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Adding a New Dimension to DNA Melting Curves.

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Abstract. - Standard DNA melting curves record the separation of the two strands versus temperature, but they do not provide any information on the location of the opening. We introduce an experimental method which adds a new dimension to the melting curves of short DNA sequences by allowing us to record the degree of opening in several positions along the molecule all at once. This adds the spatial dimension to the melting curves and allows a precise investigation of the role of the base-pair sequence on the fluctuations and denaturation of the DNA double helix. We illustrate the power of the method by investigating the influence of an AT rich region on the fluctuations of neighboring domains.

DNA melting, i.e. the separation of the two strands of the double helix, and its reverse process hybridisation are ubiquitous in biology, in vivo for instance for DNA transcription in the reading of the genetic code of a gene, as well as in vitro in biological laboratories for PCR (Polymerase Chain Reaction) or the use of DNA microarrays. This is why DNA melting has been extensively studied even in the early days of DNA structural studies [1]. An approximate understanding of the melting curves of long DNA segments, with thousands of base pairs, can be provided by simple statistical physics models, using empirical parameters because, at this large scale, the subtle effects of the base pair sequence are smoothed out. Understanding the fluctuations and melting of short DNA fragments of a few tens of base pairs with a high degree of heterogeneity is much more challenging. And it is also very important because this size is the scale at which the genetic code can be resolved. This would have some significant biological consequences to unravel the processes by which specific binding sites are recognised by proteins, drugs, mutagens and other molecules. This would also have a lot of practical importance in the design of the PCR primers which are used everyday in most of the biological laboratories [2].

The two kind of base pairs which exist in the DNA double helix have different thermal stability, the AT pair, bound by two hydrogen bonds, being weaker than the GC pair bound by three hydrogen bonds. This explains why the melting curve of a heterogeneous DNA sequence, which shows the fraction of open pairs as a function of temperature, can exhibit complex features. Those curves are easy to record experimentally because the UV absorbance of a DNA solution increases drastically when the bases are unstacked, which is

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the case in the broken regions of the molecule. But such a curve only provides an integral information on the open fraction of base pairs. Getting more local information requires involved methods. Using a clever choice of sequences such that single strands can form hairpins, and a combination of heating and quenching, Montrichok et al. [3,4] managed to get some data on the melting process of short DNA sequences, detecting whether they open at one end or by starting with an open bubble in the centre. The kinetics of proton–deuterium exchange for the protons involved in the hydrogen bonds within pairs, coupled with NMR studies to detect the location of the exchanged protons, can also provide partial information on the spatial aspect of DNA fluctuations, at the expense of heavy experiments [5]. Another approach relies on special molecular constructs which attach a fluorophore and a quencher to DNA to detect its local opening at a particular site [6, 7]. Only one position can be monitored, and one cannot exclude local perturbations of the fluctuations by the large residues attached to the DNA.

Due to their importance, the statistical and dynamical properties of DNA fluctuations and their relation to biological functions have been the subject of many theoretical studies [8–10]. These studies raised a debate on the role of statistical properties and dynamical phenomena in connection to biological function. But the validity of those theoretical approaches can only be tested if one can compare their predictions to measurements of the local fluctuations of the molecule. Moreover the study of dynamical and conformational phenomena in DNA requires a method not only able to give a precise local information, but also able to provide coupled information of events along the chain. In this letter we present an original method that can provide a mapping of the strength of the fluctuations of the double helix as function of their position along the sequence. This adds a new dimension, space, to the traditional melting curves. Our approach does not require special molecular constructs like those using dyes or fluorophores. Instead it uses DNA itself to report on its internal state and gives a snapshot of the opening of DNA at each guanine site at once.

Fig. 1: Local melting curves for different sections of a DNA sequence (S1). Symbols represent the thermal evolution of the rate $R_{Fpg}/R_{Pip}$ exhibited by two guanines $G_1$ and $G_2$, used as probes located along the molecule. This ratio reports on the closing probability of each guanine at a given temperature. The lines are the fitting curves described in the paper and the insets plot the derivative of the fitting curve for each probe. They highlight the fine structure of the melting curves and point out the existence of different local conformational transitions along the DNA sequence.

The method relies on the oxidative chemistry of the guanine bases G, and their propensity to be ionised by a two step resonance excitation [11] from a strong UV laser pulse. Two guanine modifications, oxazolone and 8-oxo-7,8-dihydro-2-oxoguanine (8-oxodG) have
been identified as the major one-electron oxidative DNA lesions. Their formation depends on the local DNA conformation and on the charge-transfer efficiency, which is affected by local fluctuations [12–14]. While oxazolone is the unique product resulting from one-electron oxidation of the free 2′-deoxyguanosine, 8-oxodG appears as soon as the nucleoside is incorporated in a helical structure. Hence the measurement of the relative yield of these photoproducts at each G site (labelled $R_{Fpg}/R_{pip}$ due to the method used for its detection [15]) tells us whether this G was in an helical structure (closed) or whether it was open when the molecule was hit by the laser pulse. As the experiment is not performed on a single molecule but in solution the results are obtained on a statistical ensemble and they give a signal representative of the probability that each G site is closed at the temperature of the study. Standard biological methods can be used to measure the relative yield of the production of oxazolone and 8-oxodG [15]. However, it is important to notice that the value of $R_{Fpg}/R_{pip}$ should not be considered as a quantitative measure of the local closing probability because it is also affected by the configuration of the DNA molecule near the probe, which depends slightly on the sequence and influence the charge transfer. Only the temperature dependence of this ratio for a given probe can be analysed quantitatively [16].

By splitting the sample into several aliquots, the measurement can be performed at different temperatures, which allows us to produce a set of melting curves for each guanine in the sequence. Figure 1 shows the results of a UV laser irradiation analysis for two guanines, labelled S1-G1 and S1-G2, belonging to a test sequence S1 (details of the sequence are provided in the figure caption and in Figure 3). It demonstrates how our method reports two complementary curves at once for the same single DNA sequence, adding a valuable information of the state of the system. The results that it provides changes the view in which DNA denaturation can be studied, adding the spatial correlations to the notion of local conformation (melted or packed helix).

In order to quantitatively analyse the measurements shown in Fig. 1 we have fitted the experimental curves by the function $f(T) = A - B_1 \tanh[C_1(T - T_1)] - B_2 \tanh[C_2(T - T_2)]$, selected according to the shape of the curves, particularly that of the probe S1-G1. Once the
optimal parameters are determined, we plot $df(T)/dT$ for each probe (right part of Fig. 1), to highlight the fine structure of each melting curve. Figure 1 clearly shows that, at the level of probe S1-G1, the melting occurs in two steps, with a precursor at $T_2 = 45.7^\circ C$ while the full melting is achieved at $55.0^\circ C$. Although a slight precursor effect can be detected for probe S1-G2 it is very weak and hardly visible on the figure. The fit of the data indicates that it occurs at $T_2' = 51.8^\circ C$, very close to the full melting detected at $T_1' = 54.4^\circ C$ for this probe. Interestingly these results show that the conformational melting can follow different paths in various parts of the same short sequence.

For longer sequences such as sequence S2 (Fig. 3), the existence of several guanine probes, located in some interesting domains, allows us to obtain a collection of snapshots for the local state of the system that can be combined to build a three-dimensional melting profile like the one shown in Figure 2. This 3D plot shows the ratio $R_{fg}/R_{pp}$ for a particular DNA sequence containing four guanines (S2-G1 to S2-G4) that are monitored as probes. The study as a function of temperature provides a melting profile for each probe, together with a view of the spatial correlations along the full sequence.

![Fig. 3: Sequences of the DNA fragments investigated in this study. S1 and S2 are artificial sequences containing a large TATA box and guanines on the 5'-3' strand, used as probes. S3 is a shorter control sequence that eliminates the TATA motif. All sequences are completed by GC-rich terminal domains (marked as dotted boxes) to stabilise them and ensure proper closing of these short DNA helices.](image)

In order to get a more precise insight on the origin of the two-step melting detected at the level of probe S1-G1, and to relate it to the effect of bubble nucleation in DNA, we have studied several artificial sequences specially designed and synthesised [17] to investigate possible non-local effects of the fluctuations. They are shown on Figure 3. Sequences S1 and S2 contain a 10 base-pair-long AT-rich fragment which is analogous to the “TATA-box”, a motif that exists in the transcription-initiation regions of the genes of various species. As the AT base pairs bound by only two hydrogen bonds are weaker than the GC pairs, these segments are expected to exhibit large fluctuations even at biological temperature because they are closer to their melting temperature.

All sequences S1, S2 and S3 contain different guanines labeled Sn-Gx, where $n$ refers to the sequence and $x$ to the particular guanine. They are spread along the strand, and according to the basis of our method, act as probes informing us about the local structural state at each temperature. As mentioned above the level of the $R_{fg}/R_{pp}$ signal that we record depends on the structure in the vicinity of the guanine of interest. This is why, in order to allow a quantitative comparison between different sequences, we have selected guanines with the same environment. For instance guanines S1-G1, S2-G1, S3-G1 are all parts of the sequence CGA, and guanines S1-G2 and S2-G2 are part of the sequence AGT.
The major difference between sequences S1 and S2 lies in the length of the domain that separates probe G1 from the AT-rich region. This “buffer region”, which is an heterogeneous domain with AT and GC pairs, has been extended from 7 base pairs in sequence S1 to 13 base pairs in sequence S2. In sequence S3 we have eliminated the large “TATA box” to keep only short AT-rich domains around probe S3-G1. In all cases these short DNA molecules are terminated by GC rich domains which act as clamps to prevent large fluctuations of the free ends of the molecules, and hold the two strands together even when we heat the sample up to 60°C. [18].

The variation versus temperature of the ratios \( \frac{R_{F,pg}}{R_{pip}} \) for all the guanine probes studied in sequences S1, S2, S3, are summarised in Fig. 4. The comparison of the various curves gives some clues to understand the differences between the two melting curves of probes S1-G1 and S1-G2 discussed above, and points out some interesting features of DNA fluctuations, which can be revealed by an experimental method able to record local melting profiles. When they are analysed in the context of the particular sequences that we studied, the curves suggest three important properties of DNA fluctuations:

i) a large AT-rich domain undergoes very large fluctuations, even at room or biological temperature, and therefore tends to easily form an “open bubble”.

ii) there is a minimum size of the AT-rich domain that allows the formation of such a bubble.

iii) the influence of such a bubble does not only affect its immediate vicinity, but extends to some distance.

Let us see how these statements are supported by our results.

As our method relies on the ionisation of the guanines, we do not directly measure the opening of the AT pairs but their fluctuations can be inferred from their influence on the adjacent guanines. Although we stressed that quantitative comparisons cannot be made between different probes because the signal that we record depends on the local structure of DNA, the very small value of \( \frac{R_{F,pg}}{R_{pip}} \) for probe S1-G2 is nevertheless a strong indication that the closing probability of this guanine, which lies next to a series of 10 AT pairs is very low even at room temperature. This can be understood as an effect of the strong tendency of the large AT rich region to open into transient bubbles, called “premelting phenomena”, starting at physiological temperatures [19–22], which certainly affects the base pair which is right next to it. Probe S3-G1 is also surrounded by AT-rich regions, but its closing probability deduced from the corresponding value of \( \frac{R_{F,pg}}{R_{pip}} \), shown on the top panel of Fig. 4, appears to be much higher that for probe S1-G2. This indicates that the fluctuations of the five-base-long AT regions which are next to probe S3-G1 are not sufficient to form open bubbles that would promote the opening of this probe. This is in agreement with the existence of a minimum size needed to allow the formation of a bubble [3].

The striking point is that the large fluctuations of the TATA box do not only perturb the adjacent geometry but also induce conformational changes that distort the closed packed helicoidal structure in regions distant from the bubble nucleation segment, giving rise to pre-melting intermediate structural states that coexist in consonance with the nucleated bubbles. This shows up in the two-step denaturation that we observed for probe S1-G1, as discussed above. Although this probe is 7 base-pairs away from the TATA box, Fig. 4 shows that precursor effects appear well below the full denaturation of this probe. Those precursors are also visible in Fig. 4. Note again how the melting at the level of probe S1-G1 differs from a simple sigmoidal curve, particularly in the temperature range from 37°C to 52°C. It is tempting to assign them to the influence of the fluctuations of the TATA, which grow when the temperature is increased and might influence the opening structure and fluctuations of the double helix even rather far away. But, to confirm such an assignment control experiments are necessary. Their results are shown on the bottom panel of Fig. 4 which shows the thermal variation of the ratio \( \frac{R_{F,pg}}{R_{pip}} \) for several guanine probes in sequence S2.

In this sequence the TATA box is present, as in sequence S1, but the length of the buffer
region that separates it from probe S2-G1, which has the same local structure as probe S1-G1, has been increased. Moreover this buffer region is strengthened because it contains several GC pairs, including two adjacent ones at the site of probe S2-G3. In this sequence the existence of the large fluctuations of the TATA box are attested by the very low closing probability of probe S2-G2, similar to what is observed for probe S1-G2. But, contrary to what was observed for probe S1-G1, the denaturation of probe S2-G1 does not show any significant precursor effect. Similar one-step transitions are also observed for probes S2-G3 and S2-G4.

In summary, these experiments clearly suggest that thermal fluctuations, which are stronger in AT-rich tracks, induce bubble nucleation phenomena and structural changes that affect not only the local geometry and dynamics of DNA in the breathing portion, but also to some distance along the helix. How this happens is highly dominated by the characteristics of the fragment sequence as well as the length of the sections involved.

Fig. 4: The top panel shows the melting profiles for the probes studied in sequences S1 and S3. Sigmoidal non-linear fits (dotted lines) have been used to describe the evolution of probes S1-G1 and S3-G1. Both fits have been performed with the same identical functional form. While the response of S3-G1 agrees with a one step standard melting transition, the result for S1-G1 suggests a two steps transition, with a premelting region, clearly shown in Fig. 1. The bottom panel shows the ratio $R_{F_{pp}}/R_{p_{pp}}$ for the guanine probes of sequence S2 versus temperature. For probe S2-G1, the premelting effect has been suppressed due to the existence of a longer more stable intermediate buffer region.
Those results are only accessible because we have introduced a method which is able to provide a spatial information that was not available until now for the study of local melting in short DNA fragments. Further studies are certainly necessary to confirm and precise the non-local effect of the fluctuations of large AT-rich regions that we have presented in this letter. They become possible with the “three-dimensional melting curves” that can now be measured.

Finally, we would like to emphasise the importance of taking into account the structural modifications induced by bubble dynamics in terms of DNA-protein binding interactions, Transcription Factor recognition or DNA-drug binding. Further studies of local fluctuations in DNA may be of significant importance in the analysis of these biological phenomena.

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[15] To measure the relative yield of the production of oxazolone and 8-oxodG we use common chemical methods used in standard analytical biology: Piperidine cleaves DNA at the sites where oxazolone is the resulting lesion whereas Fpg protein cleaves it as sites which have evolved to 8-oxodG. One end of the molecule is marked by a radioactive marker, and, after cleavage by piperidine or Fpg, performed on different aliquots at the same temperature, the lengths of the fragments that carry the marker are quantified by gel electrophoresis. The radioactively labelled fragments of length $l$ are those which were cleaved at a guanine situated at distance $l$ from the marked end. The ratio $R_{Fpg}/R_{pip}$ of the number of radioactively labeled fragments of length $l$ produced by Fpg cleavage or by piperidine cleavage indicates the probability of closing for this particular guanine, which is unambiguously identified by its distance from the marked end. As the gel image contains the data for fragments of all lengths, a single experiment measures a magnitude proportional to the opening probability at all guanine sites.
[16] The experimental errors are within the size of the data points shown in all graphs.
[17] DNA Oligos have been HPLC-RP purified and artificially synthesized by Eurogentec.
[18] We should point out that the spatial resolution near the end of the strands in the gel is usually not optimal to separate and resolve two adjacent guanines in this region. In principle the guanines in these terminal parts could also be monitored in our analysis, but such limitation makes the study inefficient.
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