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ARTICLE TYPE

# Double zipper helical assembly of deoxyoligonucleotides: mutual templating and chiral imprinting to form hybrid DNA ensembles

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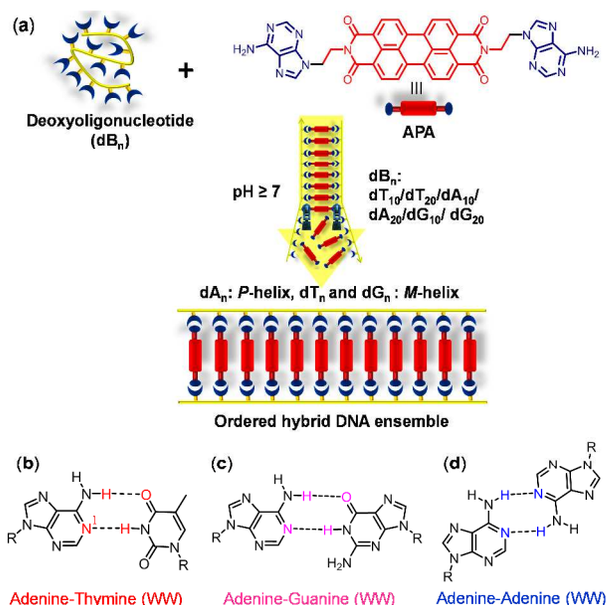
Herein, the conventional and unconventional hydrogen bonding potential of adenine in APA for double zipper helical assembly of deoxyoligonucleotides is demonstrated at ambient conditions. The quantum mechanical calculations supported the formation of hybrid DNA ensembles.

The magnificent structure-property correlations of biological systems is exemplified by the elegant molecular design and functioning of nucleic acids.<sup>1,2</sup> In particular, deoxyribonucleic acid (DNA) has structurally evolved over billions of years to effectively store and communicate the genetic information in majority of all living organisms.<sup>3-5</sup> In recent times, numerous efforts have been directed at utilizing DNA as a potential biomaterial, a biomolecular system capable of conducting electricity, single molecular wire and material building block in celebrated nanotechnological advances.<sup>6</sup> Here, we report an adenine functionalized perylene bisimide (PBI) conjugate (APA) as a promising molecular template to construct hybrid DNA ensembles through double zipper helical assembly (Fig. 1). The intriguing property of adenine to form hydrogen bond with complementary (thymine) and non-complementary (adenine and guanine) nucleobases inspired us to design APA as a double zipper template to construct new hybrid DNA structures.<sup>7,8</sup> PBI is one of the most promising aromatic  $\pi$ -conjugated systems with potential applications in organic electronics biology and supramolecular architecture.<sup>9</sup> Recently, supramolecular architecture resulting from covalent functionalization of PBI with single-stranded (ss) DNA has been reported.<sup>10</sup> In this context, APA operates through noncovalent interactions, thus, avoiding synthetic difficulties prevalent in the covalent approaches. To the best of our knowledge, this is the first report on the construction of ordered hybrid DNA ensembles through double zipper helical assembly of deoxyoligonucleotides employing versatile hydrogen bonding and  $\pi$ -stacking potential of adenine in APA.

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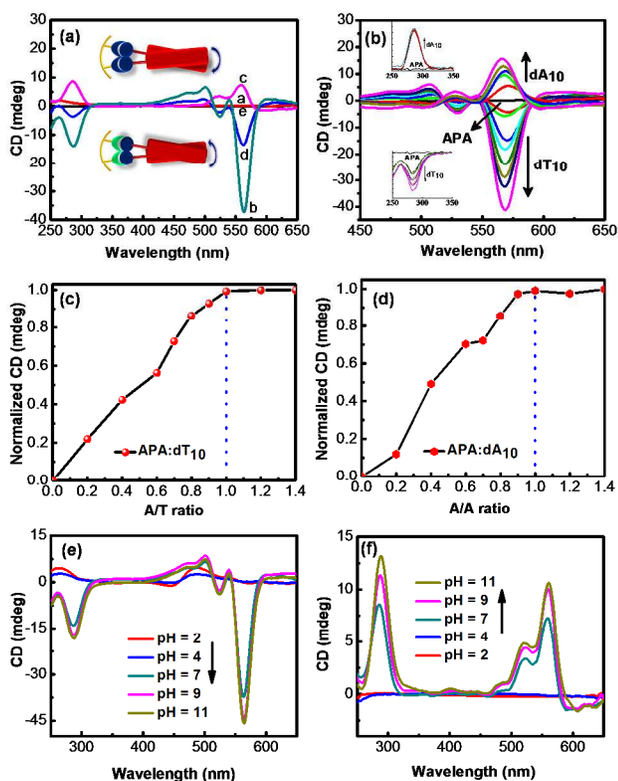
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†Electronic Supplementary Information (ESI) available: Synthesis, characterization, experimental and computational procedures. UV-vis absorption, emission, CD spectra, and simulation data of APA-oligonucleotides hybrid ensembles. See DOI: 10.1039/b000000x/



**Fig. 1** a) Molecular structure of APA and deoxyoligonucleotides (dB<sub>n</sub>) used in the present study and schematic of hybrid DNA ensemble formation. b-d) Hydrogen bonding in A-T, A-G and A-A base pairs. W = Watson-Crick hydrogen bonding site.

Herein, we present a new strategy for the construction of hybrid DNA ensembles of deoxyoligonucleotides (dB<sub>n</sub>) employing APA (Fig. 1). APA conjugate was prepared following our earlier reported procedure<sup>11</sup>, and various dB<sub>n</sub> (dA<sub>n</sub>/T<sub>n</sub>/G<sub>n</sub>/C<sub>n</sub>) were employed to construct double helical assembly of APA and dB<sub>n</sub> (Fig. S1).<sup>12</sup> First, the molecular interactions of APA were studied by evaluating its photophysical properties. The UV-vis absorption spectrum of APA (50  $\mu$ M) in DMSO exhibited three strong characteristic bands at 530, 493 and 461 nm corresponding to characteristic 0-0, 0-1 and 0-2 electronic transitions of perylene chromophore, respectively.<sup>13</sup> The absorption spectrum of APA (50  $\mu$ M) in aqueous solution (Water/DMSO = 90:10, v/v%) exhibited hypsochromic shift in the 400-550 nm region and a new band at 561 nm, which is attributed to the hydrophobic force-induced aggregation of APA (Fig. S2a). The fluorescence spectrum of APA in DMSO displayed mirror image emission bands and these bands were completely quenched in aqueous solution as a result of aggregation (Fig. S2b).<sup>12</sup>



**Fig. 2** CD spectra of APA. **a**) **a**: APA, and APA and dB<sub>n</sub> (10:2 ratio), **b**: dT<sub>10</sub>, **c**: dA<sub>10</sub>, **d**: dG<sub>10</sub> and **e**: dC<sub>10</sub> in PBS buffer (10 mM, pH = 7, 10% DMSO). *Inset*: Right and left-handed helical orientation of APA in the presence of dA<sub>10</sub> (top) and dT<sub>10</sub>/G<sub>10</sub> (bottom). **b**) Spectra of APA with variable concentration of dA<sub>10</sub> and dT<sub>10</sub>. *Inset*: Spectral features of APA and dA<sub>10</sub> or dT<sub>10</sub> in the A/T absorption region. **c**) & **d**) Plots of CD intensity monitored at 561 nm for APA and dT<sub>10</sub> or dA<sub>10</sub> different combinations of A/T and A/A ratio at fixed concentration of dT<sub>10</sub> (10 μM) and dA<sub>10</sub> (10 μM) respectively. pH dependent spectra of APA templated hybrid DNA ensembles of dT<sub>10</sub> (**e**) and dA<sub>10</sub> (**f**).

Interestingly, the absorption spectrum of APA (50 μM) in PBS buffer (10 mM, pH = 7, 10% DMSO) showed bathochromic shift with an appreciable hypochromicity (Fig. S2a). Next, we investigated photophysical properties of APA in the presence of complementary and non-complementary dB<sub>n</sub> in PBS buffer. APA (50 μM) showed hyperchromicity in the absorption with increasing concentration (1-10 μM) of dB<sub>n</sub> (Fig. S3). These results revealed the existence of mutual interactions between APA and complementary as well as non-complementary dB<sub>n</sub> through noncovalent forces to form hybrid assembly structures.

To understand the nature of molecular organizations of APA in the presence and absence of dB<sub>n</sub>, we performed circular dichroism (CD) spectroscopy studies at ambient temperature (25 °C). APA (50 μM) gave a flat CD signal, which is ascribed to the existence of equal amounts of right- and left-handed helical aggregates. APA:dT<sub>10</sub> (10:2) exhibited a bisignated CD signal in the 400-600 nm region and a negative CD signal in the 250-300 nm region corresponding to the absorption of APA and dT<sub>10</sub>, respectively (Fig. 2a). The intense negative CD signal in APA absorption region originated from the orientation of transition moments of perylene chromophores in the counter-clockwise direction (*M*-helix). The negative CD signal in 250-300 nm region revealed *M*-helical arrangement of dT<sub>10</sub> strands around the molecular template APA. To validate our proposed

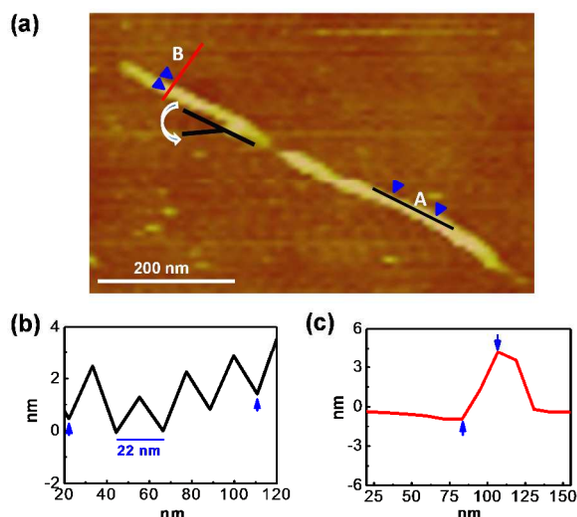
stoichiometric ratio of APA:dT<sub>10</sub> (10:2), we performed concentration dependent CD measurements by titrating increasing concentrations of APA (0 to 70 μM) to a fixed concentration of dT<sub>10</sub> (10 μM). The stoichiometry of APA:dT<sub>10</sub> in hybrid DNA ensemble is given in terms of base pairs ratio i.e A:T (A of APA:T of dT<sub>10</sub>). Plot of CD intensity at 561 nm (perylene) against the A/T ratio showed saturation at 1:1 suggesting stoichiometry of 10:2 for the formation of [dT<sub>10</sub>:(APA)<sub>10</sub>:dT<sub>10</sub>] type ensemble (Fig. 2c). We also recorded the CD spectra of fixed concentration of APA (50 μM) by adding increasing concentration of dT<sub>10</sub> (0 to 12 μM). The plot of CD intensity monitored at 561 nm against concentration of dT<sub>10</sub> showed saturation at 10 μM (Fig. S4). Thus, CD studies confirmed stoichiometric ratio of 10:2 for the complexation of APA and dT<sub>10</sub> to form [dT<sub>10</sub>:(APA)<sub>10</sub>:dT<sub>10</sub>] ensembles.

Next, CD spectra of APA in the presence of non-complementary dA<sub>10</sub>, dG<sub>10</sub> and dC<sub>10</sub> were recorded. Notably, APA:dC<sub>10</sub> (10:2) combination did not display any characteristic CD signals indicating the absence of ordered assembly between APA and dC<sub>10</sub> (Fig. 2a). Surprisingly, APA:dA<sub>10</sub> (10:2) and APA:dG<sub>10</sub> (10:2) showed unprecedented *P*-(right handed) and *M*-helical arrangements with respect to both perylene chromophore (400-600 nm) and dB<sub>n</sub> (B = A or G) (250-300 nm) (Fig. 2a). The stoichiometry study performed with APA and dB<sub>10</sub> (dA<sub>10</sub> and dG<sub>10</sub>) also suggested the formation of [dB<sub>10</sub>:(APA)<sub>10</sub>:dB<sub>10</sub>] in the ratio of 10:2 (APA:dB<sub>10</sub>) (Fig. 2d and Fig. S5). This indicated mutual templating between APA and dA<sub>10</sub>/dG<sub>10</sub> through unconventional hydrogen bonding leading to *P*- and *M*-helical imprinting in the hybrid DNA ensembles. In the control study, spectra of individual dA<sub>10</sub> showed positive and negative signal at 270 nm and 250 nm respectively (Fig. S6). Similarly, dT<sub>10</sub> showed positive and negative signals at 280 nm and 250 nm respectively. These CD signals correspond to their respective random coil structures.<sup>12</sup> Therefore, the characteristic features in the CD spectra of dT<sub>10</sub>, dA<sub>10</sub> and dG<sub>10</sub> with APA in the nucleobase and perylene absorption regions, as discussed above, signify the formation of ordered chiral assemblies. The spectra of APA as a function of added dT<sub>10</sub>/dA<sub>10</sub> exhibited an increase in the intensity of CD signals in the perylene chromophoric region (Fig. 2b). These data suggest that the dB<sub>n</sub> concentration drives the formation of ordered [dB<sub>n</sub>:(APA)<sub>n</sub>:dB<sub>n</sub>] ensembles. Further, we observed similar CD changes of APA with dA<sub>20</sub>/dT<sub>20</sub>/dG<sub>20</sub> (Fig. S7). Overall, CD studies confirmed the formation of ordered chiral ensembles of two-component systems (APA and dB<sub>n</sub>) by way of mutual templating and chiral imprinting through double zipper assembly (Fig. 1).

These results prompted us to study the effect of pH, one of the key factors that influence hydrogen bonding (nucleobases) and

**Table 1.** Melting temperatures ( $T_m$ ) of ApA templated DNA ensembles.

DNA complex	$T_m$ (at pH = 7)	$T_m$ (at pH = 9)
ds(A <sub>10</sub> +T <sub>10</sub> )	21.14 °C	---
ds(G <sub>10</sub> +C <sub>10</sub> )	62.00 °C	---
[dT <sub>10</sub> +(APA) <sub>10</sub> +dT <sub>10</sub> ]	55.25 °C	60.10 °C
[dT <sub>20</sub> +(APA) <sub>20</sub> +dT <sub>20</sub> ]	63.33 °C	64.14 °C
[dA <sub>10</sub> +(APA) <sub>10</sub> +dA <sub>10</sub> ]	67.37 °C	65.75 °C
[dA <sub>20</sub> +(APA) <sub>20</sub> +dA <sub>20</sub> ]	75.45 °C	65.75 °C
[dG <sub>10</sub> +(APA) <sub>10</sub> +dG <sub>10</sub> ]	73.83 °C	---
[dG <sub>20</sub> +(APA) <sub>20</sub> +dG <sub>20</sub> ]	78.30 °C	---



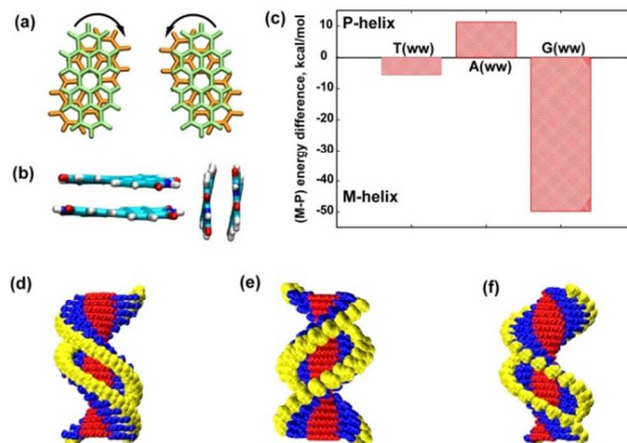
**Fig. 3** a) AFM image of left-handed helical DNA ensemble of  $[dT_{20}-(APA)_{20}:dT_{20}]$ . b) & c) Section profiles of DNA ensemble in (a) along A- and B-axis. Helical pitch and height (thickness) of DNA ensemble are found to be  $\sim 22$  nm along A-axis (b) and 5 nm along B-axis (c) respectively. For clarity, section profiles corresponding to helical pitch and thickness are shown on different ensemble-structures. Inset: Arrow shows the counter clockwise direction of DNA ensemble to the symmetry axis.

electrostatic (sugar phosphate backbone) interactions of  $dB_n$ .<sup>5</sup> In the pH range of 2-11, a flat CD signal was observed for **APA** in the absence of  $dB_n$ . **APA**: $dT_{10}$  (10:2) and **APA**: $dT_{20}$  (20:2) showed very weak CD features corresponding to perylene chromophore (400-600 nm) and  $dT_{10}/dT_{20}$  (250-300 nm) absorption regions in the acidic pH range of 2-4 (Fig. 2e and Fig. S8a).<sup>12</sup> However, strong CD signals were observed for perylene chromophore (400-600 nm) and  $dT_{10}/dT_{20}$  (250-300 nm) in the pH range of 7-11. CD spectra of **APA**: $dA_{10}/dA_{20}$  displayed a flat signal in the 400-600 nm region under acidic conditions (pH = 2-4). Remarkably, at neutral pH **APA**: $dA_{10}/dA_{20}$  exhibited strong positive CD signals in the 400-600 nm and 250-300 nm regions. The intensities of these signals were further enhanced by increasing the pH from 7 to 11 (Fig. 2f and Fig. S8b).<sup>12</sup> Interestingly, **APA**: $dA_{20}$  showed a positive CD signal while **APA**: $dA_{10}$  did not show any appreciable CD signal in the adenine absorption region (250-300 nm) under acidic conditions (pH = 2-4). These pH-dependent transformations in the CD signatures of **APA**: $dA_{10}/dA_{20}$  are mainly attributed to protonation of adenine. In acidic media (pH = 2-4), adenine ( $N^1$ ) underwent protonation with the protonated  $N^1$ -H, triggering self-complementary base pairing  $[d(AH^+):d(AH^+)]$  through reverse Hoogsteen (H) hydrogen bonding.<sup>7b</sup> This facilitates the formation of homoduplexes (A-motifs) of  $dA_{10}/dA_{20}$ , which are further stabilized by electrostatic interactions with the phosphate backbone.<sup>7b</sup> It should be noted that well-defined and characteristic CD signature for self-complementary A-motif is observed with  $dA_n$  where  $n \geq 12$ .<sup>7b</sup> However, under neutral and basic conditions the unconventional A:A hydrogen bonding interaction driven by **APA** templating of  $dA_{10}/dA_{20}$  dominates to form hybrid DNA ensembles (Fig. 1).

Next, we recorded CD spectra of **APA** in the presence of  $dG_{10}$  and  $dC_{10}$  over a pH range of 2-11. Surprisingly, **APA**: $dG_{10}$  (10:2) showed CD features similar to **APA**: $dT_{10}$  (10:2), albeit only under neutral pH conditions (Fig. S9b).<sup>12</sup> Stable unconventional hydrogen bonding-driven G-A pairing between **APA** and  $dG_{10}$  at neutral conditions led to the formation of  $[dG_{10}:(APA)_{10}:dG_{10}]$

ensemble.<sup>8</sup> The CD spectra of **APA** and  $dC_{10}$  mixture (10:2) displayed a flat signal in the pH range of 7-11, as these conditions do not favor the formation of A-C base pairing. However, under acidic conditions (pH = 4) an intense positive CD signal at 294 nm and a negative CD-signal at 265 nm were observed. These are the characteristic CD features well-documented in the literature for *i*-motifs of  $dC_{10}$  under acidic conditions (Fig. S9a).<sup>12,14</sup> Overall,  $pH \geq 7$  favored the mutual templating and chiral imprinting of **APA** and  $dB_n$  ( $B = T/A/G$ ) through conventional and unconventional hydrogen bonding to form hybrid DNA ensembles of the type  $[dB_n:(APA)_n:dB_n]$ .

In order to ascertain the thermal stability of helical ensembles of  $dB_n$  and **APA**, we carried out variable-temperature absorption and CD studies (Fig. S10 & S11). Hybrid ensembles of purine-containing  $dB_n$  and **APA** exhibited high thermal stability as indicated by the melting temperatures ( $T_m$ ) (Table 1). The thermal denaturation data revealed that all helical DNA ensembles of **APA** and  $dB_n$  were highly stable under ambient conditions. To visualize the structural morphology of hybrid helical DNA ensemble, we carried out atomic force microscopy (AFM) measurements on  $[dT_{20}:(APA)_{20}:dT_{20}]$  (A/T = 1:1) assembly structures. AFM micrograph clearly showed formation of ordered left-handed helical assembly of  $[dT_{20}:(APA)_{20}:dT_{20}]$  (Fig. 3a and Fig. S12a). The left-handed helical assembly structure is well-corroborated with the observed negative cotton effect in the CD spectrum of  $[dT_{20}:(APA)_{20}:dT_{20}]$  ensemble (Fig. S7). The AFM section profiles of  $[dT_{20}:(APA)_{20}:dT_{20}]$  structures revealed typical helical pitch of  $\sim 22$  nm along A-axis (Fig. 3b) and height (thickness) of  $5 \pm 0.5$  nm along B-axis (Fig. 3c). The observed thickness ( $5 \pm 0.5$  nm) is in agreement with the theoretically calculated value of 4.5 nm across (B-axis) hydrogen bonded **APA** and  $dT_{20}$  in  $[dT_{20}:(APA)_{20}:dT_{20}]$  (Fig. S12b). Further, the length of  $[dT_{20}:(APA)_{20}:dT_{20}]$  structures are in the range of 100-400 nm which is more than individual ensemble (Fig. S12). The observed longer helical assembly structures allowed us to consider end-to-end extension of individual ensembles through weak interactions which is very much anticipated on the surface. Overall, these studies proved the versatility of conventional and



**Fig. 4** Modeling PBI core of **APA** in dimer geometries. a) Top view of the clock and anti-clockwise twisting in PBI dimers, b) Observed bent in optimized PBI dimer structure and c) Difference between Boltzmann weighted averages of heat of formation values (in kcal/mol) of the left (*M*) and right (*P*) handed  $[dB_n-(APA)_n-dB_n]$  ( $B = A/T/G$ ) helices. d-f) Models of hybrid DNA ensembles  $[dT_{20}-(APA)_{20}:dT_{20}]$ ,  $[dA_{20}-(APA)_{20}:dA_{20}]$ , and  $[dG_{20}-(APA)_{20}:dG_{20}]$ , in their stable form.

unconventional hydrogen bonding potential of adenine as the key factor for constructing stable helical hybrid DNA ensembles.

To understand the structure and energetics of the double zipper assembly, quantum mechanical (QM) calculations were performed on carefully chosen model systems, details of which are given in the ESI.<sup>12</sup> PBI dimer, optimized using the M06 and PM7 methods, yielded a non-planar structure (Fig. 4a & b). Several model systems for APA have been built appropriately including several conformations.<sup>12</sup> High level QM calculations at the RI-MP2 level of theory indicated that both A:A and A:G prefer W:W type base pairing over other possibilities (Fig. 1c & d). The relative energies of four model systems with two base pair steps for each of the *M*- and *P*-helical forms were calculated at the semi-empirical PM7 and the Boltzmann weighted differences are presented in Fig. 4c. The energies suggest that dT<sub>n</sub> and dG<sub>n</sub> prefer *M*-helical forms, whereas dA<sub>n</sub> prefers to form the *P*-helix, which is in excellent agreement with the experimental observation discussed above. Based on these structures, three-dimensional model for the most stable helical ensemble in each of the three cases were built and are given in Fig. 4d-f. A complex combination of several factors including unique conformational preferences of the backbone increased pitch, and solvent effects are proposed to yield such structures of the assemblies.

In conclusion, we demonstrated the versatility of conventional and unconventional hydrogen bonding ability of adenine in APA as a robust double zipper molecular template to construct hybrid DNA ensembles of random coiled deoxyoligonucleotides. The formation of ordered *M*- and *P*-helical DNA ensembles was achieved by distinctive base pairing (A-T, A-A and A-G) in the processes of mutual templating and chiral imprinting of APA and deoxyoligonucleotides at pH  $\geq$  7. These experimental results were further supported by AFM analysis and computational calculations. DNA-templated studies that have been reported so far probe the helical assembly of chromophores based on their characteristic CD signatures. In the present work, we showed mutual helical assembly of functional chromophore and oligonucleotides with corresponding characteristic CD signatures in their respective absorption regions for the formation of *M*- and *P*-helical DNA ensembles. The results reported here are likely to inspire the development of new hybrid DNA ensembles of functional molecules (organic chromophores with interesting optical, electronic and biological properties) and oligonucleotides for diverse applications. The properties and applications range from electronics to nanotechnology to biomedicine. The pH dependent hydrogen bonding ability of nucleobases in the DNA ensembles can be used as tool for the development of stimuli responsive (pH-triggered) delivery systems for therapeutic small molecules and oligonucleotides.

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## 5 Graphical Abstract

**Double zipper helical assembly of deoxyoligonucleotides: mutual templating and chiral imprinting to form hybrid DNA ensembles**

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