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# Praseodymium promotes B–Z transition in self-assembled DNA nanostructures†

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Millimolar concentrations of  $\text{PrCl}_3$  can induce sequence-specific B–Z transition in various-self-assembled branched DNA (bdDNA) nanostructures. Competitive dye binding and thermal kinetics suggest that the phosphate backbone and grooves of bdDNA are wrapped with  $\text{Pr}^{3+}$  for stabilizing the Z-bdDNA. Application of EDTA can convert Z-DNA back to the B-form.

The programmable self-assembly of branched DNA (bdDNA) structures is the current trend for creating novel nanostructures with potential for diagnostic, therapeutic, and engineering applications.<sup>1</sup> Formation of functional DNA devices will potentially involve dynamic structures that can endure controllable conformational transitions. Hitherto, most switchable DNA devices have been triggered by the addition of DNA single strands and toehold-mediated strand exchange, where transitions occur on time scales of minutes.<sup>2</sup> In contrast, conformational transitions initiated by changes in ionic strength, pH, temperature, or light can be faster, and have been demonstrated to occur within seconds.<sup>3</sup> Moreover, the B-to-Z DNA transition is one of the most exceptional conversions in the biological world where the transition from right-handed B-DNA to left-handed Z-DNA with opposite chirality happens in milliseconds.<sup>4</sup> Recent studies on Z-DNA specific proteins that trigger the B-to-Z transition under physiological conditions have emphasized its role in various biological phenomena like transcription attenuation, gene regulation and participation in certain diseased conditions.<sup>5</sup> After the discovery of Z-DNA structure, it is believed that B-to-Z transition is seen specifically on  $(\text{CG})_n$  or  $(\text{GC})_n$  sequences and requires extreme ionic conditions like 4 M NaCl.<sup>6</sup> However, this vision has been changed gradually over a period of three decades. Now it is known that TA and AT repeat as well as some heterogenous-sequences can readily acquire Z-DNA in the presence of salts or metal complexes.<sup>7</sup>

Rare earth metals in this context stand as plausible candidates for inducing conformational modulations because of their strong affinity to bind protein and nucleic acids.<sup>8</sup> Nevertheless, lanthanide complexes are of increasing importance in cancer diagnosis, therapy, bioimaging and sensing, owing to

the versatile chemical and magnetic properties of the lanthanide-ion 4f electronic configuration.<sup>9</sup> Recently, we have reported the  $\text{LaCl}_3$  and  $\text{CeCl}_3$  induced B–Z transition in bdDNA nanostructures.<sup>10</sup> Praseodymium is the third element in the lanthanide series and is well used in the medicine for its radioactive isotopes  $^{59}\text{Pr}^{141}$ . Moreover, the stable form ( $^{59}\text{Pr}^{140}$ ) is used in therapeutics as well as DNA binding probes for spectroscopic studies.<sup>9</sup> However, reports regarding the interaction of  $\text{PrCl}_3$  with DNA, which involves B–Z transition, are yet to be known. On this background, the  $\text{PrCl}_3$ -induced B–Z transition was investigated in the self-assembled Y-shaped bdDNA structures in order to understand how different sequences of bdDNA affect and initiate the B–Z transition (Fig. 1). Particularly, the influence of loop length and overhang sequences on Pr-induced B–Z transition has been examined in the current communication. The details mechanism including the binding of various-dyes before and after the B–Z transition, and thermal kinetics was performed using Circular dichroism and fluorescence spectroscopy.

To understand  $\text{PrCl}_3$ -induced B–Z transition in bdDNA, a series of eight Y-shaped bdDNA structures (bdDNA US-17 to US-24) have been designed as described previously.<sup>10</sup> Four of the bdDNA (US-17, 19, 21 and 23) bear 3T in the loop whereas the other set of four bdDNAs (US-18, 20, 22 and 24) contain 5T in the loop without changing the complementary sequences. However, the overhang sequences of the bdDNA structures are different

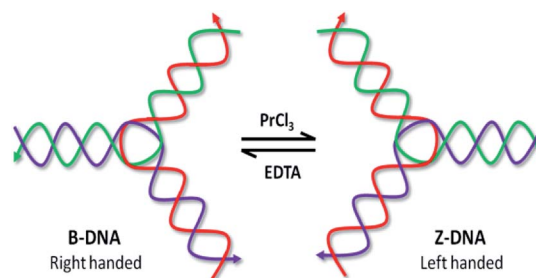


Fig. 1 Schematic presentation of Pr-induced B–Z transition in bdDNA structure and reversal transition by EDTA.

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from each other in each set of bDNA (ESI<sup>†</sup>). The interaction of bDNAs with PrCl<sub>3</sub> is examined by employing both CD and absorbance spectroscopy. The B–Z transition in DNA has distinct optical signatures in the CD spectrum due to its helicity and base stacking.<sup>11</sup> The CD spectra of bDNA US-17 is quite similar to that of bDNA US-2 with a positive peak at 280 nm and a negative peak at 250 nm.<sup>10</sup> When bDNA US-17 is treated with 2.5 to 7.5 mM of PrCl<sub>3</sub>, a decreasing in spectral trend is observed showing the loosening of base stacking (Fig. 2a). On the contrary, with 10 mM of PrCl<sub>3</sub> the CD spectra is dramatically changed showing a negative peak at 295 nm and a positive peak at 255 nm indicating a stable Z-DNA conformation. Since loop length plays an important role in the conformational flexibility,<sup>12</sup> the effect of internal loop length on the B–Z transition is explored. The loop length is increased to 5T keeping the other sequences same and the assembled product bDNA US-18 is interacted with PrCl<sub>3</sub>. With increasing concentration of PrCl<sub>3</sub>, a clear condensation is noticed with gradual decrease in the positive and negative CD without any sign of transition (Fig. 2b). These results suggest that by addition of two more thymine bases in the loop, the conformational change in bDNA is resistant to B–Z transition. Possibly, the 5T-induced structural flexibility is different from the 3T-induced conformational flexibility which is sensitive for B–Z transition. To access the generality of the observations another set of bDNA US-19 (with 3T loop) and US-20 (with 5T loop) were interacted with PrCl<sub>3</sub>. The CD spectra of bDNA US-19 and US-20 is similar and no difference is noticed in the absorbance spectra of US-19 and US-20 (Fig. S1<sup>†</sup>). While addition of 5 mM PrCl<sub>3</sub>, a significant decrease in CD spectra is observed both in US-19 and US-20. However, with 7.5 and 10 mM of PrCl<sub>3</sub> a typical Z-DNA signature is observed in bDNA US-19 unlike the progressive decrease in CD spectra of US-20. A remarkable hypochromism and bathochromic shift is also observed in the absorbance spectra of bDNA US-19 (Fig. S1<sup>†</sup>). Conversely, with 10 mM PrCl<sub>3</sub>, a partial B–Z transition is noticed in bDNA US-20 which is accompanied with a faint bathochromic shift (Fig. S1b and d<sup>†</sup>).

To move further, another set of bDNA (US-21 with 3T and US-22 with 5T) is interacted with PrCl<sub>3</sub>. When bDNA US-21 is interacted with PrCl<sub>3</sub> a small positive peak at around 260 nm is

observed despite a clear negative peak as that of Z-DNA (Fig. S2a<sup>†</sup>). This phenomenon is indicative of incomplete transition where both the B-form and Z-form co-exists which is represented by a flattened peak at 265 nm. Moreover, the hypochromism and bathochromic shift of bDNA US-21 is clearly observed in Fig. S2c,<sup>†</sup> which is an indicator of B–Z transition. When PrCl<sub>3</sub> is added to the bDNA US-22, a clear condensation is noticed similar to bDNA US-18 (Fig. S2b<sup>†</sup>). The progressive decreasing trend of both positive peak as well as negative peaks is the clear indication for the condensation of DNA. Similarly, when bDNA US-23 with 3T is interacted with 10 mM of PrCl<sub>3</sub> the CD spectra is dramatically changed to Z-bDNA with a negative peak at 295 nm and a positive peak at 255 nm (Fig. 2c). When bDNA US-24 with 5T is subjected to CD, a similar spectrum is obtained like US-23. While addition of 5 and 7.5 mM PrCl<sub>3</sub>, a significant decrease in CD spectra is observed in US-24 alike to US-23, however, with 10 mM of PrCl<sub>3</sub> a pseudo B–Z transition is noticed in bDNA US-24 (Fig. 2d). A clear bathochromic shift is also noticed in bDNA US-23 and bDNA US-24 in the presence of 10 mM PrCl<sub>3</sub>, which is comparable to the absorbance of bDNA US-17 (Fig. S3<sup>†</sup>).

From the above observations it is confirmed that the B–Z transition is clearly dependent on the loop length. The 3T variant of bDNA (US-17, US-19 and US-23) is readily exhibiting B–Z transition in presence of 10 mM of PrCl<sub>3</sub>, whereas with 5T internal loop bDNA is showing either clear condensation (US-18 and US-22) or pseudo transition (US-20 and US-24). The differential response of 5T bearing bDNA structures to B–Z transition might be due to the different overhang sequences. Another interesting observation was that when one of the overhangs AGCT was changed to AATT in bDNA US-19, the B–Z transition was observed with 7.5 mM PrCl<sub>3</sub>, whereas with three GATC overhang in bDNA US-21, a partial transition was observed. Thus, the overhangs as well as the loop length control the degree of B–Z transition. In general, 10 mM of PrCl<sub>3</sub> is sufficient to induce B–Z transition in bDNA structure. Nevertheless, among the transitions, bDNA US-23 is showing a clear B–Z transition, and its 5T variant also showing a pseudo-transition with clear bathochromic shift. Therefore, bDNA US-23 is chosen for thermal kinetics and dye binding assay.

Among the differences between B-DNA and Z-DNA, the absence of major groove in Z-DNA is an attribute which is used to differentiate the Z-DNA from the B-form of DNA.<sup>4</sup> In this regard, methyl green (MG) has been chosen that binds selectively to the major grooves of B-DNA, but unable to interact with Z-DNA. Fig. 3 showed CD spectral changes of B-bDNA and Z-bDNA in the presence of MG. When MG is bound to B-bDNA US-23, four induced CD signals (one positive peak at 650 nm and three negative peaks at 320, 425 and 620 nm) are observed which are typical characteristic of bound MG.<sup>13</sup> With incremental addition of PrCl<sub>3</sub>, the induced CD signal is gradually decreased. Interestingly, with 10 mM of PrCl<sub>3</sub> the spectrum changed to Z-bDNA with its characteristic negative peak at 298 nm and a positive peak at 262 nm. This finding supports that Pr<sup>3+</sup> could bind to the bDNA in a manner that excludes MG from the major groove. To examine the effect of MG binding to Pr-induced Z-bDNA, initially B-bDNA US-23 is interacted with

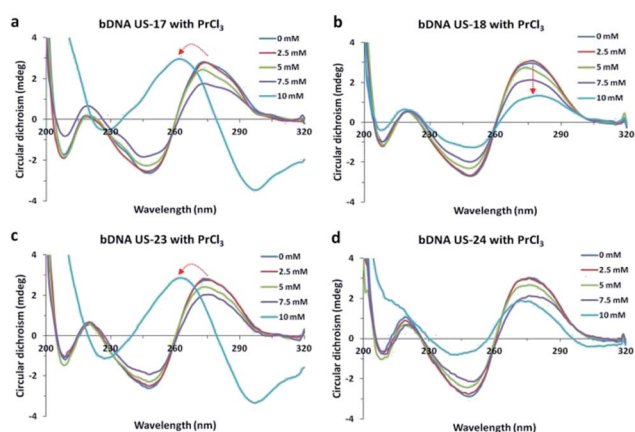


Fig. 2 (a–d) Circular dichroism spectral changes of bDNA US-17, US-18, US-23 and US-24 induced by 2.5, 5.0, 7.5 and 10 mM PrCl<sub>3</sub>.

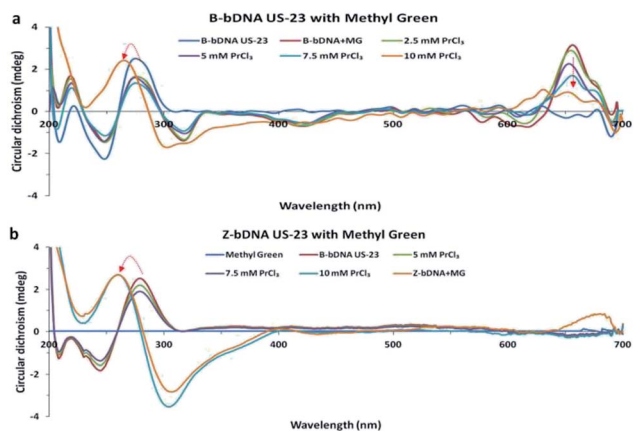


Fig. 3 (a and b) Circular dichroism spectra of B-bdDNA US-23 with MG followed by  $\text{PrCl}_3$  and (b) CD spectra of Z-bdDNA US-23 with MG.

10 mM  $\text{PrCl}_3$  to induce B–Z transition and then MG is applied to the Z-bdDNA US-23. It is observed that there is no induced negative peak at 320, 425 and 620 nm unlike Fig. 3a except a rudimentary positive peak at 650 nm. Moreover, the  $\text{PrCl}_3$ -induced Z-bdDNA conformation is unaltered in presence of MG (Fig. 3b). This result clearly supports the fact that Z-bdDNA is devoid of major grooves and therefore, MG is unable to bind to the lost major groove in  $\text{Pr}^{3+}$  induced Z-bdDNA US-23. Thus, 10 mM  $\text{PrCl}_3$  is sufficient to induce the B–Z transition in bdDNA nanostructures. Possibly, the  $\text{PrCl}_3$ -induced change in helicity leads to the drastic decrease in  $\pi$ – $\pi$  stacking between the bases and hence losing of major grooves in bdDNA.

To understand the molecular mechanism of interaction of bdDNA US-23 with  $\text{PrCl}_3$ , we employed a well-established dye binding assay.<sup>6</sup> Since nucleic acid grooves are known to interact with metal ions, two well-known minor groove binders such as DAPI and Hoechst 33342 are chosen to bind competitively with  $\text{PrCl}_3$  bound bdDNA US-23. DAPI alone did not show any CD signal, however, after interaction with bdDNA US-23, an induced peak is observed at 370 nm (Fig. S4†). Addition of  $\text{PrCl}_3$  to the bdDNA-DAPI complex led to decrease in the induced CD as well as the absorbance. With increasing concentration of  $\text{PrCl}_3$  the induced CD at 370 nm and absorbance at 260 nm is further decreased and interestingly, with  $\geq 10$  mM  $\text{PrCl}_3$ , B-to-Z transition is observed suggesting  $\text{Pr}^{3+}$  possibly replaces DAPI from bdDNA US-23. In another set of experiment, when DAPI is interacted with  $\text{Pr}$ -induced Z-bdDNA a minor induced peak is observed at 370 nm (Fig. S4c†). Similarly, Hoechst 33342 is interacted with bdDNA US-23 and an induced positive peak is observed at 340 nm and the absorbance of bdDNA is also increased at 260 nm (Fig. S5†). Addition of  $\text{PrCl}_3$  up to 5 mM did not alter the induced peak at 340 nm, however 7.5 mM led to decrease in the induced CD and with  $\geq 10$  mM  $\text{PrCl}_3$ , B-to-Z transition is observed suggesting  $\text{Pr}^{3+}$  possibly replaces Hoechst 33342. The increased absorbance is also decreased after addition of  $\text{PrCl}_3$ . On the contrary, when Hoechst 33342 is interacted with  $\text{Pr}$ -induced Z-bdDNA, an intense peak is observed at 330 nm and the usual DNA peak at 262 nm is enhanced. Collectively our results suggest that  $\text{Pr}$ -induced Z-conformation of bdDNA US-23 is stable in presence of Hoechst and DAPI. It

appears that binding affinity of  $\text{Pr}^{3+}$  towards bdDNA US-23 is higher than minor groove binders. Nevertheless, it is noticed that  $\text{PrCl}_3$  easily displaced DAPI than Hoechst 33342.

To check whether the mode of binding of  $\text{PrCl}_3$  is through intercalation we repeated the same competitive dye binding assay with ethidium bromide (EtBr). When bdDNA US-23 is interacted with EtBr, two positive peaks are observed at 270 and 300 nm and also two negative peaks are observed at 240 and 250 nm (Fig. 4a). Addition of  $\geq 10$  mM  $\text{PrCl}_3$  led to decrease in the induced CD at respective peaks without the B-to-Z transition. Possibly  $\text{Pr}^{3+}$  does not intercalate with the B-bdDNA US-23 or  $\text{PrCl}_3$  unable to displace the EtBr from bdDNA US-23. May be the EtBr bound bdDNA are in minimized energy, as a result the bases do not flip to acquire left-handed conformation even after addition of  $\text{PrCl}_3$ . In another set of reaction, when  $\text{Pr}$ -induced Z-bdDNA US-23 is interacted with EtBr, it is observed that the negative peaks at 295 nm got decreased and the positive peak at 263 nm shifted to 278 nm which correspond to the B-bdDNA (Fig. 4b). Possibly, the intercalation of EtBr to Z-bdDNA is disturbed and hence, resulting in a ruminant peak at 300 and 270 nm. Z-bdDNA–EtBr complex had a decreased absorbance at 290 nm as compared to B-bdDNA–EtBr complex which can be easily corroborated with the CD data (Fig. S6†).

EDTA, a known metal chelator is used earlier to study the reversibility of the metal ion induced B-to-Z DNA transition.<sup>10</sup> To check the  $\text{Pr}^{3+}$  chelation by EDTA, the bdDNA US-23 is pre-incubated with EDTA and then  $\text{PrCl}_3$  is added. As expected in presence of EDTA no change in B-conformation is observed suggesting that before  $\text{PrCl}_3$  being interacting with bdDNA all  $\text{Pr}^{3+}$  are trapped by EDTA, hence no B–Z transition or any other change is noticed (Fig. 4c). The absorbance spectra are also unaltered after addition of  $\text{PrCl}_3$  to the preincubated bdDNA–EDTA solution (Fig. S6c†). On the contrary, when  $\text{Pr}$ -induced Z-bdDNA is titrated with increasing concentration of EDTA the Z-conformation of bdDNA slowly moves back to the B-conformation (Fig. 4d). This suggests that EDTA-chelates the  $\text{PrCl}_3$  and Z-bdDNA are back to its native right-handed

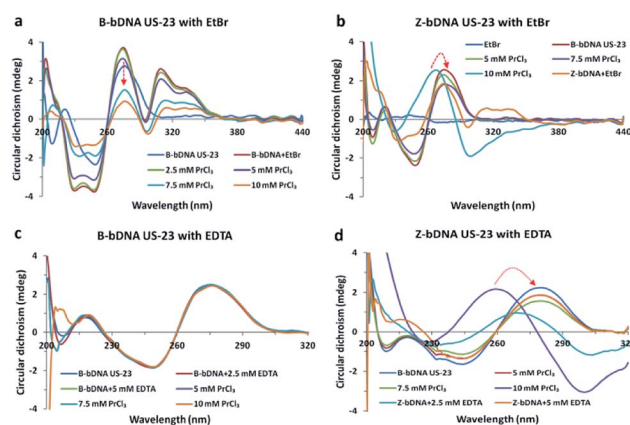


Fig. 4 (a) CD spectra of B-bdDNA US-23 with EtBr followed by  $\text{PrCl}_3$ , (b) CD spectra of Z-bdDNA US-23 with EtBr, (c) CD spectra of B-bdDNA US-23 with EDTA followed by  $\text{PrCl}_3$  and (d) CD spectra of Z-bdDNA US-23 with EDTA.

conformation. The absorbance spectra also support this observation (Fig. S6d†).

To compare these observations and understand how different fluorophores are released from B-conformation of bDNA, the fluorescence competitive binding assay is conducted after binding to  $\text{PrCl}_3$ . The fluorescence quenching experiment is widely used to establish DNA binding mode.<sup>10,14</sup> In this assay DAPI, Hoechst and EtBr dyes are interacted with B-bDNA US-23. When Hoechst 33342 (excitation wavelength, 361 nm) and DAPI (excitation wavelength, 358 nm) are interacted with bDNA US-23, an enhanced fluorescence is observed (Fig. S7†). Now, if  $\text{Pr}^{3+}$  competitively binds to the same site of bDNA where DAPI or Hoechst are bound, the fluorescence of DAPI and Hoechst would significantly decrease, because the strong binding of  $\text{Pr}^{3+}$  with bDNA should exclude these minor groove binders. As expected when  $\text{PrCl}_3$  is incrementally added to this enhanced fluorescence, invariably the emission intensity decreased than the DAPI or Hoechst bound to bDNA (Fig. S7†). With addition of 10 mM  $\text{PrCl}_3$ , fluorescence intensity of DNA declined to 30% and 60% for Hoechst 33342 and DAPI, respectively. This is suggesting that DAPI is easily released and replaced by  $\text{PrCl}_3$  similar to the spectroscopic observations. On the other hand, intercalating dye EtBr (excitation wavelength 480 nm) is widely used as a suitable fluorescence probe for DNA-binding with complexes in solution. When  $\text{PrCl}_3$  is added to the bDNA–EtBr complex, the fluorescence intensity of bDNA–EtBr complex has gradually weakened up to 50% at about 590 nm. As shown in Fig. S7,† when  $\text{PrCl}_3$  is added to US-23-dye complex, the decrease in emission is highest with DAPI and lowest with Hoechst suggesting that  $\text{Pr}^{3+}$  has DAPI like binding to the bDNA US-23. It is indicated that EtBr, DAPI or Hoechst could still bind to DNA even in the presence of  $\text{Pr}^{3+}$ . In combination with CD, competitive binding results of major and minor groove binders and fluorescence experiments further supported that  $\text{Pr}^{3+}$  could bind to major and minor grooves of bDNA. Nevertheless, the binding of  $\text{Pr}^{3+}$  is very strong towards DNA; once it is bound neither the minor groove nor major groove binders could replace and reverse the transition. The Z-conformation of the bDNA is retained even in the presence of minor and major groove binders.

Another property of left-handed Z-DNA is its thermal instability. From the thermal melting curve, it is observed that the  $T_m$  of B-bDNA US-23 is  $\sim 75^\circ\text{C}$  whereas the  $T_m$  of Z-DNA is found to be lesser than  $33^\circ\text{C}$  in both absorbance and CD spectroscopy (Fig. S8–S11†). The reason for the reduced  $T_m$  of Z-DNA can be explained as follows. The conversion of B-DNA to Z-DNA is associated with a flipping over of the base pairs so that they had an upside-down orientation relative to that of B-DNA. This flipping over resulted both in the production of a syn-conformation in purine bases and a change in the deoxyribose-ring pucker in the respective bases. Moreover, in Z-DNA, the base pairs occupy a position at the periphery instead of the centre as in B-DNA.<sup>15</sup> This may be the cause of lower  $T_m$  of Z-DNA since base stacking and base pairing are not in proper order like B-DNA. Interestingly, the  $T_m$  value of the B-bDNA is also unaltered during the process of annealing. Thus, both the CD and absorbance spectroscopy results advocate the reversible

nature of B-bDNA US-23 during the process of denaturation and renaturation unlike the  $\text{Pr}^{3+}$  bound Z-bDNA. We speculate a thorough wrapping of  $\text{Pr}^{3+}$  on the backbone of Z-DNA which hinders the formation of hydrogen bonding during the process renaturation. However, when EDTA is applied to Pr-induced Z-bDNA, the  $\text{Pr}^{3+}$  is chelated and a clear Z–B transition was observed as mentioned earlier.

In summary, for the first time our results indicate that low concentration of  $\text{PrCl}_3$  can induce a left-handed helical structure in self-assembled Y-shaped bDNA with sequence and structure selectivity. Interestingly, the bDNA having 3T in the loop are more sensitive for B–Z transition than 5T bearing bDNA. One of the most important findings in our study is that bDNA US-17, US-19, and US-23 structures with 3T are found to be more specific affinity towards  $\text{PrCl}_3$  and have considerable conformational flexibility for switching from right-handed to left-handed Z-DNA. However, the interaction between the bDNA structure and  $\text{PrCl}_3$  is different than other lanthanides discussed in earlier reports. While 5 and 7.5 mM of  $\text{LaCl}_3$  and  $\text{CeCl}_3$  respectively, are enough to induce a left-handed Z-conformation, 10 mM of  $\text{PrCl}_3$  is required for a stable Z-bDNA conformation. Thermal kinetics and dye binding experiments suggest that the B–Z transition is associated with an array of  $\text{Pr}^{3+}$  loaded on the phosphate backbone of bDNA structure as well as to the grooves of DNA and when  $\text{Pr}^{3+}$  is chelated by EDTA the bDNA transit to the original B-conformation.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

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