Detection of an *en masse* and reversible B- to A-DNA conformational transition in prokaryotes in response to desiccation

Donna R. Whelan¹, Thomas J. Hiscox², Julian I. Rood², Keith R. Bambery¹,³, Don McNaughton¹ and Bayden R. Wood¹

¹Centre for Biospectroscopy, School of Chemistry, and ²Department of Microbiology, School of Biomedical Sciences, Monash University, Victoria 3800, Australia
³Australian Synchrotron, 800 Blackburn Road, Clayton, Victoria 3168, Australia

The role that DNA conformation plays in the biochemistry of cells has been the subject of intensive research since DNA polymorphism was discovered. B-DNA has long been considered the native form of DNA in cells although alternative conformations of DNA are thought to occur transiently and along short tracts. Here, we report the first direct observation of a fully reversible *en masse* conformational transition between B- and A-DNA within live bacterial cells using Fourier transform infrared (FTIR) spectroscopy. This biospectroscopic technique allows for non-invasive and reagent-free examination of the holistic biochemistry of samples. For this reason, we have been able to observe the previously unknown conformational transition in all four species of bacteria investigated. Detection of this transition is evidence of a previously unexplored biological significance for A-DNA and highlights the need for new research into the role that A-DNA plays as a cellular defence mechanism and in stabilizing the DNA conformation. Such studies are pivotal in understanding the role of A-DNA in the evolutionary pathway of nucleic acids. Furthermore, this discovery demonstrates the exquisite capabilities of FTIR spectroscopy and opens the door for further investigations of cell biochemistry with this under-used technique.

1. Introduction

The molecular conformation of DNA *in situ* has been a point of interest ever since Watson and Crick’s X-ray diffraction studies and the ensuing analyses revealed crystal polymorphism of the double helix [1–5]. Not only did they solve the crystal structure of the assumed native, fully hydrated B-DNA, they also detected and identified a more dehydrated A-DNA polymorph in DNA crystals. As this polymorphism was detected over 60 years ago, dozens of alternative double-stranded conformations have been identified including Z-DNA [6] and Holliday junctions [7], as well as triplex [8,9] and quadruplex conformations [10,11]. The biological significance of these non-canonical conformations, however, has been hotly contested because of a lack of techniques capable of determining *in situ* DNA conformation, resulting in a heavy reliance on indirect methods such as crystallography [12,13], computational chemistry [14,15] and electron microscopy [16,17].

Nonetheless, several DNA conformations are now believed to be present in cells and to be involved in biologically relevant reactions [16]. Of the conformations now established as biologically relevant, A-DNA has been the most researched and has been identified as the most biologically active non-B-DNA conformation. In the last two decades, sequences containing 5'-CC-3' and 5'-ACT-3' repeats have been demonstrated to be predisposed to assuming an A-DNA form most easily [18]. Furthermore, the transition to A-DNA is required for many different transient interactions with enzymes [12,19,20], counter-ions [13] and ligands [21]. A-DNA has also been strongly implicated through...
protein–DNA crystal structures as a necessary structure in several polymerase reactions [22] and in all RNA/RNA [23,24] and hybrid helices [25]. In a handful of studies, A-DNA has also been demonstrated to have increased stability in response to UV radiation [26,27], chemical toxicity [28] and desiccation [29,30], though the significance of this finding has been questioned owing to the prevalence of B-DNA in live cells.

One method capable of determining DNA conformation is infrared (IR) spectroscopy and, following the discovery of the double helix, the conformation-specific IR absorptions of both B- and A-DNA were exhaustively catalogued and used to complement crystallography data [20,31–33]. In more recent years, Fourier transform infrared (FTIR) spectroscopy has been applied to intact cell and tissue samples as one of several techniques known collectively as biospectroscopy. These methods have been highlighted as both biochemical and medical diagnostic tools capable of detecting disease state [34], cell cycle [35], the onset of apoptosis and necrosis [36], as well as for investigating cell interactions with drugs, chemicals and pathogens [37]. This is because FTIR spectroscopy provides a holistic snapshot of the biochemical make-up of a sample based on the vibrational absorption of all the sample constituents. Different chemical bonds absorb at different energies and the degree of absorption is directly proportional to concentration. FTIR spectroscopy can thus be applied to live samples without any requirement for staining or targeting and with no danger of perturbation to the sample itself.

However, interpretation of the FTIR spectra of biological samples is often rendered difficult owing to the complicated overlap of absorptions from so many different molecules. Accordingly, band assignments are typically made to a set of biomolecules such as lipids, proteins or nucleic acids (see the electronic supplementary material, figure S1) [38]. Moreover, intense water absorptions in the biochemically relevant area of the IR spectrum (4000–600 cm\(^{-1}\)) continue to necessitate the use of a synchrotron source and small path lengths for sensitive measurements of hydrated, live cells because of the high throughput required to transmit light through aqueous media, the cell and IR transparent windows [39]. Over the past decade, further complications in the successful application of FTIR spectroscopy to biological samples have stalled the uptake of this technique. Cells were found to cause Mie scattering and were susceptible to refractive index mismatching, both of which manifested as nonlinear baselines and shifting absorption peaks and intensities and which led to incorrect band assignments [40]. However, recent advances in methodologies along with a better understanding of these effects have led to the development of correctional algorithms and, in the process, demonstrated the true potential of FTIR spectroscopy for the analysis of biological specimens [41–43].

Using these advances, we recently reported live-cell synchrotron–FTIR spectroscopic detection of predominantly B-DNA in eukaryotic cells [35,44]. In that study, an en masse transition of the genomic DNA from B-DNA to an A-like DNA form was observed during dehydration in all eukaryotic cells examined. Importantly, the full reversal of the B to A transition was also detected upon rehydration [44].

This transition of the cellular B- to A-DNA upon dehydration had previously only been reported once [29]. Crystallographic work by Mohr et al. identified the same reversible transition of the DNA in Bacillus subtilis upon natural desiccation. In Mohr et al.’s [29] research, the transition to A-DNA was first reported as an effect of the binding of DNA-protecting proteins to the genome during desiccation of the spores. In a second publication, the A-DNA conformation was identified as the primary mode of resistance to DNA damage, with the DNA-protecting proteins providing the mechanism by which the transition was achieved [30].

This result, in combination with other reports of A-DNA having increased resistance to various forms of damage and with the spectroscopic evidence for the surprisingly reversible B- to A-DNA transition in eukaryotic cells, highlights the potential biological significance of the genome-wide B- to A-DNA transition. Furthermore, FTIR spectroscopy has previously detected conformational change in the DNA of cells in tissue sections via the symmetric phosphate vibration [45]. In this research, conformational differences as detected using FTIR spectroscopy were observed to be strongly prescriptive of stem cell regions. This demonstrates the ability of this technique to detect otherwise undiscoverable DNA conformational changes.

Although our previous research did not endeavour to maintain cell viability during dehydration and detection of the conformational transition [44], here we report on the detection of the B-—A—B-DNA transition in desiccation-resistant bacteria with no known mechanism for DNA stabilization. The bacteria were rehydrated from a dormant state with no loss of functionality, demonstrating the B- to A-DNA transition in a functional biological system.

2. Results

Figure 1 shows the raw and second derivative spectra of partially dehydrated A-DNA (red) and the same sample after rehydration to B-DNA (blue). In the backbone fingerprint region of A-DNA, the most prominent bands are observed

Figure 1. FTIR spectra (a) and calculated second derivative spectra (b) of double-stranded DNA in the dehydrated A-DNA conformation (dashed red) and after rehydration in the B-DNA conformation (solid blue). Shading indicates, from left to right, the anti-symmetric phosphate stretch, the symmetric phosphate stretch and the backbone carbon–carbon vibration. (Online version in colour.)
to be the anti-symmetric stretching vibration ($v_{\text{asym}}(\text{PO}_2^-) = 1237 \text{ cm}^{-1}$), the symmetric phosphate stretching vibration ($v_{\text{sym}}(\text{PO}_2^-) = 1088 \text{ cm}^{-1}$) and the backbone carbon–carbon vibration ($v_{\text{C–C}} = 966 \text{ cm}^{-1}$). Upon rehydration to B-DNA, these bands all undergo significant observable changes. The $v_{\text{asym}}(\text{PO}_2^-)$ red-shifts by $14 \text{ cm}^{-1}$ to $1223 \text{ cm}^{-1}$, the $v_{\text{C–C}}$ blue shifts to $969 \text{ cm}^{-1}$ and the $v_{\text{sym}}(\text{PO}_2^-)$ decreases in intensity. These shifts have previously been associated with the respective A- and B-conformations of DNA and used to monitor polymorphism [33]. Moreover, we recently demonstrated that the change in intensity of the $v_{\text{sym}}(\text{PO}_2^-)$ was due to a change in molar extinction coefficient, also related to the B–A transition [43]. Figure S1 in the electronic supplementary material highlights the various peaks in FTIR spectra of bacteria attributed to protein, carbohydrate, lipid and nucleic acid absorptions, as well as the high absorbance of water vibrations in the approximately $3300 \text{ cm}^{-1}$ and approximately $1625 \text{ cm}^{-1}$ regions. Because of this wide distribution of absorbances and the interference from the water absorptions, we focused mainly on the phosphodiester backbone region ($1300–950 \text{ cm}^{-1}$).

The spectra of a dehydrated and subsequently rehydrated viable *Proteus vulgaris* culture along with the calculated second derivative spectra over the region $1300–950 \text{ cm}^{-1}$ are shown in figure 2. By examining the second derivative spectra overlapping absorptions can be better resolved and hence changes in relative absorption intensity and position can be detected more clearly. The IR spectrum of the dehydrated, dormant culture demonstrated the three diagnostic A-DNA absorptions: $v_{\text{asym}}(\text{PO}_2^-) = 1239 \text{ cm}^{-1}$, $v_{\text{C–C}} = 966 \text{ cm}^{-1}$ and the less intense $v_{\text{sym}}(\text{PO}_2^-) = 1084 \text{ cm}^{-1}$. By contrast, the spectrum of the hydrated culture had absorptions indicative of B-DNA: $v_{\text{asym}}(\text{PO}_2^-) = 1222 \text{ cm}^{-1}$, $v_{\text{C–C}} = 969 \text{ cm}^{-1}$ and the more intense $v_{\text{sym}}(\text{PO}_2^-) = 1086 \text{ cm}^{-1}$. It should be noted that, for the $v_{\text{asym}}(\text{PO}_2^-)$ absorption, it is particularly difficult to detect this transition owing to overlapping amide III, phospholipid and phosphorylated protein vibrations [46], all of which are observed in the second derivative. This overlap has been reported before; however, the band at approximately $1238 \text{ cm}^{-1}$, which is red-shifted by approximately $14 \text{ cm}^{-1}$ to $1224 \text{ cm}^{-1}$ is specific for the DNA conformational change and is not from other molecular vibrational modes that appear in the vicinity of this band. Accordingly, in the dormant culture, the A-DNA signal ($1239 \text{ cm}^{-1}$) is more prominent than the B-DNA signal ($1222 \text{ cm}^{-1}$). The red-shift of the $v_{\text{sym}}(\text{PO}_2^-)$ and broadening of the $v_{\text{C–C}}$ have similarly been observed previously in cell spectra and are thought to be due to overlapping absorptions from other biomolecules as well as band-broadening due to steric crowding. These spectra are representative of repeated experiments using the *P. vulgaris* strain along with a second cycle of the rehydrated culture through a dehydration cycle.

*Escherichia coli* spectra also demonstrated A- and B-DNA absorptions with representative spectra of the dehydrated and rehydrated culture shown in figure 3. Again the $v_{\text{asym}}(\text{PO}_2^-)$ was observed to be split into two absorptions in the second derivative spectra of rehydrated samples, with the dehydrated culture exhibiting a more intense peak at $1241 \text{ cm}^{-1}$ indicative of A-DNA and the rehydrated bacteria having a significantly more intense $1220 \text{ cm}^{-1}$ absorption. In the underivatized spectra, the $v_{\text{asym}}(\text{PO}_2^-)$ shifts from $1240 \text{ cm}^{-1}$ dehydrated to $1232 \text{ cm}^{-1}$ when rehydrated in accordance with the A–B transition. Furthermore, the $v_{\text{sym}}(\text{PO}_2^-)$ ($1085 \text{ cm}^{-1}$) increases dramatically in intensity upon rehydration and the $v_{\text{C–C}}$ shifts from $966$ to $969 \text{ cm}^{-1}$. Following rehydration of *P. vulgaris* and *E. coli*, the number of viable bacteria was determined to be $5.2 \times 10^8$ and $8.3 \times 10^8$ CFU ml$^{-1}$, respectively.

The spectra of dehydrated and hydrated *Salmonella typhimurium* and *Pseudomonas aeruginosa* cultures showed that the $v_{\text{sym}}(\text{PO}_2^-)$ and $v_{\text{C–C}}$ absorptions consistently undergo
the typical absorption changes indicative of the A–B transition upon rehydration (electronic supplementary material, figures S2 and S3). The expected absorption changes of the $v_{\text{asym}}(\text{PO}_2^\cdot)$ absorption were not observed in the spectra of these two cultures; however, analysis of the second derivative spectra demonstrates the presence of peak splitting here in both the hydrated and dehydrated states that can be attributed to several overlapping non-DNA bands. It seems probable, considering the presence of other backbone A- and B-DNA diagnostic absorptions, that the $v_{\text{asym}}(\text{PO}_2^\cdot)$ shift was present but was more difficult to unambiguously detect in these species.

To further confirm the A- to B-DNA conformational change during rehydration, pairs of spectra of the four bacterial strains were used in a principal component analysis (PCA) model. The scores plot of the first two principal components (PC1 and PC2) is shown in figure 4, and the corresponding loadings plots are available in the electronic supplementary material, figure 4S. The PCA was conducted using the entire 1300–950 cm$^{-1}$ supplementary material, figure 4S. The PCA was conducted using the entire 1300–950 cm$^{-1}$ supplementary material, figure 4S. The PCA was conducted using the entire 1300–950 cm$^{-1}$ supplementary material, figure 4S.

The separation of hydrated from dehydrated bacteria spectra of all four species along PC1, in particular P. vulgaris, E. coli and P. aeruginosa, which cluster completely separately, demonstrated a common difference in these spectra. Examination of the PC1 loadings plot confirmed that the correlation was indicative of the A- to B-DNA transition.

3. Discussion

The perceived biological significance of A-DNA has previously been confined to transient transitions of short stretches of DNA from the native B-form [16]. Here, we show a concerted transition of the DNA of several species of bacteria to B-DNA during rehydration from a dormant, but functional state in which the DNA is in the A-form. This result clearly demonstrates that the DNA of these prokaryotes transitions to the A-form during desiccation and then returns to the native B-form in parallel with regaining full functionality upon rehydration. These observations come from using FTIR spectroscopy, a technique capable of monitoring in situ DNA conformation directly and non-invasively [44]. This capability is in contrast to other typical methods for examining DNA conformation such as crystallography and electron microscopy, which both require rigorous sample preparation and cannot be used for live-cell measurement. FTIR spectroscopy, on the other hand, has been shown to be very sensitive to DNA conformation with changes in absorptions that enable differentiation between A- and B-DNA as well as being directly proportional to concentration in the cellular environment [43].

One limitation of FTIR is that it does not have the required sensitivity to detect conformational changes on short tracts of DNA and only en masse transitions are possible. However, because DNA typically constitutes less than 5% of bacterial cell dry weight [47], clear elucidation of these changes to the FTIR spectrum relies on a fortuitous lack of overlapping absorptions in any given sample. In an earlier study, we reported the bands diagnostic of the B–A–B transition for several eukaryotic cells [44]. The main bands indicative of the B–A transition in cell spectra include the symmetric and anti-symmetric phosphate vibrations because these bands are not as heavily overlapped by protein absorptions (amide I: approx. 1650 cm$^{-1}$, amide II: 1550 cm$^{-1}$) and because the phosphate moiety is four times more concentrated that the nucleic base moieties.

In both the E. coli and P. vulgaris samples, all the diagnostic DNA backbone absorption changes were clearly observed, whereas in the S. typhimurium and P. aeruginosa spectra the band most susceptible to overlapping bands, $v_{\text{asym}}(\text{PO}_2^\cdot)$, did not show a clear A- to B-DNA transition. PCA, however, does differentiate between hydrated and dehydrated cultures successfully based predominantly on A- and B-DNA absorptions. The lack of clear DNA bands observed in some regions of the bacterial spectra is the result of overlapping bands masking the spectral changes associated with the B–A–B transition.

With this in mind, the unequivocal detection of the A- to B-DNA transition upon rehydration in several different strains...
of desiccation-resistant bacterial cells as well as in non-functional eukaryotic cell raises questions about the role of the en masse transition to A-DNA. Particularly because the A-form of DNA has previously been implicated in several studies as the conformational central to the evolution of nucleic acids [48,49] and in resisting DNA damage. A-DNA has been documented ex vivo as having increased resistance to damage by UV [26,27], desiccation [29] and radio-sensitization [28] and is the assumed conformation taken on by *B. subtilis* spores in a concerted protein–DNA response to dehydration [30].

Because of the lack of appropriate methodology, A-DNA has not previously been considered the conformation of DNA prevalent during the desiccation and survival of prokaryotes. Here, we have used FTIR spectroscopy to directly and non-invasively monitor a B–A–B transition of the DNA in four species of bacteria, demonstrating a newly found significance of this conformation and the potential central role of A-DNA in cells under stress. Observation of the predominantly A-form DNA in prokaryotes necessitates further consideration of the B–A conformational transition and its role in defence mechanisms as well as its potential evolutionary role with regard to the ongoing debate over the origins of nucleic acids and their central role in life [50].

4. Material and methods

4.1. Bacteria culture

*Proteus vulgaris* or *E. coli* were cultured in Luria-Bertani broth at 37°C, with constant agitation, until the optical density at 600 nm (OD<sub>600</sub>) was approximately 0.1, which corresponds to approximately 3.5 × 10<sup>7</sup> CFU ml<sup>−1</sup>. The cell pellets were harvested by centrifugation at 3500×g and subsequently washed in sterile Dulbecco's phosphate-buffered saline (140 mM NaCl, 2.68 mM KCl, 4.23 mM Na<sub>2</sub>HPO<sub>4</sub>, 10 mM KH<sub>2</sub>PO<sub>4</sub>, pH 7.5). The washed cell pellets were finally resuspended in 1 ml of sterile isotonic saline. A 100 μl aliquot of the resuspended cells in isotonic saline was dehydrated onto CaF<sub>2</sub> discs at room temperature for 1 h before scratching off the CaF<sub>2</sub> windows using a clean metal spatula and then deposited onto the ATR diamond. Spectra (50 co-added scans at 4 cm<sup>−1</sup> resolution) were acquired over the region 4000–600 cm<sup>−1</sup> and ratioed to a 50-scan background recorded of the diamond window in air. The culture was then gradually rehydrated with phosphate-buffered saline as previously described and spectra were collected systematically during the rehydration [44]. Reference spectra of B- and A-DNA (from calf thymus; Sigma, St. Louis, MO, USA) were collected on the same system with the same parameters as previously described [44]. Spectra were baseline corrected using a concave rubberband method with 12 iterations and 25 baseline points and then vector normalized within experimental sets across the 1435–1140 cm<sup>−1</sup> region to correct for changes in concentration during dehydration and rehydration. For between species analysis, all spectra were vector normalized together across the 1435–1140 cm<sup>−1</sup> region. Second derivatives were calculated using nine smoothing points for the Savitsky–Golay process.

4.2. Attenuated total reflection–Fourier transform infrared spectroscopic measurement

For FTIR–attenuated total reflection (ATR) spectroscopy, *S. typhimurium* and *P. aeruginosa* cells were collected by swabbing from an agar culture and directly deposited onto the diamond window of a Golden Gate single bounce micro-ATR module coupled to a Bruker IFS Equinox FTIR spectrometer. The cell populations were dehydrated at room temperature for 1 h before measurement. *Proteus vulgaris* and *E. coli* cells were carefully scratched off the CaF<sub>2</sub> windows using a clean metal spatula and then deposited onto the ATR diamond. Spectra (50 co-added scans at 4 cm<sup>−1</sup> resolution) were acquired over the region 4000–600 cm<sup>−1</sup> and ratioed to a 50-scan background recorded of the diamond window in air. The culture was then gradually rehydrated with phosphate-buffered saline as previously described and spectra were collected systematically during the rehydration [44]. Reference spectra of B- and A-DNA (from calf thymus; Sigma, St. Louis, MO, USA) were collected on the same system with the same parameters as previously described [44]. Spectra were baseline corrected using a concave rubberband method with 12 iterations and 25 baseline points and then vector normalized within experimental sets across the 1435–1140 cm<sup>−1</sup> region. Second derivatives were calculated using nine smoothing points for the Savitsky–Golay process.

4.3. Principal component analysis

Baseline corrected and normalized spectra were converted to JCAMP files and imported into the UNSCRAMBLER software suite (v. 9.2; CAMO, Oslo, Norway) and collated into one file. PCA was conducted with five principal components over the entire sugar–phosphate backbone fingerprint region (1300–950 cm<sup>−1</sup>) and rehydration. For between species analysis, all spectra were vector normalized together across the 1435–1140 cm<sup>−1</sup> region. Second derivatives were calculated using nine smoothing points for the Savitsky–Golay process.

Funding statement. An Australian Research Council (ARC) Discovery grant (DP120100431) financially supports this work. B.R.W. is supported by ARC Future Fellowship grant no. FT120100926. D.R.W. is supported by an Australian Synchrotron Postgraduate Award. We acknowledge Mr Finlay Shanks for instrumental support.

References


