

Disruption of Energetic and Dynamic Base Pairing Cooperativity in DNA Duplexes by an Abasic Site

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Abstract

DNA duplex stability arises from cooperative interactions between multiple adjacent nucleotides that favor base pairing and stacking when formed as a continuous stretch rather than individually. Lesions and nucleobase modifications alter this stability in complex manners that remain challenging to understand despite their centrality to biology. Here we investigate how an abasic site destabilizes small DNA duplexes and reshapes base pairing dynamics and hybridization pathways using temperature-jump infrared spectroscopy and coarse-grained molecular dynamics simulations. We show how an abasic site splits the cooperativity in a short duplex into two

segments, which destabilizes small duplexes as a whole and enables metastable half-dissociated configurations. Dynamically, it introduces an additional barrier to hybridization by constraining the hybridization mechanism to a step-wise process of nucleating and zipping a stretch on one side of the abasic site and then the other.

Significance

Biological functions of nucleic acids rely on the stability and dynamics of base pairing interactions. While hydrogen-bonding contacts describe a specific base pair, base pairing stability within duplexes is primarily shaped by the continuous stretches of stacking interactions between adjoining nucleobases that give rise to highly cooperative base pair breaking and formation. Damage or epigenetic modification of a given nucleobase can disrupt this cooperativity and may change the properties of adjoining base pairs. Our investigation used time-resolved spectroscopy and computational modeling to directly reveal how an abasic site alters the stability, kinetics, and dynamics of base pairing in the duplex.

Introduction

Numerous modifications to canonical base pairs, including epigenetic chemical modifications, mismatches, and lesions, alter local base pairing interactions in double-stranded DNA which may in turn shift global duplex thermodynamic stability by interrupting cooperative interactions. Cooperativity refers to the non-additive energetic advantages of forming base pairs in a stretch of multiple adjacent nucleotides concertedly rather than individually(1). This phenomenon underlies the quantitatively accurate nearest neighbor (NN) models of DNA thermodynamic stability (2, 3). While cooperativity in base pairing is understood to be rooted in the stacking interactions between nucleobases, the molecular factors bridging local perturbations and global destabilization are not fully understood. Identifying these factors require site-specific characterization of the hybridization free energy landscape and duplex structural dynamics over wide time-windows. This motivates the combination of temperature-jump infrared (T-jump IR) spectroscopy and coarse-grained molecular dynamics (MD) simulations that are employed in this work to access thermodynamic, kinetic, and dynamic insight into how an abasic site disrupts base pairing and stacking in short oligonucleotides.

The loss of a nucleobase to form an abasic site (apurinic/apyrimidinic or AP-site) is among the most naturally abundant DNA modifications due to spontaneous or enzymatic cleavage of the

glycosidic bond.(4, 5) Previous work showed that an AP-site destabilized the DNA duplex in a sequence-dependent fashion(6-11) that was correlated with the identity of base pairs adjacent to the modified site; however, an AP-site may also disrupt base pairing well beyond nearest-neighbors.(12) Since duplexes containing an AP-site retain a B-form conformation on average,(13-15) an AP-site influences duplex stability and dynamics through disruption of cooperative interactions rather than a large-scale structural change.

We characterize the disruption of cooperativity in various sequence contexts by investigating the impact of central AP-sites in three 11-mer template oligonucleotides that are expected to exhibit wide variation in dynamic behavior due to different relative placement of G:C and A:T base pairs.(16-22) T-jump IR spectroscopy separately monitors changes in A:T and G:C base pairing from nanoseconds to milliseconds following a laser T-jump that induces duplex melting. MD simulations employing the 3-site-per-nucleotide (3SPN.2) coarse-grained model(16, 23) are used to parameterize a kinetic Markov state model (MSM)(24, 25) that allows us to interpret the experiments in terms of the temperature-induced reshaping of the duplex free energy landscape. Our results indicate that duplex destabilization primarily stems from the loss of base pairing and stacking interactions around the AP-site as well as disruption of base pairing cooperativity within the duplex. The AP-site splits the base pairing dynamics of the duplex into two segments that must each overcome a significant free energy barrier to hybridize. Although our study is limited to oligonucleotides, length-dependent trends in duplex destabilization behavior enable us to address the impact of an AP-site in biological DNA.

Results & Discussion

AP-site destabilizes the DNA duplex through disruption of base pair cooperativity

The DNA sequences shown in Fig. 1a were chosen to vary the position of three or four stabilizing G:C base pairs within an otherwise A:T sequence. Sequence “CGCcap” places them on one end to create asymmetric base pairing stability, “CCends” places them on both ends to minimize terminal base pair fraying,(19) and “GCGcore” centers them to promote frayed A:T terminal base pairs.(19, 20, 26) For each oligonucleotide, we compare the hybridization of the wild-type (WT) sequence to its complement with one containing a central AP-site at the 6th base position (AP6).

Experiments showing how a central AP-site destabilizes the duplex are presented in Figs. 1b-d and S1-S5. Duplex melting curves obtained by Fourier-transform infrared spectroscopy (FTIR) show a 20–30 °C reduction in melting temperature (T_m) that equates to a drop of 20-30 kJ/mol in the dehybridization free energy between AP6 and WT sequences ($\Delta\Delta G_{d,37}^o = \Delta G_{d,37}^o(AP) - \Delta G_{d,37}^o(WT)$), consistent with previous studies.(6, 8, 11) Calculations of $\Delta\Delta G_{d,37}^o$ using Santa Lucia's NN model(2, 27) merely removing the two central NN interaction parameters involved with the AP-site only accounts for a portion of the experimental value, suggesting that duplex destabilization is not simply additive in the NN energies (Fig. 1c, Section S1.4). An additional free energy penalty affecting both the dehybridization enthalpy (ΔH_d^o) and entropy (ΔS_d^o) must arise from the AP-site (Fig. S1). We also find that $\Delta\Delta G_{d,37}^o$ is ~ 5 kJ/mol larger for CCends-AP6 than CGCcap-AP6 even though the modified and adjacent base pairs to the AP-site are identical in each sequence, suggesting that the oligonucleotide sequence further from the AP-site contributes to $\Delta\Delta G_{d,37}^o$.

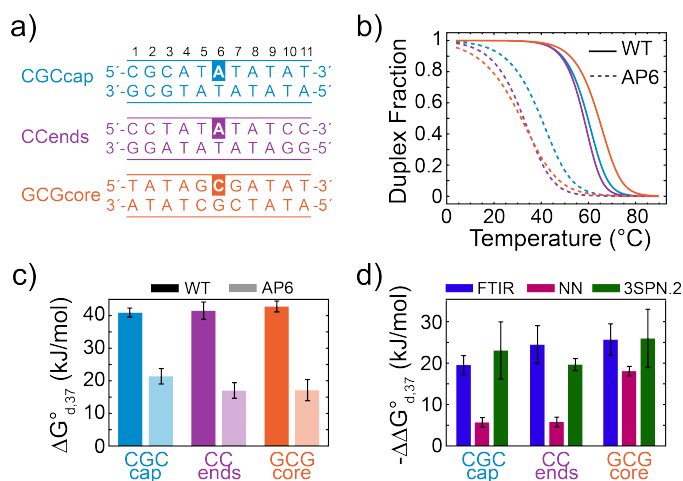


Figure 1. DNA duplex destabilization induced by an AP-site. **a**, CGCcap, CCends, and GCGcore sequences studied in this work, identifying the 6-position of the AP-site along the top strand (shaded). **b**, Duplex melting curves determined from two-state analysis of an FTIR temperature series for WT (solid lines) and AP6 (dashed lines) sequences (see Section S1.3). **c**, Duplex dehybridization free energy at 37°C ($\Delta G_{d,37}^o$) determined from FTIR data for WT (dark) and AP6 (light) sequences. **d**, Drop in $\Delta G_{d,37}^o$ for each AP6 sequence with respect to WT sequence ($\Delta\Delta G_{d,37}^o = \Delta G_{d,37}^o(AP) - \Delta G_{d,37}^o(WT)$) compared to those determined from the NN model (Section S1.4). Also shown are the change in dehybridization Helmholtz free energy ($\Delta\Delta F_{d,37}^o$) obtained from melting curves generated from 3SPN.2 MD simulations. FTIR and 3SPN.2 error bars correspond to 95% confidence intervals from fits to a two-state model (Fig. S1) while NN error bars correspond to the reported error in the model.(2)

MD simulations with the 3SPN.2 model and well-tempered metadynamics (WTMetaD) were used to gain insight into the molecular factors governing the thermodynamic stability of various oligonucleotide structures. Structurally 3SPN.2 predicts that duplexes with the AP-site are virtually unchanged from the B-form conformation of the WT duplex and the nucleobase opposite the AP-site is predominantly intrahelical, consistent with previous experimental studies and all-atom MD simulations (Figs. S6-S7).(13, 14) The extrahelical configuration of GCGcore-AP6 is negligible, yet there is minor population for CCends-AP6 (0.3%) and CGCcap-AP6 (0.8%). This sequence-dependence presumably arises from the 2-fold larger cross-stacking energy between guanine nucleobases relative to thymines in 3SPN.2.(23) The stability of each duplex was assessed using simulations as a function of temperature to compute duplex-fraction melting curves and the Helmholtz free energy for dehybridization $\Delta\Delta F_{d,37}^o$ (Figs. 1d, 2 and S8-S13). The simulations show a similar destabilization by the AP-site as in experiment, suggesting that the coarse-grained model is properly accounting for the duplex destabilization by the AP-site. We note that the simulated sequence-dependence of $\Delta\Delta F_{d,37}^o$ differs from the experimental $\Delta\Delta G_{d,37}^o$, but any differences lie within the uncertainty of the data. Previous studies have verified the accuracy of 3SPN.2 for predicting T_m in canonical duplexes with near-physiological salt concentrations ($[\text{Na}^+] = 120$ mM),(23) but we find that they are consistently 5-10 °C lower than experimental values, which is likely due to the high salt concentrations ($[\text{Na}^+] = 600$ mM) used in this work.

The computed hybridization free energy profile (FEP) as a function of the number of intact base pairs (n_{bp}) is compared for WT and AP6 sequences in Fig. 2a. These profiles overlap for the early steps in hybridization ($n_{bp} = 0-5$ for CGCcap and CCends and $n_{bp} = 0-2$ for GCGcore) and span the barrier to nucleating the duplex ($\Delta F_h^\ddagger$) at $n_{bp} = 2-3$. The free energy of duplex formation continues downhill with increasing n_{bp} in the WT sequences, as expected for a two-state nucleation and zipping process,(28-30) but the AP6 sequences display a minimum at $n_{bp} = 5$ and must pass through an additional free energy barrier to fully hybridize ($\Delta F_{h,2}^\ddagger$). The dominant base pairing configurations of AP6 sequences at $n_{bp} = 5$ are those with one side of the AP-site fully intact and the other side completely dissociated, indicating a stepwise process of hybridizing one segment and then the other (Fig. S8). The nucleobase opposite the AP-site is predominantly intrahelical at $n_{bp} = 5$. Therefore, an additional free energy penalty for base pair nucleation on the second side of the AP-site contributes to $\Delta\Delta G_{d,37}^o$, which is absent in the NN model.

We also observe that $\Delta F_{h,2}^\ddagger < \Delta F_h^\ddagger$, reflecting the difference in hybridization mechanism for the first and second segments. The temperature-dependence of $\Delta F_h^\ddagger$ and $\Delta F_{h,2}^\ddagger$ indicates each arises from an entropic penalty that is partially balanced by a favorable internal energy change for base pairing (Fig. S11). The reduction in translational and orientational entropy for bimolecular association contribute to $\Delta F_h^\ddagger$ but not $\Delta F_{h,2}^\ddagger$, where conformational entropy of the partially bound chains dominates. For a $n_{bp} = 5$ configuration with one fully hybridized segment, the separation of the unbound segment base pairs adjacent to the AP-site is more constrained than the terminal base pairs. As a result, nucleation of the second segment usually occurs by forming a base pair adjacent to the AP-site, as shown below.

Additional sequence dependent details can be observed in the free energy surfaces (FESs) in Fig. 2b, which are computed at the 3SPN.2-determined melting temperature ($T_{m,MD}$) and projected into the average base pair separation (r_{bp}) on either end of the duplex. FESs are binned and plotted on a r_{bp}^{-1} scale to better resolve different basins within the duplex state. Here 5'- and 3'-sides of the duplex refer to the top strand in Fig. 1a. Fully intact sides have $r_{bp} < 0.7$ nm whereas well-separated strand segments have $r_{bp} = 3-5$ nm. Addition of an AP6 site creates metastable configurations with only the 5'- or 3'-segment hybridized. This provides a structural rationalization for the emergence of the $\Delta F_{h,2}^\ddagger$ barriers observed in Fig. 2a. The relative stability of 5'-hybridized-to-3'-hybridized configurations qualitatively follows the expected stability of those segments predicted by the NN model (Table S4).

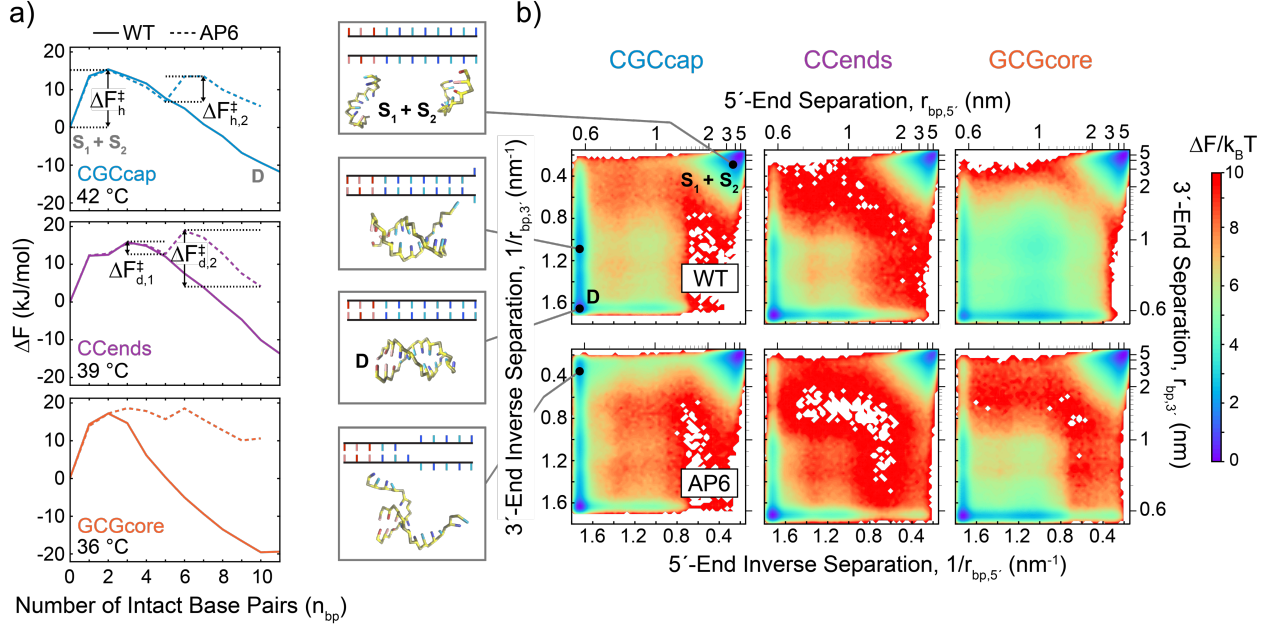


Figure 2. Influence of AP-site on base pairing free energy landscape. **a**, Free energy profiles as a function of the number of intact base pairs, $\Delta F(n_{bp}) = -RT \ln P(n_{bp})/P(0)$, for WT (solid lines) and AP6 (dashed lines) sequences extracted from 3SPN.2 MD simulations with WTMetaD at temperatures 10–20°C below $T_{m,MD}$ of each WT sequence. Base pairs are determined using a 0.7 nm radial cutoff between the center-of-mass of a nucleobase site and that of its complement. Hybridization barriers in n_{bp} for forming the first ($\Delta F_h^\ddagger$) and second ($\Delta F_{h,2}^\ddagger$) base pair segments of the AP6 sequence are indicated for CGCcap. Barriers along the reverse dehybridization direction for dissociating the second ($\Delta F_{d,2}^\ddagger$) and first ($\Delta F_{d,1}^\ddagger$) base pair segments are shown for CCends-AP6. **b**, Free energy surfaces for all sequences plotted as a function of the average separation between base pairs 1, 3, and 5 on the 5'-end ($r_{bp,5'}$) and 7, 9, and 11 on the 3'-end ($r_{bp,3'}$). 5'- and 3'-ends refer to the top strand in Fig. 1a. Free energy (ΔF) is plotted relative to the minimum of the single-strand well ($S_1 + S_2$). Surfaces were generated using 2.5 million frames from 25 unbiased 10 μ s simulation trajectories near the 3SPN.2-determined melting temperature of each sequence and are binned and plotted on a r_{bp}^{-1} scale to better resolve different base pairing configurations. Addition of an AP6-site creates local minima (top left and bottom corners) in the free energy landscape corresponding to configurations with only the 5'- or 3'-end hybridized that make up the $n_{bp} = 5$ minimum in panel **b** (Fig. S12).

Direct observation of internal nucleation barrier with T-jump spectroscopy

We experimentally test for the presence of $\Delta F_{h,2}^\ddagger$ and a half-hybridized metastable state by characterizing duplex dehybridization with time-resolved T-jump IR spectroscopy.^(19, 31, 32) We equilibrate the sample at a temperature 15°C below T_m , where most of the DNA is duplexed and use a 15°C optical T-jump to heat the solution within 7 ns and promote duplex dehybridization. Changes in base pairing interactions are probed at varying time delays after the T-jump using heterodyned dispersed vibrational echo spectroscopy (HDVE),⁽³¹⁾ which provides a transient IR

spectrum reporting on changes in nucleobase ring and carbonyl vibrational bands (Fig. 3a). The transient spectrum that we report is the change of the spectrum at a given T-jump delay relative to the initial temperature, T_i . Each nucleobase contains a distinct IR fingerprint of ring and carbonyl vibrations(33, 34) that are reshaped upon base pair formation,(20, 35) providing separate probes for G:C and A:T base pairing using the guanine band at 1550 cm^{-1} and adenine band at 1605 cm^{-1} (Figs. S14-S16).

T-jump IR measurements of WT sequences in Fig. 3b show two distinct kinetic components on $\sim 100\text{ ns}$ (τ_1) and $\sim 100\text{ }\mu\text{s}$ (τ_3) timescales, followed by thermal relaxation of the sample back to T_i at $\sim 1\text{ ms}$. The transient IR spectra are well described by global lifetime fitting with exponentially damped spectral components, and similar results are obtained from inverse-Laplace-transform rate distribution spectra (Figs. S17-S19). These processes have previously been characterized in similar sequences and they are illustrated in Scheme 1a through changes to the hybridization free energy profile induced by the T-jump.(18, 19, 36, 37) The T-jump destabilizes the duplex state, resulting in barrierless terminal base pair fraying (τ_1). The signal change observed during τ_1 increases in order from CCends-WT to CGCcap-WT to GCGcore-WT due to an increased propensity for fraying in A:T base pairs relative to G:C (Figs. S20-S21). This fast step is followed by complete dissociation into single strands by an activated barrier crossing (τ_3). The temperature-dependence of dehybridization rates obtained from τ_3^{-1} follows Arrhenius behavior with a large enthalpic barrier ($\Delta H_d^\ddagger > 100\text{ kJ/mol}$, Fig. S22) that is consistent with full strand dissociation.(30, 38)

The T-jump IR responses of CGCcap-AP6, CCends-AP6, and GCGcore-AP6 reveal an additional kinetic component that occurs on a $\sim 1\text{ }\mu\text{s}$ timescale (τ_2) and corresponds to dehybridization of the full stretch of base pairs on one side of the AP-site (half-dehybridization). Using the contribution of G (1550 cm^{-1}) and A (1605 cm^{-1}) to the spectral response during τ_2 relative to the full time window, we determine the fraction of A:T and G:C base pair loss character during τ_2 (Fig. 3c and S23-S24). The observed A:T character of 60% for CCends-AP6 and 85% for GCGcore-AP6 match the expectation for dissociating either the 5'- or 3'-half of the duplex based on the fractional A:T content of these palindromic sequences. On the other hand, the 91% A:T character for CGCcap-AP6 indicates a clear preference for dissociating the pure A:T segment. We find that τ_2^{-1} increases exponentially with temperature (Fig. 3d) and can also be described by

a Kramers-like equation (Eq. S27) as expected for an activated process. Together, the relative A:T/G:C character and activated nature of the τ_2 response directly reveal the presence of a free energy barrier in AP6 sequences for half-dehybridization ($\Delta G_{d,2}^\ddagger$), which in the reverse direction corresponds to the barrier for nucleating the remaining base pairs across the AP-site ($\Delta G_{h,2}^\ddagger$). Scheme 1b illustrates how the AP-site introduces the metastable half-hybridized state ($n_{bp} = 5$) into the free energy profile, and the resulting kinetics following the T-jump.

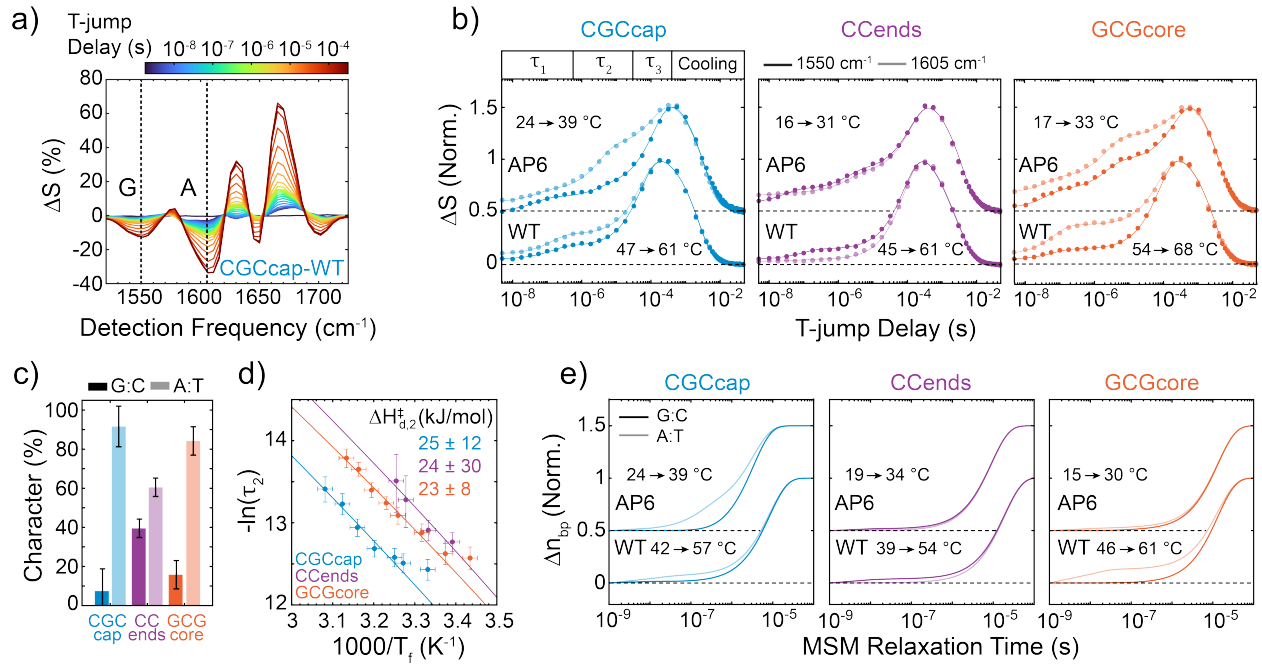
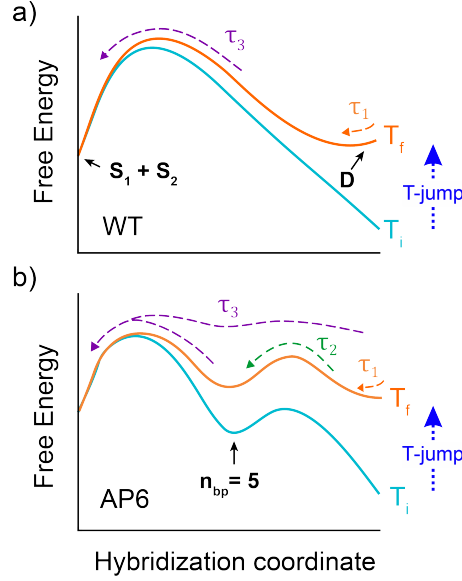


Figure 3. Direct observation of segment nucleation barrier from T-jump IR spectroscopy. **a**, T-jump IR difference (t-HDVE) spectra of CGCcap-WT from 5 ns to 1 ms delays after the T-jump, plotted as the change relative to the maximum of the spectrum before the T-jump: $\Delta S(t) = S(t)/\max(S(T_i))$. **b**, Normalized $\Delta S(t)$ time traces at 1550 cm^{-1} (dark, G:C) and 1605 cm^{-1} (light, A:T) cm^{-1} plotted for the CGCcap, CCends, GCGcore sequences, showing the initial and final temperatures for each sequence ($T_i \rightarrow T_f$). T-jumps are performed from approximately $T_m - 15^\circ\text{C}$ to T_m for each sequence. Traces are shifted vertically with respect to one another, and dashed lines indicate respective baselines. Solid lines correspond to three- or four-component fits from global lifetime analysis. **c**, Percentage of G:C (dark) and A:T (light) base pair loss character observed in half-dehybridization t-HDVE responses. Percentages are derived from t-HDVE amplitudes of the 1550 and 1605 cm^{-1} bands, respectively, determined from global lifetime fitting. Error bars indicate 95% confidence intervals propagated from global fits. **d**, Temperature-dependent observed rates for half-dehybridization responses (τ_2^{-1}) with fit to a Kramers-like equation in the high friction limit (solid lines, Eq. S27). Vertical error bars correspond to 95% confidence intervals from global lifetime fitting and horizontal error bars are the standard deviation in T-jump magnitude. The enthalpic barrier for half-dehybridization ($\Delta H_{d,2}^\ddagger$) is reported. **e**, Markov state model (MSM) T-jump simulations of the normalized change in intact G:C (dark) and A:T (light) base pairs, $\Delta n_{bp}(t) = [n_{bp}(t) - n_{bp}(T_i)]/[n_{bp}(T_f) - n_{bp}(T_i)]$. Traces are shifted vertically with respect to one another, and

dashed lines indicate respective baselines. T-jumps were initialized from a population distribution determined at $T_i = T_{m,MD} - 15^\circ\text{C}$, projected into the microstates of an MSM constructed at $T_f \approx T_{m,MD}$, and allowed to relax to the ensemble determined at T_f . The presence of a base pair is based on a $r_{bp,eq} + 0.3$ nm radial cutoff between a nucleobase and its complement, where $r_{bp,eq}$ is the equilibrium base pair separation for G:C (0.55 nm) and A:T (0.6 nm) base pairs. The relaxation kinetics are insensitive to small changes in the radial cutoff value (Fig. S28). The MSM relaxation profiles contain two to three kinetic components spanning from 1 ns to 100 μs , and CGCcap-AP6 is the only sequence that exhibits a significant amplitude of the half-dehybridization component that appears from 10 ns to 1 μs .

To provide a molecular level interpretation of the T-jump IR responses, we constructed Markov state models (MSMs) from 250 μs of unbiased 3SPN.2 simulations near $T_{m,MD}$ to furnish kinetic models of the T-jump relaxation behavior (Section S4). Using base pairing assignments for the 200 microstates, we computed the fractional change in the number of intact A:T and G:C base pairs (Δn_{bp}) as a function of relaxation time at $T_{m,MD}$ after initiating the MSM kinetics with a state occupancy obtained at $T_{m,MD} - 15^\circ\text{C}$ (Fig. 3e). The 3SPN.2 coarse-grained model is known to accelerate computational timescales by ~ 1 order of magnitude relative to experiment,⁽¹⁸⁾ but maintains a clear temporal separation between terminal fraying (~ 20 ns), half-dehybridization (~ 500 ns), and complete dehybridization (~ 10 μs). Consistent with the FEPs in Fig. 2a, we observe significant amplitude of τ_2 response for CGCcap-AP6, the only sequence for which $\Delta F_{d,2}^\ddagger < \Delta F_{d,1}^\ddagger$ at the final temperature used for the simulation. In contrast, $\Delta F_{d,2}^\ddagger > \Delta F_{d,1}^\ddagger$ for GCGcore-AP6 and CCends-AP6 sequences, leading to overlap of half-dehybridization and complete dissociation timescales and negligible τ_2 response amplitude. Experimentally, half-dehybridization responses are observed for all three AP6 sequences, so although our computational model predicts a free energy minimum at $n_{bp} = 5$ and accurately captures $\Delta F_{d,37}^o$, we surmise that small errors in the ratio of $\Delta F_{d,1}^\ddagger$ to $\Delta F_{d,2}^\ddagger$ – potentially due to inaccurate modeling of the local interactions around the AP-site for which the 3SPN.2 model was not directly parameterized – may lead to substantial changes in the kinetic predictions of the computational model. Nevertheless, the internal consistency of the model in resolving significant τ_2 response as a result of $\Delta F_{d,2}^\ddagger < \Delta F_{d,1}^\ddagger$ provides a mechanistic understanding of the origin of this observed kinetic behavior.



Scheme 1. Free energy profile reshaping by a T-jump for WT and AP6 sequences. (a) An increase in temperature promotes broadening of the duplex free energy minimum (D) by terminal base pair fraying that occurs on a ~ 100 ns timescale (τ_1 , orange arrows). This is followed by cooperative dissociation of the duplex to single-strands ($S_1 + S_2$) on a ~ 100 μ s timescale (τ_3 , purple arrows). (b) In AP6 sequences, cooperative dissociation is split into multiple steps. Following minor terminal fraying (τ_1), base pairs on one side of the AP-site dehybridize on a ~ 1 μ s timescale (τ_2 , green arrows) prior to full dissociation of the duplex (τ_3).

Impact of AP-site in longer oligonucleotides

The global impact of the AP-site across these 11-mer oligonucleotides raises the question of how an AP-site may disrupt cooperativity in longer oligonucleotides. We compared duplexes of A:T-rich sequences similar to GCGcore containing 11, 15, 19, and 27 possible base pairs (N) with and without a central AP-site using the same destabilization metrics as above (Figs. 4 and S29-S33). The observed duplex destabilization, measured with the difference in T_m between AP and WT sequences (ΔT_m), is greatest for the shortest sequences and drops with increasing N , but still remains substantial for $N = 27$. In comparison, the NN model yields values of ΔT_m that are roughly half, but qualitatively follows the same trend with N . The NN model applies only a constant free energy penalty independent of N to account for the AP-site. We find that this quantity can be corrected to nearly reproduce the experimental data by fitting the NN ΔT_m trend to that from FTIR using an entropic factor of 34.5 J/molK. This correction corresponds to a free energy penalty of -10.7 kJ/mol at 37 °C that approximately matches the difference between NN and experimental $\Delta\Delta G_{d,37}^0$ values for sequences with the GCG motif. This observation suggests that the

AP-site defect introduces a single thermodynamic penalty that is “diluted out” with increasing N . In contrast to the trends in ΔT_m , the experimental $\Delta\Delta G_{d,37}^0$ decreases with N (Fig. 4d) by an amount dependent on temperature (Fig. S29d). This could be a non-additive contribution to the AP-site destabilization, which may reflect that the fraction of $\Delta G_{d,37}^0$ lost by incorporating an AP-site ($\Delta\Delta G_{d,37}^0/\Delta G_{d,37}^0(WT)$) is much higher for short sequences than long ones.

The sizeable influence of a single lesion on the stability of a 27-mer duplex led us to investigate the predictions of the NN model for much larger oligonucleotides (Fig. S29e-f). These calculations predict that ΔT_m does not drop to an offset, but appears to follow power law behavior, $\Delta T_m \sim N^{-\nu}$, with $\nu \sim 1.1$ in A:T-rich sequences. The same power law behavior with $\nu \sim 1.4$ fits the experimental ΔT_m trend (Fig. 4c), and similar behavior is calculated for $\Delta\Delta G_{d,37}^0/\Delta G_{d,37}^0(WT)$. Of course, this NN calculation does not account for the fact that long stretches of A:T base pairs (>20 bp) may form bubbles,(39) or that an AP-site could significantly impact bubble behavior as observed for base pairing mismatches.(40) The ΔT_m trend is expected to be sensitive to sequence and A:T/G:C content. For example, NN calculations predict ΔT_m to approach zero more sharply with an exponent of $\nu \sim 3.5$ when extending a CCends template with GC steps due to the greater $\Delta G_{d,37}^0$ associated with GC steps.

For comparison, 3SPN.2 simulations employing WTMetaD were performed for the GCGcore-like sequences of length $N = 11, 15$, and 19 . We find that one central AP-site has a negligible effect on the end-to-end distance of duplex DNA, regardless of N , but does have a small lengthening effect on single-strands that becomes negligible as N increases (Fig. S32). The computed FEPs as a function of the fraction of intact base pairs have a similar free energy barrier $\Delta F_{h,2}^\ddagger$ at $n_{bp}/N = 0.5$ with a barrier height of ~ 4 kJ/mol for all lengths (Fig. 4e), indicating that the disruption of hybridization cooperativity and second nucleation penalty arising from $\Delta F_{h,2}^\ddagger$ persists even in the long duplex limit.

While measurable in the $N = 11$ AP sequence, the half-dehybridization T-jump response decreases in magnitude in the $N = 15$ AP sequence and is undetectable for $N = 19$ (Fig. S29d), indicating that $\Delta G_{d,2}^\ddagger$ increases and ultimately exceeds $\Delta G_{d,1}^\ddagger$ for large N . Similar to $\Delta\Delta G_{d,37}^0$, a decrease with N is observed in $\Delta\Delta H_d^0$ (Fig. S29a) as well as $\Delta\Delta G_{d,37}^\ddagger$ from T-jump measurements (Fig. 4d). As indicated by the FEPs in Fig. 4e, these observations indicate that the role of half-

hybridized states in duplex stability decreases as N increases, yet remains important even for oligonucleotides far longer than those investigated here.

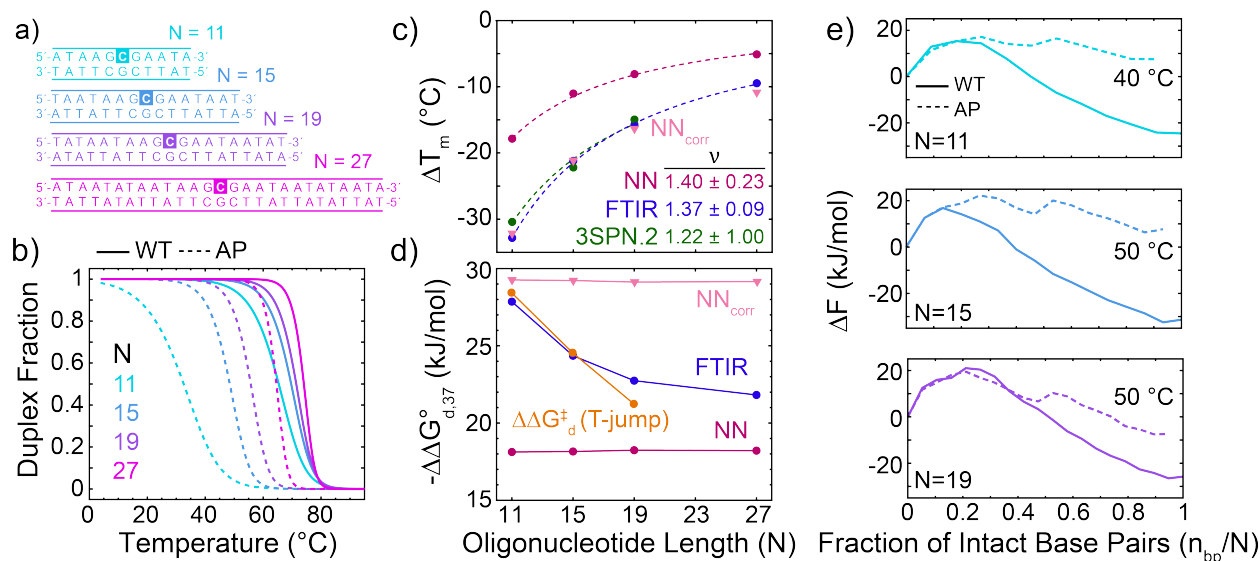


Figure 4. Influence of oligonucleotide length on duplex destabilization induced by an AP-site. **a**, GCGcore-like sequences of variable length in terms of number of possible base pairs (N), identifying the central AP-site in the top strand (shaded). Sequences contain a GCG motif with a variable number of A:T base pairs on each side leading to $N = 11, 15, 19$, and 27 . **b**, FTIR melting curves for GCGcore-like sequences where solid and dashed lines correspond to WT and AP sequences, respectively. All sequences were measured with a 1:1 strand ratio, total strand concentration of 2 mM, and in pH 7 400 mM SPB. **c**, $\Delta T_m = T_m(\text{AP}) - T_m(\text{WT})$ and **d**, $-\Delta\Delta G_{d,37}^{\circ}$ between AP and WT sequences. Values from FTIR melting curves are compared with those from NN calculations. Dashed lines indicate fits to a power law $\Delta T_m(N) = AN^{-\nu}$, and ν is listed where error corresponds to 95% confidence interval from fit. The NN ΔT_m trend was corrected to the experimental data (NN_{corr}) by fitting with an entropic factor. **e**, FEPs for WT (solid lines) and AP (dashed lines) $N = 11, 15$, and 19 sequences extracted from 3SPN.2 MD WMetaD simulations at the indicated temperatures that are approximately 20°C below $T_{m,\text{MD}}$ of each WT sequence. FEPs are plotted as a function of the fraction of intact base pairs (n_{bp}/N) within a base pair separation cutoff of 0.7 nm.

Disruption of cooperativity in hybridization dynamics

Both T-jump measurements and computed FEPs reveal the presence of free energy barriers for hybridizing each half of AP6 sequences, but they do not directly describe if or how an AP-site affects the dynamics of recognition and binding during hybridization. Even among studies of unmodified DNA many of the early events during hybridization are poorly understood, and it is clear that oligonucleotides may hybridize via multiple mechanisms.(16-18, 41) We examined the dynamics of base pairing from thousands of hybridization events generated by 3SPN.2 MD simulations at 20°C for each 11-mer sequence. Trajectories of example hybridization events for

GCGcore-WT and -AP6 sequences are shown in Fig. 5a-b, plotting the separation between all in-register base pairs as a function of time. Trajectories are trimmed to the time interval between the first base pair separation distance falling below 2.5 nm and when all distances fall below 0.7 nm. From each trajectory, we isolate the time point for first permanent base pair formation, the N1-nucleation point t_{N1} , and define a nucleation patch (N1-patch) as any intact base pairs present at that point, whether they are transient or permanent.

As illustrated in Fig. 5a-b, WT sequences exhibit a single nucleation event—here through a N1-patch of three adjacent bases—followed by concerted formation of the remaining base pairs on either side of the patch within a few nanoseconds in a classic nucleation and zipping process.(30, 38) In contrast, AP6 sequences tend to hybridize one half of the duplex and remain in the half-hybridized state with frayed termini for many nanoseconds before nucleating the second half (N2-nucleation) at t_{N2} . This is illustrated through the changes in the distribution of $t_{N1} - t_{N2}$ time-intervals between WT and AP6 sequences in Fig. 5c where t_{N2} is defined for WT sequences as the time point for nucleating a base pair on the half opposite from where t_{N1} occurs. These dynamics are consistent with the presence of two barriers to hybridization, $\Delta F_h^\ddagger$ and $\Delta F_{h,2}^\ddagger$, and suggest that n_{bp} is a good coordinate to describe hybridization.

N1- and N2-nucleation events have qualitatively different properties that strongly depend on nucleobase sequence. Quantifying the N1-nucleation position and N1-patch size shows that initial nucleation most likely occurs near G:C base pairs with a N1-patch spread over 1-5 contacts (Fig. 5d). A central AP-site localizes the N1-patch into one half of the duplex (Fig. S34), which is most disruptive to GCGcore and reshapes the nucleation free energy barrier shown in Fig 2a. The N1-patch of CCends and CGCcap are less perturbed by a central AP-site, and their corresponding WT and AP6 FEPs have nearly identical profiles from $n_{bp} = 0$ -5.

N2-nucleation typically occurs with all base pairs in the first half intact, which constrains the dissociated segment structure relative to single-stranded DNA(42, 43) and reduces the barrier to N2-nucleation. As a result, N2-nucleation is most probable adjacent to the AP-site but with characteristics that depend on the identity of the base pair adjacent to the AP-site as well as the sequence of the entire segment (Fig. 5e). GCGcore-AP6 demonstrates that an adjacent G:C base pair promotes N2-nucleation next to the AP-site relative to sequences with an adjacent A:T base pair and does not require as many base pairs in the N2-patch (Fig. S34c). Although each have an

adjacent A:T base pair, the terminal G:C base pairs of CCends-AP6 promote a higher probability of N2-nucleation away from the AP-site than in CGCcap-AP6 and exemplify that N2-nucleation dynamics depend on the sequence of the entire segment, not just the nucleobases adjacent to the AP-site.

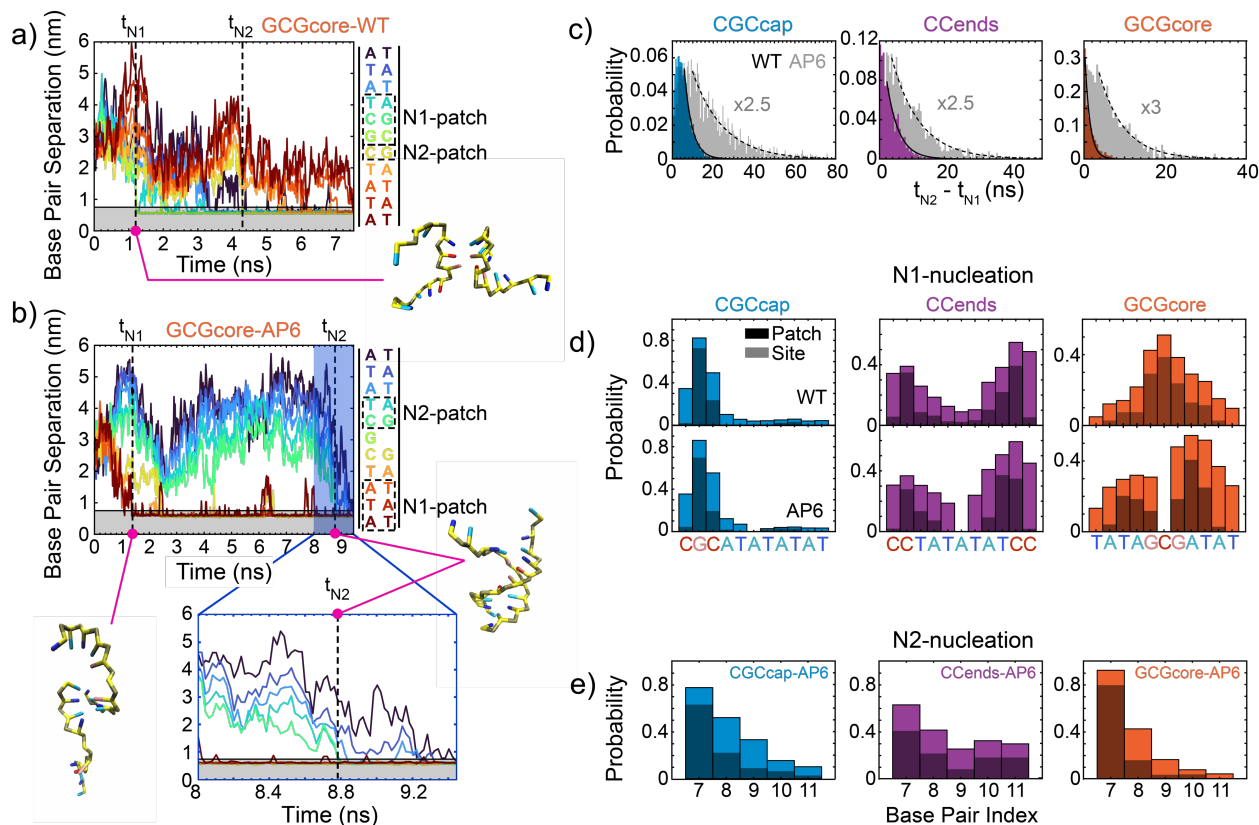


Figure 5. Splitting of hybridization dynamics into two nucleation events by an AP-site revealed from 3SPN.2 MD simulations. **a**, Example hybridization trajectory at 20 °C of GCGcore-WT plotting the separation of each in-register base pair contact (color coded). The horizontal solid line indicates the base pair separation cutoff of 0.7 nm, and vertical dashed lines indicate the time points of nucleation (t_{N1} and t_{N2}) in the example trajectories. N1-nucleation is defined as the first permanent formation of any base pair during the hybridization event and N2-nucleation is the first permanent formation of a base pair on the other half of the duplex. N1- and N2-patches correspond to all base pairs with a separation below 0.7 nm at t_{N1} and t_{N2} , respectively. Dashed boxes denote the base pairs within the N1- and N2-patches for the example trajectories. **b**, Example hybridization trajectory of GCGcore-AP6 at 20 °C. The bottom panel magnifies the portion of the trajectory near t_{N2} for GCGcore-AP6. The hybridization event is declared complete when the average base pair separation reaches 0.6 nm. The GCGcore-WT trajectory exhibits a single nucleation event where t_{N2} is defined just for comparison with GCGcore-AP6, which undergoes a distinct nucleation event for each half of the duplex. **c**, Probability distribution of time difference between N2- and N1-nucleation ($t_{N2} - t_{N1}$). Grey distributions correspond to AP6 sequences scaled up by a factor of 2.5-3 for clarity. Distribution tails are fit to an exponential decay for WT (solid) and AP6 (dashed) sequences. AP6 sequences exhibit wider $t_{N2} - t_{N1}$ distributions than WT sequences due to the added free energy barrier for

N2-nucleation. **d**, Probability that a given base pair site is part of the N1-patch (colored bars) or the N1-nucleation site (gray bars) determined from >1000 hybridization trajectories for each sequence. The AP-site reshapes the N1-patch in a sequence-dependent fashion. **e**, Probability that a base pair site is part of the N2-patch (colored bars) or the N2-nucleation site (gray bars). Base pair index refers to the distance in sequence from the terminal base pair on the duplex half where nucleation occurs (Fig. S34b). Regardless of sequence, the most probable site for N2-nucleation is the base pair adjacent to the AP-site. Nucleation properties calculated using a base pair separation cutoff of 1.0 nm are shown in Fig. S35.

Conclusions

Our study combines T-jump IR spectroscopy and coarse-grained MD simulations to reveal a hierarchy of consequences arising from a single AP-site in duplex DNA, spanning thermodynamic destabilization, hybridization pathways, and melting kinetics. An AP-site reduces the global cooperativity by effectively splitting the base pairing energetics and dynamics of the duplex into two segments. The level of impact is sequence-dependent where both the loss of base pairing interactions and height of the barrier for hybridizing across the AP-site depend on the identity of surrounding base pairs. This disruption of base pairing energetics and dynamics may be important for the recognition of AP-sites by AP endonuclease(5, 44, 45) and subsequent repair that plays a critical role in maintaining genome integrity.(46) Additionally, damage-induced AP-sites may form in clusters(47, 48) that will likely amplify the magnitude of the disruption observed from a single AP-site in this work.

The hallmark of DNA duplex stability is the nonlinearity of cooperative interactions that depend on the concerted formation of multiple continuous contacts, but we see that even one perturbation in this cooperativity can unravel duplex stability and base pairing dynamics. We have focused on an AP-site, but there are non-canonical base pairs and epigenetic and damage-induced nucleobase modifications that impose an energetic penalty similar to that of an AP-site ($-\Delta\Delta G_{d,37}^o = 5\text{-}20$ kJ/mol) and likely disrupt base pairing cooperativity within the duplex.(3, 49-53) Base pairing stability and dynamics themselves tune the interactions between DNA and other molecules yet also give rise to mechanical dynamics and distortions that facilitate protein recognition.(54, 55)

The contribution of cooperativity to stability is recognized not only among nucleic acids but for any binding partners that rely on a continuous stretch of interactions between subunits such as for protein-protein association and self-assembly of material and biological subunits. Even

though the nature of these interactions are completely different, a disruption through mutation or defect may reshape stability and dynamics, for better or for worse.

Materials and Methods

Oligonucleotide preparation

DNA oligonucleotides were purchased from Integrated DNA Technologies (IDT) at desalt-grade purity. AP-sites were incorporated as a tetrahydrofuran group (dSpacer). Samples were purified with 3 kDa centrifugal filters (Amicon). All oligonucleotides were prepared in pH* 6.8 400 mM sodium phosphate buffer (SPB, $[\text{Na}^+] = 600 \text{ mM}$). This high salt concentration was chosen to assure that the complete melting transition for all duplexes could be observed. For FTIR spectroscopy, samples were prepared in deuterated SPB solution to avoid spectral interference with the H_2O bending vibrational band. Oligonucleotide concentration was verified with UV absorbance using a NanoDrop UV/Vis spectrometer (Thermo Scientific).

Temperature-dependent FTIR spectroscopy

All samples used for FTIR spectroscopy measurements were prepared at a 1:1 ratio of complementary strands and a total strand concentration of 2 mM. This concentration was required in order to reach duplex absorbance values of 20 – 40 mOD from 1550 – 1720 cm^{-1} , which is ~6% of the D_2O solvent background absorbance. The solution of complementary strands was placed in a water bath at 90 °C for 3 min and cooled to room temperature under ambient conditions prior to each measurement to ensure oligonucleotides start in their lowest energy conformation. A discrete wavelet transform using the Mallat algorithm and symlet family was applied to the 1490 – 1750 cm^{-1} region of the FTIR spectra to separate and subtract the D_2O background absorption from the data.(56, 57)

Isothermal titration calorimetry

Isothermal titration calorimetry (ITC) measurements were performed using a MicroCal iTC200 (Malvern Panalytical). Single-stranded oligonucleotides were prepared in the cell with a concentration of 10-30 μM and the complement strand was placed in the syringe with a concentration of 100-300 μM . All samples were prepared in non-deuterated pH 6.8 400 mM SPB and degassed under vacuum for >20 min at the experimental temperature prior to each measurement. Sample conditions for each measurement are listed in Table S1 and additional

experimental details are provided in Section S1.2. The heat of dilution was measured for the titrant at each experimental temperature and subtracted from integrated thermogram of each measurement.

Temperature-jump IR spectroscopy

The temperature-jump (T-jump) IR spectrometer used in this work has been described previously.(32, 58) The T-jump magnitude ($\Delta T = T_f - T_i$) was set to ~ 15 °C for all measurements. The initial temperatures (T_i) were set using a recirculating chiller connected to a brass sample jacket and ΔT was determined from the change in mid-IR transmission following the T-jump pulse. The T-jump response of the DNA samples is probed using transient heterodyned dispersed vibrational echo spectroscopy (t-HDVE)(31) with parallel (ZZZZ) pulse polarization and fixed at a constant waiting time (t_2) of 150 fs. We report the real part of the t-HDVE spectrum, which contains similar information to an IR pump-probe spectrum.

Molecular dynamics simulations with 3SPN.2

Molecular dynamics (MD) simulations were performed using the coarse-grained 3-Site-Per-Nucleotide (3SPN.2) model.(23) Although not parameterized against sequences containing an AP-site, 3SPN.2 is directly amenable to modification for AP-site representation and simulation. Since backbone dihedral potentials are determined solely by sugar and phosphate sites, removal of one base bead does not intrinsically disrupt the backbone structure. Non-bonded interactions are constructed anisotropically as a function of the angles formed between proximate intra and inter-strand bases and their bonded sugars. The targeted removal of one base eliminates non-bonded interactions with that base without disrupting non-bonded interactions between all other intact bases. In this way, base pairing and stacking interactions in the vicinity of the AP-site are minimally perturbed.

All systems were simulated using the LAMMPS package(59) compiled with the 3SPN.2 model plugin.(23) Simulations were performed in the NVT ensemble in a periodic box of side length 8.5 nm, equivalent to an effective 5.3 mM oligomer concentration. Classical mechanical equations of motion were integrated via Langevin dynamics in the scheme developed by Bussi and Parrinello(60) with a 20 fs integration time step. Solvent was treated implicitly with an experimentally informed friction coefficient of 9.94×10^{-11} m²/s.(23, 61) A 600 mM implicit salt concentration was used and electrostatic interactions were treated using the Debye-Hückel with a

5 nm cutoff radius.(62) All simulations were initialized from WT B-DNA duplex states generated by the 3SPN.2 software. Free energies obtained from probability densities are reported as the Helmholtz free energy ΔF .

Enhanced sampling of hybridization free energy landscape

We employed different sampling methods depending on the hybridization property of interest. To efficiently sample the duplex-to-single-strand thermodynamic free energy landscape at various temperatures, we employed well-tempered metadynamics (WTMetaD) via the PLUMED plugin(63) to enhance sampling of hybridized, dissociated, and intermediate states by accelerating the dynamics in a pre-defined collective variable (CV).(64) For each system, we selected a CV that described the average distance between the beads representing native Watson-Crick-Franklin base pairs (r_{bp}). The CV was calculated by averaging over all available in-register pair distances – 11 for WT or 10 for AP sequences. Statistics were accumulated over the WTMetaD runs within the quasi-static regime where the applied bias was converged and was analytically reweighted to report unbiased thermodynamic averages and free energy landscapes in various order parameters beyond those in which sampling was conducted. Full details of the WTMetaD simulation protocol is provided in Section S2.2.

Markov state models

Markov state models (MSMs) are long-time kinetic models that provide quantitative and interpretable descriptions of the thermodynamic stability and dynamical transitions between various meta-stable states of a molecular system.(24, 25, 65-67) We constructed MSMs from 25 independent and unbiased 10 μ s molecular dynamics simulations for each of the WT and AP6 DNA systems. Simulations were performed near the sequence-dependent T_m calculated from the WTMetaD simulations ($T_{m,MD}$). Initial velocities were assigned from a Maxwell-Boltzmann distribution at $T_{m,MD}$. Each simulation was conducted for 12 μ s and frames saved to disc every 100 ps, resulting in 1-2 hybridization and dehybridization events per trajectory. The first 2 μ s of each run was discarded to produce $25 \times 10 \mu$ s = 250 μ s (2.5 million frames) of simulation data for each sequence. An additional $25 \times 1 \mu$ s of simulation data was generated for each system 15 °C below $T_{m,MD}$ in order to furnish a low-temperature initial distribution for MSM relaxation experiments. The 2D FESs shown in Fig. 2b were produced by reweighting unbiased trajectories according the

MSM microstate transition matrix. Full details of MSM construction and validation as well as T-jump relaxation simulations can be found in Section S4.

Hybridization trajectories

To efficiently sample hybridization events to characterize nucleation behavior (Fig. 5), 1000-2500 simulations of each sequence were initiated from a dissociated configuration, generated by shifting equilibrium duplexes by 2 nm in the x, y, and z directions, and were declared complete when reaching a defined duplex state cutoff of $r_{bp} = 0.6$ nm, which corresponds to a fully hybridized duplex. An umbrella potential with a force constant $k=100$ kJ/nm²mol was applied for $r_{bp} > 3$ nm prevent diffusive separation of the oligonucleotide strands. We verified that this cutoff was sufficiently large that it did not influence the binding and association events by confirming negligible differences in the hybridization mechanism when comparing trajectories harvested under restraining potentials with ($r_{bp} > 3$ nm, $k = 100$ kJ/nm²mol) and ($r_{bp} > 4$ nm, $k = 10$ kJ/nm²mol). Trajectories were trimmed to analyze the time window between when at least one in-register base pair separation distance falls below 2.5 nm to when all base pair separation distances fall below 0.7 nm.

Data availability

Python scripts for generating abasic configurations from intact 3SPN.2 files, performing metadynamics simulations, and reweighting free energy surfaces are available at <https://github.com/mrjoness/abasic-thermo/>. Scripts for running equilibrium simulations and building Markov State models are available at <https://github.com/mrjoness/abasic-kinetics/>. All PLUMED input scripts were submitted to the PLUMED-NEST public repository and are available at <https://www.plumed-nest.org/eggs/22/037/>. All unbiased and biased MD trajectories (65 GB), time- and rate-domain t-HDVE data, FTIR temperature series, and experimental and 3SPN.2-determined melting curves are uploaded to Zenodo and available at 10.5281/zenodo.7199303.

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Supporting Information

Temperature-dependent FTIR data; description of melting curve determination from two-state model; details of enhanced WMetaD 3SPN.2 simulation and melting curve determination; t-HDVE spectral assignments; description of global lifetime fitting and determination of A:T/G:C character in τ_2 response; description of two-state kinetic analysis of τ_3 response; description of MSM construction and T-jump relaxation simulations; FTIR and t-HDVE kinetic data for oligonucleotide length series.

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