Mass-resolved electronic circular dichroism ion spectroscopy

Steven Daly1, Frédéric Rosu2, Valérie Gabelica1*  

DNA and proteins are chiral: Their three-dimensional structures cannot be superimposed with their mirror images. Circular dichroism spectroscopy is widely used to characterize chiral compounds, but data interpretation is difficult in the case of mixtures. We recorded the electronic circular dichroism spectra of DNA helices separated in a mass spectrometer. We studied guanine-rich strands having various secondary structures, electrospayed them as negative ions, irradiated them with an ultraviolet nanosecond optical parametric oscillator laser, and measured the difference in electron photodetachment efficiency between left and right circularly polarized light. The reconstructed circular dichroism ion spectra resembled those of solution-phase counterparts, thereby allowing us to assign the DNA helical topology. The ability to measure circular dichroism directly on biomolecular ions expands the capabilities of mass spectrometry for structural analysis.

Two centuries ago, the interaction of polarized light with crystals revealed that many molecules come in two forms, nonsuperimposable with their mirror images (1). The activity and toxicity of natural or synthetic molecules often depend on their chirality (2). Identifying which mirror image is present in a sample is thus crucial. Furthermore, because biomolecules such as DNA or proteins consist of repetitive chiral subunits, they can form helical structures, as in the iconic DNA double helix (3, 4). Characterizing the types of helical structures formed by biomolecules is therefore also essential for structural biology.

Mass spectrometry (MS) is a widely used analytical method with expanding impact in structural biology (5, 6). MS excels at separating and quantifying complex mixtures, and additional structural characterization can be obtained using tandem MS (7), ion mobility spectrometry (8), or infrared ion spectroscopy (9–11). However, MS-based measurements are typically blind to chirality. Characterizing chiral compounds by MS currently requires a physical interaction with other chiral molecules (12), either by separation on chiral phases in front of the mass spectrometer or by the formation of complexes with chiral auxiliaries in the gas phase (13).

Another way to characterize chiral biomolecules is through interaction with chiral light. In solution, DNA and proteins are conveniently characterized by circular dichroism (CD), which measures the difference of absorption between left and right circularly polarized light. Electronic CD is particularly useful to characterize the different types of helices formed by nucleic acids (14) or the secondary structures formed by proteins (15). The structural interpretation relies on spectral databases of known structures. However, the interpretation of CD spectra becomes difficult for samples wherein multiple species or structures coexist, because the resulting spectrum is the weighted average of all contributions.

Here, we report a method for recording electronic CD spectra on DNA helices separated in a mass spectrometer. Although CD and MS have been combined before (16–18), the previous methods were based on resonant multiphoton ionization (REMPI) and were thus applicable only to volatile neutral small molecules (<200 Da). We used electrospray to produce gas-phase polyanions of intact DNA multihelices (>5000 Da) and interrogated them with circularly polarized ultraviolet light directly inside the mass spectrometer.

Our mass analyzer is a quadrupole ion trap (Paul trap) with two opposite holes (diameter 1.7 mm) in the ring electrode. We reasoned that this configuration would minimize risks of reflection of the laser beam inside the mass spectrometer, which could possibly alter the circular polarization. The electrospray source was operated in the negative mode to produce multiply charged anions. We used gentle ion transfer conditions to prevent gas-phase restructurizing and thus maximize the chances of preserving the solution-phase secondary structures.

We analyzed several DNA G-quadruplex tetrahelical structures—parallel right-handed tetramer TG4T [(dTGTTGGT)4(NH4+)3], parallel left-handed intramolecular ZG4 [dT(TGTT)TG(TGTT)5(NH4+)3] (19), antiparallel right-handed intramolecular 5′YF [d(GGGTTA)GGTTGGGG(K)x] (20)—and the silver-mediated parallel G-duplex GAgG [(dG11)2(Ag+)n] (21). These structures were chosen because they have remarkably different electronic CD spectra in solution in the wavelength range 220 to 300 nm (fig. S1). Also, these structures are stabilized by spines of central cations, which maximizes the likelihood of preserving the hydrogen-bonding and stacking arrangement in the gas phase while minimizing rearrangements due to self-solvation (21). The preservation of the known solution-phase shapes was confirmed by the good agreement between helium momentum transfer collision integrals measured in ion mobility spectrometry and those calculated for gas-phase structures obtained by molecular dynamics at the semi-empirical level (fig. S2). For TG4T, infrared ion spectroscopy also supported the preservation of G-quartets (22). We can thus safely assume that these molecular systems have the same base-stacking arrangement in the gas phase as in solution.

When coupled to MS, ion spectroscopy is an action spectroscopy (and not an absorption spectroscopy): One records the effect of the laser irradiation on the ions. Another reason for selecting DNA as the first test case is that, when irradiating DNA polyanions between 220 and 300 nm, the resulting action is electron photodetachment (ePD) (23, 24). This action has two advantages over fragmentation: (i) There are few product ions to quantify, which in terms of statistics will increase the chances

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*Corresponding author. Email: v.gabelica@iecb.u-bordeaux.fr
of detecting differences in product ion yields between the two polarizations; and (ii) ePD is monophotonic (25), and thus the normalization for fluctuations of the laser power is simply linear for all wavelengths. We verified this linear relationship, and for all spectroscopy experiments we selected the pulse energy ranges in which only linear ePD was observed (fig. S3).

Nanosecond laser pulses were generated using a wavelength-tunable optical parametric oscillator laser (nanosecond pulses, ≤100 μJ transmitted to the trap). To generate circularly polarized laser pulses, the beam passes through an air-spaced Rochon prism, which gives pure linearly polarized light. Next, the laser passes through an achromatic broadband quarter-wave plate (Fig. 1, figs. S4 to S6, and supplementary materials). The angle of the fast axis of the quarter-wave plate relative to the polarization direction is set by the Rochon prism to rotate the quarter-wave plate, and the mass spectrum (Fig. 1). Mass spectra and pulse energy are acquired for 90 s. The polarization state of the laser pulse is then changed by irradiating them with a single laser pulse of the selected wavelength, polarization, and pulse energy, and then recording their mass spectrum (Fig. 1). Mass spectra and pulse energy are averaged for 90 s. The polarization state of the laser pulse is then changed by rotating the quarter-wave plate, and the mass spectrum and pulse energy are acquired for another 90 s. This process is repeated 10 times for each polarization. The relative electron detachment yield for each mass spectrum is calculated as

\[ Y_{\text{ePD}} = \frac{I_{\text{ePD}}}{I_{\text{total}}} \]  

where \( I_{\text{ePD}} \) is the integrated intensity of the peak due to electron detachment, \( I_{\text{total}} \) is the total integrated intensity (precursor + ePD product), \( \lambda \) is the wavelength, and \( E_{\text{trans}} \) is the transmitted pulse energy determined from the reflected pulse energy by calibration (fig. S8). We then calculate the average value of the relative electron detachment yield for left- and right-handed circular polarizations. The CD monitored by ePD is expressed as an asymmetry factor, approximated by

\[ g_{\text{ePD}} = \frac{\Delta ePD}{ePD} = 2 × \left( \frac{Y_{\text{LCP}} - Y_{\text{RCP}}}{Y_{\text{LCP}} + Y_{\text{RCP}}} \right) \]

where LCP and RCP are left and right circularly polarized light, respectively. The gas-phase CD spectra are reconstructed by plotting the asymmetry factor \( g_{\text{ePD}} \) as a function of the wavelength. To facilitate the visual comparison, we plot the solution-phase CD spectra as \( \Delta A/A \) (where \( A \) is the absorbance), hence their shape differs from the ones traditionally displayed in molar CD.

To unambiguously prove that the gas-phase CD effect comes from the sample and not from an instrumental artifact, we performed the

Fig. 2. Gas-phase circular dichroism spectra (\( \Delta ePD/ePD \)) compared to solution-phase spectra (\( \Delta A/A \)). Symbols and lines denote gas-phase and solution-phase CD data, respectively. (A) TG4T: natural right-handed \([(\text{TGGGGT})_4(\text{NH}_4\text{)}^+)_3]^+ \) (blue) and its mirror image \( [(\text{TGGGGT})_4(\text{NH}_4\text{)}^+)_3]^+ \) (red). Inset: Gas-phase CD measured at 260 nm for \([(\text{TGGGGT})_4(\text{NH}_4\text{)}^+)_3]^+ \) ions prepared from 5 μM solutions with varying ratios of \( v \)-TG4T and \( l \)-TG4T. Error bars show the 95% confidence interval calculated from the standard error (three replicates, except at 260 nm with five replicates). (B to D) Gas-phase and solution-phase CD spectra for the antiparallel G-quadruplex 5YEY\(_5\) (B), the left-handed G-quadruplex ZG4\(_5\) (C), and the G-duplex GAGG\(_5\) (D). The illustrations showing helicity were generated using Chimera from Protein Data Bank coordinates (entries 2O4F, 5YEY, 6GZ6, and 4JRD, respectively); the gas-phase structures are shown in fig. S2.

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experiment on n-[TGGGGT]₄*(NH₄⁺)₃] formed from the natural DNA backbone (all D-sugars) and from its enantiomer (all L-sugars). The gas-phase CD signals have opposite signs (Fig. 2A). The magnitude is not exactly reversed, however. At 260 nm, we recorded a series of CD measurements by varying the relative concentration of the two enantiomers (ω:λ) in solution from only D to only L (Fig. 2A, inset). We obtained from the natural DNA backbone (all D-sugars) and from its enantiomer (all L-sugars). The gas-phase CD spectra result from the CD in absorption, and the differences in ePD with left and right circularly polarized light are reporting on CD; this finding suggests that the technique could be suitable for quantification. The line did not pass exactly through 0 when the proportions were 50:50, indicating the presence of residual instrumental artifacts, likely because the polarization is only ~95% circular. In the future, use of an achiral internal standard could further reduce the uncertainty. Furthermore, if such imperfections vary with the wavelength, it would induce some distortion in the CD spectra that might hamper structural assignment. We thus compared the solution-phase and gas-phase CD spectra for other typical DNA helices (Fig. 2, B to D; symbols for gas-phase CD, lines for solution-phase CD). The solution- and gas-phase CD spectral shapes are similar in terms of sign and position of the maxima and minima. This suggests that the following conditions are met: (i) The base-stacking pattern existing in solution is preserved in the gas-phase ions. (ii) The gas-phase action reflects the absorption, in terms of sensitivity to the circular polarization. In other words, the electronic excited states that are responsible for the CD effect also trigger ePD. (iii) Possible distortions due to imperfections in the polarization do not preclude assigning the base-stacking arrangement on the basis of spectral shape. As a result, the gas-phase CD spectra can unambiguously discriminate between the different guanine-rich oligonucleotide structures.

The gas- and solution-phase asymmetry factors mainly differ in their magnitude, which is consistently larger in the gas phase (up to 2%) than in solution (<0.4%). This has been observed previously for small neutral molecules measured by REMPI (17, 26–28), where magnitudes of up to 25% have been observed (28). High CD signals are also observed in photoelectron CD (PECD) (29, 30), but in that case the measured property is the angular distribution of the photoelectrons. Here, we measure the electron detachment yield, and ePD is the result of a resonant excitation above the detachment threshold (24, 25). Given that solution-phase spectra result from the CD in absorption, we conclude that the gas-phase CD effect in photodetachment yields is also due to the resonant absorption of circularly polarized light by a chiral molecule. It is thus puzzling that the CD spectral shapes are similar although the magnitudes differ.

Here we consider possible explanations. In small molecules, the solvent can affect the magnitude of CD (31), and thus a change in the dielectric environment from water to vacuum may change the magnitude of the CD spectrum. To test this hypothesis, we recorded solution-phase CD spectra of n-TG4T in mixtures of water up to 50% isopropanol (dielectric constant varying from 80.4 to 44.3). The mass spectra show that the G-quadruplex is intact, and no larger multimers are present (fig. S9). No change in the value of ΔA/A was observed (fig. S10), which suggests that the dielectric environment plays only a small role. We also recorded the gas-phase CD spectra for the 5-, 6-, and 7-ions of n-TG4T, reasoning that if intramolecular electric dipoles influence the electron ejection dynamics, the effect would increase with the charge state. We observed no significant change in either magnitude or shape of the CD spectra as a function of the charge state (fig. S11). We thus hypothesize that the origin of the larger gas-phase magnitude comes from the different definition of asymmetry factors. In the gas phase, absorption is revealed by electron detachment. If not all states that absorb result in electron detachment (25, 32) while most states that are responsible for CD do, the denominator of the asymmetry factor is smaller in gas-phase CD action spectroscopy, and thus the asymmetry factor is larger. Because the electronic states responsible for the CD effect are most likely to be delocalized on the entire DNA helices, this may result in more efficient autodetachment after resonant excitation. Thus, although future work is needed to elucidate the origins and dynamics of ePD from DNA polyanions—for example, by measuring the photoelectron CD—the ePD action serendipitously revealed itself to be especially well suited to probe DNA higher-order structures by ion spectroscopy.

All experiments above were carried out on pure samples, and tandem MS was used to select a charge state. Next, we used mass separation to obtain the CD spectra of individual components from a mixture. Human telomeric sequences, consisting of TTAGGG repeats, can form several G-quadruplex topologies (33). At submillimolar potassium concentrations, complexes with 1 K⁺ (two G-quartets) and 2 K⁺ (three G-quartets) coexist (34). The 2-K⁺ complex is fully formed at high KCl concentrations, but 1-K⁺ complexes cannot be isolated in solution. We previously reconstructed solution-phase CD data for individual K⁺ binding stoichiometries (Fig. 3A) by deconvoluting the solution-phase CD signal of a K⁺ concentration series, having determined the amount of each complex in separate MS measurements. Here we applied mass-resolved CD ion spectroscopy directly on the mixture. Because the [M+1K]⁵⁻ and [M+2K]⁶⁻ complexes...
are close in m/z, they could be co-isolated (inset of Fig. 3B), and as their m/z differs, the ePD efficiencies of both species could be monitored in a single experiment (this is also how we envisage using internal standards in the future). Each spectrum was averaged over 180 s. Although the magnitudes of gas-phase CD signals (Fig. 3B) differ from those of their solution-phase counterparts, their position and signal allow us to infer the G-quadruplex stacking topology using the same rules as in solution (35): The 2-K⁺ complex has positive CD signals at 290 and 260 nm and a negative signal at 240 nm (signature of coexisting homo- and heterostacking), whereas the 1-K⁺ complex has positive signals at 290 and 240 nm and a negative signal at 260 nm (signature of heterostacking).

Our experiments show that it is possible to measure the CD of large DNA polyanions in the gas phase while exploiting the physical separation of components inside the mass spectrometer. The similarity between solution-phase and gas-phase spectra allowed us to distinguish the secondary structures of G-rich DNA, thereby extending the scope and capabilities of structural MS. In the future, proteins and their aggregates can be studied by the same approach, but overcoming signal-to-noise challenges will likely require using internal standards and alternative circularly polarized light sources giving access to shorter wavelengths. For positive ions and small molecules in general, cold mass tagging could provide alternative monophotonic action channels that may be less dependent on the nature of the excited state. Our demonstration of feasibility thus opens new avenues to study diverse classes of chiral molecules while leveraging the separation capabilities of contemporary mass spectrometry.

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SUPPLEMENTARY MATERIALS
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