Circular dichroism calculation for natural products

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Abstract Determination of the absolute configuration (AC) is often a challenging aspect in the structure elucidation of natural products. When chiral compounds possess appropriate chromophore(s), electronic circular dichroism (ECD) may provide a powerful approach to the determination of their absolute configuration. Recently, ECD calculations by time-dependent density functional theory (TDDFT) have come to be used more commonly. In the present review, we give several examples of recent studies using TDDFT-calculated ECD spectra for the AC determination of natural products.

Keywords Circular dichroism · Absolute configuration · Density functional theory

Introduction

Determination of the absolute configuration (AC) of natural products often poses a challenging problem in structure elucidation. There are various approaches to the solution of the problem, e.g., X-ray crystallography, chiroptical methods, and NMR anisotropy methods, each with its own limitations. When chiral compounds possess appropriate chromophore(s), electronic circular dichroism (ECD) may provide a powerful approach to the determination of their AC. In general, AC determination using ECD compares the spectrum of a new compound having unknown AC to those of analogous compounds of known AC. However, AC determination by predicting the sign of one or more bands in the ECD spectrum by using empirical, semi-empirical or non-empirical rules may be an option. Another option, which has become more widely used in the past 10 years, is to compare calculated and experimental ECD spectra.

ECD calculations have been used for the determination of ACs of natural products for decades, as seen in the π-SCF studies of ECD by Mason and coworkers [1]. However, its use was quite limited until the major advancement in ECD calculations of time-dependent density functional theory (TDDFT) was introduced, which resulted in a good compromise between computational costs and accuracy [2–10]. The principle of the determination of the ACs of natural products by ECD calculation is relatively simple: basically, calculated ECD spectra are compared with experimental ECD spectra. If the two data sets are very similar to each other, then highly reliable assignment is obtained. In the present review, we briefly explain the method of TDDFT calculation of ECD spectra and give several examples of its successful application to the determination of the AC of natural products, including our own previous works. There are hundreds of publications using TDDFT-calculated ECD spectra for the AC assignment of a broad range of compounds, and examples were selected from recently published studies, especially those from 2012.

Computational methods

ECD calculations generally involve two steps, first the conformational analysis of the compound to obtain the possible conformer(s), and second, the UV/ECD TDDFT calculation of each conformer(s). The conformational analysis is often done by Monte Carlo methods using molecular mechanics (MMFF94, etc.) and/or semi-
empirical methods (AM1, etc.) for the relative energy evaluation of the conformers. The resulting conformers are then optimized further using density functional theory (DFT) methods before being subjected to the TDDFT calculations of the UV/ECD by using programs such as Gaussian [11], TURBOMOLE [12], or NWChem [13]. To obtain the calculated UV/ECD spectra of the isomers, the UV/ECD spectra of the conformers are Boltzmann averaged. The averaged UV spectra are then shifted to conform to the experimental UV spectrum and the same shifts are also applied to the corresponding calculated ECD spectra before comparing the calculated ECD spectra with the experimental ECD of the natural product in question. The calculation steps are summarized in Fig. 1.

The accuracy of TDDFT calculations itself depends mainly on the basis set and functional used for the calculations. Use of a larger basis set generally increases the accuracy, but it also increases the computational time required. In ECD calculations, basis sets with polarization and diffuse functions, such as 6-31G* or aug-cc-pVDZ, and the B3LYP functional are commonly used and give satisfactory results [14]. The ECD calculation level is often expressed as [Functional 1]/[Basis set 1]/[Functional 2]/[Basis set 2], which means that the ECD calculation was performed using Functional 1 and Basis set 1 on a geometry optimized by using Functional 2 and Basis set 2.

The results of TDDFT calculations of UV/ECD are excitation energies, and their corresponding oscillator strength and rotatory strength. The oscillator strengths are used to simulate the UV curve, and rotatory strengths to simulate the ECD curve. Both oscillator strength and rotatory strength can be calculated using either dipole–velocity gauge or dipole–length gauge, but the use of dipole–length gauge generally gives better results [3, 15]. To take the solvent effect into account, a COnductor-like continuum Solvent MOdel (COSMO) can also be employed [16, 17].

The calculated rotatory strength values are generally converted to the line-shaped ECD curve using a Gaussian distribution function:

$$\Delta \varepsilon(\sigma) = \frac{1}{2.296 \times 10^{-19}} \times \frac{1}{\Delta \sigma \sqrt{\pi}} \sum \sigma_i R_i e \left[ \left( -\frac{\sigma - \sigma_i}{\Delta \sigma} \right)^2 \right]$$

where $\sigma$ is the wavenumber (in cm$^{-1}$), $\Delta \sigma$ is half of the width of the band at $1/e$ peak-height, and $\sigma_i$ and $R_i$ the excitation wavenumber and rotatory strength for transition $i$, respectively [18]. The value of $\Delta \sigma$ can be evaluated from the corresponding UV spectrum, and is typically in the range of 0.05–0.4 eV. The use of Lorentzian distribution to simulate ECD curves has also been reported by using the function:

$$\Delta \varepsilon(\sigma) = \frac{1}{2.296 \times 10^{-19}} \times \frac{1}{\gamma \sqrt{\pi}} \sum \sigma_i R_i \left[ \frac{\gamma^2}{(\sigma - \sigma_i)^2 + \gamma^2} \right]$$

where $\gamma$ represents half the width of the band at half peak-height [18–22].

Use of TDDFT calculation of ECD spectra for the AC determination of natural products

This section gives examples where the TDDFT calculation of ECD was used for the AC determination of natural products. The examples include alkaloids, terpenoids, coumarins, anthraquinones, and other types of compounds, ranging from conformationally rigid to highly flexible compounds.
Alkaloids

Eucophylline [23]

Eucophylline (1), isolated from Leuconotis griffithii [23], has only one chiral center at C-20. Conformational analysis (MMFF94 force field, 50 kJ/mol window) generated conformations that can be divided into two groups: one group with the piperidine ring in a chair form, and the other with the ring in a boat form. Members of the same group differ in the three-dimensional (3D) location of the ethyl moiety at C-20 and the vinyl moiety at C-4. The group with the piperidine ring in chair form was predicted to be dominant, which agreed with NMR data (Fig. 2). The ECD spectra were calculated at the BP86/aug-cc-pVDZ//BP86/SVP and B3LYP/TZVPP//B3LYP/TZVPP level on two stable conformers differing in the location of the vinyl group because changes in the location of the ethyl moiety were expected to have no effect on the calculated ECD spectra. The ECD spectra for the two conformers were Boltzmann-averaged to obtain the ECD spectrum of the isomers. The two calculation levels lead to the same conclusion that the AC at C-20 of 1 was R. Of the calculation levels, the BP86/aug-cc-pVDZ//BP86/SVP calculated ECD spectrum matched the experimental spectrum well, but in the B3LYP/TZVPP//B3LYP/TZVPP calculated spectrum, the first and second cotton effects (CEs) apparently overlapped (Fig. 3).

Leucomidines A and B [24]

Leucomidines A and B (2 and 3, Fig. 4) are indole alkaloids also isolated from L. griffithii [24]. Conformational analysis (MMFF94 force field, 50 kJ/mol window) for both 2 and 3 generated conformers differing only in the orientation of the side chain at C-20. The ECD spectra of the most stable conformation for 2 and 3 were each calculated at BP86/aug-cc-pVDZ//BP86/SVP and B3LYP/TZVPP//B3LYP/TZVPP levels. Both levels gave the same conclusion that both 2 and 3 had the 20R,21S configuration. In the case of 2, however, the B3LYP/TZVPP//B3LYP/TZVPP-calculated ECD spectrum agreed with the experimental ECD spectrum better than that calculated by BP86/aug-cc-pVDZ//BP86/SVP (Fig. 5). It is to be noted that the calculated ECD spectrum for 3dis (Fig. 5)—a model compound for 3 in which the side chain at C-20 was substituted with a methyl group—reproduced the experimental ECD spectrum better.

Rupestines F–M [25]

Rupestines F–M (4–11, Fig. 6) are guaipyridine alkaloids isolated from Artemisia rupestris [25]. Of those eight compounds, 5–11 showed the CE at about 270 nm in the experimental ECD spectra, which can be attributed to transition of the pyridine moiety. To study the effect of conformation on the CE sign at 270 nm, the ECD spectra...
of possible conformations of 10, 11 and a model compound, 2-methylpyridine-cycloheptene (12), were calculated (Figs. 7, 8). The calculation results may be summarized as follows, (1) B3LYP/TZVPP calculations overestimated the excitation energy of pyridine’s first absorption band (calc. ~250 nm vs expt. ~270 nm), so that the calculated CE at around 250 nm are observed experimentally at around 270 nm; (2) when the plane of C-6, C-7, and C-8 was over the pyridine ring, a positive CE at around 215 nm and a negative CE at around 250 nm were observed regardless of the chirality at C-5. Thus, by using the CE sign at around 270 nm, the absolute structure of the molecules can be deduced or the AC at C-8 of 5–11 could be assigned accordingly; 5–11 were shown to possess the same 8S configuration.

The ECD spectrum of 4 is different from those of 5–11. By comparing the ECD spectrum calculated at B3LYP/TZVPP//B3LYP/TZVPP level with the experimental spectrum (Fig. 9), the AC of 4 was assigned as 5S,8S.

**Bisnicalaterines B and C [26]**

Bisnicalaterines B and C (13 and 14) are rotamers isolated from *Hunteria zeylanica* [26]. By changing the dihedral angle N-1–C-16–C-9–C-10, two stable conformations were obtained, one corresponding to the structure of 13 and the other to that of 14 (Fig. 11). Monte Carlo conformational analysis (MMFF94 force field, 50 kJ/mol window) of each structure generated conformers differing only in conformation at C-20 (ethyl group, C-18 and C-19) and/or at C-15’ (hydroxyethyl group, C-16’ and C-17’), which should have little or no effect on the calculated ECD spectra. Thus, the ECD calculation was performed at BP86/aug-cc-pVDZ//BP86/SVP level only for the most stable conformation of 13 and 14. The calculated and experimental ECD spectra for 13 and 14 are shown in Fig. 12, suggesting their AC to be as shown in Fig. 10.

**Schizozygine [27]**

Schizozygine (15) is an alkaloid isolated from *Schizogygia caffaeoides* [28]. Monte Carlo conformational search by using MMFF94 force field for the energy evaluation generated two stable conformations within the 20 kcal/mol window. The two conformations were then re-optimized at the B3LYP/6-31G* level, and the results were subjected to a potential energy scan (PES) at the B3LYP/6-31G* level. By varying the dihedral angle C-9–C-10–O–C-24 of both of the conformations, two additional conformations were identified. The ECD calculations for all four conformers were done at the B3LYP/aug-cc-pVDZ//B3LYP/6-31G*, B3LYP/aug-cc-pVDZ//B3LYP/TZ2P, and B3LYP/aug-cc-pVDZ//B3PW1/TZ2P levels. For each conformer, calculations at all three levels gave similar results. The ECD spectra were Boltzmann-averaged, and the resulting ECD spectrum agreed well with the experimental ECD spectrum. The AC of 15 was assigned as shown in Fig. 13 [27].

**Actinophyllic acid [29]**

The AC of actinophyllic acid (16)—an alkaloid isolated from *Alstonia actinophylla* [30]—was assigned as shown in Fig. 13 indirectly on the basis of its methyl ester derivative because of the expected complication due to the presence of the carboxylic acid group [29]. A preliminary conformational search (MMFF94) gave two stable conformers within a 10-kcal/mol window. Of the ECD spectra calculated at B3LYP/6-31G(d,p)//B3LYP/6-31G(d,p) and

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Fig. 5  Calculated (20R,21S) and experimental ECD spectra of leucomidines A (2) and B (3)

4: R¹ = α-Me, R² = C(CH₂)CO₂Me
5: R¹ = α-Me, R² = CO₂Me
6: R¹ = β-Me, R² = COCH₂CH₂OH
7: R¹ = α-Me, R² = COCH₂CH₂OH
8: R¹ = α-Me, R² = C(CH₂)CH₂OH
9: R¹ = β-Me, R² = C(CH₂)CH₂OH
10: R¹ = α-Me, R² = OH
11: R¹ = β-Me, R² = OH
12: R¹ = R² = H

Fig. 6  Structures of rupestines F–M (4–11) and 2-methylpyridine-cycloheptene (12)

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B3LYP/aug-cc-pVDZ/B3LYP/6-31G(d,p) levels, the one calculated at the B3LYP/6-31G(d,p) level generally agreed well with the experimental ECD spectrum, but did not give the negative shoulder at around 230 nm. On the other hand, the B3LYP/aug-cc-pVDZ-calculated ECD spectrum clearly reproduced the negative shoulder.
around 230 nm, but the region around 250 nm became slightly positive.

Mariline A1 and A2 [31]

Mariline A1 (17) and A2 are enantiomeric phthalimidines isolated from sponge-derived fungus *Stachylidium* sp. [31]. Detailed investigation of the highly flexible conformations of 17 was carried out at the B3LYP/SV(P) level, yielding more than 550 conformers. ECD calculations were performed at the B3LYP/SV(P) level with the COSMO model. Comparison of the calculated ECD spectra with the experimental spectrum did not allow an unambiguous assignment of the AC. Almeida and coworkers [31] associated the problem with dispersion effects caused by the highly flexible alkoxy side chain. Instead of using a dispersion-corrected functional in combination with a larger basis set to perform a DFT optimization of the conformations, they used single-point energy calculations for all previously DFT-optimized structures at the RI-SCS-MP2/TZVPP level since a very accurate valuation of the energies for the Boltzmann weighting for such a high number of conformers was indispensable. Boltzmann weighting with these more reliable energy values yielded a significantly improved overall ECD spectrum, which almost agreed with the experimental one. Thus the AC of 17 was determined to be as shown in Fig. 14.

**Chaetoglobosin Vb [32]**

Chaetoglobosin Vb (18)—a cytochalasan alkaloid—was isolated from *Chaetomium globosum* [32]. Xue and coworkers performed a TDDFT ECD calculation [B3LYP/6-311 + G(d,p)//B3LYP/6-311 + G(d,p)] using a model compound of 18 in which the 3-methylindole moiety was replaced by a hydrogen atom. They considered that the 3-methylindole ring should contribute only weakly to the observed ECD spectrum of 18 in the long-wavelength region, because (1) it is connected to the chiral skeleton by two flexible bonds; (2) it is quite distant from the other
chromophores in the molecule, especially the cyclopentenone; and (3) its transitions above 220 nm are either very weak or directed roughly perpendicular to the C-3′/C-10 bond; so its coupling with other chromophores should be near zero. The use of this model compound simplified ECD calculations, since it gives only two conformers to be considered for the calculations. Based on the calculation results, the AC of 18 was suggested to be as shown in Fig. 15.

Terpenoids

Chisomicine A [33] and sanjecumin B [34]

Chisomicine A (19) [33] and sanjecumin B (20) [34] are limonoids isolated from Chisocheton ceramicus (Meliaceae), and from Sandoricum koetjape (Meliaceae), respectively. ECD calculations by TDDFT suggested the ACs of 19 and 20 to be as shown in Fig. 16. Calculations were performed at B3LYP/SVP/B3LYP/SVP level and at BP86/aug-cc-pVDZ//BP86/SVP level for chisomicine A and sanjecumin B, respectively. The calculated ECD spectra are shown in Fig. 17 along with the experimentally obtained spectra.

Ximaolides [35]

Ximaolides, including ximaolide A (21, Fig. 18), are a series of bis-cembrane diterpenoids obtained from the Hainan soft coral Sarcophyton tortuosum [36, 37]. Kurta´n and coworker [35] assigned the AC of 21 by comparing the solid-state ECD spectrum of 21 with the TDDFT-calculated ECD spectrum of the solid-state geometry obtained from X-ray crystallography analysis. By this method, they managed to reduce the number of input conformers for the ECD calculation of this highly flexible compound to only one, and thus the AC of 21 was determined as shown in Fig. 18.

Coumarins

5-O-methyldiphysin and 5,5′-di-O-methyldiphysin [38]

Bis-dihydrocoumarins, 5-O-methyldiphysin (22) [38] and 5,5′-di-O-methyldiphysin [39] were isolated from Ormocarpum kirkii. By using the non-empirical aromatic quadrant rule [40], Dhooghe and coworkers [39] assigned the AC of 5,5′-O-dimethyldiphysin to be of the
configuration. However, Xu and coworkers [38] showed that the TDDFT ECD calculation of (3R,4S,3′R,4′S)-22 in both gas and solution phases [(B3LYP/6-31G(d,p)//B3LYP/6-31G(d,p)] produced an ECD spectrum quite similar to the reported experimental spectrum of 5,5′-di-O-methyldiphysin, indicating that the non-empirical aromatic quadrant rule is not appropriate for defining the AC of C-4 of this class of compounds (Fig. 19).

Anthraquinones

**Altersolanol N [41]**

Altersolanol N (23) is an anthraquinone derivative isolated from *Sternphilyum globuliferum*—an endophytic fungus isolated from *Mentha pulegium* (Lamiaceae) [41]. First, DFT minimizations were executed at B3LYP/6-31G(d) level, and all the resulting DFT minima with relative internal energies within 10 kcal/mol were reoptimized at the B3LYP/6-31G + (d,p) level with the polarizable continuum model (PCM) for acetonitrile. For the TDDFT ECD calculations, a preliminary screening of various functionals (B3LYP, CAM-B3LYP, BH&HLYP, PBE0) and basis sets (SVP, TZVP, aug-TZVP) showed that CAM-B3LYP provided the best result, and the results obtained by the use of the smallest basis set used (SVP) was compatible with those obtained by the use of larger ones. Therefore, the CAM-B3LYP/SVP combination was chosen for the final calculations, which determined the AC of 23 to be as shown in Fig. 20.

**Coniothyrrinone A [42]**

Coniothyrrinone A (24), an anthraquinone derivative, was isolated from *Coniothyrium* sp.—an endophytic fungus isolated from *Salsola oppositifolia* [42]. The TDDFT ECD calculations were performed using the TZVP basis set and three functionals (B3LYP, BH&HLYP, CAM-B3LYP) and all of them produced ECD spectra that agreed well with the experimental ECD spectrum; the use of CAM-B3LYP gave the best agreement. Thus, the AC of 24 was assigned to be as shown in Fig. 20.

**Others**

**Phomopsinones A–D [43]**

Phomopsinones A–D (25–28) were obtained from an endophytic strain of *Phomopsis* sp., isolated from the halotolerant plant *Santolina chamaecyparissus* [43]. Hussain and coworkers used (S)-4-methoxy-7-methyl-7,8-dihydropyrano[4,3-b]pyran-2(5H)-one (29) as a model compound to find the best functional and basis set level for ECD calculations. The DFT optimized geometry [B3LYP/6-31G(d)] with equatorial 7-methyl was used as input in TDDFT ECD calculations with various combinations of hybrid functionals (CAM-B3LYP, B3LYP, PBE0) and basis sets (SVP, TZVP, aug-TZVP). The calculated ECD spectra for 29 were then compared with the experimental ECD spectrum of 25. The long-range functional CAM-B3LYP gave the best agreement, and all the basis sets performed similarly. Thus, the CAM-B3LYP/SVP combination was employed in subsequent ECD calculations of 25, 26, and 28 to assign their ACs to be as shown in Fig. 21.
Conclusion

TDDFT calculation of ECD simplified the interpretation of the ECD–AC relationship and is a promising tool for the AC determination of natural products with chiral centers. Successful application of TDDFT calculations of ECD spectra to the AC determination of various type of natural products, ranging from conformationally rigid to highly flexible compounds, are exemplified in this review. Further improvements both in the computer system technologies and TDDFT, may make the TDDFT calculation of ECD an integral part of the AC determination of natural products.

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