrm{O}_2$	$^3\Sigma_{\rm g}^-  o {}^2\Pi_{\rm g}$	-0.11	-0.03	0.04	-0.09	-0.02	-0.04	$12.0697\pm0.0002$
$\mathrm{F}_2$	$^{1}\Sigma_{\mathrm{g}}^{+} \rightarrow ^{2}\Pi_{\mathrm{g}}$	0.06	0.06	0.04	0.08	0.01	-0.03	$15.697 \pm 0.003$
NO	$^2\Pi \rightarrow ^1\Sigma^+$	-0.15	-0.05	-0.05	-0.05	-0.02	-0.09	$9.26438 \pm 0.00005$
OF	$^2\Pi  ightarrow ^3\Sigma^-$	0.11	0.12	-0.10	-0.03	-0.02	-0.11	$12.77 \pm 0.01 ^{\mathrm{b}}$
HNC	$^{1}\Sigma_{\mathrm{g}}^{+} \rightarrow ^{2}\Sigma^{+}$	0.27	0.14	-0.12	-0.14	-0.08	-0.04	$12.04\pm0.01^{\rm \ c}$
HOF	$^{1}\text{A}' \rightarrow {}^{2}\text{A}''$	0.20	0.17	-0.10	-0.03	-0.04	-0.07	$12.71 \pm 0.01$
FNO	$^{1}\mathrm{A'} \rightarrow ^{2}\mathrm{A''}$	0.51	0.10	-0.02	-0.02	-0.00	0.04	$12.63 \pm 0.03$
$\mathrm{F_2N}$	$^2\mathrm{B}_1  ightarrow {}^1\mathrm{A}_1$	0.07	0.10	0.07	0.01	0.03	-0.08	$11.63 \pm 0.01$
$F_2O$	$^{1}\mathrm{A}_{1}  ightarrow ^{2}\mathrm{B}_{1}$	0.49	0.37	-0.01	0.05	0.04	-0.04	$13.11 \pm 0.01$
	$\Delta_{\mathrm{MAE}}$ :	0.21	0.13	0.06	0.05	0.03	0.06	
	$\Delta_{\mathrm{SD}}$ :	0.22	0.12	0.08	0.06	0.04	0.04	

<sup>&</sup>lt;sup>a</sup> Reference 72.

<sup>&</sup>lt;sup>b</sup> Reference 73.

<sup>&</sup>lt;sup>c</sup> Reference 74.

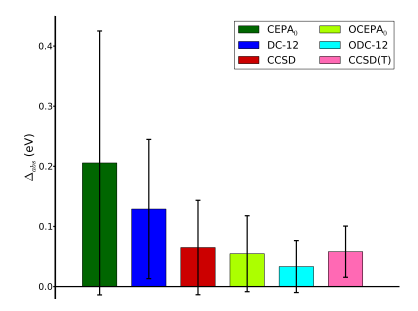


Figure 2.4: Mean absolute deviations ( $\Delta_{MAE}$ , eV) and the standard deviations from the mean signed error ( $\Delta_{SD}$ , eV) of adiabatic ionization energies for 10 diand triatomic molecules computed using six methods with the cc-pCVQZ basis set. The errors are relative to experimental values.<sup>71–74</sup> The  $\Delta_{MAE}$  value is represented as a height of each colored box, while the  $\Delta_{SD}$  value is depicted as a radius of the black vertical bar. See table 2.4 for data on individual molecules.

di- and triatomic electron-dense molecules (table 2.4), i.e. those that are composed of elements with small atomic radius, high electron affinity, and high electronegativity (N, O, F), in order to increase the magnitude of electron correlation effects. We use experimentally measured ionization energies reported to high precision  $(\sim 0.01 \text{ eV})^{71-74}$  as reference values for our benchmark (IE<sub>ref</sub>, table 2.4). The AIEs were computed using the cc-pCVQZ basis set, with harmonic ZPVE corrections applied to each neutral and cationic system.

The  $\Delta_{\text{MAE}}$  and  $\Delta_{\text{SD}}$  values for our computed AIEs relative to experiment are plotted in fig. 2.4. Of the six methods, CEPA<sub>0</sub> and DC-12 exhibit the largest  $\Delta_{\text{MAE}}$  values (0.21 and 0.13 eV, respectively). The closest agreement with experiment is given by ODC-12, with  $\Delta_{\text{MAE}} = 0.03$  eV. OCEPA<sub>0</sub>, CCSD, and CCSD(T) show somewhat poorer performance ( $\Delta_{\text{MAE}} = 0.05$ , 0.06 and 0.06 eV, respectively). The  $\Delta_{\text{SD}}$  for ODC-12 matches that of CCSD(T) (0.04 eV). For the other methods, the  $\Delta_{\text{SD}}$  values decrease in the order CEPA<sub>0</sub> (0.22 eV) > DC-12 (0.12) > CCSD (0.08) > OCEPA<sub>0</sub> (0.06).

Individual errors for each system are shown in table 2.4. Both DC-12 and CEPA<sub>0</sub> exhibit large deviations for  $F_2O$  (0.49 and 0.37 eV), and CEPA<sub>0</sub> also gives a large error for FNO (0.51 eV) which is the maximum error for this dataset. Both DC-12 and CEPA<sub>0</sub> give errors exceeding 0.1 eV for seven of the ten systems, whereas CCSD exhibits errors in excess of 0.1 eV for only three systems (OF, HNC, and HOF). CCSD(T) has only one such error (0.11 eV for OF), as does OCEPA<sub>0</sub> (0.14 eV for HNC). ODC-12 does the best of the methods considered, with a maximum error of 0.08 eV, found for the AIE of HNC.

#### 2.5.5 Covalent Bond Stretching in Diatomic Molecules

Finally, we benchmark DCFT methods for covalent bond stretching. Although accurate description of bond stretching demands the use of multireference methods, our aim here is to explore the limits of DCFT away from equilibrium. For this purpose, we compute the energy as a function of bond distance for diatomic molecules with single (HF and BH), double (BeO), and triple ( $N_2$ ) bonds using the

Figure 2.5: Error in the total energy  $(mE_h)$ , relative to full CI, as a function of B–H internuclear separation  $(\mathring{A})$  computed using six methods with the DZP basis set. The full CI reference is depicted with a horizontal dotted line. The dashed vertical line indicates the full CI equilibrium bond distance.

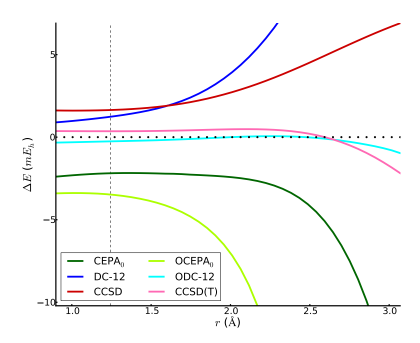


Figure 2.6: Error in the total energy  $(mE_h)$ , relative to full CI, as a function of H–F internuclear separation  $(\mathring{A})$  computed using six methods with the DZP basis set. The full CI reference is depicted with a horizontal dotted line. The dashed vertical line indicates the full CI equilibrium bond distance.

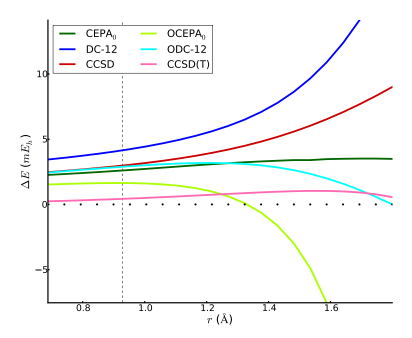


Figure 2.7: Error in the total energy (m $E_{\rm h}$ ), relative to full CI, as a function of Be–O internuclear separation (Å) computed using six methods with the 6-31G basis set. The full CI reference is depicted with a horizontal dotted line. The dashed vertical line indicates the full CI equilibrium bond distance.

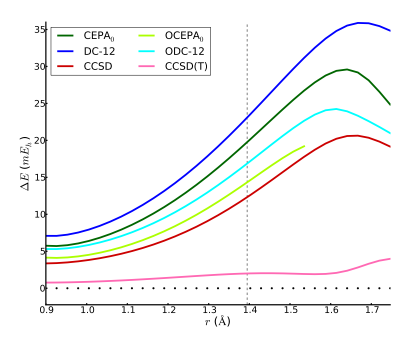
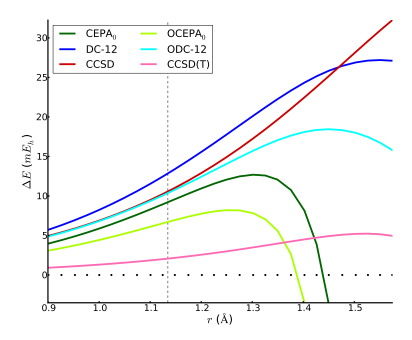


Figure 2.8: Error in the total energy (m $E_{\rm h}$ ), relative to full CI, as a function of N–N internuclear separation (Å) computed using six methods with the 6-31G basis set. The full CI reference is depicted with a horizontal dotted line. The dashed vertical line indicates the full CI equilibrium bond distance.



CEPA<sub>0</sub>, OCEPA<sub>0</sub>, DC-12, ODC-12, CCSD, and CCSD(T) methods. We restrict ourselves to modest basis sets in order to use full CI (FCI) as a reference, and plot the errors with respect to FCI ( $\Delta E$ ) as a function of internuclear distance for each molecule. The relative performance of the methods is described below using non-parallelity errors (NPE =  $\Delta E_{\rm max} - \Delta E_{\rm min}$ , m $E_{\rm h}$ ) computed for specific bond distance ranges.

BH Figure 2.5 shows errors relative to FCI for the BH molecule. DC-12 and CCSD increasingly overestimate the energy at larger internuclear distances, whereas the CEPA<sub>0</sub> error curve is concave down. Orbital optimization lowers the binding energy for OCEPA<sub>0</sub> even further compared to CEPA<sub>0</sub>, leading to large errors with respect to FCI for r(B-H)  $\downarrow$  1.5  $r_{\rm e}$ , where  $r_{\rm e}$  is the FCI equilibrium bond distance ( $r_{\rm e}=1.244~{\rm Å}$ ). At 1.87  $r_{\rm e}$ , OCEPA<sub>0</sub> encounters convergence problems, which originate from numerical instabilities due to the method's deficiencies in the description of N-representability. The ODC-12 method exhibits much more stable behavior with respect to bond stretching in this case, fortuitously showing smaller errors and better parallelity than CCSD(T). For the range [0.72  $r_{\rm e}$ , 2.47  $r_{\rm e}$ ], the NPEs decrease in the order DC-12 (24 m $E_{\rm h}$ ) > CEPA<sub>0</sub> (15) > CCSD (5) > CCSD(T) (3) > ODC-12 (1).

HF Errors for HF bond stretching are plotted in fig. 2.6. The  $\Delta E$  values of CCSD and DC-12 increase as a function of r(H-F), while CEPA<sub>0</sub> fortuitously maintains parallelity similar to CCSD(T) over the range [0.74  $r_{\rm e}$ , 1.94  $r_{\rm e}$ ] ( $r_{\rm e} = 0.929$  Å). OCEPA<sub>0</sub> increasingly overestimates the HF binding energy away from equilibrium,

failing to converge past 1.82  $r_{\rm e}$ . The ODC-12 method exhibits larger NPE than was observed for BH, and encounters convergence problems past 1.94  $r_{\rm e}$ . CCSD(T) shows the best overall performance, with errors between 0 and 1 m $E_{\rm h}$ . In the range [0.74  $r_{\rm e}$ , 1.94  $r_{\rm e}$ ] the computed NPE values are: DC-12 (15 m $E_{\rm h}$ ) > CCSD (7) > ODC-12 (3) > CEPA<sub>0</sub> (1)  $\approx$  CCSD(T) (1). Recently, the orbital-optimized variants of CCSD(T) have been shown to yield good performance for HF bond stretching.<sup>75</sup>

BeO The double bond of BeO presents a more challenging test for the single-reference methods under consideration (fig. 2.7). All methods but CCSD(T) show qualitatively similar error curves, with inflection points near the FCI equilibrium ( $r_{\rm e}=1.394~{\rm \AA}$ ) and valleys/peaks around 0.6  $r_{\rm e}/1.2~r_{\rm e}$ . OCEPA<sub>0</sub> encounters convergence problems past 1.10  $r_{\rm e}$ . The ODC-12 method performs similarly to CCSD. Overall, the NPEs for the range [0.65  $r_{\rm e}$ , 1.10  $r_{\rm e}$ ] decrease in the following order: DC-12 (29 m $E_{\rm h}$ ) > CEPA<sub>0</sub> (24) > ODC-12 (19) > CCSD (17) > CCSD(T) (3).

 $N_2$  Figure 2.8 depicts the errors relative to FCI for triple bond stretching in  $N_2$ . Here, OCEPA<sub>0</sub> fails to converge past 1.24  $r_{\rm e}$  ( $r_{\rm e}=1.135$  Å). The ODC-12 method significantly overestimates the binding energy, possibly due to the lack of three-body correlation effects, but shows much more stable performance compared to methods other than CCSD(T). NPEs in the range [0.79  $r_{\rm e}$ , 1.39  $r_{\rm e}$ ] decrease in the order: CEPA<sub>0</sub> (802 m $E_{\rm h}$ )<sup>‡</sup>> CCSD (27) > DC-12 (21) > ODC-12 (14) > CCSD(T) (4).

 $<sup>^{\</sup>ddagger}\text{CEPA}_0$  exhibits a vertical asymptote at 1.36  $r_{\rm e}$  for  $N_2$  stretching.

### 2.6 Conclusions

We have presented the benchmark study of four density cumulant functional theory (DCFT) methods (DC-06, DC-12, ODC-06, and ODC-12) developed recently in our group. <sup>2;4;51;52</sup> Specifically we have compared the performance of DCFT to that of coupled electron pair methods (CEPA<sub>0</sub> and OCEPA<sub>0</sub>), as well as coupled-cluster theory [CCSD and CCSD(T)] for predicting a variety of chemical properties relevant to thermochemistry and kinetics, with a particular focus on open-shell, electron-dense, and non-equilibrium systems.

Our results indicate that among the four DCFT methods, the best agreement with available reference data is obtained for the ODC-12 method. While all four DCFT formulations yield similar results for the description of noncovalent interactions, DC-06, DC-12, and ODC-06 exhibit worse performance than ODC-12 for thermodynamic and kinetic properties of reactions involving open-shell molecules. In particular, DC-06 and ODC-06 frequently encounter convergence problems that originate from poor description of N-representability. In comparing ODC-12 to other methods, several trends can be observed:

- (i) For all benchmark datasets, ODC-12 outperforms CCSD with errors smaller by almost a factor of two, on average. ODC-12 is also superior to CCSD for the description of single bond stretching in BH and HF, although it does not converge for all bond distances.
- (ii) The performance of ODC-12 and OCEPA<sub>0</sub> is comparable. In particular, for hydrogen-transfer reaction barrier heights, the OCEPA<sub>0</sub> method yields smaller percent errors than ODC-12, whereas, for the radical stabilization energies (RSE)

and adiabatic ionization energies (AIE) in electron-dense molecules, the ODC-12 method smaller standard deviations than OCEPA<sub>0</sub>. For AIEs, ODC-12 gives smaller mean absolute deviations by almost a factor of two. ODC-12 also shows significantly smaller non-parallelity errors than OCEPA<sub>0</sub> for covalent bond stretching, and can be converged for a larger range of distances for all diatomic molecules studied.

(iii) For the two most challenging datasets, RSE and AIE, the standard deviation of ODC-12 and CCSD(T) are similar. While CCSD(T) yields smaller mean absolute errors for the RSE database, the ODC-12 method significantly outperforms CCSD(T) for the AIE test case. However, for bond stretching ODC-12 is competitive with CCSD(T) only for the BH dissociation and shows worse results for other molecules.

Overall, the data presented herein indicates that the ODC-12 method can be used as an efficient  $\mathcal{O}(n^6)$  alternative to CCSD, capable of predicting thermodynamic and kinetic quantities that are competitive in accuracy with the "gold-standard"  $\mathcal{O}(n^7)$  CCSD(T). Although our current implementation of ODC-12 is far from optimal, the ODC-12 equations have reduced non-linearities compared to CCSD, which makes them more amenable to parallel implementation. The efficiency of ODC-12 can also greatly benefit from spin-adaptation, <sup>44;76;77</sup> local approximations, <sup>37;78–80</sup> and density fitting. <sup>78;81–83</sup> Another important advantage of ODC-12 over CCSD is its stationarity, which makes the computation of first-order properties and analytic gradients more efficient and easily accessible. In particu-

lar, ODC-12 has potential to be used for computing accurate response properties which do not suffer from a lack of gauge-invariance.  $^{84;85}$ 

# Chapter 3

Linear-Response Density Cumulant Theory for Excited States: First Implementation and Benchmark Calculations\*

### 3.1 Abstract

We present a linear-response formulation of density cumulant functional theory (DCT) that provides accurate access to many electronic states. DCT expresses the electronic energy as a Hermitian, size-extensive, and stationary functional of the one-particle density matrix and the two-particle density cumulant. In the original DCT formulation only the information about a single electronic state (usually, the ground state) is obtained. In this research, we combine DCT with linear response theory to obtain information about many electronic states simultaneously. We discuss the derivation of linear-response DCT, present its implementation for the ODC-12 method (LR-ODC-12), and benchmark its performance against highly accurate equation-of-motion coupled cluster theory with up to full triple excitations (EOM-CCSDT). Our results for a set of small molecules demonstrate that LR-ODC-12 vertical excitation energies are in closer agreement with EOM-CCSDT

<sup>\*</sup>A. V. Copan and A. Yu. Sokolov (to be submitted in J. Chem. Theory Comput).

than those obtained from equation-of-motion coupled cluster theory with up to double excitations (EOM-CCSD). In addition, we report a linear-response formulation of the orbital-optimized linearized coupled cluster theory with double excitations (LR-OLCCD), which we obtain by neglecting the non-linear terms in the LR-ODC-12 equations.

### 3.2 Introduction

Accurate simulation of excited electronic states remains one of the major challenges in modern electronic structure theory. Ab initio methods for excited states can be divided into single-reference and multi-reference categories, based on their ability to treat static electron correlation. Multi-reference methods can correctly describe static correlation in near-degenerate valence orbitals and electronic states with multiple-excitation character, but often lack accurate treatment of important dynamic correlation effects (e.g., multi-configurational self-consistent field or multi-reference perturbation theories) <sup>86–95</sup> or become very costly when the number of near-degenerate orbitals is large (e.g., multi-reference configuration interaction or coupled cluster theories). <sup>96–108</sup> Meanwhile, single-reference methods <sup>25;27;109–121</sup> often provide a compromise between the computational cost and accuracy, and can be used to reliably compute properties of molecules in low-lying electronic states near the equilibrium geometries. In these situations, single-reference equation-of-motion coupled cluster theory (EOM-CC) <sup>25;27;111–114</sup> is usually the method of choice, especially when high accuracy is desired.

The EOM-CC methods yield size-intensive excitation energies <sup>116;117</sup> and can be systematically improved by increasing the excitation rank of the cluster operator in the exponential parametrization of the wavefunction. Although EOM-CC is usually formulated in the context of a similarity-transformed Hamiltonian, its excitation energies are equivalent to those obtained from linear-response coupled cluster theory (LR-CC). <sup>115–117</sup> Both EOM-CC and LR-CC are based on non-Hermitian eigenvalue problems, complicating the computation of molecular properties (e.g., transition dipoles) by requiring evaluation of left and right eigenvectors. <sup>122–125</sup> Several Hermitian alternatives to EOM-CC and LR-CC have been proposed to avoid these problems, such as algebraic diagrammatic construction <sup>126–130</sup>, unitary and variational LR-CC, <sup>131–133</sup> similarity-constrained CC, <sup>134</sup> and propagator-based LR-CC. <sup>135;136</sup>

In this work, we present the development of linear-response density cumulant functional theory (LR-DCT), a size-intensive approach for excited electronic states. In density cumulant functional theory (DCT), <sup>1–4;51;52;137</sup> the electronic energy is obtained by optimizing the energy functional directly in terms of the one-particle reduced density matrix and the two-particle density cumulant, a fully connected part of the two-particle reduced density matrix (2-RDM). <sup>41–44;46;50;138–141</sup> In this regard, DCT is related to approaches that are based on the variational optimization <sup>43;142–147</sup> or parametrization <sup>38–40</sup> of 2-RDM. On the other hand, DCT has a close relationship <sup>2;4</sup> with wavefunction-based electronic structure theories, such as linearized, unitary, and variational coupled cluster theory. <sup>9;148–155</sup> In contrast to variational 2-RDM theory <sup>47–49</sup> and traditional coupled cluster methods [e.g.,

CCSD and CCSD(T)], <sup>25;27</sup> DCT naturally combines size-extensivity and a Hermitian energy functional. In addition, the DCT electronic energy is fully relaxed with respect to all of its parameters, which greatly simplifies computation of the first-order molecular properties. <sup>156–159</sup> We have successfully applied DCT to a variety of chemical systems with different electronic structure effects (e.g., open-shell, symmetry-breaking, and multi-reference). <sup>3;4;137;160;161</sup> One limitation of the original DCT formulation is that it can only obtain information about the lowest-energy state of a particular symmetry (usually, the ground state). By combining DCT with linear response theory, we remove this limitation, providing access to many electronic states simultaneously.

We begin with a brief overview of linear response theory (section 3.3.1). We then discuss the derivation of linear-response theory for the ODC-12 method (LR-ODC-12, section 3.3.2). In section section 3.3.3, we derive equations for the linear-response orbital-optimized linearized coupled cluster theory with double excitations (LR-OLCCD), which we obtain by neglecting the non-linear terms in the LR-ODC-12 equations. We outline the computational details in section 3.4. In section section 3.5, we demonstrate that the LR-ODC-12 excitation energies are size-intensive (section 3.5.1), test the performance of LR-ODC-12 for the dissociation of  $H_2$  (section 3.5.2), and benchmark the accuracy of LR-ODC-12 for vertical excitation energies of small molecules (section 3.5.3).

### 3.3 Theory

## 3.3.1 Linear Response Theory

We now briefly review linear response theory in the quasi-energy formulation.  $^{162}$  The quasi-energy of a system perturbed by a time-dependent interaction  $\hat{V}f(t)$  is defined as

$$Q(t) = \langle \Psi(t) | \hat{H} + \hat{V}f(t) - i\frac{\partial}{\partial t} | \Psi(t) \rangle$$
 (3.1)

where  $\Psi(t)$  is the phase-isolated wavefunction, from which the usual Schrödinger wavefunction can be recovered as  $e^{-i\int_0^t dt' Q(t')} \Psi(t)$ . Assuming that the perturbation is periodic

$$f(t) = \sum_{\omega} f(\omega)e^{-i\omega t} \tag{3.2}$$

the time average of the quasi-energy over a period of oscillation, denoted as  $\{Q(t)\}$ , is variational with respect to the exact dynamic state. <sup>163</sup> The independent parameters  $\mathbf{u}(t)$  that define such a state can be written using a Fourier expansion

$$\mathbf{u}(t) = \sum_{n=0}^{\infty} \sum_{\omega_1 \cdots \omega_n} \mathbf{u}(\omega_1, \dots, \omega_n) e^{-i(\omega_1 + \dots + \omega_n)t}$$
(3.3)

where the outer sum runs over polynomial orders in f(t). The stationarity of the time-averaged quasi-energy then implies the following relationship <sup>164</sup>

$$0 = \frac{d}{df(\omega)} \frac{\partial \{Q(t)\}}{\partial \mathbf{u}^{\dagger}(\omega)} \bigg|_{f=0} =$$

$$\frac{\partial^{2} \{Q(t)\}}{\partial \mathbf{u}^{\dagger}(\omega) \partial \mathbf{u}(\omega)} \frac{\partial \mathbf{u}(\omega)}{\partial f(\omega)} \bigg|_{f=0} + \frac{\partial^{2} \{Q(t)\}}{\partial \mathbf{u}^{\dagger}(\omega) \partial f(\omega)} \bigg|_{f=0}$$
(3.4)

which constitutes a linear equation for the first-order response of the system to the perturbation. When the frequency  $\omega$  is in resonance with an excitation energy of the system, eq. (3.4) will result in an infinite first-order response  $\frac{\partial \mathbf{u}(\omega)}{\partial f(\omega)}$ . From eq. (3.4), we find that these poles occur when the Hessian matrix of the quasi-energy with respect to the wavefunction parameters  $\mathbf{u}(\omega)$  becomes singular. We can express this Hessian matrix in the form

$$\left. \frac{\partial^2 \{Q(t)\}}{\partial \mathbf{u}^{\dagger}(\omega) \partial \mathbf{u}(\omega)} \right|_{f=0} \equiv \mathbf{E} - \omega \,\mathbf{M}$$
(3.5)

where **E** is the Hessian of the time-averaged electronic energy  $\{\langle \Psi(t)|\hat{H}|\Psi(t)\rangle\}$  and  $\omega \mathbf{M}$  is the Hessian of the time-derivative overlap  $\{\langle \Psi(t)|i\dot{\Psi}(t)\rangle\}$ . The excitation energies of the system  $\omega_k$  can therefore be determined by solving the following generalized eigenvalue equation:

$$\mathbf{E}\mathbf{z}_k = \omega_k \mathbf{M}\mathbf{z}_k \tag{3.6}$$

where **M** serves as the metric matrix. Equation (3.6) allows the determination of excitation energies for an arbitrary parametrization of  $|\Psi(t)\rangle$ .

The generalized eigenvectors  $\mathbf{z}_k$  can be used to compute transition properties for excited states. In particular, in the exact linear response theory <sup>165</sup> the transition strength of the perturbing interaction,  $|\langle \Psi | \hat{V} | \Psi_k \rangle|^2$ , is equal to the complex residue of the following quantity at  $\omega \to \omega_k$ :

$$\langle\!\langle \hat{V}; \hat{V} \rangle\!\rangle_{\omega} \equiv \mathbf{v}'^{\dagger} \cdot \frac{\partial \mathbf{u}(\omega)}{\partial f(\omega)} \Big|_{f=0}$$
 (3.7)

This quantity is known as the linear response function and  $\mathbf{v}'$  is termed the property gradient vector, <sup>166</sup> which is defined as follows:

$$\mathbf{v}' \equiv \left. \frac{\partial^2 \{Q(t)\}}{\partial \mathbf{u}^{\dagger}(\omega) \partial f(\omega)} \right|_{f=0}$$
 (3.8)

Substituting eqs. (3.5) and (3.8) into eq. (3.4) and decomposing the quasi-energy Hessian as

$$\mathbf{E} - \omega \mathbf{M} = (\mathbf{Z}^{\dagger})^{-1} (\mathbf{Z}^{\dagger} \mathbf{M} \mathbf{Z}) (\Omega - \omega \mathbf{1}) (\mathbf{Z})^{-1}$$
(3.9)

where  $\Omega$  is a diagonal matrix of generalized eigenvalues and  $\mathbf{Z}$  is a matrix of generalized eigenvectors that simulataneously diagonalizes  $\mathbf{E}$  and  $\mathbf{M}$ , we obtain the general formula for the transition strengths:

$$\lim_{\omega \to \omega_k} (\omega - \omega_k) \langle \langle \hat{V}; \hat{V} \rangle \rangle_{\omega} = \frac{|\mathbf{z}_k^{\dagger} \mathbf{v}'|^2}{\mathbf{z}_k^{\dagger} \mathbf{M} \mathbf{z}_k}$$
(3.10)

In section 3.3.2, we will use the quasi-energy formalism to derive equations for the linear-response ODC-12 method (LR-ODC-12).

### 3.3.2 Linear-Response ODC-12

In the ODC-12 method, the electronic energy Hessian can be written in the following form

$$\mathbf{E} = \begin{pmatrix} \mathbf{A}_{11} & \mathbf{A}_{12} & \mathbf{B}_{11} & \mathbf{B}_{12} \\ \mathbf{A}_{21} & \mathbf{A}_{22} & \mathbf{B}_{21} & \mathbf{B}_{22} \\ \mathbf{B}_{11}^* & \mathbf{B}_{12}^* & \mathbf{A}_{11}^* & \mathbf{A}_{12}^* \\ \mathbf{B}_{21}^* & \mathbf{B}_{22}^* & \mathbf{A}_{21}^* & \mathbf{A}_{22}^* \end{pmatrix}$$
(3.11)

where the submatrices are defined in general as

$$\mathbf{A}_{nm} = \left. \frac{\partial^2 E}{\partial \mathbf{t}_n^{\dagger} \partial \mathbf{t}_m} \right|_{f=0}, \ \mathbf{B}_{nm} = \left. \frac{\partial^2 E}{\partial \mathbf{t}_n^{\dagger} \partial \mathbf{t}_m^*} \right|_{f=0}.$$
 (3.12)

These complex derivatives relate to the second derivatives of the electronic energy with respect to variations of the orbitals  $(\mathbf{A}_{11}, \mathbf{B}_{11})$  and cumulant parameters  $(\mathbf{A}_{22}, \mathbf{B}_{22})$ . Similarly, the mixed second derivatives couple variations in the orbitals and cumulant parameters  $(\mathbf{A}_{12}, \mathbf{B}_{12})$ . The metric matrix  $\mathbf{M}$  has a block-diagonal structure, as a consequence of the linear parametrization of the wavefunction in eq. (1.11):

$$\mathbf{M} = \begin{pmatrix} \mathbf{S}_{11} & \mathbf{0} & \mathbf{0} & \mathbf{0} \\ \mathbf{0} & \mathbf{1}_{2} & \mathbf{0} & \mathbf{0} \\ \mathbf{0} & \mathbf{0} & -\mathbf{S}_{11}^{*} & \mathbf{0} \\ \mathbf{0} & \mathbf{0} & \mathbf{0} & -\mathbf{1}_{2} \end{pmatrix}$$
(3.13)

where  $\mathbf{1}_2 = \langle \Phi | \mathbf{a}_2^{\dagger} \mathbf{a}_2 | \Phi \rangle$  is an identity matrix over the space of unique two-body excitations and the orbital metric is defined as follows:

$$\omega \mathbf{S}_{11} = \frac{\partial^2 \{ \langle \Psi(t) | i\dot{\Psi}(t) \rangle \}}{\partial \mathbf{t}_1^{\dagger}(\omega) \partial \mathbf{t}_1(\omega)} \bigg|_{f=0}$$
(3.14)

Equations for all blocks of  $\mathbf{E}$ ,  $\mathbf{M}$ , and the property gradient vector  $\mathbf{v}'$  are shown explicitly in the Supporting Information (section 3.B). We note that, due to the Hermitian nature of the DCT energy functional (1.7), the ODC-12 energy Hessian  $\mathbf{E}$  is always symmetric. As a result, in the absence of instabilities (i.e., as long as the Hessian is positive semi-definite), the LR-ODC-12 excitation energies are guaranteed to have real values.

To illustrate the derivation of the LR-ODC-12 energy Hessian, let us consider the diagonal two-body block of  $\mathbf{E}$ . Expressing the energy (1.7) using the cumulant expansion (1.8) and differentiating with respect to  $\mathbf{t}_2$ , we obtain:

$$\mathbf{A}_{22} = \frac{\partial^{2} E}{\partial \mathbf{t}_{2}^{\dagger} \partial \mathbf{t}_{2}} = f_{p}^{q} \frac{\partial^{2} \gamma_{q}^{p}}{\partial \mathbf{t}_{2}^{\dagger} \partial \mathbf{t}_{2}} + \overline{g}_{pr}^{qs} \frac{\partial \gamma_{q}^{p}}{\partial \mathbf{t}_{2}^{\dagger}} \frac{\partial \gamma_{s}^{r}}{\partial \mathbf{t}_{2}} + \frac{1}{4} \overline{g}_{pq}^{rs} \frac{\partial^{2} \lambda_{rs}^{pq}}{\partial \mathbf{t}_{2}^{\dagger} \partial \mathbf{t}_{2}}$$

$$(3.15)$$

where we have introduced the generalized Fock matrix  $f_p^q \equiv h_p^q + \overline{g}_{pr}^{qs} \gamma_s^r$ . The derivatives of the one-body density matrix can be expressed in terms of the derivatives of the density cumulant

$$\mathbf{A}_{22} = \mathcal{F}_{p}^{q} \frac{\partial^{2} \lambda_{qt}^{pt}}{\partial \mathbf{t}_{2}^{\dagger} \partial \mathbf{t}_{2}} + \mathcal{G}_{pr}^{qs} \frac{\partial \lambda_{qt}^{pt}}{\partial \mathbf{t}_{2}^{\dagger}} \frac{\partial \lambda_{su}^{ru}}{\partial \mathbf{t}_{2}} + \frac{1}{4} \overline{g}_{pq}^{rs} \frac{\partial^{2} \lambda_{rs}^{pq}}{\partial \mathbf{t}_{2}^{\dagger} \partial \mathbf{t}_{2}}$$

$$(3.16)$$

where the intermediates  $\mathcal{F}_p^q$  and  $\mathcal{G}_{pr}^{qs}$  can be computed using a transformation of the one- and two-electron integrals to the natural spin-orbital basis (see section 3.A for details). These cumulant derivatives are straightforward to evaluate from eqs. (1.10) and (1.11) using either algebraic or diagrammatic techniques.

Next, let us outline the derivation of the one-body metric. Substituting eq. (1.11) into eq. (3.14) gives

$$\omega \mathbf{S}_{11} = \frac{1}{2} \frac{\partial^2 \{ \langle \Psi | [i\dot{T}_1^{\dagger}(t), T_1(t)] | \Psi \rangle \}}{\partial \mathbf{t}_1^{\dagger}(\omega) \partial \mathbf{t}_1(\omega)} \bigg|_{f=0}$$

$$- \frac{1}{2} \frac{\partial^2 \{ \langle \Psi | [T_1^{\dagger}(t), i\dot{T}_1(t)] | \Psi \rangle \}}{\partial \mathbf{t}_1^{\dagger}(\omega) \partial \mathbf{t}_1(\omega)} \bigg|_{f=0}$$
(3.17)

where we have assumed that we are working in the variational orbital basis so that  $T_1(t)|_{f=0} = 0$ , and  $\Psi = \Psi(t)|_{f=0}$  denotes the ground state wavefunction. Using the Fourier expansion of the  $\mathbf{t}_1(t)$  parameters (eq. (3.3)), the gradients of the time derivatives can be evaluated as:

$$\frac{\partial i \dot{T}_{1}^{\dagger}(t)}{\partial \mathbf{t}_{1}^{\dagger}(\omega)} \bigg|_{t=0} = -\omega \mathbf{a}_{1}^{\dagger} e^{+i\omega t}$$
(3.18)

$$\left. \frac{\partial i \dot{T}_1(t)}{\partial \mathbf{t}_1(\omega)} \right|_{t=0} = +\omega \mathbf{a}_1 e^{-i\omega t}$$
(3.19)

Substituting eqs. (3.18) and (3.19) into eq. (3.17) and evaluating the gradients of  $T_1$  and  $T_1^{\dagger}$  similarly gives the final working equation for the one-body metric:

$$\omega(\mathbf{S}_{11})_{ia,jb} = \omega \langle \Psi | [a_a^i, a_j^b] | \Psi \rangle$$

$$= \omega(\delta_a^b \gamma_j^i - \delta_j^i \gamma_a^b)$$
(3.20)

### 3.3.3 Linear-Response OLCCD

The orbital-optimized linearized coupled cluster doubles method (OLCCD) can be considered as an approximation to the ODC-12 method where all of the non-linear

 $\mathbf{t}_2$  terms are neglected. Similarly, we can formulate the linear-response OLCCD method (LR-OLCCD) by linearizing the LR-ODC-12 equations. This simplifies the expressions for the electronic Hessian blocks that involve the second derivatives with respect to  $\mathbf{t}_2$ . For example, for the  $\mathbf{A}_{22}$  block, we obtain:

$$\mathbf{A}_{22} = (f_0)_i^j \frac{\partial^2 \lambda_{jr}^{ir}}{\partial \mathbf{t}_1^b \partial \mathbf{t}_2} - (f_0)_a^b \frac{\partial^2 \lambda_{br}^{ar}}{\partial \mathbf{t}_1^b \partial \mathbf{t}_2} + \frac{1}{4} \overline{g}_{pq}^{rs} \frac{\partial^2 \lambda_{rs}^{pq}}{\partial \mathbf{t}_2^b \partial \mathbf{t}_2}$$
(3.21)

where  $(f_0)_p^q = h_p^q + \overline{g}_{pi}^{qi}$  is the usual (mean-field) Fock operator. Comparing eq. (3.21) with eq. (3.16) from the LR-ODC-12 method, we observe that the former equation can be obtained from the latter by replacing the  $\mathcal{F}_p^q$  intermediates with the mean-field Fock matrix elements and ignoring the term that depends on  $\mathcal{G}_{pr}^{qs}$ . These simplifications arise from the fact that the  $\mathcal{F}_p^q$  and  $\mathcal{G}_{pr}^{qs}$  intermediates contain high-order  $\mathbf{t}_2$  contributions that are not included in the linearized LR-OLCCD formulation (see section 3.A and Ref. 2 for details). For the  $\mathbf{B}_{22}$  block, we find that all of the Hessian elements are zero. A complete set of working equations for LR-OLCCD is given in the Supporting Information (section 3.B).

## 3.4 Computational Details

The LR-ODC-12 and LR-OLCCD methods were implemented as a standalone Python program, which was interfaced with Psi4<sup>167</sup> and PyscF<sup>168</sup> to obtain the one- and two-electron integrals. Our implementation of the energy Hessian was validated by computing the static response function for a dipole perturbation (i.e.,

the dipole polarizability):

$$\langle\langle \hat{V}; \hat{V} \rangle\rangle_0 = -\mathbf{v}'^{\dagger} \mathbf{E}^{-1} \mathbf{v}'$$
 (3.22)

This quantity can be evaluated numerically as a derivative of the ground state energy

$$\langle\langle \hat{V}; \hat{V} \rangle\rangle_0 = \frac{d^2E}{df^2} \bigg|_{f=0}$$
 (3.23)

by perturbing the one-electron integrals  $h_p^q \leftarrow h_p^q + f v_p^q$  with the integrals of the perturbing dipole operator,  $v_p^q$ , and solving the ODC-12 or OLCCD equations for different values of f. In this way, we have matched our response calculation to  $10^{-9}$  a.u. for the electric dipole polarizability along the  $C_2$  symmetry axis of water.

We used Q-CHEM  $4.4^{169}$  to obtain results from equation-of-motion coupled cluster theory with single and double excitations (EOM-CCSD) and EOM-CCSD with triple excitations in the EOM part [EOM-CC(2,3)]. The MRCC program <sup>170</sup> was used to obtain results for equation-of-motion coupled cluster theory with up to full triple excitations (EOM-CCSDT). All electrons were correlated in all computations. We used tight convergence parameters in all ground-state ( $10^{-8}$   $E_h$ ) and excited-state computations ( $10^{-5}$   $E_h$ ). In sections 3.5.2 and 3.5.3, the augmented aug-cc-pVTZ and d-aug-cc-pVTZ basis sets of Dunning and co-workers were employed. <sup>60</sup> To compute vertical excitation energies in section 3.5.3, geometries of molecules were optimized using ODC-12 (for LR-ODC-12), OLCCD (for LR-OLCCD), or CCSD [for EOM-CCSD, EOM-CC(2,3), and EOM-CCSDT].

#### 3.5 Results and Discussion

## 3.5.1 Size-Intensivity of the LR-ODC-12 Energies

Table 3.1: Ground-state energies (in  $E_{\rm h}$ ) and vertical excitation energies (in eV) for the four lowest-energy excited states of the CO molecule and noninteracting systems of CO with Ne atoms (CO + nNe, n=1,2,3) computed using the ODC-12 and LR-ODC-12 methods (cc-pVDZ basis set). The noninteracting systems were separated from each other by 10000 Å and the C-O bond distance was set to 1.12547 Å. Results demonstrate size-intensivity of the LR-ODC-12 excitation energies.

	CO	CO + Ne	CO + 2Ne	CO + 3Ne
$X^1\Sigma_q^+$	-113.051282	-241.730913	-370.410543	-499.090174
$^3\Pi$	6.48596	6.48596	6.48596	6.48596
$^3\Sigma^+$	8.41225	8.41225	8.41225	8.41225
$^{1}\Pi$	8.90866	8.90866	8.90866	8.90866
$\Delta$	9.33189	9.33189	9.33189	9.33189

All DCT methods are by construction size-extensive, meaning that their electronic energies scale linearly with the number of electrons. In this section, we demonstrate that the LR-ODC-12 excitation energies are size-intensive, i.e. they satisfy the following property:  $E(A^* + B) = E(A^*) + E(B)$ , where A and B are two noninteracting fragments in their corresponding ground states and  $A^*$  is the fragment A in an excited state. Table 3.1 shows the ODC-12 ground-state energies and the LR-ODC-12 excitation energies for the CO molecule and noninteracting systems composed of CO and the neon atoms separated by 10000 Å (CO + nNe, n = 1, 2, 3). The scaling of the ODC-12 energies for the ground  $X^1\Sigma_g^+$  electronic state is perfectly linear up to  $10^{-8}$   $E_h$ , which is the convergence parameter used in our ODC-12 computations. Upon the addition of the neon atoms, the excita-

tion energies of the CO molecule remain constant up to the convergence threshold set in LR-ODC-12 ( $10^{-6}$  eV). These results provide numerical evidence that the LR-ODC-12 excitation energies are size-intensive.

### 3.5.2 H<sub>2</sub> Dissociation

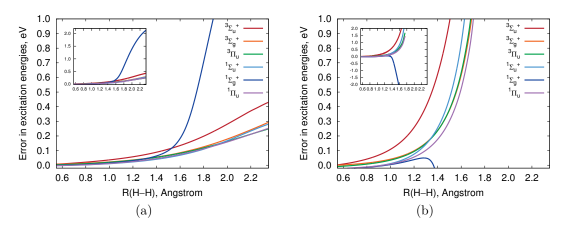


Figure 3.1: Errors in vertical excitation energies (eV) for six lowest-lying electronic states of  $\rm H_2$  computed using LR-ODC-12 (3.1a) and LR-OLCCD (3.1b) as a function of the H–H bond length, relative to full configuration interaction. All methods employed the d-aug-cc-pvtz basis set. In each figure, the inset shows the same plot for a larger range of errors.

One of the desirable properties of an electronic structure method is exactness for two-electron systems. While the ODC-12 method is not exact for two-electron systems, it has been shown to provide a very good description of the ground-state  $H_2$  dissociation curve, with errors of  $\sim 1$  kcal mol<sup>-1</sup> with respect to full configuration interaction (FCI) near the dissociation limit.<sup>4</sup> Here, we investigate the performance of LR-ODC-12 for the excited states of  $H_2$ . fig. 3.1a shows the errors in vertical excitation energies for six lowest-lying electronic states as a function of the H–H distance, relative to FCI. The FCI energies were computed using

the EOM-CCSD method, which is exact for two-electron systems. At the equilibrium geometry ( $r_{\rm e}=0.742~{\rm \AA}$ ) the errors in excitation energies for all states do not exceed 0.02 eV. Between 0.6 and 1.45 Å ( $r\approx 2r_{\rm e}$ ), the LR-ODC-12 excitation energies remain to be in a good agreement with FCI, with errors less than 0.1 eV for all states. In this range, the largest error is observed for the  $^3\Sigma_u^+$  state. For  $r\geq 1.5~{\rm \AA}$ , the error in the  $^1\Sigma_g^+$  excited state energy rapidly increases from 0.10 eV (at 1.5 Å) to 2.13 eV (at 2.35 Å), while for other states the errors increase much more slowly. Analysis of the FCI wavefunction for the  $^1\Sigma_g^+$  state shows a significant contribution from the  $(1\sigma_g)^2 \to (1\sigma_u)^2$  double excitation already at  $r=1.55~{\rm \AA}$ . This contribution becomes dominant for  $r\geq 1.75~{\rm \AA}$ . Thus, the large LR-ODC-12 errors observed for the  $^1\Sigma_g^+$  state are likely due to the double-excitation character of this electronic state at long H–H bond distances. The second largest error near the dissociation is observed for the  $^3\Sigma_u^+$  state (0.43 eV). For other electronic states, smaller errors of  $\sim 0.25$  eV are observed near the dissociation.

We now compare the performance of LR-ODC-12 to that of the LR-OLCCD method, which can be considered as an approximation to LR-ODC-12 where all of the non-linear terms are removed in the description of electron correlation. Figure 3.1b shows the errors in the LR-OLCCD vertical excitation energies as a function of the H–H bond length. Although near the equilibrium geometry the performance of LR-OLCCD and LR-ODC-12 is similar, the LR-OLCCD errors increase much faster with increasing H–H distance compared to LR-ODC-12. At r = 1.3 Å, the LR-OLCCD error for the  $^3\Sigma_u^+$  state (0.4 eV) is almost six times larger than the corresponding error from LR-ODC-12 (0.07 eV). For  $r \geq 1.35$  Å, the LR-

OLCCD errors for all excitation energies show very steep increase in magnitude, ranging from 1.5 to 4.7 eV already at r=1.75 Å. We were unable to converge the LR-OLCCD equations for  $r\geq 1.80$  Å. Overall, our results demonstrate that the non-linear terms in LR-ODC-12 significantly improve the description of excited states at long H-H distances when the electron correlation effects become particularly strong.

### 3.5.3 Benchmark: Small Molecules

Here we investigate the performance of our method for vertical excitation energies in several small molecules:  $N_2$ , CO, HCN, HNC,  $H_2C_2$ , and  $H_2CO$ . The reference values have been computed with EOM-CCSDT and we compare all-electron calculations with the same aug-cc-pVTZ basis in order to exclude basis set incompleteness error from the comparison. As an aggregate measure of the tendency for each method to deviate from the reference values, we calculate their mean absolute errors ( $\Delta_{MAE}$ ). As a measure of the degree to which the errors are systematically clustered above or below the reference values, we look at their standard deviation from the average signed error ( $\Delta_{SD}$ ).

For the singlets, LR-ODC-12 has errors above 0.15 eV for three states, with a maximum error of 0.20 eV. All three of these occur for  ${}^{1}\Pi$  states of linear molecules, which are dominated by  $\sigma \to \pi^{*}$  excitations. EOM-CCSD has eleven errors greater than 0.15 eV, with a maximum error of 0.26 eV and no predictable pattern – the largest three errors occur for three qualitatively unrelated states:  ${}^{1}\Delta$ ,  ${}^{1}\Sigma^{-}$ , and  ${}^{1}\Sigma^{+}$ . Interestingly, the EOM-CCSD errors are relatively tightly clustered around

Table 3.2: Errors in vertical excitation energies (eV) for singlet states computed using LR-OLCCD, LR-ODC-12, and EOM-CCSD, relative to EOM-CCSDT (aug-cc-pVTZ basis set). All electrons were correlated in all computations. Also shown are mean absolute errors ( $\Delta_{\rm MAE}$ ) and standard deviations ( $\Delta_{\rm SD}$ ) computed for each method.

		$\Delta \text{EOM-CCSD}$	$\Delta$ LR-OLCCD	$\Delta$ LR-ODC-12	EOM-CCSDT
$\overline{\mathrm{N}_{2}}$	$^{1}\Pi_{g}$	0.18	0.08	0.20	9.29
	$^{1}\Sigma_{u}^{-}$	0.23	0.15	0.09	9.84
	$^1\Delta_u$	0.26	0.14	0.10	10.26
CO	$^{1}\Pi$	0.16	0.09	0.17	8.46
	$^{1}\Sigma^{-}$	0.19	-0.10	-0.01	9.89
	$^{1}\Delta$	0.19	-0.22	-0.05	10.03
HCN	$^1\Sigma^-$	0.16	0.05	0.00	8.25
	$^{1}\Delta$	0.17	0.04	0.01	8.61
	$^{1}\Pi$	0.17	0.05	0.20	9.12
HNC	$^{1}\Pi$	0.15	-0.01	0.10	8.13
	$^{1}\Sigma^{+}$	0.24	0.05	0.12	8.46
	$^1\Sigma^-$	0.15	-0.09	0.04	8.67
	$^{1}\Delta$	0.15	-0.18	-0.03	8.84
$\mathrm{H_2C_2}$	$^{1}\Sigma_{u}^{-}$	0.12	0.06	0.02	7.11
	$^{1}\Delta_{u}$	0.10	0.07	0.03	7.45
${\rm H_2CO}$	$^{1}\mathrm{A}_{2}$	0.10	-0.07	0.02	3.95
$\Delta_{ m MAE}$		0.09	0.08	0.17	
$\Delta_{\mathrm{SD}}$		0.11	0.08	0.05	

Table 3.3: Errors in vertical excitation energies (eV) for triplet states computed using LR-OLCCD, LR-ODC-12, and EOM-CCSD, relative to EOM-CCSDT (aug-cc-pVTZ basis set). All electrons were correlated in all computations. Also shown are mean absolute errors ( $\Delta_{MAE}$ ) and standard deviations ( $\Delta_{SD}$ ) computed for each method.

		$\Delta  ext{EOM-CCSD}$	$\Delta$ LR-OLCCD	$\Delta$ LR-ODC-12	EOM-CCSDT <sup>a</sup>
$\overline{\mathrm{N}_{2}}$	$^3\Sigma_u^+$	0.11	0.04	-0.02	7.63
	$^3\Pi_g$	0.15	0.06	0.11	8.00
	$^3\Delta_u$	0.17	0.08	0.03	8.82
	$^3\Sigma_u^-$	0.28	0.03	0.01	9.63
	$^3\Pi_u$	0.14	-0.01	0.10	11.18
CO	$^3\Pi$	0.12	0.06	0.08	6.27
	$^3\Sigma^+$	0.05	-0.03	-0.03	8.38
	$^3\Delta$	0.11	-0.07	-0.03	9.21
	$3\Sigma^-$	0.19	-0.18	-0.06	9.72
HCN	$3\Sigma^+$	0.05	-0.04	-0.10	6.40
	$^3\Delta$	0.13	-0.02	-0.06	7.40
	$^3\Pi$	0.10	0.08	0.06	8.01
	$3\Sigma$	0.16	-0.10	-0.05	8.15
HNC	$^3\Pi$	0.09	0.00	0.03	6.06
	$3\Sigma^+$	0.04	-0.09	-0.11	7.20
	$^3\Delta$	0.10	-0.14	-0.11	8.02
	$3\Sigma^+$	0.22	-0.05	0.04	8.38
	$3\Sigma^-$	0.15	-0.02	0.11	8.56
$\mathrm{H_2C_2}$	$^{3}\Sigma_{u}^{+}$ $^{3}\Delta_{u}$	0.01	-0.02	-0.08	5.52
	$^3\Delta_u$	0.08	-0.02	-0.05	6.41
	$^3\Sigma_u^-$	0.10	-0.03	-0.05	7.10
$\mathrm{H_{2}CO}$	$^3A_2$	0.04	-0.02	0.01	3.56
	$^3A_1$	0.02	-0.06	-0.14	6.06
$\Delta_{ m MAE}$		0.17	0.09	0.08	
$\Delta_{ m SD}$		0.05	0.11	0.08	

<sup>&</sup>lt;sup>a</sup> For CO, HCN, HNC, and  $\mathrm{H_2C_2}$ , the  $^3\Sigma^-$  ( $^3\Sigma_u^-$ ) excitation energies were obtained from EOM-CC(2,3), which energies were shifted to reproduce the EOM-CCSDT energy for the  $^1\Sigma^-$  ( $^1\Sigma_u^-$ ) state.

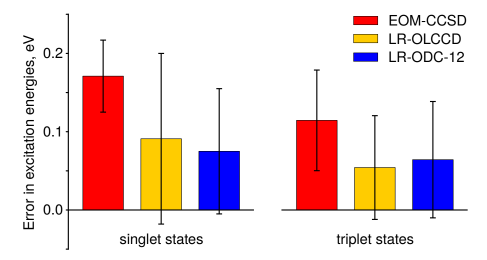


Figure 3.2: Mean absolute deviations ( $\Delta_{MAE}$ ) and standard deviations from the mean signed error ( $\Delta_{SD}$ ) in vertical excitation energies for the states in tables 3.2 and 3.3 for OLCCD, LR-ODC-12, and EOM-CCSD relative to EOM-CCSDT (augcc-pVTZ basis set). The height of each colored box represents the  $\Delta_{MAE}$  value of the method and the height of the black bar above each box represents its  $\Delta_{SD}$  value.

their mean error of 0.17 eV, with a standard deviation of 0.05 eV compared to 0.08 eV for LR-OCD-12. For the triplets, LR-ODC-12 has no errors at all above 0.15 eV, whereas EOM-CCSD has six, with a maximum error of 0.28 eV for the  $^3\Sigma_u^-$  state of N<sub>2</sub>. Here the EOM-CCSD description of  $^3\Sigma^-$  states is consistently poor, but some of the larger errors occur for  $^3\Sigma^+$  and  $^3\Delta$  states as well. In aggregate, we see that LR-ODC-12 cuts the mean absolute errors of EOM-CCSD roughly in half for both singlet and triplet states. Notably, whereas the EOM-CCSD errors are exclusively positive, the LR-ODC-12 errors are more centered on 0 eV, suggesting a more balanced treatment of the ground and excited states.

Comparing LR-ODC-12 to LR-OLCCD we see that most of the errors are within 0.1 eV of each other, suggesting that the non-linear terms are relatively

unimportant for many of these states. Significant differences occur for the  $^{1}\Delta$  states of CO and HCN, where LR-OLCCD shows some deterioration, and for the  $^{1}\Pi$  state of HCN, where the linearization leads to a significant error cancellation.

# 3.A Derivatives of the One-Body Density Matrix in Density Cumulant Theory

Repeated differentiation of the one-body n-representability condition in DCT (eq. (1.9)) gives the following formulas for the first and second derivatives of the cumulant partial trace:

$$\frac{\partial \lambda_{qr}^{pr}}{\partial y} = \gamma_s^p \frac{\partial \gamma_q^s}{\partial y} + \frac{\partial \gamma_s^p}{\partial y} \gamma_q^s - \frac{\partial \gamma_q^p}{\partial y}$$
(3.24)

$$\frac{\partial^2 \lambda_{qr}^{pr}}{\partial x \partial y} = \gamma_s^p \frac{\partial \gamma_q^s}{\partial x \partial y} + \frac{\partial \gamma_s^p}{\partial x \partial y} \gamma_q^s - \frac{\partial \gamma_q^p}{\partial x \partial y} + \frac{\partial \gamma_s^p}{\partial x} \frac{\partial \gamma_q^s}{\partial y} + \frac{\partial \gamma_q^s}{\partial x} \frac{\partial \gamma_s^p}{\partial y}$$
(3.25)

Transforming to the natural spin-orbital basis (NSO, denoted by prime indices) where the one-body density matrix is diagonal, the first and second derivatives of the one-body density matrix can be determined from the cumulant derivatives as follows:

$$\frac{\partial \gamma_{q'}^{p'}}{\partial u} = \theta_{p'q'} \frac{\partial \lambda_{q'r}^{p'r}}{\partial u} \tag{3.26}$$

$$\frac{\partial^{2} \gamma_{q'}^{p'}}{\partial x \partial y} = \theta_{p'q'} \frac{\partial^{2} \lambda_{q'r}^{p'r}}{\partial x \partial y} - \delta_{r'}^{s'} \theta_{p'q'} \theta_{p's'} \theta_{q'r'} \frac{\partial \lambda_{s't}^{p't}}{\partial x} \frac{\partial \lambda_{r'u}^{r'u}}{\partial y} - \delta_{r'}^{s'} \theta_{p'q'} \theta_{p's'} \theta_{q'r'} \frac{\partial \lambda_{q'u}^{r'u}}{\partial x} \frac{\partial \lambda_{s't}^{p't}}{\partial y}$$

$$-\delta_{r'}^{s'} \theta_{p'q'} \theta_{p's'} \theta_{q'r'} \frac{\partial \lambda_{q'u}^{r'u}}{\partial x} \frac{\partial \lambda_{s't}^{p't}}{\partial y}$$
(3.27)

Here, we have defined the following matrix:

$$\theta_{p'q'} \equiv \begin{cases} (\gamma_{p'} + \gamma_{q'} - 1)^{-1} & \text{if } p', q' \in \text{occ or vir} \\ 0 & \text{otherwise} \end{cases}$$
 (3.28)

where  $\gamma_{p'}$  denotes an eigenvalue of the one-body density matrix (i.e., an occupation number). The natural spin-orbital p' is occupied if  $\gamma_{p'} > 0.5$ .

Equations (3.26) and (3.27) can be used to derive expression for the two-body energy Hessian in eq. (3.15). Simplifying the resulting equations allows us to determine the intermediates defined in eq. (3.16). In the NSO basis, these intermediates are given by

$$\mathcal{F}_{p'}^{q'} \equiv \theta_{p'q'} f_{p'}^{q'} \tag{3.29}$$

$$\mathcal{G}_{p'r'}^{q's'} \equiv \theta_{p'q'}\theta_{r's'}(\overline{g}_{p'r'}^{q's'} - \mathcal{F}_{p'}^{s'}\delta_{r'}^{q'} - \mathcal{F}_{r'}^{q'}\delta_{p'}^{s'})$$
(3.30)

Although these intermediates are computed in the NSO basis, they must be backtransformed to the original spin-orbital basis using the eigenvectors of the oneparticle density matrix before being used in eq. (3.16) (see Ref. 2 for more details).

### 3.B Supporting Information

Here, we present the working equations for the LR-ODC-12 and LR-OLCCD methods. We refer the reader to the main article for definitions and notation, and note that prime indices refer to the natural spin-orbital basis (NSO) where the one-particle density matrix is diagonal,  $\gamma_{q'}^{p'} = \delta_{q'}^{p'} \gamma_{q'}$ . Quantities computed in the NSO

basis can be back-transformed to the original basis using the eigenvectors of the one-particle density matrix.

For the property gradient vectors, we define the block structure as follows

$$\mathbf{v}^{\prime\dagger} \equiv (\mathbf{p}_1 \ \mathbf{p}_2 \ \mathbf{p}_1^* \ \mathbf{p}_2^*)^{\dagger} \qquad \mathbf{p}_m \equiv \frac{\partial \langle \Psi | \hat{V} | \Psi \rangle}{\partial \mathbf{t}_m^{\dagger}}$$
(3.31)

where  $\hat{V}$  can be any one-particle operator, defined through its spin-orbital integrals,  $v_p^q = \langle \psi_p | \hat{v} | \psi_q \rangle$ , as follows.

$$\hat{V} = v_p^q a_q^p \tag{3.32}$$

A typical example would be an electric dipole operator.

## 3.B.1 Working Equations for LR-ODC-12

## **Density Matrices**

$$\gamma_j^i = \frac{1}{2} (\mathbf{1} + \sqrt{\mathbf{1} + 4\mathbf{d}})_j^i \qquad (\mathbf{d})_j^i \equiv -\frac{1}{2} t_{cd}^{ik} t_{jk}^{cd*}$$
 (3.33)

$$\gamma_a^b = \frac{1}{2} (\mathbf{1} - \sqrt{\mathbf{1} + 4\mathbf{d}})_a^b \qquad (\mathbf{d})_a^b \equiv -\frac{1}{2} t_{ac}^{kl} t_{kl}^{bc*}$$
 (3.34)

$$\gamma_{kl}^{ij} = \frac{1}{2} t_{cd}^{ij} t_{kl}^{cd*} + P_{(k/l)} \gamma_k^i \gamma_l^j \tag{3.35}$$

$$\gamma_{ab}^{cd} = \frac{1}{2} t_{ab}^{kl} t_{kl}^{cd*} + P^{(c/d)} \gamma_a^c \gamma_b^d$$
 (3.36)

$$\gamma_{ja}^{ib} = -t_{ac}^{ik}t_{jk}^{bc*} + \gamma_j^i\gamma_a^b \tag{3.37}$$

$$\gamma_{ab}^{ij} = t_{ab}^{ij} \tag{3.38}$$

## Blocks of the Energy Hessian

$$(\mathbf{A}_{11})_{ia,jb} = h_j^i \gamma_a^b + h_a^b \gamma_j^i - (\bar{\mathbf{F}})_j^i \delta_a^b - (\bar{\mathbf{F}})_a^b \delta_j^i + \overline{g}_{nj}^{mi} \gamma_{ma}^{nb} + \overline{g}_{ma}^{nb} \gamma_{nj}^{mi} + \overline{g}_{jf}^{ie} \gamma_{ae}^{bf} + \overline{g}_{ae}^{bf} \gamma_{jf}^{ie} + \overline{g}_{me}^{ib} \gamma_{ja}^{me} + \overline{g}_{ja}^{me} \gamma_{me}^{ib}$$

$$(3.39)$$

$$\bar{\mathbf{F}} \equiv \frac{1}{2} (\mathbf{F} + \mathbf{F}^{\dagger}) \qquad (\mathbf{F})_{p}^{q} \equiv h_{p}^{t} \gamma_{t}^{q} + \frac{1}{2} \overline{g}_{pt}^{uv} \gamma_{uv}^{qt}$$
(3.40)

$$(\mathbf{B}_{11})_{ia,jb} = \overline{g}_{be}^{im} \gamma_{ma}^{je} + \overline{g}_{ma}^{je} \gamma_{be}^{im} + \overline{g}_{mb}^{ie} \gamma_{ae}^{jm} + \overline{g}_{ae}^{jm} \gamma_{mb}^{ie} + \frac{1}{2} \overline{g}_{mn}^{ij} \gamma_{ab}^{mn} + \frac{1}{2} \overline{g}_{ab}^{mn} \gamma_{mn}^{ij} + \frac{1}{2} \overline{g}_{ab}^{ef} \gamma_{ef}^{ij}$$

$$(3.41)$$

$$\begin{aligned} (\mathbf{A}_{22})_{ijab,klcd} &= -P_{(a/b|k/l)}^{(c/d)} \mathcal{F}_{a}^{c} \delta_{b}^{d} \delta_{k}^{i} \delta_{l}^{j} - P_{(k/l)}^{(i/j|c/d)} \mathcal{F}_{k}^{i} \delta_{l}^{j} \delta_{a}^{c} \delta_{b}^{d} + P_{(k/l)} \overline{g}_{ab}^{cd} \delta_{k}^{i} \delta_{l}^{j} + P^{(c/d)} \overline{g}_{kl}^{ij} \delta_{a}^{c} \delta_{b}^{d} \\ &- P_{(a/b|k/l)}^{(i/j|c/d)} \overline{g}_{la}^{jc} \delta_{b}^{d} \delta_{k}^{i} + P_{(a/b)}^{(c/d)} \mathcal{G}_{af}^{ec} t_{eb}^{ij} t_{kl}^{fd*} + P_{(a/b|k/l)} \mathcal{G}_{ka}^{me} t_{eb}^{ij} t_{ml}^{cd*} \\ &+ P^{(i/j|c/d)} \mathcal{G}_{me}^{ic} t_{ab}^{mj} t_{kl}^{ed*} + P_{(k/l)}^{(i/j)} \mathcal{G}_{mk}^{in} t_{ab}^{mj} t_{nl}^{cd*} \end{aligned}$$

$$(3.42)$$

$$\mathcal{F}_{p'}^{q'} \equiv \theta_{p'q'} f_{p'}^{q'} \qquad \mathcal{G}_{p'r'}^{q's'} \equiv \theta_{p'q'} \theta_{r's'} (\overline{g}_{p'r'}^{q's'} - \mathcal{F}_{p'}^{s'} \delta_{r'}^{q'} - \mathcal{F}_{r'}^{q'} \delta_{p'}^{s'})$$
(3.43)

$$\theta_{p'q'} \equiv \begin{cases} (\gamma_{p'} + \gamma_{q'} - 1)^{-1} & \text{if } p, q \in \text{occ or vir} \\ 0 & \text{otherwise} \end{cases}$$
 (3.44)

$$(\mathbf{A}_{12})_{ia,klcd} = P_{(k/l)} \overline{g}_{la}^{cd} \delta_{k}^{i} + P^{(c/d)} \overline{g}_{kl}^{id} \delta_{a}^{c} + P_{(k/l)} (\mathcal{I}_{a}^{i})_{k}^{m} t_{ml}^{cd*} + P^{(c/d)} (\mathcal{I}_{a}^{i})_{e}^{c} t_{kl}^{ed*}$$

$$+ P_{(k/l)}^{(c/d)} \overline{g}_{ac}^{mc} t_{ml}^{ed*} \delta_{k}^{i} + P_{(k/l)}^{(c/d)} \overline{g}_{ke}^{im} t_{ml}^{ed*} \delta_{a}^{c} + \frac{1}{2} P_{(k/l)} \overline{g}_{la}^{mn} t_{mn}^{cd*} \delta_{k}^{i}$$

$$+ \frac{1}{2} P^{(c/d)} \overline{g}_{ef}^{id} t_{kl}^{ef*} \delta_{a}^{c}$$

$$(3.46)$$

$$(\mathbf{B}_{12})_{ia,klcd} = P^{(k/l)} (\mathcal{I}_{a}^{i})_{m}^{k} t_{cd}^{ml} + P_{(c/d)} (\mathcal{I}_{a}^{i})_{c}^{e} t_{ed}^{kl} + P_{(c/d)}^{(k/l)} \overline{g}_{ad}^{le} t_{ce}^{ki} + P_{(c/d)}^{(k/l)} \overline{g}_{md}^{il} t_{ca}^{km} + \overline{g}_{ma}^{kl} t_{cd}^{im} + \overline{g}_{cd}^{ie} t_{ae}^{kl}$$

$$(3.47)$$

$$(\mathcal{I}_a^i)_{k'}^{l'} \equiv \theta_{k'l'}(I_a^i)_{k'}^{l'} \qquad (I_a^i)_k^l \equiv +f_a^l \delta_k^i - \overline{g}_{ka}^{ml} \gamma_m^i + \overline{g}_{ke}^{il} \gamma_a^e$$
 (3.48)

$$(\mathcal{I}_a^i)_{c'}^{d'} \equiv \theta_{c'd'}(I_a^i)_{c'}^{d'} \qquad (I_a^i)_c^d \equiv -f_c^i \delta_a^d + \overline{g}_{ac}^{md} \gamma_m^i - \overline{g}_{ec}^{id} \gamma_a^e$$
 (3.49)

## Blocks of the Metric Matrix

$$(\mathbf{S}_{11})_{ia,jb} = (\delta_a^b \gamma_j^i - \delta_j^i \gamma_a^b) \tag{3.50}$$

# Blocks of the Property Gradient Vector

$$(\mathbf{p}_1)_{ia} = v_a^k \gamma_k^i - v_c^i \gamma_a^c \tag{3.51}$$

$$(\mathbf{p}_2)_{ijab} = -P_{(a/b)} \mathcal{V}_a^e t_{eb}^{ij} - P^{(i/j)} \mathcal{V}_m^i t_{ab}^{mj}$$
(3.52)

$$\mathcal{V}_{p'}^{q'} \equiv \theta_{p'q'} v_{p'}^{q'} \tag{3.53}$$

## 3.B.2 Working Equations for LR-OLCCD

# **Density Matrices**

$$\gamma_{j}^{i} = \delta_{j}^{i} - \frac{1}{2} t_{cd}^{ik} t_{jk}^{cd*} \tag{3.54}$$

$$\gamma_a^b = \frac{1}{2} t_{ac}^{kl} t_{kl}^{bc*} \tag{3.55}$$

$$\gamma_{kl}^{ij} = \frac{1}{2} t_{cd}^{ij} t_{kl}^{cd*} + P_{(k/l)} \delta_k^i \delta_l^j + \frac{1}{2} P_{(k/l)}^{(i/j)} \delta_k^i t_{cd}^{ij} t_{lm}^{cd*}$$
(3.56)

$$\gamma_{ab}^{cd} = \frac{1}{2} t_{ab}^{kl} t_{kl}^{cd*} \tag{3.57}$$

$$\gamma_{ja}^{ib} = -t_{ac}^{ik}t_{jk}^{bc*} + \frac{1}{2}\delta_{j}^{i}t_{ac}^{kl}t_{kl}^{bc*}$$
(3.58)

$$\gamma_{ab}^{ij} = t_{ab}^{ij} \tag{3.59}$$

## Blocks of the Energy Hessian

$$(\mathbf{A}_{11})_{ia,jb} = \text{Equation } (3.39) \text{ with OLCCD density matrices.}$$
 (3.60)

$$(\mathbf{B}_{11})_{ia,jb} = \text{Equation (3.41)} \text{ with OLCCD density matrices.}$$
 (3.61)

$$(\mathbf{A}_{22})_{ijab,klcd} = P_{(a/b|k/l)}^{(c/d)}(\mathbf{f}_{0})_{a}^{c}\delta_{b}^{d}\delta_{k}^{i}\delta_{l}^{j} - P_{(k/l)}^{(i/j|c/d)}(\mathbf{f}_{0})_{k}^{i}\delta_{l}^{j}\delta_{a}^{c}\delta_{b}^{d} + P_{(k/l)}\overline{g}_{ab}^{cd}\delta_{k}^{i}\delta_{l}^{j} + P^{(c/d)}\overline{g}_{kl}^{ij}\delta_{a}^{c}\delta_{b}^{d} - P_{(a/b|k/l)}^{(i/j|c/d)}\overline{g}_{la}^{jc}\delta_{b}^{d}\delta_{k}^{i}$$

$$(3.62)$$

$$(\mathbf{f}_0)_p^q = h_p^q + \overline{g}_{pi}^{qi} \tag{3.63}$$

$$(\mathbf{B}_{22})_{ijab,klcd} = 0 \tag{3.64}$$

$$(\mathbf{A}_{12})_{ia,klcd} = P_{(k/l)} \overline{g}_{la}^{cd} \delta_{k}^{i} + P^{(c/d)} \overline{g}_{kl}^{id} \delta_{a}^{c} + P_{(k/l)} (I_{a}^{i})_{k}^{m} t_{ml}^{cd*} - P^{(c/d)} (I_{a}^{i})_{e}^{c} t_{kl}^{ed*}$$

$$+ P_{(k/l)}^{(c/d)} \overline{g}_{ae}^{mc} t_{ml}^{ed*} \delta_{k}^{i} + P_{(k/l)}^{(c/d)} \overline{g}_{ke}^{im} t_{ml}^{ed*} \delta_{a}^{c} + \frac{1}{2} P_{(k/l)} \overline{g}_{la}^{mn} t_{mn}^{cd*} \delta_{k}^{i}$$

$$+ \frac{1}{2} P^{(c/d)} \overline{g}_{ef}^{id} t_{kl}^{ef*} \delta_{a}^{c}$$

$$(3.65)$$

$$(I_a^i)_k^l \equiv +f_a^l \delta_k^i - \overline{g}_{ka}^{il} \qquad (I_a^i)_c^d \equiv -f_c^i \delta_a^d + \overline{g}_{ac}^{id}$$
(3.66)

$$(\mathbf{B}_{12})_{ia,klcd} = P^{(k/l)} (I_a^i)_m^k t_{cd}^{ml} - P_{(c/d)} (I_a^i)_c^e t_{ed}^{kl} + P_{(c/d)}^{(k/l)} \overline{g}_{ad}^{le} t_{ce}^{ki} + P_{(c/d)}^{(k/l)} \overline{g}_{md}^{il} t_{ca}^{km} + \overline{g}_{ma}^{kl} t_{cd}^{im} + \overline{g}_{cd}^{ie} t_{ae}^{kl}$$

$$(3.67)$$

## Blocks of the Metric Matrix

$$(\mathbf{S}_{11})_{ia,jb} = \text{Equation (3.50)} \text{ with OLCCD density matrices.}$$
 (3.68)

## Blocks of the Property Gradient Vector

$$(\mathbf{p}_1)_{ia} = \text{Equation (3.51)} \text{ with OLCCD density matrices.}$$
 (3.69)

$$(\mathbf{p}_2)_{ijab} = P_{(a/b)} v_a^e t_{eb}^{ij} - P^{(i/j)} v_m^i t_{ab}^{mj}$$
(3.70)

# Chapter 4

Linear-Response Density Cumulant Theory for Excited States: Better Algorithms, Bigger Systems

#### 4.1 Abstract

We have recently proposed the LR-ODC-12 model for electronic excited states, a linear response extension of density cumulant theory with linearized double excitations and variationally optimized orbitals. Our benchmark calculations have shown that LR-ODC-12 is consistently more accurate than its popular equation-of-motion coupled-cluster counterpart, EOM-CCSD. Here we propose improved algorithms for solving the LR-ODC-12 equations in order to extend its range of application to larger systems. We use the new implementation to study the valence spectra of ethylene, butadiene, and hexatriene.

#### 4.2 Introduction

The success of ground state coupled-cluster theory comes not from CCSD itself, which has a relatively poor cost-accuracy ratio, but from the non-iterative (T) correction to CCSD, which affords truly exceptional accuracy relative to its cost.<sup>57</sup> Coupled-cluster theory has also been extremely popular in the study of electronic excited states through the equation-of-motion coupled-cluster (EOM-CC) method, <sup>114</sup> because it is relatively black-box compared to multi-reference methods like CASPT2<sup>92</sup> which require artfully chosen active spaces and empirical shift parameters. <sup>171</sup> Unfortunately, despite ongoing efforts to develop perturbative triples corrections for EOM-CCSD excitation energies, none of these attempts have been nearly as successful as CCSD(T). <sup>172</sup> As a result, one must resort to non-iterative triples methods like EOM-CC3<sup>173</sup> or EOM-CCSDT<sup>174</sup> to achieve quantitative (< 0.1 eV) accuracy, which are prohibitive even for modest-sized systems and basis sets. The LR-ODC-12 method improves on EOM-CCSD without including triples parameters.

#### 4.3 Algorithms

The LR-ODC-12 eigenvalue equation has a two-by-two block structure which describes the independent variation of the state parameters and their complex conjugates.

$$\mathbf{E}(\mathbf{z}_k) = \omega_k \mathbf{M}(\mathbf{z}_k), \quad \mathbf{E} = \begin{pmatrix} \mathbf{A} & \mathbf{B} \\ \mathbf{B}^* & \mathbf{A}^* \end{pmatrix}, \quad \mathbf{M} = \begin{pmatrix} \mathbf{S} & \mathbf{0} \\ \mathbf{0} & -\mathbf{S}^* \end{pmatrix}, \quad \mathbf{z}_k = \begin{pmatrix} \mathbf{x}_k \\ \mathbf{y}_k \end{pmatrix}$$
 (4.1)

This block symmetry leads to a paired system of eigenvalues,  $\omega_{\pm k} = \pm \omega_k$ . The submatrices in eq. (4.1) are further blocked according to whether they describe

**Algorithm 1** Canonical multiroot Davidson algorithm for a generalized non-symmetric eigenvalue problem,  $\mathbf{L}\mathbf{v}_k = \lambda_k \mathbf{G}\mathbf{v}_k$ , with periodic subspace collapse. Requires linear transformation functions and diagonal approximations (indicated by tildes) for  $\mathbf{L}$  and  $\mathbf{G}$  and solves for the lowest n eigenvalues and eigenvectors.

```
    procedure Davidson(L(·), G(·), L, G, U<sup>(0)</sup>, n, d<sub>max</sub>, i<sub>max</sub>, r<sub>tol</sub>)
    Initialize the expansion space with a set of guess vectors, U ← U<sup>(0)</sup>.
```

3: **for**  $1 \le i \le i_{\text{max}}$  **do** 

4: Construct subspace representation and solve the lowest n eigenvalues.

$$\mathbf{L}^{\mathrm{sub}} = \mathbf{U}^{\dagger} \mathbf{L}(\mathbf{U})$$

$$\mathbf{G}^{\mathrm{sub}} = \mathbf{U}^{\dagger} \mathbf{G}(\mathbf{U})$$

$$\mathbf{L}^{\mathrm{sub}}\mathbf{v}_{k}^{\mathrm{sub}}=\lambda_{k}\mathbf{G}^{\mathrm{sub}}\mathbf{v}_{k}^{\mathrm{sub}}$$

5: Calculate the eigenvector residuals over the full space.

$$\mathbf{r}_k = (\mathbf{L}(\mathbf{U}) - \lambda_k \mathbf{G}(\mathbf{U})) \mathbf{v}_k^{\mathrm{sub}}$$

- 6: **if**  $\max(\mathbf{r}_k) < r_{\text{tol}}$  for all k **then**
- 7: Set  $\mathbf{v}_k \leftarrow \mathbf{U}\mathbf{v}_k^{\mathrm{sub}}$  and quit the loop. The eigenvectors are converged.
- 8: end if
- 9: Determine new direction vectors by a quasi-Newton-Raphson step.

$$\mathbf{d}_k^{(i)} = -(\tilde{\mathbf{L}} - \lambda_k \tilde{\mathbf{G}})^{-1} \mathbf{r}_k$$

10: Project out the span of U and orthogonalize via SVD compression.

$$\widehat{\mathbf{U}}^{(i)} = (\mathbf{1} - \mathbf{U}^{\dagger} \mathbf{U}) \mathbf{D}^{(i)}$$

$$\widehat{\mathbf{U}}^{(i)} pprox \mathbf{U}^{(i)} \mathbf{\Sigma}^{(i)} \mathbf{W}^{(i)\dagger}$$

- 11: if  $\operatorname{rank}(\mathbf{U}) + \operatorname{rank}(\mathbf{U}^{(i)}) < d_{\max}$  then
- 12: Extend the expansion space,  $\mathbf{U} \leftarrow (\mathbf{U} \ \mathbf{U}^{(i)})$
- 13: else
- 14: Collapse the expansion space,  $\mathbf{U} \leftarrow (\mathbf{U}\mathbf{v}_1^{\text{sub}} \cdots \mathbf{U}\mathbf{v}_n^{\text{sub}})$ .
- 15: **end if**
- 16: end for
- 17: return  $\lambda_k$ ,  $\mathbf{v}_k$
- 18: end procedure

variations with respect to one-body  $(\mathbf{t}_1)$  or two-body  $(\mathbf{t}_2)$  parameters.

$$\mathbf{A} = \begin{pmatrix} \mathbf{A}_{11} & \mathbf{A}_{12} \\ \mathbf{A}_{21} & \mathbf{A}_{22} \end{pmatrix} \quad \mathbf{B} = \begin{pmatrix} \mathbf{B}_{11} & \mathbf{B}_{12} \\ \mathbf{B}_{21} & \mathbf{B}_{22} \end{pmatrix} \quad \mathbf{S} = \begin{pmatrix} \mathbf{S}_{11} & \mathbf{0} \\ \mathbf{0} & \mathbf{1}_{2} \end{pmatrix} \quad \mathbf{x}_{k} = \begin{pmatrix} \mathbf{x}_{k,1} \\ \mathbf{x}_{k,2} \end{pmatrix}$$
(4.2)

The dimensions of the diagonal two-body blocks ( $\mathbf{A}_{22}$  and  $\mathbf{B}_{22}$  and  $\mathbf{1}_s$ ) in these matrices scale as  $(o^2v^2)^2$  with the number of occupied (o) and virtual (v) orbitals, rapidly overwhelming the storage space on a given computer as the number of electrons and basis functions increases. This problem can be circumvented by using so-called "direct" matrix algorithms, which avoid the explicit representation of a matrix as an array of coefficients  $\mathbf{L} = [L_{\mu\nu}]$  by instead defining a function  $\mathbf{L}(\mathbf{U})$  which returns the output of the linear transformation on an arbitrary set of vectors  $\mathbf{U} = (\mathbf{u}_1 \cdots \mathbf{u}_m)$ . The standard direct algorithm for extremal eigenvalues of large, diagonally-dominant matrices is the Davidson algorithm,  $^{175;176}$  which solves the eigenvalue problem by progressively growing an expansion space for the n lowest (or highest) generalized eigenvectors. The procedure is described in Algorithm 1.

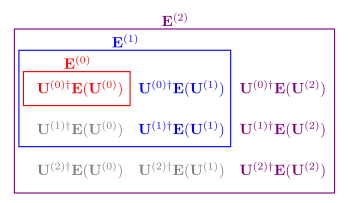
The full set of linear transformation formulas for the LR-ODC-12 Hessian and metric blocks is given in the appendix (section 4.A), but let us consider the diagonal two-body Hessian as an example. The image of an arbitrary two-body vector  $\mathbf{u}_{\mu,2} = [u^{ij}_{\mu,ab}]$  under this transformation is given by

$$\begin{split} (\mathbf{A}_{22}(\mathbf{u}_{\mu,2}))_{ijab} &= -P_{(a/b)}\mathcal{F}_{a}^{c}u_{\mu,cb}^{ij} - P^{(i/j)}\mathcal{F}_{k}^{i}u_{\mu,ab}^{kj} + \frac{1}{2}\overline{g}_{ab}^{cd}u_{\mu,cd}^{ij} + \frac{1}{2}\overline{g}_{kl}^{ij}u_{\mu,ab}^{kl} \\ &- P_{(a/b)}^{(i/j)}\overline{g}_{la}^{jc}u_{\mu,cb}^{il} + \frac{1}{2}P_{(a/b)}\mathcal{G}_{af}^{ec}t_{eb}^{ij}t_{kl}^{fd*}u_{\mu,cd}^{kl} + \frac{1}{2}P_{(a/b)}\mathcal{G}_{ka}^{me}t_{eb}^{ij}t_{ml}^{cd*}u_{\mu,cd}^{kl} \\ &+ \frac{1}{2}P^{(i/j)}\mathcal{G}_{me}^{ic}t_{ab}^{mj}t_{kl}^{ed*}u_{\mu,cd}^{kl} + \frac{1}{2}P^{(i/j)}\mathcal{G}_{mk}^{in}t_{ab}^{mj}t_{cd*}^{cd*}u_{\mu,cd}^{kl} \end{split} \tag{4.3}$$

where the  $i, j, k, l, \ldots$  run over occupied spin-orbitals,  $a, b, c, d, \ldots$  run over virtual (un-occupied) spin-orbitals, and summation over repeated indices is implied. The important thing to note is that we need only two v<sup>4</sup>-sized intermediates ( $\overline{g}_{ab}^{cd}$  and  $\mathcal{G}_{ab}^{cd}$ ) to define our linear transformations, and that the image of the two-body Hessian only has  $\mathcal{O}(d_{\mu}o^2v^2)$  elements where  $d_{\mu}$  is the dimension of the  $\mu$  index. In general the Davidson algorithm requires no more than 10-20 subspace vectors per root, sometimes as few as 2 or 3, so that even for small systems it is easy to satisfy  $d_{\mu} \ll o^2v^2$ , resulting in substantial memory savings. Equation (4.3) also includes the timing bottleneck for the LR-ODC-12, the contraction of the  $v^4$  integrals with the expansion vector,  $g_{ab}^{cd}u_{\mu,cd}^{ij}$ , which scales as  $\mathcal{O}(d_{\mu}o^2v^4)$  in the number of floating point operations. This term is the rate limiting step in EOM-CCSD as well.

A direct application of algorithm 1 to eq. (4.1) is complicated by the fact that the desired (small, positive) eigenvalues are in the middle of the spectrum, rather than at the extremes. We make use of two strategies.

Figure 4.1: A schematic illustrating the sequence of reduced representations of **E** in a blocked Davidson algorithm. The colored entries must be explicitly computed, whereas the gray entries are determined by Hermitian symmetry.



**Strategy 1** One approach is to invert the eigenvalue equation, solving for the *largest* positive inverse eigenvalues, which is easily done by selecting the highest rather than the lowest roots at each step in algorithm 1.

$$\mathbf{M}(\mathbf{z}_k) = \omega_k^{-1} \mathbf{E}(\mathbf{z}_k) \tag{4.4}$$

This has the added advantage that the energy Hessian  $\mathbf{E}$  is generally positive definite, which allows us to treat the subspace diagonalizations as standard Hermitian eigenvalue equation with real eigenvalues and orthonormal eigenvectors. A simple, effective choice of guess vectors in this approach is to determine the lowest eigenvectors of the following diagonal approximation to eq. (4.3)

$$(\tilde{\mathbf{A}}_{22}(\mathbf{u}_{\mu}))_{ijab} \equiv -(\mathcal{F}_i^i + \mathcal{F}_j^j + \mathcal{F}_a^a + \mathcal{F}_b^b)u_{\mu,ab}^{ij}$$

$$(4.5)$$

which are simply the standard unit vectors associated with the smallest diagonal entries. The virtual coefficients  $\mathcal{F}_a^a$ ,  $\mathcal{F}_b^b$  in this equation are generally negative, so this has the form of an orbital energy difference like we see in Hartree-Fock theory. For larger systems, we manage the memory requirements of the method by keeping successive groups of expansion vectors and images,  $\mathbf{U}^{(i)}$ ,  $\mathbf{E}(\mathbf{U}^{(i)})$ ,  $\mathbf{M}(\mathbf{U}^{(i)})$ , on disk and reading them in as needed. These can be further subdivided into even smaller blocks if needed. Figure 4.1 illustrates which entries in the subspace representation need to be explicitly computed as we add vectors to the expansion space.

**Strategy 2** Assuming real matrices, we can add and subtract the block rows of eq. (4.1) to arrive at the following pair of equations.

$$(\mathbf{A} \pm \mathbf{B})(\mathbf{x}_k \pm \mathbf{y}_k) = \omega_k \mathbf{S}(\mathbf{x}_k \mp \mathbf{y}_k) \tag{4.6}$$

Substituting one into the other leads to a non-symmetric eigenvalue equation for the squares of the excitation energies.

$$\mathbf{H}(\mathbf{x}_k - \mathbf{y}_k) = \omega_k^2(\mathbf{x}_k - \mathbf{y}_k), \quad \mathbf{H} \equiv \mathbf{S}^{-1}(\mathbf{A} + \mathbf{B})\mathbf{S}^{-1}(\mathbf{A} - \mathbf{B})$$
 (4.7)

By halving the dimension of the problem and eliminating the metric, this reduces the memory requirements by a factor of three, provided we do not need to recover the full eigenvector  $\mathbf{z}_k$  to compute transition properties. From eq. (4.2) we can see that computing the inverse of  $\mathbf{S}$  requires us to invert the orbital block of the metric. This matrix is given by

$$(\mathbf{S}_{11})_{ia,jb} = \gamma_i^i \delta_a^b - \delta_i^i \gamma_a^b \tag{4.8}$$

where  $\gamma_j^i$  and  $\gamma_a^b$  are occupied and virtual blocks of the one-body density matrix. By expanding these matrices in the natural spin-orbital basis (indicated by prime indices) where the one-body density matrix is diagonal

$$\gamma_j^i = (\mathbf{Y})_j^{j'} (\mathbf{Y}^\dagger)_{i'}^i \delta_{j'}^{i'} \gamma_{j'} \qquad \gamma_a^b = (\mathbf{Y})_a^{a'} (\mathbf{Y}^\dagger)_{b'}^b \delta_{a'}^{b'} \gamma_{a'}$$
(4.9)

we can derive the following simple and inexpensive formula for the inverse of the orbital metric.

$$(\mathbf{S}_{11}^{-1})_{ia,jb} = \frac{(\mathbf{Y}^{\dagger})_{j'}^{i}(\mathbf{Y})_{a}^{b'}}{\gamma_{j'} - \gamma_{b'}} (\mathbf{Y}^{\dagger})_{b'}^{b} (\mathbf{Y})_{j}^{j'}$$

$$(4.10)$$

The appropriate diagonal approximation to  ${\bf H}$  can simply be computed as the square of  $\tilde{\bf A}$ , since in low order we have  ${\bf B}\approx {\bf 0}$  and  ${\bf S}\approx {\bf 1}$ 

#### 4.4 Computational Details

The LR-ODC-12 and LR-OLCCD methods were implemented as a standalone Python program, which was interfaced with Psi4<sup>167</sup> and PyscF<sup>168</sup> to obtain the one- and two-electron integrals. We used several iterations of Strategy 1, as described in section 4.3, to assign states based on the transition dipole moments and used Strategy 2 to fully converge the energies. We used the Q-CHEM program, version 4.4, <sup>169</sup> to calculate EOM-CCSD values for comparison. The ANO-L-pVXZ (X = D, T) basis sets<sup>177</sup> were used in the excitation energy computations as in Ref. 178. Frozen-core MP2/cc-pVQZ geometries were used as in Refs. 178 and 179.

#### 4.5 Results and Discussion

We apply the LR-ODC-12 method to challenging excited states of ethylene ( $C_2H_4$ ), butadiene ( $C_4H_6$ ), and hexatriene ( $C_6H_8$ ). A reliable description of these electronic states requires accurate treatment of electron correlation. <sup>178;179;181–195</sup> All three molecules feature a dipole-allowed  $1^1B_u$  (or  $1^1B_{1u}$ ) state that is well de-

Table 4.1: Vertical excitation energies computed using LR-OLCCD, LR-ODC-12, and EOM-CCSD for the low-lying electronic states of ethylene ( $C_2H_4$ ), butadiene ( $C_4H_6$ ), and hexatriene ( $C_6H_8$ ). Computations employed the ANO-L-pVDZ (for  $C_4H_6$  and  $C_6H_8$ ) and ANO-L-pVTZ (for  $C_2H_4$ ) basis sets and the MP2/cc-pVQZ optimized geometries. For LR-OLCCD and LR-ODC-12, oscillator strengths of the allowed transitions are given in parentheses. All electrons were correlated in all computations.

		EOM-CCSD	LR-OLCCD	LR-ODC-12	SHCIa
$C_2H_4$	$1^3 \mathrm{B_{1u}}$	4.46	4.66	4.52	4.59
	$1^1\mathrm{B}_{1\mathrm{u}}$	8.14	8.20(1.8)	8.14(1.9)	8.05
$C_4H_6$	$1^3 \mathrm{B_u}$	3.20	3.58	3.43	3.37
	$1^1\mathrm{B_u}$	6.53	6.76(4.2)	6.67(4.4)	6.45
	$2^1 A_g$	7.28	7.14	6.81	6.58
$C_6H_8$	$1^3 B_u$	2.64	3.01	2.83	2.77
0 0	$1^1\mathrm{B_u}$	5.60	5.89(6.5)	5.74(8.1)	5.59
	$2^1 A_g$	6.55	$4.\overline{21}$	5.73	5.58

<sup>&</sup>lt;sup>a</sup> Also shown are the excitation energies from the semistochastic heat-bath CI (SHCI) method, extrapolated to full CI limit. <sup>180</sup> The 1s orbitals of carbon atoms were not included in the SHCI correlation treatment. The SHCI computations used the same basis sets and optimized geometries as those used for LR-OLCCD, LR-ODC-12, and EOM-CCSD.

scribed by a  $\pi - \pi^*$  excitation, but requires a very accurate description of dynamic correlation between the  $\sigma$  and  $\pi$  electrons. In butadiene and hexatriene, the  $1^1B_u$  state is near-degenerate with a dipole-forbidden  $2^1A_g$  state that has a substantial double-excitation character, requiring the description of static correlation in the  $\pi$  and  $\pi^*$  orbitals. <sup>189–191</sup> For this reason, the relative energies and ordering of the  $1^1B_u$  and  $2^1A_g$  states are very sensitive to various levels of theory. For example, single-reference methods truncated to single and double excitations are usually more accurate in describing the  $1^1B_u$  state than the  $2^1A_g$  state, while

multi-reference methods are more reliable for the  $2^1A_g$  state, missing important dynamic correlation for the  $1^1B_u$  state. Very recently, Chien et al.  $^{180}$  reported accurate vertical excitation energies for the low-lying states of ethylene, butadiene, and hexatriene computed using semistochastic heat-bath configuration interaction (SHCI) extrapolated to the full CI limit. In this section, we will use the SHCI results to benchmark the accuracy of the LR-ODC-12 method.

Table 4.1 reports the vertical excitation energies of ethylene, butadiene, and hexatriene computed using the EOM-CCSD, LR-OLCCD, and LR-ODC-12 methods, along with the SHCI results from Ref. 180. All methods employed the same optimized geometries and basis sets (see table 4.1 for details). We refer to the  $B_{1u}$  states of  $C_2H_4$  as  $B_u$  for brevity. All excitation energies decrease as the number of double bonds increases. For butadiene and hexatriene, the (1<sup>1</sup>B<sub>u</sub>; 2<sup>1</sup>A<sub>g</sub>) excitation energies computed using the SHCI method are (6.45; 6.58) and (5.59; 5.58) eV, respectively, indicating that the two states are nearly degenerate for the longer polyene. This feature is not reproduced by the EOM-CCSD method, which predicts the 1<sup>1</sup>B<sub>u</sub> state energies in close agreement with SHCI, but significantly overestimates the energies for the doubly-excited 2<sup>1</sup>A<sub>g</sub> state. As a result, the EOM-CCSD method overestimates the energy spacing between the 1<sup>1</sup>B<sub>u</sub> and 2<sup>1</sup>A<sub>g</sub> states by  $\sim 0.6$  eV and 1.0 eV for butadiene and hexatriene, respectively.

On the contrary, the LR-ODC-12 method correctly describes the relative energies and ordering of the  $1^1B_u$  and  $2^1A_g$  states, predicting their energy spacing to be 0.14 and 0.01 eV for butadiene and hexatriene, respectively, in an excellent agreement with the SHCI results (0.13 and 0.01 eV). For the singlet excited states,

the LR-ODC-12 method consistently overestimates the SHCI excitation energies by  $\sim 0.1-0.2$  eV. For the  $1^3B_u$  state, the LR-ODC-12 errors are smaller in magnitude ( $\sim 0.06$  eV). Importantly, these results suggest that the LR-ODC-12 method provides a balanced description of the excited states with different electronic structure effects, as illustrated by its consistent performance for the  $1^3B_u$ ,  $1^1B_u$ , and  $2^1A_g$  states in ethylene, butadiene, and hexatriene.

Comparing the LR-ODC-12 and the LR-OLCCD results suggests that including the non-linear terms in the description of electron correlation significantly improves the agreement with the reference SHCI data. In particular, for the  $1^3B_u$  and  $1^1B_u$  states, the LR-OLCCD errors are larger than the LR-ODC-12 errors by  $\sim 0.15$  eV. For the doubly-excited  $2^1A_g$  state, LR-OLCCD shows very large errors of 0.56 and -1.37 eV for butadiene and hexatriene, respectively. These results indicate that the non-linear terms included in LR-ODC-12 are very important for the description of excited states with double-excitation character.

#### 4.A LR-ODC-12 Linear Transformation Formulas

$$(\mathbf{A}_{11}(\mathbf{u}_{\mu,1}))_{ia} = h_{j}^{i} \gamma_{a}^{b} u_{\mu,b}^{j} + h_{a}^{b} \gamma_{j}^{i} u_{\mu,b}^{j} - (\bar{\mathbf{F}})_{j}^{i} u_{\mu,a}^{j} - (\bar{\mathbf{F}})_{a}^{b} u_{\mu,b}^{i} + \bar{g}_{nj}^{mi} \gamma_{ma}^{nb} u_{\mu,b}^{j} + \bar{g}_{ma}^{nb} \gamma_{nj}^{mi} u_{\mu,b}^{j} + \bar{g}_{jf}^{ie} \gamma_{ae}^{bf} u_{\mu,b}^{j} + \bar{g}_{ae}^{bf} \gamma_{jf}^{ie} u_{\mu,b}^{j} + \bar{g}_{me}^{ib} \gamma_{ja}^{me} u_{\mu,b}^{j} + \bar{g}_{ja}^{me} \gamma_{me}^{ib} u_{\mu,b}^{j}$$

$$(4.11)$$

$$(\mathbf{B}_{11}(\mathbf{u}_{\mu,1}))_{ia} = \overline{g}_{be}^{im} \gamma_{ma}^{je} u_{\mu,j}^{b} + \overline{g}_{ma}^{je} \gamma_{be}^{im} u_{\mu,j}^{b} + \overline{g}_{mb}^{ie} \gamma_{ae}^{jm} u_{\mu,j}^{b} + \overline{g}_{ae}^{jm} \gamma_{mb}^{ie} u_{\mu,j}^{b} + \frac{1}{2} \overline{g}_{mn}^{ij} \gamma_{ab}^{mn} u_{\mu,j}^{b} + \frac{1}{2} \overline{g}_{ab}^{mn} \gamma_{mn}^{ij} u_{\mu,j}^{b} + \frac{1}{2} \overline{g}_{ef}^{ij} \gamma_{ab}^{ef} u_{\mu,j}^{b} + \frac{1}{2} \overline{g}_{ab}^{ef} \gamma_{ef}^{ij} u_{\mu,j}^{b}$$

$$(4.12)$$

$$(\mathbf{A}_{22}(\mathbf{u}_{\mu,2}))_{ijab} = -P_{(a/b)}\mathcal{F}_{a}^{c}u_{\mu,cb}^{ij} - P^{(i/j)}\mathcal{F}_{k}^{i}u_{\mu,ab}^{kj} + \frac{1}{2}\overline{g}_{ab}^{cd}u_{\mu,cd}^{ij} + \frac{1}{2}\overline{g}_{kl}^{ij}u_{\mu,ab}^{kl}$$

$$-P_{(a/b)}^{(i/j)}\overline{g}_{la}^{jc}u_{\mu,cb}^{il} + \frac{1}{2}P_{(a/b)}\mathcal{G}_{af}^{ec}t_{eb}^{ij}t_{kl}^{fd*}u_{\mu,cd}^{kl} + \frac{1}{2}P_{(a/b)}\mathcal{G}_{ka}^{me}t_{eb}^{ij}t_{ml}^{cd*}u_{\mu,cd}^{kl}$$

$$+ \frac{1}{2}P^{(i/j)}\mathcal{G}_{me}^{ic}t_{ab}^{mj}t_{kl}^{ed*}u_{\mu,cd}^{kl} + \frac{1}{2}P^{(i/j)}\mathcal{G}_{mk}^{in}t_{ab}^{mj}t_{cd}^{cd*}u_{\mu,cd}^{kl}$$

$$(4.13)$$

$$(\mathbf{B}_{22}(\mathbf{u}_{\mu,2}))_{ijab} = \frac{1}{2} P_{(a/b)} \mathcal{G}_{ac}^{ef} t_{eb}^{ij} t_{fd}^{kl} u_{\mu,kl}^{cd} + \frac{1}{2} P_{(a/b)} \mathcal{G}_{na}^{ke} t_{eb}^{ij} t_{cd}^{nl} u_{\mu,kl}^{cd} + \frac{1}{2} P^{(i/j)} \mathcal{G}_{mc}^{if} t_{ab}^{mj} t_{fd}^{kl} u_{\mu,kl}^{cd} + \frac{1}{2} P^{(i/j)} \mathcal{G}_{mn}^{ik} t_{ab}^{mj} t_{cd}^{nl} u_{\mu,kl}^{cd}$$

$$(4.14)$$

$$(\mathbf{A}_{12}(\mathbf{u}_{\mu,2}))_{ia} = \frac{1}{2} \overline{g}_{la}^{cd} u_{\mu,cd}^{il} + \frac{1}{2} \overline{g}_{kl}^{id} u_{\mu,ad}^{kl} + \frac{1}{2} (\mathcal{I}_{a}^{i})_{k}^{m} t_{ml}^{cd*} u_{\mu,cd}^{kl} + \frac{1}{2} (\mathcal{I}_{a}^{i})_{e}^{c} t_{kl}^{ed*} u_{\mu,cd}^{kl}$$

$$+ \overline{g}_{ae}^{mc} t_{ml}^{ed*} u_{\mu,cd}^{il} + \overline{g}_{ke}^{im} t_{ml}^{ed*} u_{\mu,ad}^{kl} + \frac{1}{4} \overline{g}_{la}^{mn} t_{mn}^{cd*} u_{\mu,cd}^{il}$$

$$+ \frac{1}{4} \overline{g}_{ef}^{id} t_{kl}^{ef*} u_{\mu,ad}^{kl}$$

$$(4.15)$$

$$(\mathbf{B}_{12}(\mathbf{u}_{\mu,2}))_{ia} = \frac{1}{2} (\mathcal{I}_{a}^{i})_{m}^{k} t_{cd}^{ml} u_{\mu,kl}^{cd} + \frac{1}{2} (\mathcal{I}_{a}^{i})_{c}^{e} t_{ed}^{kl} u_{\mu,kl}^{cd} + \overline{g}_{ad}^{le} t_{ce}^{kl} u_{\mu,kl}^{cd} + \overline{g}_{md}^{il} t_{ca}^{km} u_{\mu,kl}^{cd} + \frac{1}{4} \overline{g}_{ma}^{kl} t_{cd}^{im} u_{\mu,kl}^{cd} + \frac{1}{4} \overline{g}_{cd}^{ie} t_{ae}^{kl} u_{\mu,kl}^{cd}$$

$$(4.16)$$

$$(\mathbf{A}_{21}(\mathbf{u}_{\mu,1}))_{ijab} = P^{(i/j)} \overline{g}_{ab}^{jc} u_{\mu,c}^{i} + P_{(a/b)} \overline{g}_{kb}^{ij} u_{\mu,a}^{k} + P^{(i/j)} (\mathcal{I}_{k}^{c})_{m}^{i} t_{ab}^{mj} u_{\mu,c}^{k}$$

$$+ P_{(a/b)} (\mathcal{I}_{k}^{c})_{a}^{e} t_{eb}^{ij} u_{\mu,c}^{k} + P_{(a/b)}^{(i/j)} \overline{g}_{ma}^{ce} t_{eb}^{mj} u_{\mu,c}^{i} + P_{(a/b)}^{(i/j)} \overline{g}_{km}^{ie} t_{eb}^{mj} u_{\mu,a}^{k}$$

$$+ \frac{1}{2} P^{(i/j)} \overline{g}_{mn}^{jc} t_{ab}^{mn} u_{\mu,c}^{i} + \frac{1}{2} P_{(a/b)} \overline{g}_{kb}^{ef} t_{ef}^{ij} u_{\mu,a}^{k}$$

$$(4.17)$$

$$(\mathbf{B}_{21}(\mathbf{u}_{\mu,1}))_{ijab} = P^{(i/j)}(\mathcal{I}_{c}^{k})_{m}^{i} t_{ab}^{mj} u_{\mu,k}^{c} + P_{(a/b)}(\mathcal{I}_{c}^{k})_{a}^{e} t_{eb}^{ij} u_{\mu,k}^{c} + P_{(a/b)}^{(i/j)} \overline{g}_{cb}^{je} t_{ae}^{ik} u_{\mu,k}^{c} + P_{(a/b)}^{(i/j)} \overline{g}_{kj}^{mb} t_{ac}^{im} u_{\mu,k}^{c} + \overline{g}_{mc}^{ij} t_{ab}^{km} u_{\mu,k}^{c} + \overline{g}_{ab}^{ke} t_{ce}^{ij} u_{\mu,k}^{c} + P_{(a/b)}^{(i/j)} \overline{g}_{cb}^{mb} t_{ae}^{im} u_{\mu,k}^{c} + \overline{g}_{ab}^{ij} t_{ce}^{im} u_{\mu,k}^{c}$$

$$(4.18)$$

$$(\mathbf{S}_{11}(\mathbf{u}_{\mu,1}))_{ia} = \gamma_i^i u_{\mu,a}^j - \gamma_a^b u_{\mu,b}^i$$
(4.19)

$$(\mathbf{S}_{11}^{-1}(\mathbf{u}_{\mu,1}))_{ia} = \frac{(\mathbf{Y}^{\dagger})_{j'}^{i}(\mathbf{Y})_{a}^{b'}}{\gamma_{j'} - \gamma_{b'}} (\mathbf{Y}^{\dagger})_{b'}^{b} (\mathbf{Y})_{j'}^{j'} u_{\mu,b}^{j}$$
(4.20)

$$(\mathbf{Y}^{\dagger})_{q'}^{p} \gamma_q^p (\mathbf{Y})_p^{p'} = \delta_{q'}^{p'} \gamma_{q'} \tag{4.21}$$

$$\tilde{\mathbf{S}}_{11} \equiv \tilde{\mathbf{S}}_{11}^{-1} \equiv \mathbf{1}_1 \tag{4.22}$$

$$(\tilde{\mathbf{A}}_{11})_{ia,ia} \equiv -f_i^i + f_a^a \tag{4.23}$$

$$(\tilde{\mathbf{A}}_{22})_{ijab,ijab} \equiv -\mathcal{F}_i^i - \mathcal{F}_j^j - \mathcal{F}_a^a - \mathcal{F}_b^b \tag{4.24}$$

## Chapter 5

### Conclusion

We have presented comprehensive benchmarks to demonstrate that orbital-optimized density cumulant theory with double excitations (ODC-12) consistently outperforms coupled-cluster theory with singles and doubles (CCSD) for the description of noncovalent interactions, hydrogen-transfer barrier heights, radical stabilization energies, ionization energies, and covalent bond stretching. Thus having established the promising performance of this model for ground state calculations, we have extended the theory for the calculation of excitation energies and transition properties through the use of linear response theory. After numerically demonstrating that our initial working equations and implementation are correct, we have empirically shown this method to be size-intensive, i.e. displaying the correct qualitative behavior with respect to excited states of independent systems. Next, we have shown that this method is more stable with respect to strong electron correlation than its linearized variant, LR-OLCCD, which often achieves impressive error cancellation in the absence of strong correlation. This demonstrates that the infinite order one-particle n-representability conditions defining the ODC-12

method contribute to a robust description of the electron distribution for more challenging states. Our initial benchmark study of the vertical excitation energies predicted by this method shows that it reduces the mean absolute error by roughly a factor of two relative to the popular equation-of-motion coupled-cluster with singles and doubles (EOM-CCSD) method, similar to our findings for ground states. For well-behaved systems we find that the linearized model, LR-OLCCD, is an effective approximation to LR-ODC-12 with a lower cost prefactor. Finally, we develop some improvements to the algorithms used for solving the LR-ODC-12 equations using disk-based direct matrix algorithms (variants of the Davidson algorithm). These developments allow us to study polyene systems as large as hexatriene with a natural orbital basis of double-zeta quality. This calculation involves 44 electrons and 124 spatial orbitals with nearly 20 million unique wavefunction parameters, which would not be feasible without the new algorithms. The advantages of LR-ODC-12 over EOM-CCSD and LR-OLCCD for these polyene systems are even more stark than for our previous benchmarks. Whereas EOM-CCSD overestimates the energy of the challenging 2<sup>1</sup>A<sub>g</sub> state of hexatriene and its gap with the neighboring 1<sup>1</sup>B<sub>u</sub> state by close to 1 eV each, LR-ODC-12 matches its energy to within 0.15 eV and matches the energy gap to within 0.01 eV. Given the relative sparsity of inexpensive alternatives to EOM-CCSD, we believe that these results merit further development of algorithms for the LR-ODC-12 method to expand our toolkit for studying excited electronic states.

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