Probing Molecular Chirality of Ground and Electronically Excited States in the UV–vis and X-ray Regimes: An EOM-CCSD Study

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Abstract

We present several strategies for computing electronic circular dichroism (CD) spectra across different frequency ranges at the equation-of-motion coupled-cluster singles and doubles level of theory. CD spectra of both ground and electronic excited states are discussed. For selected cases, the approach is compared with coupled-cluster linear response results as well as time-dependent density functional theory. The extension of the theory to include the effect of spin–orbit coupling is presented and illustrated by calculations of X-ray CD spectra at the L edge.

1 Introduction

An object is chiral if its mirror images cannot be brought into coincidence by rotation. Chirality allows one to distinguish between “right-handed” and “left-handed” forms of the
same object. In molecules, chirality typically arises due to the presence of “chiral centers” such as asymmetrically substituted tetrahedral centers (most often carbon atoms, but other types of atoms are also possible) or due to the handedness of a helical structure; the latter is also known as axial chirality. In axially chiral systems, the clockwise and counterclockwise structures are non-superimposable. The isomers with non-superimposable mirror structures are called enantiomers, whereas those that are non-superimposable and are not mirror images are named diastereomers (or diastereoisomers).¹

Chirality can be mathematically characterized by means of rotationally invariant pseudoscalars (mixed scalar–vector products). Pseudoscalars do not depend on the orientation of the object and change sign upon inversion,¹ which means that they have opposite sign for opposite enantiomers. The interaction between two chiral objects depends on their relative handedness, so the observable outcome of the interaction of a chiral probe with the different enantiomers of the same molecular species is different, which is at the foundation of chiral recognition.

Chirality is a common and important molecular property with broad consequences. At molecular level, many biological objects, such as sugars, amino acids, and secondary structures of proteins and nucleic acids, are chiral. Even more fascinating, life is homochiral—in living organisms, biological molecules exist almost exclusively as single enantiomers (enantiomerically pure). For example, all naturally occurring amino acids are left-handed and all naturally occurring sugars are right-handed. This is critical for molecular recognition and replication processes. Hence, chirality is essential for understanding the origin of life and its molecular-level operation. The origin of the homochirality in life remains an unsolved mystery: when synthesized in the lab, left- and right-handed molecules of a compound most often form a racemic mixture, implying that primordial processes in Nature must have been driven by a directing (chiral) template.² Chirality makes living beings enantio-sensitive. This is critically important for the pharmaceutical and food industry because different enantiomers have drastically different biological effects: e.g., left-handed sugars taste bitter; different
enantiomers of the same drug may have different pharmacological effects, the thalidomide tragedy\textsuperscript{3} being a never-fading reminder. Any prospective new drug based on a chiral structure must have clearly assigned absolute configuration.

Opposite enantiomers have the same energy spectrum; hence, they cannot be distinguished by simple absorption of linearly polarized light. A pseudoscalar effect is needed, such as the differential interaction with the left or right circularly polarized components of linearly polarized light. Differential transmission results in oppositely signed rotation of the plane of polarization, which is exploited in optical rotation (OR) measurements. Even more powerful is the technique of measuring the differential absorption of the two circularly polarized components, which is the basis of circular dichroism spectroscopies.\textsuperscript{4} In electronic circular dichroism (ECD),\textsuperscript{4,5} the probing light activates electronic transitions in the sample, most typically in the UV–visible frequency region, where valence excitations occur. X-ray radiation can also be used, in principle, to probe chirality via excitations of core electrons. In vibrational circular dichroism (VCD),\textsuperscript{1,4,6} infrared light is used to probe dichroism effects in vibrational transitions of the enantiomers in their ground states. An emerging chiral rotational spectroscopy exploits dichroism effects using microwave (MW-CD) radiation.\textsuperscript{7–9} Chirality can also be probed by transitions to the continuum, as done in photoelectron circular dichroism.\textsuperscript{10,11}

The key step in contemporary use of spectroscopy for chiral discrimination is the simulation of the spectra for an \textit{a priori} chosen absolute configuration of the chiral species.\textsuperscript{12,13} Indeed, experimental measurements of optical activity alone are not sufficient to establish the absolute configuration, since there is no direct correspondence between the enantiomer’s configuration and the measured sign of the chiroptical property. However, if the simulated spectrum of the chosen configuration matches the sign pattern of the measured spectrum, one can conclude that the experimental sample contains the same enantiomer as in the simulation,\textsuperscript{5,12–14} provided, of course, that the computational method is reliable.

The appeal of this strategy has stimulated the development of reliable computational
methods to simulate ORD, ECD, and VCD spectra. Time-dependent density functional approaches (TDDFT) are undoubtedly playing the leading role in this endeavour,\textsuperscript{5,12} but the need for alternative, wave-function-based methods has been highlighted on many occasions—at the very least, to provide a reliable benchmark against which the validity and accuracy of TDDFT can be checked.\textsuperscript{15–19}

Equation-of-motion coupled-cluster theory (EOM-CC)\textsuperscript{20–23} and the closely related coupled-cluster response theory (CC-RSP)\textsuperscript{18,24,25} are popular and robust theoretical frameworks for simulating spectroscopy. The portfolio of spectroscopies that can be treated using EOM-CC/CC-RSP methods is vast and expanding. Spanning linear and non-linear regimes, as well as different frequency ranges of probing radiation, it comprises UV–vis one- and two-photon absorption\textsuperscript{26–28} and dichroism,\textsuperscript{29} magnetic circular dichroism,\textsuperscript{30,31} X-ray absorption,\textsuperscript{32} dichroism\textsuperscript{33} and photoemission,\textsuperscript{32,34} resonant inelastic X-ray scattering,\textsuperscript{33,35,36} and many more. The non-Hermitian (bi-variational) nature of the underlying formalism has initially posed some complications (compared to Hermitian variational methods) for addressing effects related to natural optical activity, but solutions have been proposed as how to deal with issues such as gauge-invariance and gauge-origin dependency.\textsuperscript{37,38}

In this work, we focus on the EOM-CC theory, specifically, the EOM-CC model with singles and doubles (EOM-CCSD),\textsuperscript{20} as the theoretical framework for computing electronic circular dichroism spectra. We consider the calculation of ECD both in the UV–vis and in the X-ray frequency regimes. The latter requires using specific techniques to treat the core-level transitions—for that purpose, we employ our frozen-core Core–Valence-Separated (fc-CVS) EOM-CC approach.\textsuperscript{32} In addition to the ECD of molecules in their ground states, we also simulate excited-state circular dichroism, i.e., the ECD spectra of electronically excited species, as needed for the simulation of time-resolved/pump–probe experiments employing circular dichroism as the probe.\textsuperscript{39–43} So far, theoretical studies of excited-state ECD have been limited to the TDDFT\textsuperscript{43,44} and algebraic diagrammatic construction (ADC) levels of theory.\textsuperscript{45}
While the fundamental equations for simulating ECD using EOM-CCSD have been known for decades, only CC-RSP appears to have been used.\textsuperscript{16,17,46–49} One probable reason may have been a fear of the “lack of size-intensivity” of the EOM-CCSD transition moments.\textsuperscript{50} However, whether this feature of the theory poses any serious practical problems in calculations of ECD spectra has not been explored; and we consider this issue here. An advantage of EOM-CC over CC-RSP is that EOM-CC does not require additional response equations (for the excited-state multipliers) to be solved and additional matrix transformations to be performed, thus resulting in a computationally slightly less expensive approach. For excited-state properties, EOM-CC is also less prone to divergences, which may become particular important in the high-energy frequency regime, and does not have artificial poles arising due to the $T$-amplitude response.\textsuperscript{51}

The paper is organized as follows. In the theory section we recapitulate the general definitions of the ECD spectroscopic parameters and observables, and then discuss alternative strategies for computing them within EOM-CCSD and CCSD-RSP. This encompasses both the calculation of “stick spectra” (i.e., excitation energies and ECD rotatory strengths) and the direct calculation of the ECD cross sections via damped response theory. We then present illustrative results for the UV ECD of the ground and excited states of methyloxirane, norcamphor, and binol. We also discuss the exemplary X-ray CD results for methyloxirane and L-alanine at the $K$ edge and chloroethanol at the $L$ edge of chlorine, with and without the inclusion of spin–orbit splitting. Concluding remarks are given at the end.

2 Theory

2.1 Definitions

Circular dichroism spectra are typically obtained by calculating the rotatory strengths $R_{nf}$ for the electronic transitions $n \rightarrow f$ (most often $n = 0$, i.e. the ground state). The rotatory strengths are defined, in the length gauge (label lg), as scalar products of the electric and
magnetic transition dipole moments (Rosenfeld equation),

\[ R_{nf}^{lg} = \sum_{\alpha=x,y,z} \Im \langle n| \hat{\mu}_\alpha | f \rangle \langle f | \hat{m}_\alpha | n \rangle = \frac{1}{2} \sum_{\alpha=x,y,z} \Im \langle n| \hat{r}_\alpha | f \rangle \langle f | \hat{L}_\alpha | n \rangle, \]  

(1)

where \( \Im \) stands for the imaginary part and \( \hat{\mu}_\alpha \) and \( \hat{m}_\alpha \) are the Cartesian components of the electric dipole and magnetic dipole operators, respectively. Note that, in atomic units (a.u.), \( \hat{\mu} = -\hat{r} \) and \( \hat{m} = -\frac{1}{2} \hat{L} \), where \( \hat{r} \) is the position operator, \( \hat{L} = \hat{r} \times \hat{p} = -i\hat{r} \times \hat{\nabla} \) is the orbital angular momentum, and \( \hat{p} = -i\hat{\nabla} \) is the linear momentum. By exploiting Ehrenfest’s off-diagonal hypervirial relationship (in a.u.)\(^{15,52,53}\)

\[ i \langle n| \hat{p} | f \rangle = \langle n| [\hat{r}, \hat{H}] | f \rangle = \omega_{fn} \langle n| \hat{r} | f \rangle, \]  

(2)

where \( \omega_{fn} \) is the energy difference between the two states, one obtains the rotatory strength in the velocity gauge (abbreviated vg)

\[ R_{nf}^{vg} = \frac{1}{\omega_{fn}} \sum_{\alpha=x,y,z} \Re \langle n| \hat{p}_\alpha | f \rangle \langle f | \hat{p}_\alpha | n \rangle = -\frac{1}{2\omega_{fn}} \sum_{\alpha=x,y,z} \Re \langle n| \hat{p}_\alpha | f \rangle \langle f | \hat{L}_\alpha | n \rangle, \]  

(3)

where \( \Re \) indicates the real part.

The dipole oscillator strength \( f \) of one-photon absorption can also be expressed in different gauges—length, velocity, and mixed:\(^{54}\)

\[ f^{lg} = \frac{2\omega_{fn}}{3} \sum_{\alpha} \langle n| \hat{\mu}_\alpha | f \rangle \langle f | \hat{\mu}_\alpha | n \rangle, \]  

(4)

\[ f^{vg} = \frac{2}{3\omega_{fn}} \sum_{\alpha} \langle n| \hat{p}_\alpha | f \rangle \langle f | \hat{p}_\alpha | n \rangle, \]  

(5)

\[ f^{mx} = \frac{2i}{3} \sum_{\alpha} \langle n| \hat{\mu}_\alpha | f \rangle \langle f | \hat{p}_\alpha | n \rangle. \]  

(6)
2.2 EOM-CC theory for valence and core-level states

The (ground-state) coupled-cluster wave function ($\Psi_{\text{CC}}$) is given in terms of the exponential of the cluster operator $\hat{T}$ acting on the reference (usually, Hartree–Fock) Slater determinant $\Phi_0$: \(^5^5\)

$$|\Psi_{\text{CC}}\rangle \equiv |\Psi_0\rangle = e^{\hat{T}}|\Phi_0\rangle. \quad (7)$$

At CCSD level, the cluster operator is expressed in terms of electron-annihilation ($\hat{a}_i, \hat{a}_j$) and electron-creation ($\hat{a}^\dagger_a, \hat{a}^\dagger_b$) operators as follows:

$$\hat{T} = \hat{T}_1 + \hat{T}_2; \quad \hat{T}_1 = \sum_{ia} t^a_i \hat{a}^\dagger_a \hat{a}_i; \quad \hat{T}_2 = \frac{1}{4} \sum_{ijab} t^{ab}_{ij} \hat{a}^\dagger_a \hat{a}^\dagger_b \hat{a}_j \hat{a}_i, \quad (8)$$

where $t^a_i$ and $t^{ab}_{ij}$ are cluster amplitudes associated with singles and doubles excitations. We adopt the standard notation, where the indices $i, j, k, \cdots$ refer to occupied, $a, b, c, \cdots$ to virtual, and $p, q, r, \cdots$ to general molecular spin-orbitals; the separation between the occupied and virtual orbital spaces is determined by the choice of the reference determinant ($\Phi_0$). The cluster amplitudes satisfy the CC equations

$$\langle \Phi_v | \hat{H} | \Phi_0 \rangle = 0, \quad (9)$$

where, for CCSD, $\Phi_v$ spans the singles and doubles excitation manifold. $\hat{H} = e^{-\hat{T}} \hat{H} e^{\hat{T}}$ is the similarity-transformed Hamiltonian.

The EOM-CCSD target state $n$ is expressed using the (right) EOM operator $\hat{R}^n$ acting on the CCSD wave function as follows:

$$|\Psi_n\rangle = \hat{R}^n \Psi_{\text{CC}} = \hat{R}^n e^{\hat{T}}|\Phi_0\rangle. \quad (10)$$

The choice of target states determines the set of configurations spanned by the EOM-CCSD operator $\hat{R}^n$ (also called the sector of the Fock space). Within EOM-CCSD for excitation
energies (EOM-EE-CCSD), $\hat{R}^n$ spans the reference and valence singly and doubly excited configurations:

$$\hat{R} = r_0 + \sum_{ia} r_i^a a^\dagger_a \hat{a}_i + \frac{1}{4} \sum_{ijab} r_{ij}^{ab} a^\dagger_a \hat{a}_i^\dagger a^\dagger_b \hat{a}_j \hat{a}_i,$$

(11)

where $r_0$, $r_i^a$, and $r_{ij}^{ab}$ are the EOM-EE-CCSD amplitudes satisfying the following eigenvalue equation:

$$\hat{H}\hat{R}^n\lvert\Psi_0\rangle = E_n\hat{R}^n\lvert\Psi_0\rangle.$$

(12)

The state with $r_0 = 1$ and $r_i^a = r_{ij}^{ab} = 0$ \forall \{i,j,a,b\} is the reference CCSD state ($n = 0$, usually but not always the ground state) and the rest ($n > 0$) are EOM-CC target states (e.g., valence excited states). In a similar fashion, target core-excited states can be computed from the EOM-CCSD eigenvalue equation by restricting the EOM-CCSD operator to accessing only the core-excited configurations only. In the core–valence separated EOM-EE-CCSD approach with frozen–core approximation (fc-CVS-EOM-EE-CCSD), this corresponds to:

$$\hat{R} = \sum_{ia} r_i^a a^\dagger_a \hat{a}_I + \frac{1}{2} \sum_{ljab} r_{lj}^{ab} a^\dagger_a \hat{a}_j \hat{a}_I + \frac{1}{4} \sum_{IJab} r_{IJ}^{ab} a^\dagger_a \hat{a}_b \hat{a}_J \hat{a}_I,$$

(13)

where capital labels $I$ and $J$ denote core orbitals. In contrast to the CVS scheme employed by Coriani and Koch, in the fc-CVS-EOM-EE-CCSD approach of Vidal et al., the core orbitals are frozen (uncorrelated) in the ground-state calculation.

Within the non-Hermitian EOM-CC theory, the left and right eigenfunctions of $\hat{H}$ are not complex conjugates of each other. The left eigenfunctions are obtained from the left EOM-CC eigenvalue equation,

$$\langle\Psi_0|\hat{L}^n\hat{H} = \langle\Psi_0|\hat{L}^n E_n,$$

(14)

where $\hat{L}^n$ is the left EOM-CC operator. The left EOM-EE-CCSD and fc-CVS-EOM-EE-
CCSD (de-excitation) operators are defined according to
\begin{equation}
\hat{L} = l_0 + \sum_{ia} l_i^a \hat{a}_i^\dagger \hat{a}_a + \frac{1}{4} \sum_{ijab} l_{ij}^{ab} \hat{a}_i^\dagger \hat{a}_j^\dagger \hat{a}_b \hat{a}_a
\end{equation}
and
\begin{equation}
\hat{L} = \sum_{ia} l_i^a \hat{a}_i^\dagger \hat{a}_a + \frac{1}{2} \sum_{Ijab} l_{IJ}^{ab} \hat{a}_I^\dagger \hat{a}_J^\dagger \hat{a}_b \hat{a}_a + \frac{1}{4} \sum_{IJab} l_{IJ}^{ab} \hat{a}_I^\dagger \hat{a}_J^\dagger \hat{a}_b \hat{a}_a,
\end{equation}
respectively. In the context of the left ground-state reference, \( \hat{L} \) is historically referred to as \( 1 + \hat{\Lambda} \) according to
\begin{equation}
\langle \tilde{\Psi}_n | = \langle \Phi_0 | (1 + \hat{\Lambda}) e^{-\hat{T}} ; \quad \hat{\Lambda} = \sum_{\mu \neq \Phi_0} \lambda_\mu \hat{\tau}_\mu .
\end{equation}
Note that \( l_0 = 0 \) for the EOM-CC states and \( l_0 = 1 \) for the reference (\( n = 0 \)) CCSD state.

In the literature, the ground-state left amplitudes are also often denoted as \( \tilde{t}_\mu \) to highlight that they are Lagrangian multipliers of the right amplitudes \( t_\mu \).\(^{25}\)

The EOM-CC left and right eigenvectors can be normalized to form a biorthonormal set, such that
\begin{equation}
\langle \hat{L}^n | \hat{R}^m \rangle = \delta_{nm}
\end{equation}
where \( \delta_{nm} \) is the Kronecker delta.

### 2.3 EOM-CCSD ECD rotatory strengths: Density-based implementation

Within the non-Hermitian coupled-cluster theory, the right (\( \langle \hat{O}^J \rangle \)) and the left (\( \langle \hat{O}^n \rangle \)) transition moments between different states are not equal; here, \( \hat{O} \) is a general operator. As a result, the expressions for rotatory strengths (and, similarly, for the oscillator strengths)
involve explicit symmetrization with respect to complex conjugation as follows:

\[
R_{ng}^{\text{lg}} = \frac{1}{2} \sum_{\alpha} \left\{ \frac{1}{2} \left( \langle \hat{r}_{\alpha}^{nf} \hat{L}_{\alpha}^{fn} \rangle + \langle \hat{r}_{\alpha}^{nf} \hat{L}_{\alpha}^{fn} \rangle^* \right) \right\} = \frac{1}{4} \sum_{\alpha} \left\{ \langle \hat{r}_{\alpha}^{nf} \hat{L}_{\alpha}^{fn} \rangle - \langle \hat{L}_{\alpha}^{fn} \hat{r}_{\alpha}^{nf} \rangle \right\} \quad (19)
\]

\[
R_{nf}^{\text{vg}} = -\frac{1}{4\omega_{nf}} \sum_{\alpha} \left\{ \langle \hat{p}_{\alpha}^{nf} \hat{L}_{\alpha}^{fn} \rangle + \langle \hat{L}_{\alpha}^{nf} \hat{p}_{\alpha}^{fn} \rangle \right\}. \quad (20)
\]

The right and left transition moments are given by

\[
\langle \hat{O}^f \rangle = \langle \Phi_0 \hat{L}^f e^{-\hat{T}} | \hat{O} | e^T \hat{R}^n \Phi_0 \rangle = \sum_{pq} \gamma_{fpq}^{f-n} O_{pq} \quad (21)
\]

and

\[
\langle \hat{O}^n \rangle = \langle \Phi_0 \hat{L}^n e^{-\hat{T}} | \hat{O} | e^T \hat{R}^f \Phi_0 \rangle = \sum_{pq} \gamma_{npq}^{n-f} O_{pq}, \quad (22)
\]

respectively. Here, \( \gamma_{fpq}^{f-n} \) and \( \gamma_{npq}^{n-f} \) are the right and left reduced transition one-particle density matrices (1PDMs), respectively; \( O_{pq} \) is the matrix representation of the operator in the MO basis (i.e., property integrals). The programmable expressions for the transition 1PDMs between different EOM-CCSD target states used in this study are presented in the Appendix and also reported elsewhere.\(^{20,32}\)

### 2.4 EOM-CCSD rotatory strengths: Alternative implementation

Within coupled-cluster response theory (CCRSP), the left and transition moments are often expressed as contractions of fundamental CC building blocks\(^{24,25}\)

\[
\langle \hat{O}^0 \rangle_{\text{rsp}} = \sum_{\mu} f_{\mu}^{\text{C}} O_{\mu}, \quad (23)
\]

\[
\langle \hat{O}^f \hat{O}^0 \rangle_{\text{rsp}} = \sum_{\mu} \eta_{\mu}^C r_{\mu}^{\text{C}} + \sum_{\mu} M_{\mu}^f (\omega_f) \xi_{\mu}^C, \quad (24)
\]

\[
\langle \hat{O}^k \hat{O}^j \rangle_{\text{rsp}} = \sum_{\mu\nu} (k_{\mu} A_{\mu\nu}^C r_{\nu}^{\text{C}}) + \sum_{\mu} \tilde{N}_{\mu}^{kj} (\omega_k, \omega_j) \xi_{\mu}^C, \quad (25)
\]
where

\[
\xi^{\text{O}}_{\mu} = \langle \Phi_{\mu} | e^{-\hat{T}} \hat{O} e^{\hat{T}} | \Phi_0 \rangle, \tag{26}
\]
\[
A^{\text{O},\text{rsp}}_{\mu\nu} = \langle \Phi_{\mu} | [e^{-\hat{T}} \hat{O} e^{\hat{T}}, \hat{\tau}_{\nu}] | \Phi_0 \rangle, \tag{27}
\]
\[
\eta^{\text{O},\text{rsp}}_{\mu} = \langle \Phi_0 | [e^{-\hat{T}} \hat{O} e^{\hat{T}}, \hat{\tau}_{\mu}] | \Phi_0 \rangle + \sum_{\lambda} \bar{t}_\lambda A^{\text{O},\text{rsp}}_{\lambda\mu}, \tag{28}
\]

and \( \tilde{M}_f(\omega_f) \) and \( \tilde{N}_{\mu}^{kj}(\omega_k, \omega_j) \) are excited-state Lagrangian multipliers.\(^\text{25}\) Greek indices refer to the excitation levels (here either singles or doubles) and \( \hat{\tau}_{\mu} \) is the corresponding excitation operator. Note the superscripts \( f \) and \( k \) to distinguish between the excited state amplitudes \( r_{\mu} \) and \( l_{\mu} \).

Analogous expressions can be written for the EOM-CCSD transition moments:\(^{35,50,57}\)

\[
\langle \hat{O}^f \rangle = \sum_{\mu} l^f_{\mu} \xi^{\text{O}}_{\mu}, \tag{29}
\]
\[
\langle \hat{O}^0 \rangle = \sum_{\mu} \eta^{\text{O}}_{\mu} r^{f}_{\mu}, \tag{30}
\]
\[
\langle \hat{O}^{kj} \rangle = \sum_{\mu\nu} l^k_{\mu\nu} A^{\text{O}}_{\mu\nu} r^{j}_{\nu} - \left( \sum_{\rho} \bar{t}_\rho \xi^{\text{O}}_{\rho} \right) \delta_{kj} - \left( \sum_{\rho} l^k_{\rho} \xi^{\text{O}}_{\rho} \right) \left( \sum_{\rho} \bar{t}_\rho r^{j}_{\rho} \right), \tag{31}
\]

where the EOM-CCSD building blocks are:\(^{35}\)

\[
\eta^{\text{O}}_{\mu} = \langle \Phi_{0} | e^{-\hat{T}} \hat{O} e^{\hat{T}} | \Phi_{\mu} \rangle + \sum_{\lambda} \bar{t}_\lambda \langle \Phi_{\lambda} | e^{-\hat{T}} \hat{O} e^{\hat{T}} | \Phi_{\mu} \rangle - \langle \hat{O} \rangle_{\text{CC}} \bar{t}_{\mu}, \tag{32}
\]
\[
\eta^{\text{O}}_{\mu} = \langle \Phi_{0} | e^{-\hat{T}} \hat{O} e^{\hat{T}} | \Phi_{\mu} \rangle + \sum_{\lambda} \bar{t}_\lambda A^{\text{O}}_{\lambda\mu}, \tag{33}
\]
\[
A^{\text{O}}_{\lambda\mu} = \langle \Phi_{\lambda} | e^{-\hat{T}} \hat{O} e^{\hat{T}} | \Phi_{\mu} \rangle - \delta_{\lambda\mu} \langle \Phi_{0} | e^{-\hat{T}} \hat{O} e^{\hat{T}} | \Phi_{0} \rangle, \tag{34}
\]

and \( \left( \sum_{\rho} \bar{t}_\rho r^{j}_{\rho} \right) = -r^{j}_{0}. \)
Simple manipulations connect the CC response and the EOM building blocks

\[ \eta^\text{O} = \eta^\text{O, rsp} + \sum_{\mu > \nu} \tilde{t}_\mu \left( \Phi^\mu | \hat{\tau}_\nu e^{-\hat{T} \hat{O} e^{\hat{T}}} | \Phi^0 \right) - \left( \sum_{\rho} \tilde{t}_\mu^\rho \eta^\rho \right) \tilde{t}_\mu, \]  
\[ A^\text{O, C} = A^\text{O, rsp} + \langle \Phi^\mu | \hat{\tau}_\nu e^{-\hat{T} \hat{O} e^{\hat{T}}} | \Phi^0 \rangle (1 - \delta_{\mu \nu}). \]  

### 2.5 EOM-CCSD ECD spectra from the damped optical rotation tensor

For isotropic samples, ECD spectra can also be obtained directly from the absorbing component of the electric dipole-magnetic dipole optical rotation tensor, \( G'_{\alpha \beta}(g) \),

\[ \Delta \epsilon \propto -\frac{1}{3} \omega \mu \lambda \lambda G'_{\alpha \alpha}(g), \]  

which, within damped linear response theory, corresponds to

\[ G'_{\alpha \beta}(-\omega, \omega) \propto \Re \langle \hat{m}_\alpha ; \hat{\mu}_\beta \rangle_\omega + i \gamma \]  

in the length gauge. In the velocity gauge, this becomes the imaginary part of the optical rotation tensor for the linear momentum and the magnetic dipole operator

\[ G'_{\alpha \beta} \propto \Im \langle \hat{m}_\alpha ; \hat{p}_\beta \rangle_\omega + i \gamma, \]  

where \( \gamma \) is the damping term, which can be regarded as a common inverse lifetime of all excited states.

The general sum-over-states expression of a complex linear response function in a.u. is

\[ \langle \hat{O}_1 ; \hat{O}_2 \rangle_\omega + i \gamma = \sum_{k>0} \left\{ \frac{\langle \hat{O}_1^{0k} \rangle \langle \hat{O}_2^{k0} \rangle}{E_k - E_0 + \omega + i \gamma} + \frac{\langle \hat{O}_2^{0k} \rangle \langle \hat{O}_1^{k0} \rangle}{E_k - E_0 - \omega + i \gamma} \right\} \]  

12
In our present implementation, we adopted an asymmetric form for the CC damped rotation tensor. In the vector-matrix notation of Section 2.4 this reads

\[
\langle\langle X; x \rangle\rangle_{\omega+i\gamma} = \frac{1}{2} \hat{C}^{\pm\omega} \left\{ \bar{t}^x(\omega+i\gamma)\xi^x + \eta^x t^x(\omega+i\gamma) \right\}, \tag{41}
\]

where the permutator \( \hat{C}^{\pm\omega} \) only acts on the real frequency \( \omega \). Eq. (41) requires the solution of the damped linear equations for the right response amplitudes, \( t^x(\omega+i\gamma) \), and the left response multipliers \( \bar{t}^x(\omega+i\gamma) \) of the same operator \( x \)

\[
\begin{align*}
[A - (\omega + i\gamma)1] t^x(\omega+i\gamma) &= -\xi^x, \tag{42} \\
\bar{t}^x(\omega+i\gamma) [A + (\omega+i\gamma)1] &= -\eta^x. \tag{43}
\end{align*}
\]

where \( A \) is a sub-block of similarity transformed Hamiltonian matrix, often referred to as the CC Jacobian

\[
\bar{H}_{\mu\nu} = \begin{pmatrix}
E_{CC} & \eta_{\nu} \\
\Omega_{\mu} & A_{\mu\nu}
\end{pmatrix}
\]

(44)

Note that, in contrast to the case of CC linear response, the EOM equations for the response multipliers are decoupled from those for the response amplitudes. The damped response equations Eqs. (42)–(43) can be solved iteratively using damped variants of standard procedures such as the Direct Inversion in the Iterative Subspace (DIIS) or generalized Davidson.

3 Computational details

We implemented the above expressions for the calculation of the electronic circular dichroism within the EOM-CCSD approach in Q-Chem. The theory is applicable for valence excitations (EOMEE-CCSD) and well as core excitations (fc-CVS-EOMEE-CCSD) of both ground and excited states. py-CCRSP was used for code profiling and testing and for
the CPP calculations. To illustrate the methodology, we computed various spectra for the following systems (shown in Figure 1): \textit{R}-methyloxirane, \textit{1R}-norcamphor, \textit{S}_\text{a}- and \textit{R}_\text{a}-binol, \textit{L}-alanine, and \textit{L}-chloroethanol.

![Figure 1: Structures of molecules considered in this work: a) \textit{R}-methyloxirane, b) \textit{1R}-norcamphor, c) \textit{S}_\text{a}-binol, d) \textit{R}_\text{a}-binol, e) \textit{L}-alanine, and f) \textit{L}-chloroethanol. Atom labeling is given when relevant for the discussion of the results.](image)

\textit{R}-methyloxirane has previously been used as model system to computationally investigate excited-state CD (ES-ECD) at the TDDFT level of theory by Rizzo and Vahtras.\textsuperscript{44} We use the same geometry as in Ref. 44. We investigated basis-set effects by employing various correlation-consistent basis sets\textsuperscript{68–70} with different augmentation and polarization levels, as well as Pople’s 6-311++G** set. The EOM-CCSD results were compared to linear response theory results obtained with Dalton.\textsuperscript{71} The ES-ECD for the 15 lowest valence excitations was computed with d-aug-cc-pVDZ and for the eight lowest states with d-aug-cc-pVTZ.

For consistency, we used the ground-state geometry of (\textit{1R})-norcamphor optimized at the DFT/CAM-B3LYP/aug-cc-pVTZ level from Ref. 45. Ground-state and \textit{S}_1 excited-state absorption and CD spectra including 12 excited states were simulated using the d-aug-cc-pVDZ basis set.

For binol, we considered two structures, namely the \textit{S}_0 structure of the \textit{S}_\text{a}-isomer from Ref. 43 and the \textit{S}_0 structure of the \textit{R}_\text{a}-isomer of Ref. 45 (including a symmetrized variant). Calculations with the former structure were carried out using the 6-31G* basis set, as was done in Ref. 43 at the TDDFT level. The 10 lowest excitations in each irrep (\textit{A} and \textit{B} irreps of \textit{C}_s point group) were computed. The ES-ECD was calculated for the two lowest excited states in each irrep. In the case of the \textit{R}_\text{a}-isomer, we used the cc-pVDZ basis set,
as was done in Ref. 45 at the ADC(2) and TDDFT levels, and computed the GS-ECD from the 10 lowest excitations in each irrep, as well as the ES-ECD of the four lowest states. These calculations were carried out using single-precision implementation of CCSD and EOM-CCSD equations.\textsuperscript{72}

In the X-ray regime, (XCD) we considered, once again, methylxirane, employing the same geometry for the $R$-isomer as above, as well as the MP2/cc-pVTZ geometry from Ref. 73. We considered different basis sets, namely augmented Dunning basis sets as well as standard and uncontracted variants of the 6-311++G** set, which have been shown to be well-suited for calculations of XAS spectra.\textsuperscript{74} The core-excited states of the carbon $K$-edge were computed using the fc-CVS-EOMEE-CCSD scheme in Q-Chem. The linear response results were obtained with the CVS-CCSD scheme implemented in Dalton.\textsuperscript{56,71,75}

The XCD of $L$-alanine was investigated with the d-aug-cc-pVDZ basis set. We computed the $K$-edges of carbon, nitrogen, and oxygen for the neutral and zwitterionic forms of the amino acid. Coordinates for the neutral form were taken from Ref. 73; the zwitterion structure was obtained from neutron diffraction experiment in Ref. 76.

Finally, inspired by Ref. 77, we considered the chlorine $L$-edge of chloroethanol. XAS and XCD spectra with and without spin–orbit coupling (SOC) were computed with the 6-311(2+,+)G** basis uncontracted to describe the $2p$ orbitals of the Cl atom.\textsuperscript{66}

Unless otherwise specified, the gauge origin was located at the center of charge (coc), which is the default setup in Q-Chem. RSP-CCSD and TDDFT/CAM-B3LYP results used for comparison were obtained using DALTON.\textsuperscript{71,75}
4 Results and discussion

4.1 UV–vis ECD spectra

4.1.1 Methyloxirane

Methyloxirane (also known as propylene oxide or 1, 2-epoxypropane) is a popular test case for calculations of optical activity both in the UV–vis and X-ray regimes. This chiral organic molecule is small enough for high-level \textit{ab initio} calculations and is also manageable for experimental UV absorption and circular dichroism studies available in the gas phase.\textsuperscript{78,79}

We investigated the basis-set dependence of the ground-state one-photon absorption (OPA) and ECD obtained from the EOM-CCSD and CCSD-RSP calculations. The resulting spectra are shown in Figure 2. The corresponding numerical data for the rotatory and oscillator strengths are listed in Tables S1 and S2 in the SI, respectively. We observe good agreement between the EOM-CCSD rotatory and oscillator strengths obtained from the length and the velocity formulations for all considered basis sets. The maximum absolute difference between the EOM rotatory strength in the two gauges is 0.004 a.u. for the Pople basis set, 0.003 a.u. for aug-cc-pVDZ and d-aug-cc-pVDZ, and 0.002 a.u. for aug-cc-pVTZ and d-aug-cc-pVTZ. Increasing the basis-set size does not seem to affect this difference, at least not within the sequence of the bases investigated here. However, the quality of the basis moderately affects the peak positions and intensities, although the overall sign pattern is the same for all basis sets. The EOM-CCSD and RSP-CCSD results are nearly identical; the only noticeable difference seen for the d-aug-cc-pVTZ case is due to a smaller number of roots considered in the RSP calculation.

Next, we examined the performance of EOM-CCSD for calculation of excited-state CD (ES-ECD), using the doubly augmented double-zeta and triple-zeta basis sets, see Tables S3 and S4 in the SI. We considered only the transitions from the four lowest excited states to the eight lowest states. We immediately note larger deviations between the two gauges, now up to 0.04 a.u. in absolute values of the rotatory strength for d-aug-cc-pVDZ and 0.03
Figure 2: \textit{R}-methyloxirane. EOM-CCSD (left) and CCSD-RSP (right) basis-set study of ground-state transition properties. ECD (top) and OPA (bottom) spectra in length (solid line) and velocity (dashed line) gauge. Lorentzian broadening with HWHM $= 0.124$ eV.

It is remarkable that for a few high-lying excited-state transitions, a negative value of the velocity-gauge oscillator strength is obtained with both basis sets, in one case ($6 \rightarrow 7$) also accompanied by a difference in sign and size of the rotatory strengths (see Tables S3 and S4).

The simulated absorption and CD spectra of the ground state ($S_0$) and first two excited states ($S_1$ and $S_2$) obtained using the d-aug-cc-pVDZ basis set are shown in Figure 3, along with the results of TDDFT/CAM-B3LYP calculations. The experimental spectra for the ground state taken from Ref. 79 are also shown. The CAM-B3LYP excitation energies are slightly lower than the CCSD ones for all three cases; otherwise, the spectra are qualitatively rather similar. No energy shift was applied. The $S_0$ spectra agree reasonably well with the available experimental results for \textit{R}-methyloxirane, at least up to 8 eV. Above 8 eV, both computational methods yield a much more intense negative peak than in the experiment, in line with previous findings.\cite{19} Adding Rydberg-type functions to the basis set partly re-distributes the intensity over several electronic transitions, yet the third band remains more intense than observed experimentally (see S2 in the SI). Note that the first ionization energy is 10.13 eV (EOM-IP-CCSD/d-aug-cc-pVDZ).

Concluding this section, Figure 4 shows proof-of-principle illustration of our damped-response implementation of the ECD cross section. The trace of the real component of
the optical rotation tensor in the length gauge was computed at the frequency grid points indicated by the red X marks and then splined to yield the cross section. The spectrum is practically identical to the one obtained from a Lorentzian broadening of the individual rotatory strengths from the standard EOM-CCSD theory.

4.1.2 Norcamphor

Moving on to a larger organic molecule, we considered the bicyclic ketone (1R)-norcamphor (a.k.a. 2-norbornanone), which has also been investigated in previous theoretical\textsuperscript{19,45,80} and experimental\textsuperscript{80} gas-phase studies. The ground-state ECD spectra, see right panel of Figure 5, from EOM-CCSD and TDDFT/CAM-B3LYP are rather similar in peak positions, intensities, and sign patterns up to 7.7 eV. Above 7.7 eV, CAM-B3LYP yields a strong negative band, whereas CCSD has a weak positive peak. The corresponding ADC(3) spectrum of Ref. 45, also shown in Figure 5, is blue-shifted and has a negative peak at around 8 eV. Most electronic transitions have been previously assigned ns and np characters.\textsuperscript{80} When Rydberg-type functions are included, a negative peak at around 8.0 eV emerges, but two positive peaks are still present between 7 and 8 eV; see Figure S3. The experimental ab-

Figure 3: \textit{R}-methyloxirane. Absorption (bottom panel) and CD (upper panel) spectra for \textit{S}_0, \textit{S}_1, and \textit{S}_2. EOM-CCSD (15 states) and TDDFT/CAM-B3LYP (15 states) with the d-aug-cc-pVDZ basis set. Dashed line is velocity gauge. Experimental spectra (dotted line) were digitized from Ref. 79. Lorentzian broadening with HWHM = 0.124 eV.
Figure 4: R-Methyloxirane. EOM-CCSD/aug-cc-pVDZ ground-state ECD (sticks and broadened spectrum, HWHM = 0.005 a.u.) and CPP (grid points and cubic fit) in length gauge.

The shapes of the ECD spectra of the lowest electronic excited state are again similar for all three methods up to 3.5 eV, with an overall blue shift of the ADC(3) spectrum. We note here that we recomputed the CAM-B3LYP ES-ECD spectrum and obtained intensity twice as large as the one reported by Scott et al.\(^{45}\) Above 3.5 eV the three methods clearly differ, because a different number of final states was computed by the three methods (i.e., only 10 states were reported in the ADC(3) study). CAM-B3LYP yields an additional strong negative band and CCSD yields a weaker positive one, reflecting the differences observed in
the ground-state spectrum. The raw data of Figure 5 are given in Tables S7 and S9 in the SI.

Figure 5: (1R)-norcamphor. Excited-state (ES) and ground-state (GS) ECD (upper panels) and absorption (bottom panels) spectra from EOM-CCSD (this work, 16 states), TDDFT/CAM-B3LYP (this work, 15 states), and ADC(3) (Ref. 45, 10 states) using the d-aug-cc-pVDZ basis. Dashed lines correspond to velocity gauge. The EOM-CCSD and TDDFT/CAM-B3LYP OPA and ESA spectra are shown in the two bottom panels. Experimental spectra (dotted line) from Ref. 80. Lorentzian broadening with HWHM = 0.124 eV.
Binol is an axially chiral compound composed of two naphthol moieties connected by a single bond. The two atropoisomers are labeled either $R_a/S_a$ or $P/M$. Here we considered two structures: the ground-state optimized structure ($S_0$) of the $S_a$ enantiomer taken from Ref. 43, where the dihedral angle between the two moieties is $\approx 90^\circ$ (“orthogonal” conformation) and the $S_0$ of the $R_a$ enantiomer taken from the recent computational study at the ADC level of theory, which is a DFT/CAM-B3LYP/cc-pVTZ optimized structure. A symmetrized version of the latter was also employed for computational efficiency, after validating that the two structures yield equivalent results. The steady-state and time-resolved ECD spectra of binol have been measured by Hache and coworkers in different solvents. Niezborala and Hache report the steady-state spectrum of $R$-binol in ethanol, whereas the one of $S$-binol in cyclohexane was reported by Schmid et al.

Electronic transitions in binol are often labeled using the Platt nomenclature relative to the 2-naphthol precursor. The first two bright transitions are called $L_a$ and $L_b$, where $L_a$ is the more intense one (HOMO$\rightarrow$LUMO character). The relative ordering of the $L_a$ and $L_b$ transitions is method dependent. At the EOM-CCSD/6-31G(d) level, the lowest excited state of isolated 2-naphthol is $L_b$ at 4.45 eV ($f_{lg} = 0.02$), whereas the second, at 5.39 eV ($f_{lg} = 0.06$), is $L_a$. They are followed by a third state at 6.28 eV ($f_{lg} = 0.04$). An intense $B$ state (according to Platt’s notation) comes as fourth at 6.60 eV ($f_{lg} = 1.29$). With the cc-pVDZ basis, we obtain 4.40 eV (0.02); 5.26 eV (0.047); 6.19 eV (0.033); 6.49 eV (1.21). At the TDDFT level, the ordering of the two states is reversed, with $L_a$ dropping below $L_b$.

This different ordering carries over to binol. Exciton pairs are obtained from the combination of the $L_a$ and $L_b$ states of the two naphthol moieties. The lowest EOM-CCSD exciton pair comprises the two states, respectively at 4.42 eV ($f_{lg} = 0.05$) and 4.44 eV ($f_{lg} = 0.012$) in the 6-31G(d) basis, which are labelled $L_b$-1 (of B symmetry) and $L_b$-2 (of A symmetry). The states of the second exciton pair, of $L_a$ character, are at 5.29 eV ($f_{lg} = 0.20$, $L_a$-1, B symmetry) and 5.33 eV ($f_{lg} = 0.01$, $L_a$-2, A symmetry). At the TD-DFT/CAMB3LYP
level the first exciton pair corresponds to $L_a$-1 (4.4359 eV, $f^{lg} = 0.1436$) and $L_a$-2 (4.4825 eV, $f^{lg} = 0.0200$); the second pair is $L_b$-1 (4.8550 eV; $f^{lg} = 0.1003$) and $L_b$-2 (4.8649 eV; $f^{lg} = 0.0001$). We summarize in Table 1 our results for energies, oscillator strengths, and rotatory strengths of the $L_a$ and $L_b$ states of the $R_a$ and $S_a$ structures considered here, using EOM-CCSD and CAM-B3LYP and two different basis sets. Thus, in comparing the

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ES-ECD of the excited states of binol for the two methods, one should keep in mind that the character of the first excited state is different at the EOM-CCSD and CAM-B3LYP levels.

Figure 6 shows the EOM-CCSD GS- and ES-ECD spectra of $S_a$-binol, together with the experimental data from Ref. 43 (measured in cyclohexane). The ES-ECD were computed for all four lowest excitations in Table 1, plus states 5A and 5B ($B_b$-1 and $B_b$-2). Raw data are given in Tables S10–S17. Table S18 presents the data from the gauge dependence study.
Table S11 contains natural transition orbitals (NTOs) and transition properties of the four lowest GS transitions. The corresponding data for the \( R_a \) conformation are given in the SI.

The EOM-CCSD GS-ECD spectra of binol agree well with the experimental results of Refs. 81 (\( R_a \)) and 43 (\( S_a \)), despite the neglect of solvent effects. Curiously, the split-band at 6 eV is present in the experimental spectrum of \( R_a \) (reproduced by the calculations), is absent in the experimental spectrum of \( S_a \). Both CAM-B3LYP (this work, as well as Ref. 43) and ADC(2) (Ref. 45) also agree with the experimental data.

Turning our attention to the ES-ECD spectra, the first general observation is that we could not reproduce the CAM-B3LYP results for the ES-ECD of \( S_a \)-binol reported in the SI of Ref. 43. Our CAM-B3LYP results are shown in Figure S1. Since our results for methyloxirane and norcamphor are consistent with CAM-B3LYP data reported by other authors, we suspect there may be a problem in the results reported by Schmid et al.

According to our calculations, the A and B symmetry excitation pairs have ES-ECD that are almost mirror images of each other across the \( x \)-axis. In the lower-energy region, the agreement between the length (origin at center-of-charge) and velocity gauges is somewhat poor. Our TD-DFT results show the same trend, at least for the \( S_3 \) (\( L_b^{-1} \)) and \( S_4 \) (\( L_b^{-2} \)) transitions.

Given the poor agreement between length and velocity ES rotatory strengths at lower energies, we examined the gauge-origin dependence of the ES-ECD of the \( L_b^{-1} \) and \( L_b^{-2} \) states of \( S_a \)-binol. The gauge origin was manually located at an arbitrary carbon atom. The data reported in Table S18 shows that it is only the rotatory strengths for transitions between different symmetries that are affected by the change of gauge origin. Overall, the effect of this shift of origin is fairly small, and it did neither significantly improve nor worsen the agreement between the two gauges. It could be interesting to test the effect of locating the gauge origin at the center-of-mass (or -charge) of the two naphthol moieties, respectively, but we have not pursued it here.
Figure 6: $S_a$-binol. Ground-state (label GS) and excited-state ($L_a$, $L_b$ and $B_b$ states) absorption (bottom panels) and ECD (upper panels) spectra in the length gauge computed at the EOM-CCSD/6-31G(d) level. Twenty excitations were considered in each irrep of the $C_2$ point group. Experimental values from Ref. 43. The CCSD energies have been shifted by $-0.9$ eV to align the main peaks in the experimental OPA spectrum. Lorentzian broadening with HWHM = 0.2 eV.
4.2 X-ray Circular Dichroism

4.2.1 Methyloxirane

The experimental XCD spectrum of methyloxirane in vapor phase at the carbon $K$-edge has been reported by Turchini et al.,$^{82}$ and by Alberti et al.$^{83}$ Computational results at different levels of theory have also been reported.$^{33,84,85}$ A later study by Piancastelli et al.$^{86}$ focused on the non-chiral core-level photoelectron spectroscopy, XAS at both C and O $K$-shells, resonant Auger and normal Auger spectroscopy.

As explained by Turchini et al., a complicating factor in XCD measurements is that, to the first approximation, the magnetic dipole transition element of $K$-edge excitations is zero, since magnetic dipole transitions arising from $s$ orbitals are forbidden. Consequently, the only possible source of magnetic dipole intensity involves $1s$–$np$ (particularly $1s$–$2p$) mixing. Theoretical studies$^{84,87}$ had indicated that this mixing could be sufficient to produce observable CD for simple organic molecules.

In the experimental measurement of Turchini et al., a nonlinear sloping background was present in the XCD of the two isomers, as well as their racemate $[S(-)+R(+)]$. To obtain a monosignate signal, the measured spectra of the two enantiomers were halved and algebraically subtracted. This yielded the composite spectrum of $S(-)\text{-methyloxirane}$ without the baseline background, since $S(-)_c = S(-)_m - \frac{1}{2}[S(-) + R(+)]_m = \frac{1}{2}[S(-)_m - R(+)_m]$. Turchini et al. assigned the CD spectrum to the methyne and methylene carbon atoms (i.e. the oxygen-bound ones), and suggested that the methyl carbon’s CD signal would essentially cancel out to zero because of the small energy splitting and the resolution of the experiment. A photoemission experiment revealed three binding energies of 291.2, 292.2, and 292.6 eV. With the d-aug-cc-pVDZ basis set, the three lowest ionization energies are 293.5, 294.8, and 294.9 eV, whereas with the d-aug-cc-pVTZ basis set the IEs are 291.3, 292.6, and 292.7 eV.

The simulation of the XCD (and XAS) spectra of methyloxirane proved to be rather
challenging. We explored different basis sets, including d-aug-cc-pVDZ and d-aug-cc-pVTZ on all atoms, aug-cc-pVTZ on the C and O atoms and aug-cc-pVDZ on the H atoms, as well as Pople’s 6-311++G** and 6-311(2+,+)G**, also uncontracting the inner functions. Two different optimized structures were also considered. Comparisons between EOM-CCSD and CCSD-RSP results were carried out. Note that we report results for the $R$ enantiomer, whereas the XCD spectra of Ref. 11 refer to the $S$ enantiomer.

The simulated spectra for two different structures (by EOM-CCSD and CC-RSP using the 6-311++G** basis set) are shown in Figure 7, along with experimental results. EOM and RSP results are consistent, though with a few noticeable differences in XCD intensity in the length gauge. The moderate differences between velocity and length XCD results depend, to a certain extent, on the absolute size of the rotatory strength, as transitions with extremely small rotatory strengths sometimes have oppositely signed strengths.

The spectra stem from a number of closely lying excitations, whose rotatory strengths may significantly vary with the basis set. The spectra computed for the two optimized equilibrium geometries considered here show large differences. For the structure labeled ‘Geometry 2’ in Figure 7, which is the one from Ref. 44 also used for ESCD, the excited states are less clustered, so the XAS spectral features are more smeared out. The most dramatic difference is, however, observed in the XCD spectra, where the first negative and weak broad band is replaced by a positive one, and the second negative one is far more pronounced. Changing the basis also affects the spectra, as illustrated in Figure 8 for Geometry 1 (the MP2 optimized structure from Ref. 73).

The computed XAS spectra agree with the experiments, with overall energy shifts that vary between 0.06 eV and 2.3 eV, depending on the basis set. The shift was computed with respect to the first peak of the spectrum of Piancastelli et al.. Clearly identifiable XCD spectral patterns are reproduced by all basis sets and for both geometries, yet strikingly different from the measured ones. Given the results, we cannot conclude whether there is a problem with the measured XCD spectrum or whether the differences are due to deficiencies
in our simulation. One possible reason for discrepancy is that our calculations do not include dynamical sampling of the structures. We note in passing, though, that the computed rotatory strengths reported by Turchini et al.\textsuperscript{82} as taken from Ref. 85 do not appear to correspond to the values tabulated in Ref. 85. Also, to the best of our understanding, the calculation of Ref. 85 were carried out on the \textit{R} isomer and not the \textit{S} one.

It is important to identify the origin of the main peaks in our spectra. Inspection of the NTOs reveals that all three carbon atoms are optically active, in agreement with Ref. 84. In the 6-311++G** basis, the first XAS transition at Geometry 1 (287.5 eV) is due to the methyl C ("C3"), which is dark in XCD. The second and third electronic transitions are responsible for the broad band at around 288.5 eV, i.e. the one changing sign in the two different geometries, and they both come from C3. Three excitations are responsible for the following (negative) band, the one with the largest (negative) \( R \) coming from the non-chiral oxygen-bound C ("C2"). The next, positive, peak (at around 289.5 eV) stems from two transitions, both with positive \( R \), the first one from the chiral center C ("C1"), and the next one from C3. The following negative feature originates mainly from the net balance of two closely lying and oppositely signed sticks at Geometry 1, one positive from C2 and one negative from C1, whereas at Geometry 2 this feature comes from two negative sticks, the strongest due to C2. The positive band at around 290 is a convolution of several transitions, from all three carbon atoms.

The first broad band in the XCD spectra of both geometries is, as mentioned, due to two excitations from only the methyl carbon. This is also the band that changes sign between the two geometries. It is therefore possible that a conformational average (which takes into account the group’s free rotation) would result in a net zero contribution from these C3 excitations. However, at higher energies, excitations from C3 do not appear to cancel out.
Figure 7: R-methyloxirane. Carbon K-edge XCD (upper subpanels) and XAS (bottom subpanels) spectra at two different ground-state optimized geometries. Basis set 6-311++G**. Length gauge pictured in solid line, velocity in dashed. Experimental XAS\textsuperscript{82,83,86} and XCD\textsuperscript{82,83} are also shown. The experimental XCD of the S enantiomer is here reported as its mirror image. Lorentzian broadening with HWHM= 0.2 eV.
Figure 8: $R$-methylxirane. EOM-CCSD Carbon $K$-edge XCD (upper subpanels) and XAS (bottom subpanels) spectra with different basis sets (Geometry 1). The short notation dD stands for d-aug-cc-pVDZ; aT/aD/aD means aug-cc-pVTZ on the C atoms and aug-cc-pVDZ on the remaining atoms. Length gauge pictured in solid line, velocity in dashed. Experimental XAS$^{82,83,86}$ and XCD$^{82,83}$ are also shown. The experimental XCD of the $S$ enantiomer is here reported as its mirror image. Lorentzian broadening with HWHM = 0.2 eV.
4.2.2 L-alanine

The XCD of alanine has been theoretically investigated previously using the STEX approach, the RPA method, as well as at the complex polarization propagator (CPP) approach at the TDDFT/CAM-B3LYP level of theory. A basis set study from Ref. 89 highlighted a strong basis-set dependence of the rotatory strengths, especially for the oxygen and nitrogen K-edges.

Our calculated XAS and XCD spectra at the carbon, oxygen, and nitrogen K-edges are illustrated in Figures 9 (C and O), and 10 (N). Raw spectral data for all figures can be found in the SI. Figure S6 shows the oxygen K-edge XCD at two different gauge origins calculated using different basis sets on the different atoms.

The computed carbon XAS of the neutral structure shows one dominating peak (third excited state) at 290.75 eV ($f_{lk}=0.076$). This corresponds to a transition from the 1s orbital of C2 into the $\pi^*$ of the carboxyl group (see Figure 1 for atom labeling). The majority of the computed states are transitions from the methyl carbon. It is the equivalent C 1s $\rightarrow$ $\pi^*$ transition that dominates the carbon XAS spectrum of the zwitterion at 291.19 eV ($f_{lk}=0.066$).

The oxygen XAS spectrum of the neutral structure has a simple profile with one major peak as the first transition at 536.45 eV ($f_{lk}=0.038$) with the second transition at 3 eV higher in energy. The lowest transition occurs between the 1s orbital of the double-bound oxygen (O2) into the $\pi^*_{\text{COOH}}$ orbital. The second, higher-lying peak, corresponding to the transition from O1 1s orbital into $\pi^*_{\text{COOH}}$, was not reached with the chosen number of computed states. In the zwitterion, the carboxyl group is deprotonated, making the two oxygen atoms nearly equivalent. The oxygen K-edge XAS spectrum of the zwitterion shows more features, with two close-lying peaks of equal magnitude belonging to the O 1s $\rightarrow$ $\pi^*_{\text{COO}^-}$ transitions. The first transition at 536.55 eV ($f_{lk}=0.028$) is from the O1 1s orbital, and the fourth transition (second peak) at 536.89 eV ($f_{lk}=0.029$) originates from the O2 1s orbital. Two almost dark excitations lie between the O 1s $\rightarrow$ $\pi^*_{\text{COO}^-}$ transitions.
The XCD results indicate that all carbon atoms are optically active for both structures, where the majority of transitions are from the methyl carbon 1s orbital. At the nitrogen and oxygen K-edges, pronounced disagreement between the two gauges is observed, especially in the nitrogen case where the velocity gauge rotatory strengths are orders of magnitude smaller than the corresponding strengths in the length gauge. The large difference was attributed to strong origin dependence of the rotatory strengths. To investigate this further, we tried moving the gauge origin to the nitrogen atom and to the center of mass of the carboxyl moiety for the nitrogen and oxygen K-edges, respectively. The nitrogen XCD spectra at the default and shifted origins are depicted in Figure 10. The effect of the origin shift from the center of charge to the N atom is striking, leading to almost complete overlap of the length and velocity gauge XCD spectra. A closer look at the NTOs of the N 1s transitions reveals that these transitions are highly localized on the N atom. We note, however, once more, that the obtained rotatory strengths are very small, so it remains as a general concern that the resulting agreement was merely a consequence of having computed equally small residual errors. As an additional test, we carried out calculations on the neutral D-isomer (geometry produced by mirroring the L-structure in the xy plane). This produced the exact mirror image of the L-alanine N K-edge XCD spectra at both gauge origins.

Applying a gauge shift does have an impact on the oxygen K-edge XCD spectrum too, but it is not as satisfying as for the Nitrogen K-edge XCD. The particle NTOs of the O(1s) transitions are not as localized as for the N 1s, which we try to compensate for by placing the origin in the center of mass of the carboxyl moiety (since the main XAS peaks arise from the O(1s) → π*_{COO} transitions).

Comparing the problematic XCD spectra of this work with the ones simulated with the CPP/CAM-B3LYP/d-aug-cc-pVDZ method reported in Ref. 90, we observe that the N K-edge spectra of the neutral structures have similar features for the two methods, if one considers the gauge-shifted EOM-CCSD one. Meanwhile, the corresponding spectra of the zwitterion are quite different, although slight similarity can be found between the TD-DFT
spectrum and the CCSD one with center-of-charge origin. The computed XAS spectra are similar. The oxygen XCD spectra of the zwitterionic structure are very different at any gauge origins. We note that the conformer considered in Ref. 90 is different from the one of this work.

Figure 9: L-alanine in its neutral and zwitter-ionic forms. fc-CVS-EOM-CCSD/d-aug-cc-pVDZ carbon (left, 10 states) and oxygen (right, 10 states for neutral, 8 states for zwitterion) \( K \)-edge XAS (bottom panels) and XCD (upper panels). Length gauge in solid lines, velocity in dashed. Lorentzian broadening with HWHM = 0.2 eV.
Figure 10: L-alanine in its neutral and zwitter-ionic form. fc-CVS-EOM-CCSD/d-aug-cc-pVDZ Nitrogen K-edge XAS (bottom panels) and XCD (upper and middle panels) spectra. Length gauge in solid lines, velocity in dashed. The top row shows the XCD with gauge origin in the center of charge, the middle row the ones with gauge origin at the position of the nitrogen atom. The spectra in gray (top panels) have been magnified by $10^2$. Lorentzian broadening with HWHM = 0.125 eV.
4.2.3 Chloroethanol

Inspired by Ref. 77, where chloroethanol was used as an illustrative case for XCD of larger chloro-compounds, we considered the chlorine $L_{2,3}$-edge XCD, focusing on the effect of spin–orbit coupling.

At the $L$-edge, the core transitions occur from the $2p$ orbitals of the targeted atom. The degeneracy of the $3p$-orbitals is split by the SOC, which is clearly seen in the X-ray absorption spectra. This splitting is expected to be seen in the XCD spectra as well.

Recently, an extension of EOM-CCSD for computation of X-ray L-edge spectra was reported. The authors employed a state-interaction two-step scheme. First, a set of non-relativistic EOM-CC states is computed and used to evaluate SOCs using Breit-Pauli operator and mean-field approximation of the two-electron part, as described in Refs. 91 and 92. Then the resulting SOC-perturbed Hamiltonain matrix is diagonalized, giving rise to SOC-mixed states. Using the computed transformation between zero-order and SOC-perturbed states, the non-Hermitian electric dipole transition strengths are the transformed into a basis of SO-split states, according to

$$\bar{S}_{nf}^{sf} = U^{\dagger}\langle \mu_{nf}^{f} \rangle \langle \mu_{fn}^{f} \rangle U,$$

where the matrix $U$ contains the eigenvectors of the spin–orbit mean-field Hamiltonian matrix. In the present work the property of interest is the rotatory strength of circular dichroism, so we considered the (anti)symmetrized product of the electric dipole and magnetic dipole transition moments (Eq. (19)):

$$\bar{R}_{nf}^{sf} = \frac{1}{2} U^{\dagger} \left\{ \langle \hat{r}_{nf}^{sf} \rangle \langle \hat{L}_{fn}^{f} \rangle - \langle \hat{L}_{nf}^{sf} \rangle \langle \hat{r}_{fn}^{f} \rangle \right\} U$$

and, likewise, for the velocity formulation (Eq.(20)). Eight singlet excited states were computed for non-relativistic transition properties, and eight singlet and eight (spin-flip) triplet
states were computed for the SOC calculations.

As for the nitrogen and oxygen $K$-edge XCD signals of alanine, we observe strong origin dependence of the $L$-edge XCD spectrum of chloroethanol. The simulated signals in the length gauge with gauge origin in the center of charge are an order of magnitude larger than the velocity gauge equivalents. Shifting the origin from the center of charge to the chlorine atom improves the agreement between the two gauges significantly, although not completely. The numerical values of the computed signals are very small, especially in the velocity gauge and in the length gauge with origin on Cl. Figure 11 shows the non-relativistic and SOC spectra for gauge origin at the Cl atom. Figure S7 in the SI illustrates the gauge effect. Raw data are given in Table S34 in the SI.

![Figure 11: Chloroethanol. fc-CVS-EOM-CCSD/6-311(2+:+)G** non-relativistic (NR, 8 states) and SOC Cl $L_{2,3}$-edge XCD (upper panels) and XAS (bottom panels). Gauge origin on the Cl atom, dashed lines are velocity gauge. Geometry from Ref. 77. Lorentzian broadening with HWHM= 0.124 eV.](image)

The non-relativistic XAS spectrum shows two peaks separated by 1.3 eV. The first peak is a convolution of two near-degenerate excited states at 201.68 eV of same magnitude. The NTOs reveal that these transitions are of $2p \rightarrow \sigma^*_{C-Cl}$ type for the $2p$ orbitals perpendicular
to the C-Cl bond. Their rotatory strengths have opposite signs and nearly the same magnitude. The second peak comprises three states; two near-degenerate at 202.97 and 202.98 eV, both with weak CD signals (note the sign disagreement between length and velocity gauge results for one of the transitions), and a single state at 203.11 eV with a positive CD signal. The near-degenerate states are transitions from the two perpendicular 2p-orbitals into diffuse orbitals, and the single state is a transition between the 2p orbital lying along the C-Cl bond into diffuse orbitals. The bisignate feature in the XCD spectrum just below 204 eV belongs to two transitions at 203.84 eV and 203.87 eV with rotatory strengths of same magnitude and opposite sign. They are practically dark in the absorption spectrum.

The effect of SOC on XAS is a splitting of the spectrum into three peaks, where the middle one contains contributions from both the up-shifted splitting of the lowest NR peak and and down-shifted splitting from highest NR peak. Introducing SOC into XCD results in a richer spectrum, with a clear bisignate feature for the middle band and two positive peaks in correspondence to the absorption peak at 204 eV.

5 Conclusion

We implemented a computational scheme for simulating optical rotation and circular dichroism spectra based on the EOM-CCSD framework in the Q-Chem package. We applied these new tools to calculate ground-state and excited-state circular dichroism spectra of the selected compounds. The circular dichroism spectra were computed for both valence excitations and for core excitations at the K- and L-edges. In the calculations of L-edge X-ray circular dichroism, spin–orbit effects were included via the state-interaction procedure.

We observed generally good agreement between the simulated spectra obtained from EOM-CCSD with those from response theory at the CCSD level, and from TDDFT/CAM-B3LYP calculations. The assessment of the accuracy of the XCD results proved challenging. For methyloxirane, the computed spectra at the C K-edge do not bear clear resemblance
with the experimental ones. A strong origin dependence of the XCD in length gauge was observed at the oxygen and nitrogen $K$-edges of L-alanine, as well as the chlorine $L_{2,3}$-edge. When the transition is highly localized, locating the gauge origin at the atom of excitation improves the agreement between the spectra from origin-dependent length gauge and those from the velocity gauge. In the X-ray regime, all computed signals, especially for the ones with strong gauge dependence, are extremely small and possibly at the limit of what the methodology can reliably reproduce.
6 Appendix: Expressions for transition $1PDM$s

For brevity, in the following subsections, the left and right amplitudes of target EOM-EE-CCSD and fc-CVS-EOM-EE-CCSD states do not carry the state indices $n$ and $f$. The CCSD $\Lambda$ amplitudes do not carry the state index 0.

6.1 EOM-EE-CCSD state $\leftarrow$ CCSD state transition

\begin{equation}
\gamma_{ia}^{f \leftarrow 0} = \sum_{jb} \left( t_{ij}^{ab} - t_{i,j}^{ba} \right) l_j^b - \sum_{j} \tilde{l}_{ji} t_j^a - \sum_{b} \tilde{l}_{ba} t_i^b
\end{equation}

(47)

\begin{equation}
\gamma_{ai}^{f \leftarrow 0} = l_i^a
\end{equation}

(48)

\begin{equation}
\gamma_{ji}^{f \leftarrow 0} = -\tilde{l}_{ij} - \sum_{a} l_i^a t_{j}^a
\end{equation}

(49)

\begin{equation}
\gamma_{ab}^{f \leftarrow 0} = \tilde{l}_{ab} + \sum_{i} l_i^a t_i^b
\end{equation}

(50)

6.2 CCSD state $\leftarrow$ EOM-EE-CCSD state transition

Here, $l$ represents the amplitudes of CCSD $\Lambda$.

\begin{equation}
\gamma_{ia}^{0 \leftarrow f} = r_i^a + \sum_{jb} \left( \tilde{r}_{ij}^{ab} - \tilde{r}_i^a l_j^b - \tilde{r}_j^a l_i^b \right) l_j^b + \sum_{jb} \left( t_{ij}^{ab} - t_{i,j}^{ba} \right) Y_{jb}
\end{equation}

\begin{equation}
- \sum_{k} \tilde{l}_{ki} t_k^a - \sum_{c} \tilde{r}_i^c - \sum_{j} \tilde{l}_{ji} t_j^a - \sum_{b} \tilde{l}_{ba} t_i^b
\end{equation}

(51)

\begin{equation}
\gamma_{ai}^{0 \leftarrow f} = r_0 l_i^a + Y_{ia}
\end{equation}

(52)

\begin{equation}
\gamma_{ji}^{0 \leftarrow f} = -\tilde{l}_{ij} - \sum_{a} l_i^a \tilde{r}_j^a - \sum_{a} Y_{ia} t_j^a
\end{equation}

(53)

\begin{equation}
\gamma_{ab}^{0 \leftarrow f} = \tilde{l}_{ab} + \sum_{i} l_i^a \tilde{r}_i^b + \sum_{i} Y_{ia} t_i^b
\end{equation}

(54)
6.3 EOM-EE-CCSD state $f \leftarrow$ EOM-EE-CCSD state $n$ transition

$$
\gamma_{ia}^{f\leftarrow n} = \sum_{jb} (\tilde{r}_{ij} - r_{ij}^a - r_{ij}^b) l_j^b + \sum_{jb} (\tilde{r}_{ij}^a - t_{ij}^b) Y_{jb} 
- \sum_k \tilde{l}_{ik} r_k^a - \sum_c \tilde{l}_{ca}^c r_i^c - \sum_j \tilde{l}_{ji} t_j^a - \sum_b \tilde{l}_{ba} t_b^b
$$

(55)

$$
\gamma_{a_i}^{f\leftarrow n} = r_0 l_i^a + Y_{ia}
$$

(56)

$$
\gamma_{ji}^{f\leftarrow n} = -\tilde{l}_{ij} - \sum_a l_i^a \tilde{r}_j^a - \sum_a Y_{ia} t_j^a
$$

(57)

$$
\gamma_{ab}^{f\leftarrow n} = \tilde{l}_{ab}^a + \sum_i l_i^a \tilde{r}_i^b + \sum_i Y_{ia} t_i^b
$$

(58)

6.4 fc-CVS-EOM-EE-CCSD state $\leftarrow$ CCSD state transition

$$
\gamma_{jI}^{f\rightarrow 0} = -\tilde{l}_{Ij} - \sum_a l_i^a t_j^a
$$

(59)

$$
\gamma_{aI}^{f\rightarrow 0} = l_i^a
$$

(60)

6.5 CCSD state $\leftarrow$ fc-CVS-EOM-EE-CCSD state transition

Here, $l$ represents the amplitudes of CCSD $\Lambda$.

$$
\gamma_{ji}^{0f\rightarrow} = -\tilde{l}_{Ij} - \sum_a l_i^a r_j^a
$$

(61)

$$
\gamma_{Ia}^{0f\rightarrow} = r_i^a + \sum_{jb} \left( r_{ij}^a - r_{ij}^b t_i^b \right) l_j^b - \sum_c \tilde{l}_{ca}^c r_i^c - \sum_j \tilde{l}_{ji} t_j^a
$$

(62)

6.6 Intermediates

$$
Y_{ia} = \sum_{jb} l_{ij}^a r_j^b
$$

(63)

$$
\tilde{r}_{ij}^{ab} = r_{ij}^{ab} + r_0 t_{ij}^{ab}
$$

(64)

$$
\tilde{r}_i^a = r_i^a + \frac{1}{2} r_0 t_i^a
$$

(65)
\[
\tilde{l}_{ij} = \frac{1}{2} \sum_{kab} l_{ik} l_{jk} \quad (66)
\]
\[
\tilde{l}_{ab} = \frac{1}{2} \sum_{ijc} l_{ik} l_{ij} \quad (67)
\]
\[
\tilde{l}_{ij} = \frac{1}{2} \sum_{kab} l_{ik} \tilde{l}_{jk} \quad (68)
\]
\[
\tilde{l}_{ab} = \frac{1}{2} \sum_{ijc} l_{ik} \tilde{l}_{ij} \quad (69)
\]
\[
\tilde{l}_{ij} = \frac{1}{2} \sum_{kab} l_{ik} l_{jk} \quad (70)
\]
\[
\tilde{l}_{ab} = \frac{1}{2} \sum_{ijc} l_{ij} l_{ij} \quad (71)
\]
\[
\tilde{l}_{iJ} = \frac{1}{2} \sum_{kab} l_{ik} l_{jk} \quad (72)
\]

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**Notes**

The authors declare the following competing financial interest(s): A. I. K. is the president and a part-owner of Q-Chem, Inc.
Supporting Information Available

Additional information: Tables of raw data, Additional spectra, CAM-B3LYP results for Binol, Geometrical parameters.

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