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Exciton Delocalization and Scaffold Stability in Bridged Nucleotide-Substituted, DNA Duplex-Templated Cyanine Aggregates

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ABSTRACT: Molecular excitons play a foundational role in chromophore aggregates found in light-harvesting systems and offer potential applications in engineered excitonic systems. Controlled aggregation of chromophores to promote exciton delocalization has been achieved by covalently tethering chromophores to deoxyribonucleic acid (DNA) scaffolds. Although many studies have documented changes in the optical properties of chromophores upon aggregation using DNA scaffolds, more limited work has investigated how structural modifications of DNA via bridged nucleotides and chromophore covalent attachment impact scaffold stability as well as the configuration and optical behavior of attached aggregates. Here we investigated the impact of two types of bridged nucleotides, LNA and BNA, as a structural modification of duplex DNA-templated cyanine (Cy5) aggregates. The bridged nucleotides were incorporated in the domain of one to four Cy5 chromophores attached between adjacent bases of a DNA duplex. We found that bridged nucleotides increase the stability of DNA scaffolds carrying Cy5 aggregates in comparison with natural nucleotides in analogous constructs. Exciton coupling strength and delocalization in Cy5 aggregates were evaluated via steady-state absorption, circular dichroism, and theoretical modeling. Replacing natural nucleotides with bridged nucleotides resulted in a noticeable increase in the coupling strength (≥10 meV) between chromophores and increased H-like stacking behavior (i.e., more face-to-face stacking). Our results suggest that bridged nucleotides may be useful for increasing scaffold stability and coupling between DNA templated chromophores.

INTRODUCTION

Molecular (Frenkel) excitons arise from the collective interaction among coupled chromophores that allows creation of excited states that are shared across participating chromophores. The shared excitation energy extends in a wavelike manner over an aggregate (i.e., excitons are delocalized).1−7 Excitons have garnered considerable research interest arising from their function in natural light-harvesting systems8−11 and their potential applications in artificial light harvesting, organic optoelectronics,12,13 and nanoscale computing.14,15 It follows that the realization of devices using excitonic systems requires controlled delocalization of excitons along a specific path (e.g., exciton wire). A variety of chromophores have been shown to spontaneously aggregate in solution,16−19 with reported delocalization over 60+ chromophores in self-assembled aggregates (i.e., -mers)19. In the absence of additional structural support or scaffolding, however, further control over the relative placement and orientations of individual dyes to influence how aggregation occurs remains elusive. Natural photosynthetic systems use proteins as a scaffold for chromophores to facilitate controlled delocalization for efficient energy transfer, but protein scaffolds present design challenges due to complex folding mechanisms and large number of possible amino acid combinations.20−23

Deoxyribonucleic acid (DNA)-based nanotechnology has emerged as an effective and accessible method to design scaffolded chromophore aggregate systems at subnanometer scales because of the simple design rules, sequence selectivity, small number of components, and commercial availability of custom-sequenced chromophore-conjugated DNA oligomers. DNA has been used successfully to induce chromophore aggregation by covalently attaching chromophores to specific bases and allowing strands to self-assemble to facilitate aggregation.24,25,34−43,26,44−53

Electronic interactions between aggregated chromophores are well approximated by an augmented Frenkel Hamiltonian54−58 of the following form:

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where $\epsilon_m$ is the monomer transition energy, $J_{m,n}$ is the exchange energy associated with transition dipole coupling between chromophores from a single exciton, $K_{m,n}$ is associated with the average energy of a pair of singly excited chromophores at sites $m$ and $n$, $\Delta_m$ is associated with the energy of a single chromophore with two simultaneous excitations, and the operators $\hat{B}_i^\dagger$ and $\hat{B}_i$ are exciton creation and annihilation operators, respectively. Note that this Hamiltonian does not contain the vibrational aspect of chromophores or impact of the bath (environment) on the individual chromophores or aggregates.

Of particular interest for this study is the magnitude of $J_{m,n}$, as this parameter quantifies the interaction that allows the single exciton to transfer, or “hop”, from the chromophore, $m$, to an aggregate to the adjacent chromophore, $n$. Because of this inherent transfer of excitation energy associated with $J_{m,n}$, we refer to this parameter as the excitonic hopping parameter. Note that Mass et al. recently provided an extensive discussion of previous terminology regarding $J_{m,n}$.28 The coupling arising from the $J_{m,n}$ term in the Hamiltonian is responsible for exciton delocalization54,56−61 and is therefore of interest for designing an exciton wire.

Transfer of energy predominately occurs through a dipole−dipole interaction between the induced transition dipole moments (TDMs) of participating chromophores, and thus the magnitude of $J_{m,n}$ is sensitive to interchromophore separation. Weak coupling has been observed for chromophores with base-pair (bp) separations of 2 bp or more,14,32,62,63 whereas strong coupling—resulting in exciton delocalization—has been reported with chromophore separations of 1 bp or less.24,26,32,37,64 Although attempts have been made to influence aggregation by changing separation distances between chromophores on a single strand33 and with chromophores positioned on opposing strands in a duplex,26,33 little information is available about how changes to the scaffold strands themselves affect exciton delocalization.
DNA, thus restricting the orientations available to the attached linkers. As such, a chromophore is attached more rigidly to the which both ends of the dye are attached to the DNA via short chromophores in the excitonic wire, we chose Cy5 attachment chromophores in a wire. To ensure the desired number of several chromophores is more likely to result in "".

In contrast, the postmodi are present because of the excitonic wires on DNA, as it ensures that all dyes in the wire tiding carrying an anchor group for attachment (e.g., NHS-ester method or as a postmodi solid-state synthesis of oligonucleotides via a phosphoramidite technique). The Journal of Physical Chemistry B pubs.acs.org/JPCB

The pentose sugar ring is “locked” into the 3′-endo conformation (N-type; Figure 1a). The N-type sugar pucker is typical of an A-form DNA duplex, which is associated with higher hybridization stability. The sugar is forced into this conformation by a bridge spanning the 2′ oxygen and the 4′ carbon of the deoxyribose sugar ring. The bridge restricts the sugar from adopting the 2′-exo conformation (S-type) found in a typical B-form DNA duplex and thus eliminates a flexural degree of freedom in the oligomer backbone. It follows that the resulting perturbation of the scaffold helix would influence the relative geometry between dyes that are covalently bonded to the scaffold strand in the vicinity of a modified nucleotide; however, to the authors’ knowledge, no studies to date have reported the influence of bridged nucleotides on exciton delocalization in covalently bonded DNA-templated chromophore aggregates.

Another important consideration in designing DNA-templated chromophore networks is the choice of conjugation. There have been successful attempts to use DNA to template chromophores to form aggregates by noncovalent binding to DNA; however, these types of chromophore aggregation are difficult to control in regard to the precise location and number of chromophores. Covalent bonding to the DNA scaffold offers a higher degree of control over chromophore placement and the number of chromophores in an aggregate or network. Chromophores can be attached covalently to DNA during the solid-state synthesis of oligonucleotides via a phosphoramidite method or as a postmodification of a synthesized oligonucleotide carrying an anchor group for attachment (e.g., NHS-ester method). The former method is more beneficial for creating excitonic wires on DNA, as it ensures that all dyes in the wire are present because of the “capping” step in the synthesis cycle. In contrast, the postmodification of an oligonucleotide with several chromophores is more likely to result in “missing” chromophores in a wire. To ensure the desired number of chromophores in the excitonic wire, we chose Cy5 attachment via a synthetically available dual-phosphoramidite method in which both ends of the dye are attached to the DNA via short linkers. As such, a chromophore is attached more rigidly to the DNA, thus restricting the orientations available to the attached chromophores. In comparison, however, with a single linker attachment to an internal nucleobase which is often done via a postmodification, the dual-phosphoramidite method requires an interruption of the continuous DNA backbone, thus disrupting the base-stacking interactions that help stabilize the duplex.

In this study, we investigate exciton delocalization in cyanine (Cy5) chromophore aggregates templated sequentially along modified DNA duplex scaffolds to form linear aggregates (i.e., dimer, trimer, and tetramer) that serve as an exciton wire. We chose to conjugate the Cy5 chromophores to the DNA template with a rigid two-linker attachment between adjacent phosphate groups (see Figure 1). We report on the influence on exciton delocalization and thermodynamic stability arising from inclusion of either of two types of bridged nucleotides in DNA scaffold strands in the domain of attached chromophores. The first type is locked nucleic acid (LNA), which is a first-generation bridged nucleotide that contains a single carbon linker across the deoxyribose sugar to restrict the sugar conformation.

Comparison with the DNA-only, i.e., DNA containing only natural bases because of less conformational flexibility in the phosphate backbone and more efficient base stacking, which, we hypothesize, may mitigate the occurrence of low activation energy breathing events. Conversely, the perturbation of the helix near the chromophores may instigate nucleation of breathing events, which, in principle, could lead to changes in relative position and orientation of chromophores within the templated aggregate, as evidenced by changes in optical properties in aggregates ranging from monomers (one chromophore) up to tetramers (four chromophores). As a result, we want to know if the inclusion of bridged nucleotides near attached chromophores can be used to alter the relative orientations of the chromophores within the aggregates to enhance exciton delocalization (i.e., increase j uptake) to facilitate use of the aggregate as an exciton wire as well as to evaluate the ability of bridged nucleotides to counteract the destabilizing effect of inserting chromophores into the phosphate–sugar backbone of a DNA duplex.

### MATERIALS AND METHODS

#### Sample Preparation.

Nucleic acid strands in lyophilized form were obtained from two different vendors. All initial dye-labeled, strands with only DNA, and strands containing LNA were obtained from Integrated DNA Technologies (Coralville, IA). Strands containing BNA and additional (duplicate) dye-labeled strands containing three and four Cy5 chromophores were obtained from Bio-Synthesis Inc. (Lewisville, TX). All Cy5-labeled strands, and unlabeled strands with LNA or BNA modifications, were purified by dual high-performance liquid chromatography (HPLC). Unlabeled strands containing only DNA were purified by standard desalting methods.

All strands were rehydrated with ultrapure water (UPW) (Barnstead Nanopure, Thermo Scientific) to a stock concentration of 100 μM. We verified concentrations by measuring the absorbance via a NanoDrop One microvolume ultraviolet–visible (UV–vis) spectrometer (Thermo Scientific) using the theoretical extinction coefficients provided by the manufacturers. Forty-two base-pair (bp) duplexes were
prepared by combining equimolar amounts of complementary strands with UPW and a concentrated buffer (10× Tris-borate-EDTA [TBE], 150 mM MgCl₂) to obtain 150 μL samples in 1× TBE 15 mM MgCl₂. Except where noted, samples were prepared at a nominal 5 μM concentration (verified from absorbance values at 260 nm and the DNA scaffold extinction coefficient at 260 nm). All duplexes were annealed by using a Mastercycler Nexus polymerase chain reaction (PCR) cycler (Eppendorf). Samples were held at 98 °C for 15 min and then slowly cooled at a rate of 0.4 °C/min until they reached 25 °C. Samples were then allowed to equilibrate to room temperature (∼20 °C).

**Melting Curves.** Melting curves were obtained by using a Cary-5000 UV–vis–near-infrared (NIR) spectrophotometer (Agilent Technologies). Unlabeled 42 bp duplexes were prepared at 2 μM, whereas chromophore-labeled strands were prepared at 5 μM. All samples were prepared in 1× TBE with 15 mM MgCl₂ and transferred to a 1 cm path length low-headspace quartz cuvette. The absorbance at 260 nm was monitored as temperature was increased from 25 up to 98 °C at a rate of 1 °C/min until they reached 25 °C. Samples were then allowed to equilibrate to room temperature (∼20 °C).

**Steady-State Absorption Spectroscopy.** Steady-state, ensemble absorption spectroscopy was used to characterize the electronic structure of chromophore aggregates. Absorption spectra were recorded by using the Cary-5000 (Agilent) in dual-beam mode at 22 °C. Samples (50 μL, 5 μM, 1× TBE, 15 mM MgCl₂) were transferred into a 1 cm path length quartz cuvette (50 μL capacity, Starna Cells). Absorption was monitored from 230 to 800 nm in 1 nm steps.

**Circular Dichroism.** The CD spectra were recorded by using a JASCO J-810 CD spectrophotometer (JASCO). Samples (100 μL, 5 μM, 1× TBE 15 mM MgCl₂) were measured in a 1 cm path length quartz cuvette (100 μL capacity, JASCO) at room temperature (∼22 °C). The CD signal was monitored from 230 to 800 nm for three consecutive scans at 200 nm/min and averaged.

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**Figure 2.** (a) Example of an experimental melting curve (middle) and the derivative with respect to temperature (right) with Gaussian curves identifying the main transition (red) and the minor transition (green) for the dimer on DNA:DNA. (b) Melting temperatures for scaffolds containing zero to four covalently attached chromophores (2–5 μM samples in 1× TBE 15 mM MgCl₂; absorbance at 260 nm monitored as temperature increased from 25 to 98 °C at 1 °C/min). Melting transition temperatures were determined to be inflection points in the melting curves. The solid data points indicate the main melting point, whereas the open data points refer to the minor "premelting" point. “0” chromophores denote duplex DNA without chromophores. The lines were added to highlight trends.

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The Supporting Information for additional melting data, including hysteresis.

**Nondenaturing Polyacrylamide Gel Electrophoresis (PAGE).** Samples (5 μM, 25 μL) were combined with 5 μL loading buffer [20% v/v Ficoll (Sigma-Aldrich) and 20% v/v bromophenol blue (Sigma-Aldrich)] and loaded into wells of 10% native PAGE gel (1.5 mm) casted with 1× TBE buffer. Samples were allowed to migrate through the gel for 90 min with 150 V of applied voltage at 15 °C in 1× TBE, 15 mM MgCl₂ running buffer. The resultant gel was imaged in a FluoroChem Q imager (Alpha Innotech, San Leandro, CA) under epi-UV illumination.

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KRM Model Simulation Tool. Theoretical modeling was performed, based on the work of Kühn, Renger, and May (KRM),\textsuperscript{74} to estimate the relative positions and orientations of chromophores within the templated aggregates. Our in-house developed software\textsuperscript{26−28,31,75} was used to extract coupling information ($J_{m,n}$) and relative chromophore positions and orientations by simultaneously fitting experimental absorbance and CD data. The KRM Model Simulation Tool evaluates the energy eigenstates and eigenvalues of a Holstein-like Hamiltonian. An extended dipole approximation is employed to estimate the excitonic hopping parameter(s), $J_{m,n}$. Eigenstates and eigenvalues are used to compute theoretical absorbance and CD spectra which are then compared with experimental data. A stochastic gradient search method is employed to iteratively generate theoretical spectra to minimize the deviation from experimental results (see section S3 of the Supporting Information).

■ RESULTS
To address our research questions and compare the effects of replacing natural nucleotides with LNA or BNA near chromophore aggregates on exciton delocalization, we obtained DNA oligomers that included from zero up to four chromophores on one DNA-only strand of a duplex to be hybridized with three different complementary strands resulting in a total of 15 samples. We chose to use commercially available cyanine-5 (Cy5) as the study chromophore given its high extinction coefficient and its already extensive characterization of spectral properties.

Construct Design. Each nucleic acid sample is a 42 bp duplex with identical base pair sequences. The Cy5 chromophores were internally bound in the DNA backbone by using a double-link attachment to phosphate groups on adjacent nucleotides (nt) near the center of a 42 nt DNA strand. Complementary strands are 42 nt with a sequence designed to hybridize with the dye-labeled strands by using Watson−Crick base pairing rules to form a duplex. Sequences were determined by using in-house developed sequence optimization software, and final designs were checked for the probability of forming unwanted secondary structures by using the Nucleic Acid Package (NUPACK; California Institute of Technology).\textsuperscript{76} Three types of complementary strands were prepared containing (1) DNA only, (2) seven LNA nucleotides, or (3) seven BNA nucleotides spanning the domain of the chromophores on the labeled strand (Figure 1b).

Characterization of the Scaffold. To determine the influence of bridged nucleotides on the stability and conformation of a DNA duplex scaffold, we used two approaches. We used melting curves to evaluate the duplex...
stability of each sample and a series of polyacrylamide gel electrophoresis (PAGE) experiments to investigate conformational changes of the DNA helix when substituting DNA bases with LNA or BNA.

Thermal Denaturation.

We applied DNA thermal denaturation to quantify the change in thermodynamic stability upon substitution of DNA with BNA or LNA. Melting curves quantify thermodynamic stability in nucleic acid scaffolds as a function of melting temperature \( T_m \). The melting temperature in this context is defined as the temperature at which half of the base pairs in a DNA duplex are dissociated. In our analysis, the \( T_m \) for the 42 bp DNA-only duplex was 82.4 °C (Figure 2). Replacing seven consecutive nucleotides on one strand of the duplex with LNA increased \( T_m \) to 90.9 °C. Conversely, replacing the same seven nucleotides with BNA increased \( T_m \) to 89.5 °C. The higher melting temperatures suggest greater hybridization affinity between complementary strands and greater thermodynamic stability of the scaffold. In addition, we observed that constructs containing chromophores show multiple melting transitions, evident in the inflection points in the melting curves. By fitting Gaussian curves to the first derivative of our experimental melting curves, we can determine one or more melting points to compare \( T_m \) of scaffolds with only DNA to those with LNA or BNA substituted into the DNA duplex. In addition, melting curves also can reveal how chromophore interactions with the preceding may alter the results.

Inclusion of chromophores led to more complicated melting behaviors. In all cases, Gaussian peak fitting suggested that the chromophore-labeled scaffolds had a minor “premelting” transition at a lower temperature than the main melting transition (Figure 3 and section S1). The theoretical \( T_m \) of 82.2 °C for the unlabeled, unmodified DNA sequence (Table 1, entry 1) strongly agreed with our experimental melting temperature of 82.4 °C. While insertion of one chromophore to the DNA:DNA duplex did not affect the temperature of the main transition, an additional transition at 79.5 °C was observed. The addition of a second and third chromophore to the unmodified duplex resulted in further decrease of both main and minor melting transitions to 80.2 and 76.7 °C, respectively, for two chromophores and 79.2 and 74.7 °C, respectively, for three chromophores. Adding a fourth chromophore further reduced the temperature of the main melting transition to 77.9 ± 0.3 °C for all samples. Therefore, for this last attachment scheme, we observed no appreciable difference in melting temperature between natural DNA samples and those modified with LNA or BNA. To estimate if the major and minor transitions were due to independent melting of DNA duplex arms on each side of the chromophore(s), we calculated the theoretical melting points.

| Entry | \(^a\)Sequence | \(^b\)Theoretical \( T_m \) (°C) |
|-------|----------------|-------------------------------|
| 1     | CGATGTAAGGCCAGAGTTCG-GCTACATTCCGTTCTCAAGGGG-GCTACATTCCGTTCTCAAGGGGC | 82.2 |
| 2     | CGATGTAAGGCCAGAGTTCG-GCTACATTCCGTTCTCAAGGGG | 66.1 |
| 3     | CGATGTAAGGCCAGAGTTCG-GCTACATTCCGTTCTCAAGGGG | 68.4 |
| 4     | CGATGTAAGGCCAGAGTTCG-GCTACATTCCGTTCTCAAGGGG | 70.8 |
| 5     | CGCCCATGAGCATAGTTCG-GCTACATTCCGTTCTCAAGGGGC | 73.2 |
| 6     | CGCCCATGAGCATAGTTCG-GCTACATTCCGTTCTCAAGGGGC | 71.0 |
| 7     | GCCCATGAGCATAGTTCG-GCTACATTCCGTTCTCAAGGGGC | 68.8 |

\(^a\)Entry 1 contains the full 42 bp sequence used in this study. Bases at the midpoint of the sequence are shown in violet for reference. Entries 2–7 are partial fragments of the sequence in entry 1. \(^b\)Theoretical melting temperatures were obtained by using an online software program provided by European Molecular Biology Laboratory\(^7\) (Cambridgeshire, UK) using the nearest-neighbor method “Allawi et al. 1997”, 15 mM MgCl2, and 89 mM TRIS. Salt correction was included by using the “SantaLucia et al. 1998” option.
of the corresponding duplex fragments (Table 1, entries 2–7). Observed minor melting temperatures are considerably higher than temperatures of duplex arms, suggesting that the major and minor transitions were not due to independent melting of the DNA duplex arms on either side of the chromophore(s).

**Polyacrylamide Gel Electrophoresis (PAGE).** To further investigate changes to the DNA scaffold induced by inclusion of LNA/BNA, or resulting from chromophore addition and aggregation behavior, we used the relative migration rates of samples through the polyacrylamide gels to evaluate changes to the tertiary structure of the DNA scaffolds. We observed that in general the migration rate of each sample was unaffected by the inclusion of LNA or BNA, indicating that LNA and BNA did not significantly affect the tertiary structure of the duplexes (Figure 3b,c). When chromophores were added, however, each added chromophore resulted in slower gel migration. The slower migration rate was attributed to the increased bulk of dye–DNA constructs upon attaching chromophores to DNA.

To determine if additional factors, such as chromophore position or interchromophore attractive forces, contributed to the slower migration rate caused by the addition of chromophores, we used a control sample composed of an unlabeled duplex equipped with terminal charged phosphoryl (PO₄⁻) groups. This design helped to determine if the change in migration rate could be attributed to the change in net charge introduced by adding positively charged Cy5 molecules. The control sample migrated at a slightly slower rate than the unmodified duplexes, which indicated that the additional negative charge on the chromophores did not alter the speed of gel migration (Figure 3a). To test the possible role of chromophore position, we designed another experiment using a total of four chromophores in which the gel migration rate with two chromophores that were located at each end of the duplex was contrasted with samples containing four chromophores at the center of the duplex. Specifically, we wanted to know if interchromophore attractive forces due to aggregation distorted the DNA scaffold and caused a significant change in the linearity of the DNA helix which would result in slower gel migration. We again observed no difference in migration rates (Figure 3c). This result suggested that the slower observed migration rate from inclusion of Cy5 chromophores was largely independent of chromophore position and interchromophore attractive forces.

**Characterization of Aggregate Behavior.** We used a combination of steady-state spectroscopy and theoretical modeling to investigate the influence of LNA and BNA on...
the optical behavior and geometry of Cy5 aggregates templated via modified duplex scaffolds.

**Absorption Spectroscopy.** Each scaffold type was characterized first with one attached Cy5 molecule (monomer). The Cy5 monomer sample on DNA:DNA exhibited an absorbance maximum at 647 nm. The maximum is assigned to the 0–0 energy transition, that is, the excitation of the monomer from the ground state to the lowest energy excited state (Figure 4a). A second absorbance band around 600 nm is assigned to the 0–1 vibronic transition, that is, the excitation from the ground state to the first electronic excited state including one quanta of vibration. Monomers on DNA:LNA and DNA:BNA scaffolds had similar and slightly higher energy absorbance maxima at 644 nm (0–0) and a similarly shifted 0–1 transition near 600 nm. All three monomer structures had an extinction coefficient of about 30000 M\(^{-1}\) cm\(^{-1}\) at the absorbance maximum, which is generally in agreement with published\(^{75}\) and manufacturer-stated values.

The Cy5 dimer sample on the DNA:DNA scaffold showed a hypsochromic (blue) shift in the absorbance maximum: that is, the redistribution of the oscillator strength was toward shorter wavelengths, relative to the DNA:DNA monomer (compare Figure 4b to Figure 4a). The shift in absorbance to higher energy transitions was consistent with an H-like (i.e., face-to-face stacking) molecular packing arrangement.\(^{77}\) The absorbance spectrum showed two main bands, with the most intense feature now at 601 nm. The other major band, which was slightly broader and of lower intensity, was centered around 643 nm. Dimers on DNA:LNA and DNA:BNA both showed a similar hypsochromic shift, with the maximum absorbance feature centered at 599 and 600 nm, respectively, and the broader, lower intensity bands centered at 645 and 644 nm, respectively. For all three scaffold types, the intensities of the two main absorbance features were comparable, and each contained an enhanced minor feature centered near 580 nm.

The trimer sample on DNA:DNA had two main absorbance bands with similar hypsochromic shifts relative to the corresponding dimer, with the most intense peak at 603 nm and a second feature around 645 nm (Figure 4c). The intensities of the two most prominent features were similar, as with the DNA:DNA dimer. Trimers on DNA:LNA and DNA:BNA showed absorption spectra that were nearly identical with each other, with the most intense peak blue-shifted (i.e., greater redistribution of oscillator strength) by 11 nm relative to the DNA:DNA trimer. In both cases, the shorter wavelength peak was significantly more intense than the lower energy band at 647 nm (Figure 4c). The comparable shift in peak wavelength intensity and the similar increase in peak intensity ratio between higher energy and lower energy absorbance bands suggested that the addition of either LNA or BNA into the DNA duplex scaffold near attached chromophores had a similar impact (when compared with each other) on the electronic structure of the attached chromophore aggregates.

Tetramers had absorbance spectra similar to the trimer series (Figures 4c and 4d). The DNA:DNA-scaffolded tetramer had a maximum absorbance band at 598 nm, with the second most intense band around 645 nm. The DNA:LNA and DNA:BNA-scaffolded tetramers again showed nearly identical absorbance spectra compared with each other, and both were distinct from the DNA:DNA version. Tetramers on both scaffold types had an absorbance maximum at 592 nm, which is blue-shifted by about 6 nm from the DNA:DNA tetramer maximum. The lower energy band was centered at 647 nm. As with the trimer series, the intensity of the lower energy feature was significantly lower than the higher energy feature (Figure 4c).

The greater blue-shift in absorbance maximum of the DNA:LNA and DNA:BNA templated aggregates than the DNA:DNA templated aggregates was consistent with the addition of LNA or BNA in the scaffold altering the orientations of the attached chromophores toward a more H-like configuration, suggesting the chromophores may have been more strongly coupled (i.e., greater exciton delocalization).

**Circular Dichroism Spectroscopy.** Circular dichroism (CD) spectroscopy measures the differential absorption of right- versus left-hand circularly polarized light. The CD rotational strength for a pair of coupled chromophores is proportional to a quadrupole product\(^{80}\) with this form:

$$\text{rotational strength} = \frac{\mu_m \times \mu_n}{|\mathbf{R}_{mn}|}$$

where \(\mu_m\) and \(\mu_n\) are the TDM vectors for chromophore \(m\) and \(n\), respectively, and \(\mathbf{R}_{mn}\) is the separation vector between the TDMs. From this expression it is clear that the CD signal depends not only on the coupling strength \((J_{mn})\) but also on the proximity \((|\mathbf{R}_{mn}|)\) and mutual orientation of the participating TDMs. For Cy5, a single chromophore is achiral (i.e., little to no CD signal; see Figure 4b); therefore, any CD signal obtained from Cy5 aggregates in this study is a signature of exciton delocalization.

The CD spectrum from the DNA:DNA dimer showed a well-defined positive feature at 660 nm (Figure 4b). Below 650 nm, the signal was negative, but weak (low intensity) and noisy (complex line shapes), which we attributed to the presence of multiple configurations of dimer orientations within the sample solution (aggregate heterogeneity). The DNA:LNA and DNA:BNA dimers showed similar spectral features, with both structures having a positive feature at 679 nm that was red-shifted by 19 nm from the DNA:DNA dimer and a negative feature at 601 nm that was not present in the DNA:DNA spectrum.

The Cy5 trimer on DNA:DNA showed the strongest and most well-defined CD signal (Figure 4c). A significant negative feature occurred at 684 nm and a strong positive feature at 655 nm that were both lower in energy than the 0–0 monomer transition. The negative band at 599 nm corresponded to the aggregate absorbance maximum. Trimers on DNA:LNA and DNA:BNA scaffolds showed CD spectra that were nearly identical when compared with each other and distinct from the DNA:DNA trimer. A negative CD feature corresponded to the absorbance maxima at 591 nm, and a positive feature appeared around 667 nm in both spectra, which was significantly red-shifted from the 0–0 monomer peak.

The DNA:DNA tetramer showed a negative feature at 686 nm and a positive feature at around 655 nm, similar to the DNA:DNA trimer CD spectrum (Figure 4d). We observed a negative feature at 599 nm corresponding to the absorbance maximum of the aggregate. Although quite noisy, DNA:LNA and DNA:BNA versions both showed similar features when compared with each other, with a negative signal at 576 nm and two positive features at 620 and \(\sim 665\) nm.
Theoretical Modeling. We used our in-house KRM Model Simulation Tool\textsuperscript{27,28,31} to extract relative orientations of chromophores within each aggregate. Our model employs the Hamiltonian in eq 1 which is truncated to consider the case for single excitations ($K$ and $\Delta_m$ terms are ignored) and augmented with terms to capture vibrational effects (see section S3). Each system Hamiltonian is diagonalized, and the allowed vibronic transitions and corresponding transition amplitudes (eigenvalues of the Hamiltonian) are convolved with Gaussian line shapes and summed to form a theoretical spectrum. This modeled spectrum enables comparison with experimental data to extract information related to coupling strength and chromophore spatial proximity and orientation relative to neighboring chromophores.

In Table 2, $J_{m,n}$ indicates the electronic coupling strength between each pair of chromophores. In general, for H-like aggregates, smaller center-to-center distances ($R_{m,n}$ in Table 2) between chromophores, short of introducing charge-transfer effects, will lead to larger values of $J_{m,n}$. The deviation from parallel between each pair of TDMs is given by the oblique angle, $\alpha$, which we will call $\alpha$—that describes the absolute angle between TDMs in three dimensions (see Figure 5). A smaller oblique angle implies that the packing is closer to parallel, which we expect to increase $J_{m,n}$ for H-like packing arrangements. An example of simulated spectra and associated molecular geometry leading to the results stated in Table 2 are shown in Figure 6. For complete modeling results, see section S3.

Inclusion of LNA and BNA generally led to higher modeled values of $J_{m,n}$, regardless of the number of chromophores within an aggregate (Table 2). Dimers on all three scaffold types adopted an H-like packing arrangement. For the dimer on the DNA:DNA scaffold, the oblique angle $\alpha$ between the long axes

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Table 2. KRM Model Simulation Tool Fitting Results for Oblique Angle $\alpha$ (deg), Center-to-Center Distance $R$ (Å), and Exciton Hopping Parameter $J$ (meV) for Each Pair of TDMs within Each Aggregate for (a) Dimers, (b) Trimers, and (c) Tetramers	extsuperscript{a}

| Scaffold   | $\alpha_{(1,2)}$ (deg) | $R_{(1,2)}$ (Å) | $J_{(1,2)}$ (meV) |
|-----------|------------------------|-----------------|-----------------|
| DNA:DNA   | 37.0                   | 8.3             | 62.7            |
| DNA:LNA   | 14.1                   | 5.3             | 88.9            |
| DNA:BNA   | 26.3                   | 6.9             | 72.6            |

| Scaffold   | $\alpha_{(m,n)}$ (deg) | $R_{(m,n)}$ (Å) | $J_{(m,n)}$ (meV) |
|-----------|------------------------|-----------------|-----------------|
| DNA:DNA   | 35.1                   | 42.7            | 48.5            |
| DNA:LNA   | 12.5                   | 36.2            | 40.1            |
| DNA:BNA   | 36.9                   | 36.6            | 25.2            |

| Scaffold   | $\alpha_{(m,n)}$ (deg) | $R_{(m,n)}$ (Å) | $J_{(m,n)}$ (meV) |
|-----------|------------------------|-----------------|-----------------|
| DNA:DNA   | 3.3                    | 12.9            | 14.1            |
| DNA:LNA   | 9.5                    | 6.8             | 5.4             |
| DNA:BNA   | 5.0                    | 21.5            | 21.8            |

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\textsuperscript{a}See section S3 for expanded modeling results, including diagrams that show the spatial arrangement of each modeled aggregate.

Figure 5. Diagram depicting relative orientation parameters for a pair of TDMs in an H-like stacking arrangement, including oblique angle, $\alpha$, the center-to-center separation vector, $R$, the relative slip angles, $\theta_s$, and the twist angle, $\theta_t$ (see section S3). Note that, here, $\alpha$ is shown in two dimensions for the special case of zero twist angle, but in general the angle spans three dimensions. The inset shows the twist angle looking along the separation vector from $m$ to $n$. The slip angle for nonparallel TDMs will depend on which TDM is chosen as the reference.
of the chromophores was 37°, whereas dimers with LNA and BNA showed a more parallel arrangement with oblique angles of 14.1° and 26.3°, respectively. The excitonic hopping parameter $J_{in}$ was 62.7 meV for the DNA:DNA dimer. The $J_{in,n}$ parameter for LNA and BNA increased to 88.9 and 72.6 meV, respectively, which reflected stronger coupling due to enhanced H-packing (i.e., closer to parallel and face-to-face).

Trimers showed consistently smaller oblique angles between chromophores in the constructs containing LNA and BNA compared to DNA-only. The smaller angles led to more parallel H-packing arrangements and higher $J_{in,n}$ values. Modeling results for all trimer scaffold types indicated that all three chromophores interacted with each other and suggested that exciton delocalization occurred over all three chromophores.

Results for tetrarmers on all three scaffolds indicated that three of the chromophores may have been in close proximity and significantly interacting, whereas the fourth was farther away. Results suggested that coupling between adjacent chromophores as indicated by larger $J_{in,n}$ values was significantly stronger for constructs containing LNA or BNA (see section S3 for expanded KRM Model Simulation Tool results).

**DISCUSSION**

First, we discuss scaffold behavior, in particular, how attachment of chromophores affected the stability and structure of the DNA:DNA, DNA:LNA, and DNA:BNA scaffolds. Incorporation of LNA and BNA was shown to be accompanied by increased duplex stability and changes in its tertiary structure. The increased thermal stability of LNA and BNA is believed to arise from the restricted 3’-endo sugar conformation that is associated with the A-form of the DNA helix rather than the B-form helix that is typically associated with natural DNA. The thermodynamic factors contributing to the stability provided by bridged nucleotides such as LNA remains an open question. Stability has previously been attributed to improved base stacking associated with the A-form (enthalpic) and to decreased entropy loss upon aggregation due to preorganization of the phosphate–sugar backbone (entropic).

Our experimental CD results (see section S5) suggest that the B-form DNA helix acquired some features of the A-form in the region where seven bridged LNA or BNA nucleotides were incorporated. In particular, we noted a small blue-shift in the CD feature associated with the DNA (UV region) in constructs containing bridged nucleotides relative to otherwise identical constructs containing only natural DNA.

Previous work suggests that the contribution to $T_m$ per substituted LNA and BNA nucleotide differs significantly depending on sequence and experiment design. In our experimental melting results for unlabeled duplex scaffold, we also observed that the inclusion of LNA or BNA led to stabilization of duplex scaffold manifested in a higher $T_m$ compared to DNA-only scaffold.

Experimental melting results for chromophore-labeled samples confirmed that the attached chromophores destabilized the DNA duplex. Analysis of the derivative of the melting data revealed a more complex dissociation process compared to unlabeled duplexes. In all of the chromophore-labeled samples, we observed a minor “premelting” transition occurring at a lower temperature than the main melting transition. We suggest that the instability introduced by inserting chromophores into the backbone of one component strand in the duplex without accommodating for the size of the chromophore on the complementary strand disturbed the integrity of the duplex. We suggest that this disruption resulted in the dissociation of the chromophore domain at a lower
temperature than the full duplex possibly by allowing nucleation of breathing events. Breathing is known to be a minor effect in a well-formed duplex, with only a tiny fraction of base pairs (~10^{-5} or less) dissociated at any one time. We note, however, that disturbances to the helix are thought to increase the probability of local fluctuations (i.e., breathing). We propose that the destabilization from chromophore insertion can promote breathing at a lower temperature than the main melting transition. Although thermal stability in each construct was generally increased by the inclusion of LNA and BNA, as evidenced by a higher temperature for the main melting transitions (Figure 2) for up to three chromophores, the instability from chromophore insertion appeared to counteract the stabilizing effect of the bridged nucleotides. With the addition of the fourth chromophore, thermal stability gains from the bridged nucleotides were no longer observed.

Theoretical melting temperatures for DNA fragments (Table 1) suggest that although the templates showed some disruption, likely due to dye insertion, the duplexes appeared to melt as a single unit rather than as two separate fragments. The fragments representing the unmodified duplex sections on either side of the aggregates were found to have theoretical melting temperatures well below the observed premelting transitions, suggesting that base hybridization was occurring to a certain extent between the base pairs within the domain containing the chromophores.

Incorporation of the chromophores, indeed, might influence DNA tertiary structure (DNA conformations). For example, Mass et al. showed that interchromophore attractive forces are strong enough to influence scaffold conformation and serve to stabilize 4-arm DNA Holliday junctions with attached squaraine aggregates. To access the changes in the tertiary structure of the DNA scaffolds upon insertion of chromophores, we performed PAGE experiments. The results of the PAGE experiments indicated that each additional chromophore inserted into the scaffold caused a noticeable incremental decrease in the migration rate (see section S2), which we attribute to the added bulk of each chromophore. In contrast, the migration rate of the duplex scaffolds through the gel was insensitive to the position of the attached chromophores along the duplex (Figure 3c), suggesting that interchromophore interactions did not significantly contribute to the tertiary structure of the scaffolds. Additionally, the change in the construct charge due to the attachment of positively charged chromophores did not appear to be responsible for a slower migration rate of the duplex. Finally, it appears that the replacement of the natural nucleotides with the LNA or BNA did not appear to disturb the tertiary structure of the DNA compared with DNA-only samples.

Next, we proceed with a discussion of aggregate behavior. Absorbance results from all dimer, trimer, and tetrameter aggregates all exhibited exciton delocalization behavior in a similar manner—they all showed a redistribution of oscillator strength toward shorter wavelengths compared with the monomer spectrum. The hypsochromic shift in absorbance, along with substantially reduced fluorescence (see section S6), was indicative of H-like packing geometries within the aggregates. Inclusion of either LNA or BNA induced greater hypsochromic shifts in the absorbance spectrum, suggesting that the presence of LNA/BNA in the scaffold favorably influenced relative chromophore orientation within each aggregate toward an enhanced H-like packing arrangement. Furthermore, inclusion of LNA or BNA had comparable impacts on the spectra relative to DNA-only constructs. We suggest that the observed spectral shifts arose predominantly from the locked conformation of the sugar ring in the backbone of LNA and BNA.

In general, the CD spectra of the aggregates showed complex line shapes with relatively low intensity, with the notable exception of the trimer on a purely DNA duplex. The complexity of the CD spectra may arise because (1) aggregates were not adopting stable configurations—due to local scaffold instability—or (2) a sample could contain a mix of aggregate orientations (heterogeneity) with competing CD signals. The low intensity in the CD signals could arise either from the reasons mentioned above or from geometrical factors. In eq 2, the rotational strength depends on the cross-product of the TDM vectors and the subsequent dot product with the separation vector, R_{m,n}. If one considers the case of coplanar TDM vectors, the cross-product will yield a vector that is normal to the plane of the TDMs. The separation vector in this case would necessarily lie in the same plane as the TDM vectors; therefore, the dot product would be zero and no CD signal is produced. As a result, one can expect the CD signal to go to zero as the twist angle (θt; see Figure 5) decreases to zero (chromophores become coplanar), even if the chromophores are strongly coupled (large J_{m,n}). To add further complexity, J_{m,n} also depends on the relative TDM orientations (see section S3). The CD signal is further convoluted by coupling to vibrational modes that lead to a mixing of electronic and vibronic states (see section S4). Given all these considerations, one must use caution when interpreting CD data. Although one might expect that strongly coupled chromophores (large J_{m,n}) would exhibit an intense CD signal, the geometric terms prevent direct interpretation. The intensity of the CD signal alone is not an exclusive indicator of the magnitude of J_{m,n} or the extent of exciton delocalization.

Consistent with the absorbance results, aggregates with either LNA or BNA showed similar CD features to each other that were distinct from those in DNA-only duplexes. Additionally, all aggregates with LNA or BNA showed a larger split between positive and negative features compared with corresponding aggregates on DNA-only scaffolds. All aggregates exhibited CD features at longer wavelengths (lower energies) than the lowest energy electronic monomer transition (0–0) that is consistent with the idea that additional energy states are accessed upon chromophore aggregation—a signature of molecular exciton delocalization.

Although the (suspected) presence of multiple aggregate configurations in sample solutions prevented the KRM Model Simulation Tool from completely describing the observed spectra, theoretical modeling results supported the finding that LNA and BNA influence the packing arrangement of chromophores toward a more H-like arrangement and strengthened coupling. Modeling constructs with LNA and BNA returned smaller oblique angles (α) between transition dipole moments and enhanced J_{m,n} values compared with the same constructs on DNA-only scaffolds. The inability to completely describe aggregate spectra may be a result of the instability introduced in each scaffold duplex by inserting chromophores without accommodating for their physical size on the complementary strand, which contributed to variations in aggregate configurations. Alternatively, the difficulty in fitting the absorbance spectra of the higher order aggregates may suggest that our model should be refined to include additional coupling-related terms to account for charge
transfer\textsuperscript{54,58} that may become significant at short interchromophore distances where wave function overlap is significant.\textsuperscript{58}

Although our modeling supported the general finding of increased coupling strength with the inclusion of LNA/BNA, we acknowledge that the model cannot account for multiple potential orientations still present in a sample solution. The inability to fully reproduce the data with a single aggregate orientation, combined with the overall complexity of the aggregate CD signals, indicated that the presence of multiple aggregate orientations is likely. As a result, the modeling outputs represented a possible average orientation within each aggregate. The suspected variation in chromophore orientations may arise from the scaffold instability introduced from chromophore insertion that counteracted the stabilizing effect of LNA/BNA inclusion in the scaffold.

\section{CONCLUSIONS}

Steady-state optical characterization results suggested that our DNA/LNA/BNA-templated Cy5 chromophore aggregates supported exciton delocalization over as many as four chromophores. Furthermore, inclusion of bridged nucleotides (LNA or BNA) in the domain of the chromophores led to enhancements in coupling strength ($I_{n,n}$) as evidenced by additional shifts in absorbance and CD spectra and extracted from theoretical modeling (Table 2). Stronger coupling and a more rigid scaffold may lead to applications as an exciton wire; however, melting data suggested that the inclusion of chromophores in the DNA backbone had a destabilizing effect that partially counteracted the additional stability gained from inclusion of the synthetic nucleotides.

Realizing rational control of aggregate packing geometry and exciton delocalization for use as an exciton wire and gaining the full benefits of including LNA or BNA in the scaffold template will require a reduction in heterogeneity in aggregate orientations. Possible paths to reduced heterogeneity include decreasing the induced strain in the scaffold that resulted from the manner in which chromophores are inserted in the scaffold backbone or by altering the attachment method to include a shorter linker from the scaffold to the chromophore. The former may provide a more stable scaffold on which to template an aggregate network with reduced scaffold fluctuations, and the latter may, in principle, restrict the degrees of freedom and possible orientations available to the chromophores within the aggregate, leading to fewer possible aggregate configurations.

\section{ASSOCIATED CONTENT}

\textbullet Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcc.1c07602.

Melting study, polyacrylamide gel electrophoresis, KRM Model Simulation Tool description and results, vibrational influence in steady-state optical spectra, ultraviolet circular dichroism, steady-state fluorescence, and supplemental references (PDF)

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