

Synthesis and Properties of Size-Expanded DNAs: Toward Designed, Functional Genetic Systems

ANDREW T. KRUEGER, HAIGE LU,
ALEX H. F. LEE, AND ERIC T. KOOL*

Department of Chemistry, Stanford University,
Stanford, California 94305

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ABSTRACT

We describe the design, synthesis, and properties of DNA-like molecules in which the base pairs are expanded by benzo homologation. The resulting size-expanded genetic helices are called xDNA (“expanded DNA”) and yDNA (“wide DNA”). The large component bases are fluorescent, and they display high stacking affinity. When singly substituted into natural DNA, they are destabilizing because the benzo-expanded base pair size is too large for the natural helix. However, when *all* base pairs are expanded, xDNA and yDNA form highly stable, sequence-selective double helices. The size-expanded DNAs are candidates for components of new, functioning genetic systems. In addition, the fluorescence of expanded DNA bases makes them potentially useful in probing nucleic acids.

Introduction

Over the past decade there has been a substantial research effort focused on designing replacements for the bases in

Andrew T. Krueger received his B.S. in chemistry from the University of Wisconsin—Madison in 2003. While at Wisconsin, he was engaged in research under Professor Howard E. Zimmerman, where he worked on the synthesis and photochemistry of substituted 4,4-diphenylcyclohexenones. He then moved to Stanford University to pursue graduate work under Professor Eric Kool. His research interests include the synthesis, evaluation, and structural studies of novel DNA base analogues.

Haige Lu received his B.Sc. degree from Peking University in 2001. He then moved to Stanford University, where he is carrying out his Ph.D. study under the supervision of Professor Eric Kool. His research concentrates on the synthesis, structure, and DNA polymerase kinetics of yDNA (wide DNA).

Alex H. F. Lee was born and brought up in Hong Kong. He pursued his B.Sc. and M.Phil. degrees from The Chinese University of Hong Kong and obtained his Ph.D. from The Hong Kong Polytechnic University in 2003 with Professor Albert S. C. Chan. Then he was awarded the Croucher Foundation Fellowship and undertook his postdoctoral training with Professor Eric T. Kool at Stanford University (2003–2006). His research interests include antitumor antibiotics, free radical-mediated DNA cleavage mechanistic studies, and modified nucleoside, expanded DNA, and novel nucleic acid research and development.

Eric T. Kool is the George and Hilda Daubert Professor in the Department of Chemistry at Stanford University. He received his undergraduate degree from Miami University (Ohio) in 1982. He pursued graduate studies as an NSF Fellow at Columbia University with Ronald Breslow and received his postdoctoral training at Caltech with Peter Dervan. Kool started his independent research career in 1990 as Assistant Professor at the University of Rochester. He moved to Stanford in 1999, where he teaches and directs his research group. His research interests include the design of molecular mimics for biological molecules and pathways and development of new chemical approaches for biosensing and bioimaging.

DNA. Until recently, DNA base replacements have been studied mainly for their effects in the context of natural DNA. For example, studies are often aimed at testing local effects of new bases and base pairs on the stability of the helix or on local biomolecular interactions with individual bases and pairs in natural DNA.^{1–8}

With the present project we are taking a different approach: we are asking whether *all* of the pairs of DNA can be replaced with a new geometry. Our strategy has been to design new base pairs that are larger than natural ones and assemble them to make enlarged DNA-like helices, called xDNA and yDNA. The aim of this Account is to outline progress made in assembling and studying these helices and to point out some of our future aims.

One of the long-term goals of research on expanded DNAs is the design and exploration of the properties of a nonnatural genetic system. This line of research could be classified as “biomimetic chemistry”⁹ or “synthetic biology”,^{10,11} both of which involve assembly of systems from smaller components (atoms and molecules, biochemical pathways) with the purpose of behaving like natural biomolecules, natural pathways, and, ultimately, living organisms. If chemists are to design a new genetic system, this might well start with a design of a molecule analogous to DNA. However, DNA alone is not a genetic system in itself, but is rather the information-encoding part of the genetic system. Thus, to make a functioning system, one would need, at a minimum, not only an analogue of DNA, but also its monomeric building blocks and polymerase enzymes that replicate it.

By working to recapitulate the functions of Nature’s genetic system, we can test our knowledge of how it operates, and how tightly constrained the natural design is. A second goal of our research in size-expanded DNAs is the development of tools for use in biology and medicine. For example, a set of expanded-size nucleotides might be useful as probes of steric effects in enzymes that recognize the natural nucleotides.¹² In addition, size-expanded fluorescent oligomers might have applied utility in genetic reporting and imaging.

Background and Related Studies

The formalism of Watson and Crick, who noted the analogous structures of hydrogen-bonded purine–pyrimidine pairs in 1953,¹³ had a strong influence on the DNA chemistry field in its first three or four decades. The main examples of non-AT/GC pairs prior to the mid-1990s were cases that still adhered to the paradigmatic purine–pyrimidine rule (Figure 1). Among the most important early examples in that vein were the designed purine–pyrimidine base pairs of Benner and of Rappaport.^{14,15}

The concept of nucleotides having increased size originated with Leonard, who in 1974 synthesized an

* To whom correspondence should be addressed. E-mail: kool@stanford.edu.

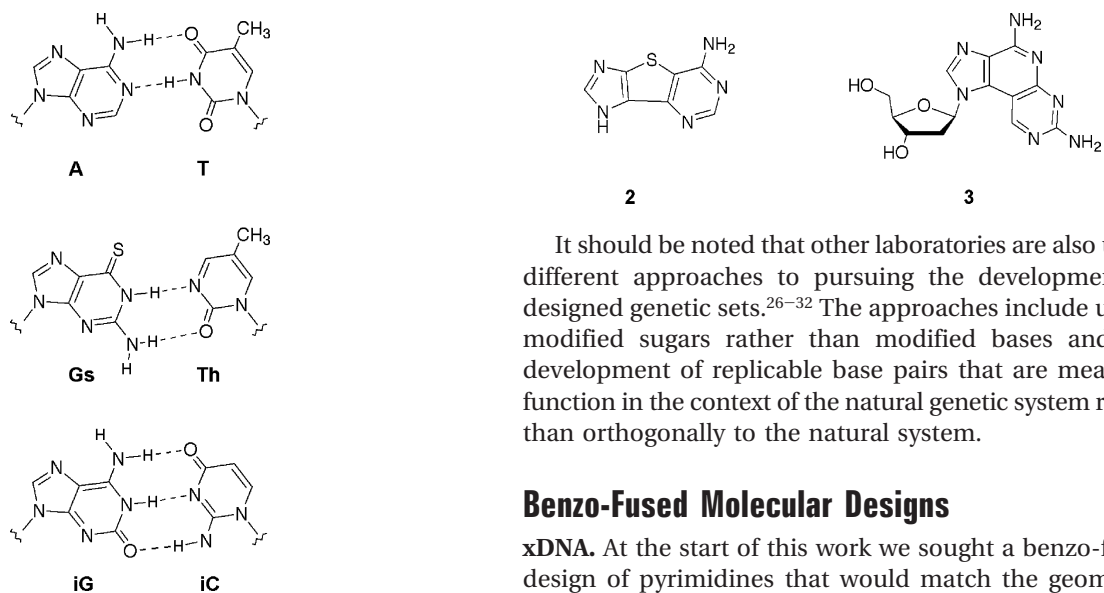
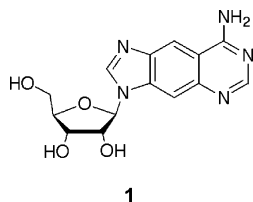


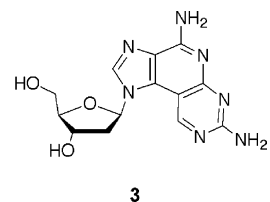
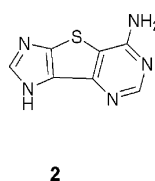
FIGURE 1. Examples of natural and nonnatural purine–pyrimidine pairs: A–T, Gs–Th,¹⁵ isoG–isoC.¹⁴ All of these conform to Watson and Crick’s purine–pyrimidine structural paradigm for the DNA double helix.

adenine ribonucleoside analogue (**1**) in which the base was “stretched out” by benzo fusion.¹⁶ The benzo-



sine nucleotide was studied as a substrate for several ATP-dependent enzymes, including firefly luciferase and a cyclic AMP-dependent protein kinase.¹² Subsequently, a similar GTP analogue was also synthesized,¹⁷ and a 2'-deoxy variant of benzoA was later prepared from the ribonucleoside.¹⁸ At the time of Leonard’s work, it was not practical to synthesize oligonucleotides. However, he predicted that benzoA was too large for a natural DNA or RNA helix.¹⁹

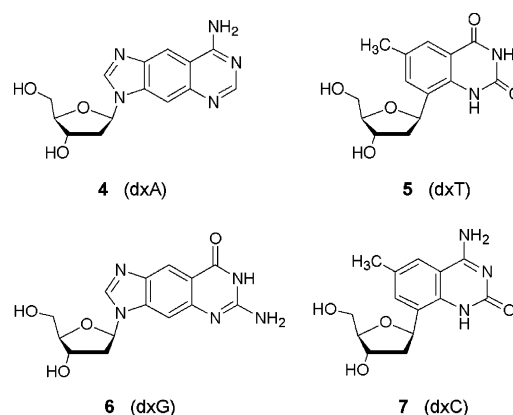
Only a few examples exist of studies in DNA involving bases larger than natural ones. Benzo-fused DNA bases were prepared by Moreau and by Saito.^{20,21} However, their point of attachment was at the natural position; thus, base pairs with the natural partner would adopt the normal rather than stretched geometry. The same is true of a large cytosine derivative reported by Matteucci.²² In studies more closely related to our strategy, Seley prepared a thiophene-fused adenine-like nucleobase (**2**);²³ this molecule was expanded in its potential pairing size, but it has not been incorporated into DNA to test this possibility. Finally, Matsuda has prepared DNA bases with extra rings (see **3**) and has examined their pairing properties.²⁴ Some of these were intended to increase the number of hydrogen bonds to four per pair, without changing the overall DNA geometry; however, a recent report describes base pairs (inserted into the natural DNA context) that could take on a stretched glycosidic distance.²⁵



It should be noted that other laboratories are also using different approaches to pursuing the development of designed genetic sets.^{26–32} The approaches include use of modified sugars rather than modified bases and the development of replicable base pairs that are meant to function in the context of the natural genetic system rather than orthogonally to the natural system.

Benzo-Fused Molecular Designs

xDNA. At the start of this work we sought a benzo-fused design of pyrimidines that would match the geometric expansion of purines conceived by Leonard.¹² A generalized design would, in principle, allow for the combination of expanded versions of A, G, C, and T with the natural base complements and might allow them to form a regular helix. This concept led to the new deoxyribosides dxT and dxC and to the complete set of expanded DNA (xDNA) nucleosides (**4–7**). All of the expanded bases have their



pairing edges shifted outward by 2.4 Å (the width of benzene), with similar vectors for this extension. Preliminary AM1 calculations predicted the structural stability and tautomeric preferences of the xDNA bases,³³ and later high-level calculations by Fuentes-Cabrera et al. added support.³⁴

The base pair designs for xDNA (Figure 2) are in some respects closely analogous to those of DNA, but are also different in important ways. Natural DNA sequences are composed of one of four letters at each position and involve purines paired with pyrimidines. In xDNA, *benzopurines* are paired with pyrimidines and *benzopyrimidines* with purines, and there are eight components of xDNA,³⁵ with four types of ring systems.

Pairing selectivity in xDNA comes (in principle) not only from complementary hydrogen bonding, but also from complementary size. For example, adenine paired with thymine is a mismatch in xDNA, because (although it has complementary hydrogen-bonding groups) the pair

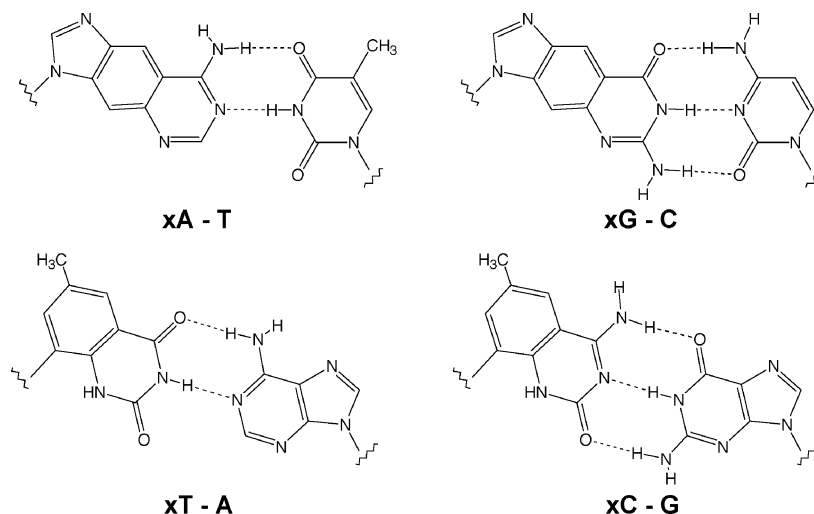


FIGURE 2. Four base-pairing schemes of xDNA. xDNA is comprised of eight bases, while natural DNA has four bases with two pairing schemes.

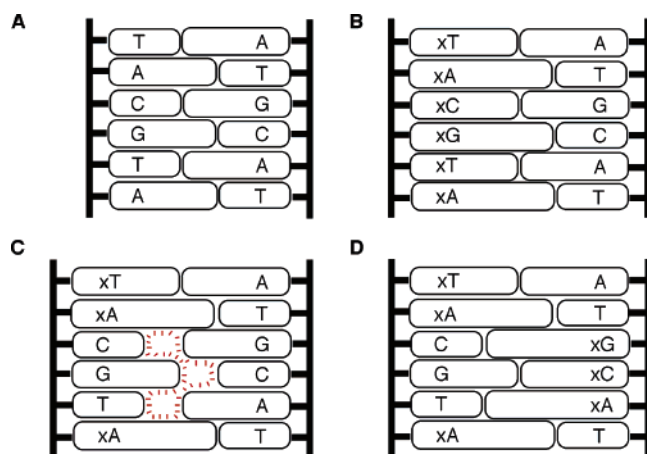
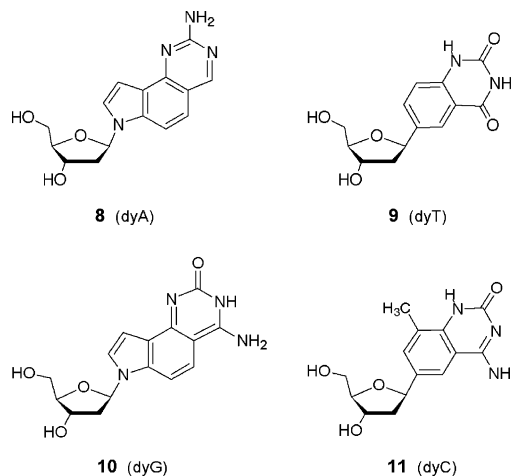


FIGURE 3. Illustration of nucleobase size differences that lead to two classes of expanded DNA helices. (A) Watson–Crick DNA strands are complementary in size. (B) An all-expanded xDNA strand is size complementary to natural DNA. (C) Mixing expanded and nonexpanded bases on a strand (eight-base xDNA) results in a size mismatch (orthogonality) with natural DNA. (D) Eight-base xDNA can be complementary to another eight-base xDNA strand.

is too small for the xDNA helix by approximately 2.4 Å. For that reason, the eight-base xDNA genetic set and the natural DNA genetic set are “orthogonal”, in that they are not expected to be able to interact with one another (Figure 3). However, it also occurred to us that a *subset* of xDNA designs might allow the modified xDNA bases to recognize natural DNA or RNA. In that subset, all the xDNA bases would be segregated onto one strand, entirely composed of xA, xG, xT, and xC.³⁵ Such size-expanded strands, in the proper sequence, might bind natural nucleic acids and still form a large helix (Figure 3B). In principle, one might use this recognition, and the inherent fluorescence of xDNA bases (see below), to sense natural DNAs or RNAs.

yDNA. Modeling studies suggested that there was more than one way to increase the size of a base pair by benzo fusion. This led to the “yDNA” (an abbreviation of “wide

DNA”) design.^{36–38} Four benzo-widened nucleoside components of yDNA are shown below (8–11). Simple analysis

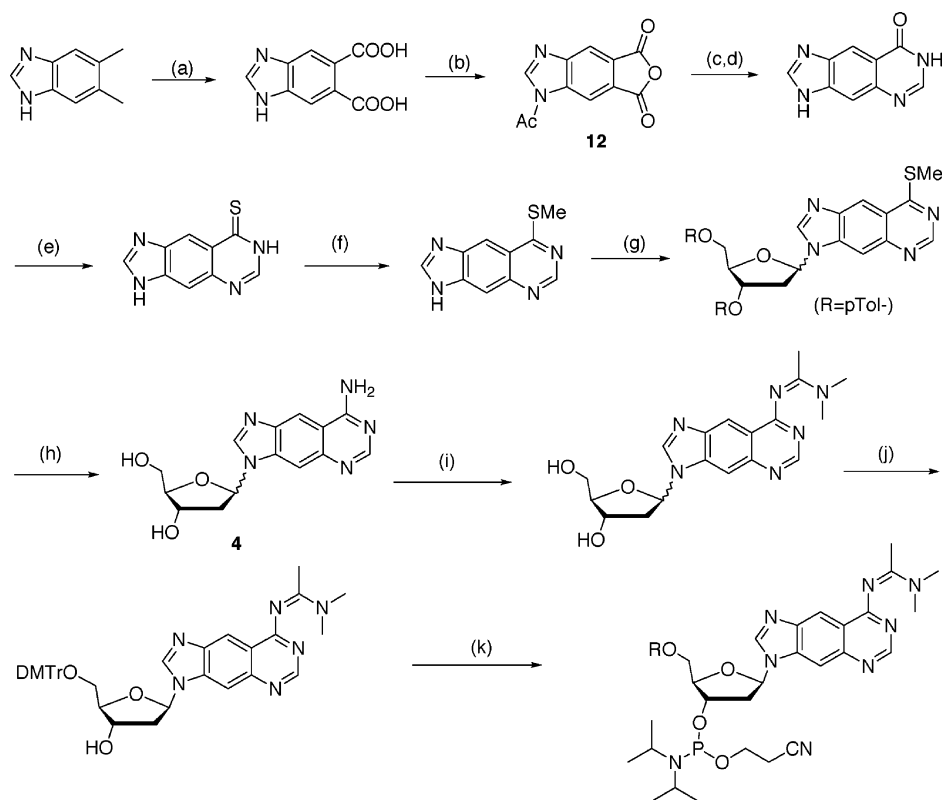


of base pair geometry suggested that yDNA would have about the same degree of helix expansion as xDNA, and analysis of possible base stacking geometries (assuming a right-handed helix) suggested that yDNA is at least as favorable in area of overlap as xDNA.

A genetic pairing system based on the yDNA design would (aside from the geometry of extension) be analogous to xDNA. However, AM1 calculations of the yDNA bases suggested that tautomeric preferences might be more complex than for the xDNA cases. The yT and yA bases were predicted to prefer the desired tautomers, but yC and yG were calculated to prefer tautomers different from those needed for pairing as in the original design.^{38,39} Of course, aqueous solvation might change the relative tautomeric preferences. As a result of these calculations we focused our initial synthetic efforts only on yT, yC, and yA and postponed studies aimed at yG.

Synthesis of Size-Expanded Nucleosides

The syntheses of seven xDNA and yDNA deoxynucleosides have been completed to date; in general, the expanded

Scheme 1. Synthetic Route for Preparation of the dxA Phosphoramidite Derivative^a

^a Conditions: (a) KMnO_4 , tBuOH , H_2O , 70°C , 1.5 h, 60%; (b) Ac_2O , 155°C , 3 h, 90%; (c) $(\text{TMS})\text{N}_3$, $90\text{--}95^\circ\text{C}$, 3 h; (d) formamidine acetate, DMF , 155°C , 3 h, 55% (two steps); (e) P_2S_5 , Py , 140°C , 36 h, 100%; (f) CH_3I , KOH , 1 h, 64%; (g) NaH , CH_3CN , Hoffer's chlorosugar, 6 h, 37%; (h) NH_3 , EtOH , 140°C , 36 h, 61%; (i) N,N -dimethylacetamide dimethyl acetal, MeOH , 80°C , 72 h, 69%; (j) DMTrCl , DIPEA , Py , 23°C , 6 h, 54%; (k) 2-cyanoethyl N,N -diisopropylchlorophosphoramidite, CH_2Cl_2 , DIPEA , 23°C , 5 h, 90%. Reprinted from ref 33. Copyright 2004 American Chemical Society.

pyrimidines (dxT, dxC, dyT, dyC) are readily prepared in good yields, while the expanded purines (dxA, dxG, dyA) are more challenging.

The synthesis of dxA started with the approach of Leonard⁴⁰ and then deviated to be complete in fewer steps and higher yields (Scheme 1).⁴¹ The deoxyribonucleoside **4** was subsequently converted to an amidine-protected phosphoramidite (Scheme 1) which can be used to make synthetic DNA oligonucleotides on an automated synthesizer. The synthesis of dxG⁴² began with intermediate **12** from the dxA synthesis. Starting from **12**, the dxG phosphoramidite was synthesized in 12 steps and 4.6% overall yield and is the most challenging of the expanded nucleosides to prepare.

In contrast to the expanded purines, the dxT and dxC deoxyribonucleosides were made much more efficiently, in up to 34% overall yield.^{41,42} The key step for these C-glycosides was glycoside bond formation (Scheme 2); in the presence of $\text{Pd}(\text{OAc})_2$ and AsPh_3 , the iodinated base analogues were added via Heck reaction onto a silyl-protected dihydrofuran (see Scheme 2).⁴³

The syntheses of three yDNA nucleosides have also been reported recently.^{36,38} The preparation of the expanded deoxyadenosine analogue dyA was achieved in 10 steps and 5.7% overall yield. The yC and yT variants were prepared in good yields in a manner closely analogous to the earlier syntheses of their xDNA analogues.³⁸

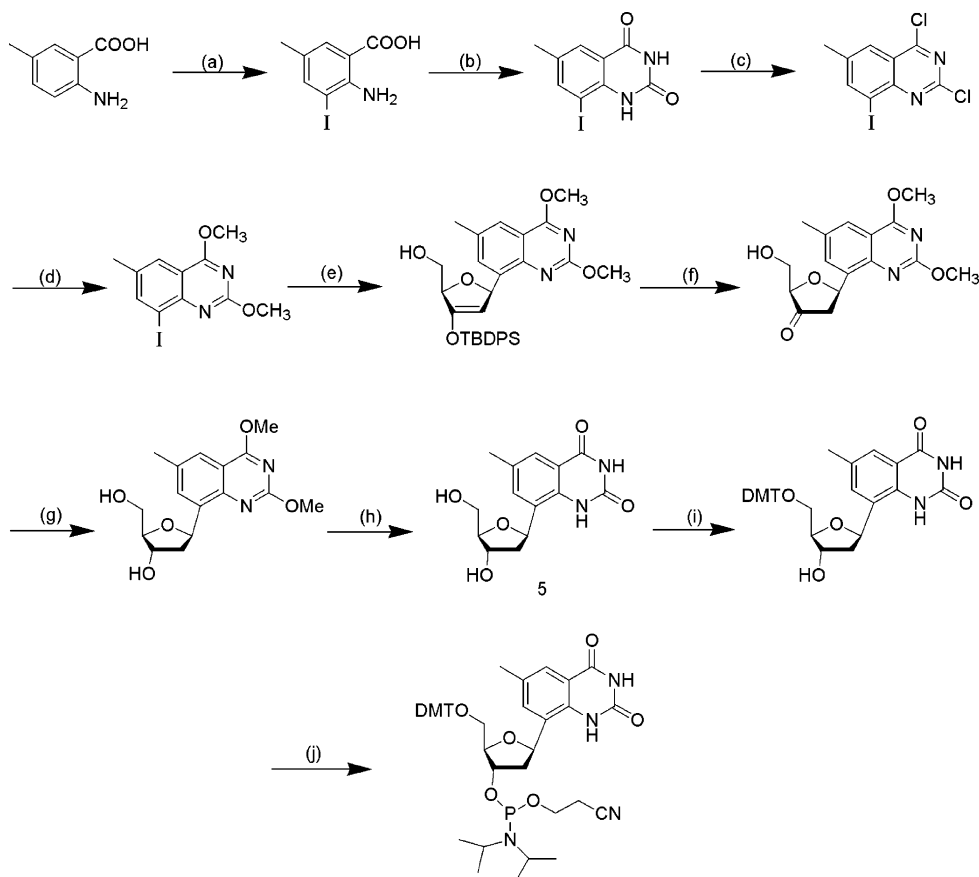
Fluorescence of Expanded DNA Bases

The additional conjugation rendered by the benzo fusion in the expanded DNA bases makes them inherently fluorescent, a useful property that is practically absent in the native DNA bases. In conjunction with the abilities of xDNA and yDNA to pair and assemble (below), this "built-in" fluorescence lends itself to possible applications in probing DNA/RNA hybridization and protein–DNA interactions.

The expanded DNA bases fluoresce in the blue-violet to violet range with conveniently large Stokes shifts of 50–80 nm (Figure 4).^{36,38,41,42} They are efficient fluorophores, with quantum yields between 0.3 and 0.6. Studies are currently under way to evaluate interaction of x,yDNA bases in the context of common fluorescence assays involving static fluorescence, time-resolved fluorescence, energy transfer, quenching, etc. Thus, it appears that expanded DNA bases will join other recent fluorescent DNA base analogues that can report on biophysical and biochemical phenomena that occur inside the helix.^{21,44–52}

Base Pairing and Stacking in Natural DNA

Before studying fully expanded helices, we tested the effects of benzo-expanding a single pair in otherwise natural DNA. Expanded-size analogues were incorporated into short double helices, and their stability was measured (by thermal denaturation) relative to that of fully natural

Scheme 2. Synthetic Route for Preparation of the dxT Phosphoramidite Derivative^a

^a Conditions: (a) ICl, HCl(aq), 96%; (b) urea, 150 °C, 92%; (c) POCl₃, *N,N*-diethylaniline, reflux, 90%; (d) NaOCH₃/CH₃OH, reflux, 87%; (e) 1,2-dehydro-3-*O*-(*tert*-butyldiphenylsilyl)-5-(hydroxymethyl)furan, Pd(OAc)₂, AsPh₃, N(Bu)₃, 75 °C, 64%; (f) TBAF, THF, 0 °C, 84%; (g) NaB(OAc)₃H, THF, AcOH, -15 °C, 93%; (h) NaI, AcOH, 60 °C, 100%; (i) 4,4'-dimethoxytrityl chloride, DIPEA, pyridine, 84%; (j) *N,N*-diisopropylammonium tetrazolid, 2-cyanoethyl tetraisopropylphosphoramidite, CH₂Cl₂, 82%. Reprinted from ref 33. Copyright 2004 American Chemical Society.

helices. The results showed consistently that a single expanded pair in the middle of a natural DNA duplex was destabilizing to the double helix. This is not too surprising considering the size mismatch of the expanded base pair in natural DNA, which would impose strain on the backbone. For the xDNA analogues, the destabilization penalty ranged from 0.3 to 1.7 kcal/mol relative to the stability of natural base pairs.⁵³ For yDNA pairs, the range was 0.6–2.2 kcal/mol.^{36,38} Interestingly, while single expanded DNA analogues decreased the thermal stability of the duplex, they were still selective for their Watson–Crick partners. For example, xA formed a more stable duplex paired opposite T than it did when paired opposite C, G, A, or a sugar lacking a base.⁵³

Stacking interactions between the aromatic DNA bases play a dominant role in duplex stability,^{54,55} and these interactions appear to be driven largely by van der Waals forces and, for hydrophobic molecules, by solvophobic effects.^{55,56} Through experiments with xDNA bases dangling at the ends of duplex DNAs, we observed strong stacking of the benzo-fused bases (Figure 5). In comparison with the natural bases as overhanging moieties, xA and xT are better stackers by 1.2 and 1.0 kcal per base.⁵³ For the expanded analogues xC and xG the advantage in stacking over the natural bases was 0.5 and 0.7 kcal per base, respectively.⁴² Subsequent studies with the yDNA

bases showed that most stack nearly as strongly as the xDNA bases.^{36,38} These exceptionally strong stacking abilities apparently result from a larger surface area (lending greater polarizability) and greater hydrophobicity. Similar observations with bases having increased aromatic surfaces have been noted by a number of laboratories.^{24,57–60}

Size-Expanded Double Helices

In a functioning genetic system, not only must isolated bases pair correctly, but they must also recognize their pairing partners when strung together in information-encoding sequences. In natural DNA, there are four monomeric components forming four pair orientations (A–T, T–A, C–G, G–C), which are assembled by polymerases (or by a DNA synthesizer). With xDNA, by contrast, there exist eight pairing possibilities; thus, the assembly of xDNA could be considerably more complex.

As mentioned above, a single expanded base pair is destabilizing when incorporated into natural DNA. However, one can envision a double helix composed entirely of expanded-size base pairs. By accommodating this size in all pairs, the backbone distortion created by a lone expanded pair might, in principle, be relieved.

Base Pairing in xDNA. In 2003 our laboratory demonstrated that a self-complementary oligomer composed

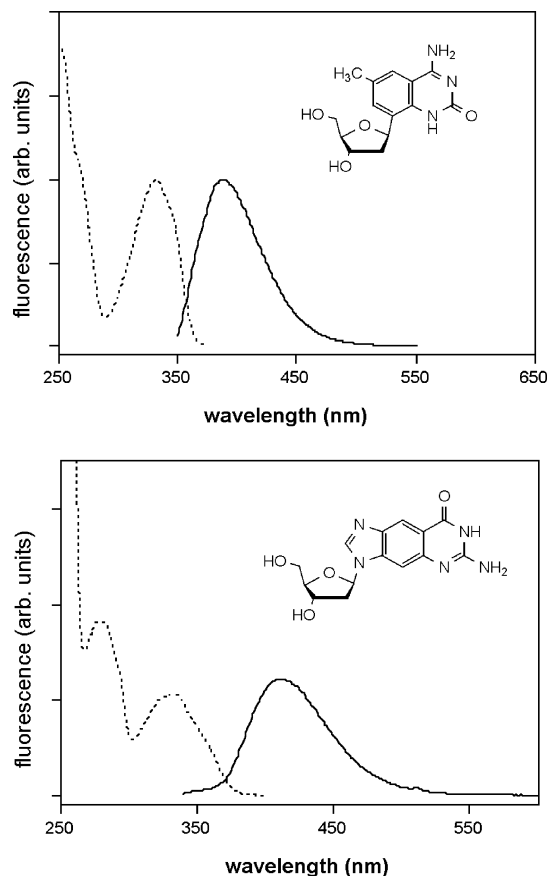


FIGURE 4. Fluorescence spectra of two xDNA nucleosides (dxC and dxG) in methanol.⁴² Emission spectra are shown with solid lines and excitation spectra with dashed lines. All xDNA and yDNA nucleobases synthesized to date are efficient fluorophores.

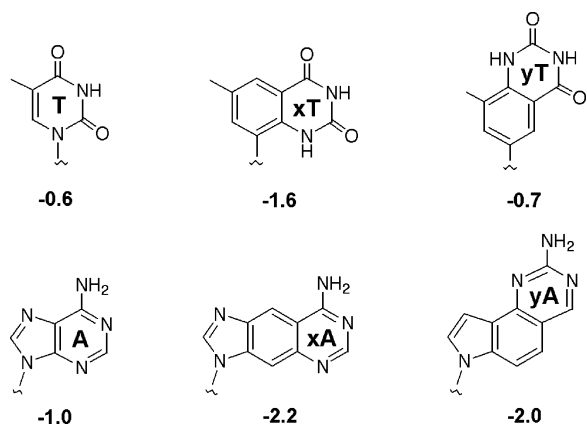


FIGURE 5. High stacking proficiency of xDNA and yDNA bases.^{36,38,42,53} Free energies for stacking (kcal/mol, 37 °C) were measured at the end of a short DNA duplex. The strong stacking of the expanded bases is likely due to both stronger van der Waals forces and the hydrophobic effect.⁶²

entirely of xA–T expanded base pairs (a 10-mer) showed a cooperative, sigmoidal transition in its UV absorbance as the temperature was raised, a property characteristic of double-stranded DNA.⁶¹ Interestingly, this complex denatured at a temperature higher than that of a natural duplex of identical sequence (56 °C vs 21 °C (Figure 6)), which might be explained by the expanded bases' enhanced stacking.

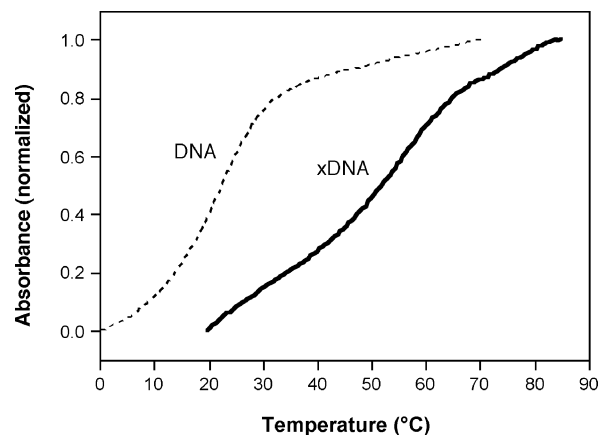


FIGURE 6. Thermal denaturation curves showing the high stability of an xDNA helix compared to DNA of the same sequence.⁶¹

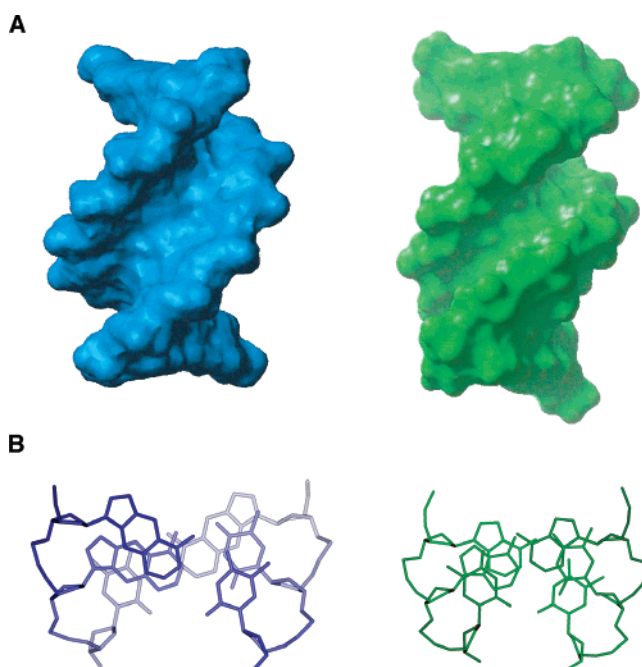


FIGURE 7. Structures of xDNA measured by NMR, with comparison to DNA (right).⁶² (A) space-filling models showing wider grooves of xDNA, but similar right-handed helical turns; (B) view of three adjacent xDNA pairs, showing a large surface area of overlap compared to that of DNA.

Subsequently, we determined a high-resolution NMR solution structure of this complex consisting of xA–T and T–xA base pairs.⁶² xDNA, like native B-DNA, exhibits a right-handed turn, antiparallel orientation, Watson–Crick hydrogen-bonding patterns, *anti* glycosidic bond conformations, and standard 2'-*endo* deoxyribose sugar conformations (Figure 7). However, the duplex diameter is larger by 3.0 Å, contributing to a smaller helical twist requiring more base pairs per turn (12 vs 10 in natural DNA). In addition, the major and minor grooves are wider than those of natural DNA by 2.5 and 2.2 Å, respectively (Figure 7A).

One of the most important hallmarks of the natural genetic system is that DNA hybridizes not only with high affinity, but also with robust sequence selectivity. We tested this with xDNA strands that contained a single

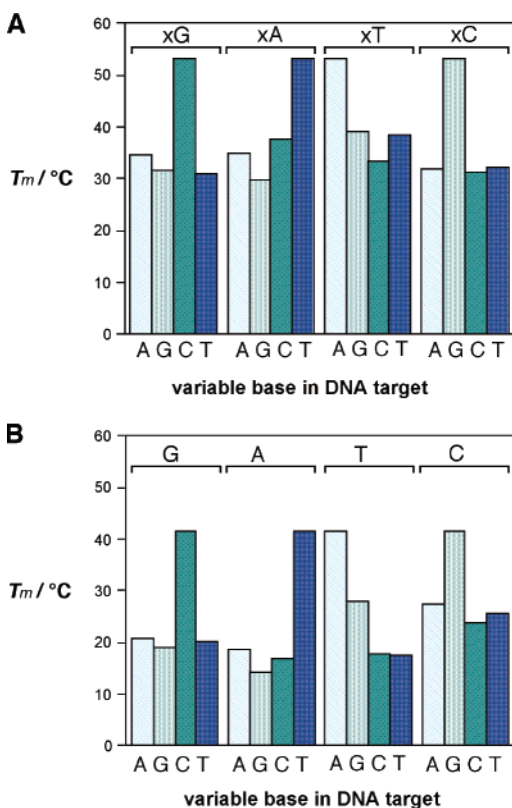


FIGURE 8. Graphs showing base-pairing selectivity of xDNA (A) compared to DNA (B), as measured by melting temperatures of short duplexes containing a single mismatched pair as shown. Reprinted with permission from ref 35. Copyright 2005 Wiley-VCH.

mismatch. These expanded duplexes also exhibited high mismatch selectivity for the expected hydrogen-bonding partners, with a selectivity profile (Figure 8) remarkably analogous to that of natural DNA.³⁵ This reinforces the possibility that an artificial genetic system other than the natural one may one day be used to selectively encode information. On a practical level, such selectivity is also important if xDNA is to be applied in hybridization-based assays with RNA or DNA.

Pairing in yDNA. The development of yDNA as part of a second designed genetic system is also furthered by data on its mismatch selectivity toward complementary hydrogen-bonding partners.^{36,38} We demonstrated that short oligomers composed entirely of yC/yT bases exhibited greater selectivity against mismatched targets than did natural DNA of analogous sequence. For example, for mismatches opposite yT, T_m values dropped 15–27 °C with a single mismatch, representing sizable selectivity. By comparison, natural DNA showed smaller T_m drops of 9–16 °C.

Early Studies of Polymerase Replication of Expanded DNAs

As was discussed in the Introduction, one of the long-term goals of this work is to determine whether size-expanded DNAs can mimic the biochemical functions of the natural genetic system. One of the most important functions of natural DNA is its ability to template its own synthesis, catalyzed by DNA polymerase enzymes. Thus,

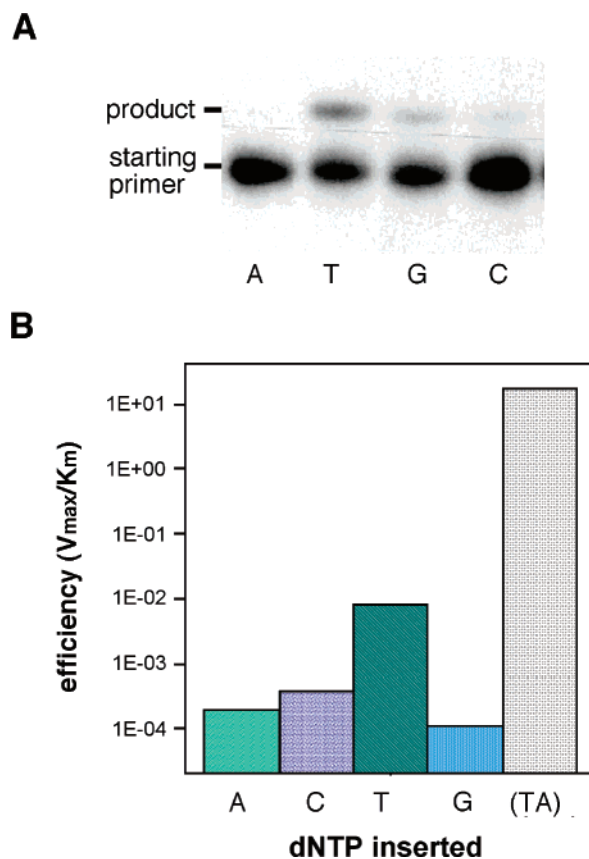


FIGURE 9. Preliminary data showing the ability of DNA polymerase I (Klenow fragment) to function with the xA base:⁶⁶ (A) polyacrylamide gel showing the product addition of dxA to the end of a primer opposite A, T, G, or C; (B) histogram showing the relative efficiency of insertion of dTTP opposite xA. Note the log scale of efficiency; the natural TA base pair efficiency is given for comparison at the right.

we ask the same question of xDNA and yDNA: Is it possible to find, modify, design or evolve an enzyme to replicate these expanded helix-forming polymers?

We expect that discovery or development of such an enzyme—an xDNA or yDNA replicase—will not be a simple task, because of the highly specialized functions of natural DNA polymerase enzymes. Many DNA polymerases work efficiently (up to hundreds of nucleotides per second) and with high specificity, making an error in nucleotide selection only once per 10000 to 100000 events.⁶³ Data suggest that this selectivity is governed chiefly by the enzyme tightly surrounding the base pair being synthesized, using steric effects to reject incorrect nucleotides.⁶⁴ Even small steric differences can cause large kinetic effects.⁶⁵ For this reason, we might expect most DNA polymerases to accept xDNA base pairs reluctantly.

Early experiments with xDNA bases in template strands of DNA add some support to this expectation (Figure 9). DNA polymerase I from *Escherichia coli* (Klenow fragment) inserts natural DNA nucleotides opposite xDNA bases relatively poorly, with an efficiency ca. 1000-fold lower than that of natural pairs.⁶⁶ However, the fact that there is any activity at all could be considered promising, and interestingly, some selectivity remains in this activity. For example, thymine (rather than C, G, or A) is preferentially

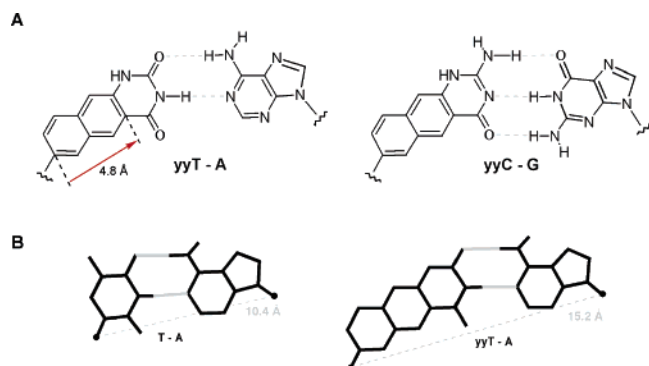


FIGURE 10. Pairing scheme of double-wide DNA (yyDNA): (A) expected base pair structures; (B) comparison of the DNA pair size and geometry with those of the yyDNA pair. The proposed double helix has an expected diameter nearly 5 Å wider than that of natural DNA.

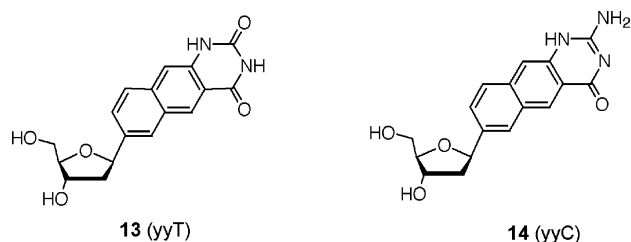
inserted opposite benzo adenine (Figure 9B), and similar selectivity is also seen with other expanded bases.⁶⁶

Addition of a single nucleotide is not replication, so there remains much work to be done in finding replicases for expanded DNA. The longer term goals will require the synthesis of entire complementary xDNA or yDNA strands and demonstration that two complementary strands can template enzymatic synthesis. Possible strategies for this might include searching for naturally occurring enzymes that are more forgiving in their substrate specificity⁶⁷ and mutagenesis of existing enzymes. A combination of such approaches may eventually yield enzymes capable of full replication of expanded DNAs.

Is Further Expansion Possible? The Naphtho Fusion Approach

The self-assembly of the xDNA and yDNA systems leads naturally to a new question: How far can base pairs be expanded and still allow for this assembly? A homologous way to test this would be to add yet another benzene ring to the pairs. This would yield 4.8 Å of expansion, giving hypothetical xxDNA and yyDNA designs (Figure 10).⁶⁸

A recent study began testing this question in the yyDNA (“double-wide DNA”) system. It proved straightforward to synthesize the two naphtho-T and -C analogues dyyT (**13**) and dyyC (**14**).⁶⁸ They are both fluorescent (like their



smaller yDNA analogues), but with red-shifted absorption and emission, as expected for the more greatly conjugated π systems. A preliminary series of helix assembly studies has been carried out in the yyDNA system. Results showed complex melting behavior consistent with strong stacking of the component bases.⁶⁸ Mixing plots, CD, and

fluorescence studies all gave evidence for double-stranded helix formation. Quenching and fluorescence resonance energy transfer experiments also pointed to double-stranded complexes of yyDNA. Further studies will be needed to evaluate the structure and stability of these complexes.

There are a number of possible ways to assemble naphtho-expanded systems. For example, it may be possible to pair xDNA bases with other xDNA bases, and one could imagine a similar yyDNA design as well. Thus, there is more work to be done in testing the limits of expanded helices.

Conclusions and Future Prospects

Our experiments confirm that natural DNA is not the only base-pairing framework that allows for stable and selective helix assembly. Size-expanded DNA sequences can spontaneously assemble themselves into right-handed, double-stranded helices with antiparallel orientation. They form base pairs with sequence selectivity and stability at least as high as those of natural DNA. Thus, it is clear that new base pair structures and geometries can be grafted onto the natural DNA backbone, with retention of helix assembly properties.

The research to date also suggests that xDNAs and yDNAs may have useful applications. The fluorescent monomers may be useful probes of DNA-modifying enzymes and nucleotide-dependent enzymes in general. When expanded DNA monomers are assembled into oligomeric strands, such strands can bind natural DNAs or RNAs sequence selectively, forming expanded helices. This binding occurs with high affinity, suggesting use of xDNAs or yDNAs in binding natural RNAs or DNAs even where they are partially blocked or folded in existing structures.

These experiments also give us new perspectives about the evolved design of the natural genetic system. Clearly, DNA base pairs did not evolve their structure because that was the only framework that could properly function in self-assembly. Our data show that size-expanded base pair designs appear to function as well. This suggests that it is more important to have a regular helix with consistent base pair geometries than it is to conserve the absolute size of DNA. Perhaps the most convincing reason for the evolution of DNA's native structure (rather than something like xDNA) is its chemical simplicity. The natural DNA bases are smaller and have fewer carbon-carbon bonds than our expanded designs, which makes their prebiotic synthesis easier. Although the greater structural complexity does not prevent chemists from making molecules such as xDNA, it does represent an advantage for natural DNA in a primordial soup where base assembly first occurred.

There remains much to be discovered with size-expanded genetic sets. In the biological realm, how far can the biomimicry be taken: Can expanded DNAs be replicated? Can they be transcribed? Can they evolve? Can they function inside native or modified cells? And beyond

the basic biological implications, are there other useful applications of expanded DNAs in biotechnology or nanotechnology? Such questions will keep us and others in the field of base pair design occupied for some time to come.

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