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N2′f**P3**′ **Phosphoramidate Glycerol Nucleic Acid as a Potential Alternative Genetic System**

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Glycerol nucleic acid (GNA), which is based upon an acyclic, three-carbon backbone (Figure 1A), is a promising starting point for the construction of an artificial genetic system owing to its ability to form stable, antiparallel duplexes.^{1,2} As proposed by Orgel, 3 we imagine that a nonenzymatic, template-directed polymerization process mediated replication during the evolution of early genetic systems,^{4,5} in place of enzymatic replication as in the modern genetic system. To recapitulate this process, our group has focused on the chemistry of polymerization facilitated by an imidazoleactivated phosphate group and an amino group as the nucleophile (Figure 1B and 1C, highlighted in red).⁵ This approach requires the newly synthesized strand with $N\rightarrow P$ phosphoramidate linkages to form stable, Watson-Crick base pairs with the template strand. The base-pairing properties of the GNA analogue with $N2'\rightarrow P3'$ phosphoramidate linkages (npGNA, Figure 1A) have not been fully explored.⁶ In this study, we demonstrate that npGNA can form stable duplexes with itself and with GNA. We also show that GNA oligonucleotides containing $N2' \rightarrow P3'$ phosphoramidate linkages can be assembled via a nonenzymatic, template-directed ligation of 3′ imidazole-activated-2′-amino GNA dinucleotides (Figure 1C). These results suggest that npGNA is a potential candidate for a selfreplicating system based on phosphoramidate linkages.

Figure 1. GNA-based genetic system. (A) Structures of GNA and npGNA. (B and C) Chemistry of templated synthesis of npGNA using imidazoleactivated monomer (B) or dimer (C). The nucleophile and leaving group are highlighted in red.

npGNA oligomers were synthesized via a solid-phase, oxidative amination coupling method^{7,8} using $2'$ -amino glycerol nucleosides as the monomers (**1t**-**1g**, Scheme 1). Compared with the lengthy synthesis $(7-9$ steps) of phosphoramidites used in the previously reported amine-exchange method,⁶ $1t-1g$ can be prepared in $2-3$ steps from the readily available 3′-dimethoxytrityl (DMT) protected glycerol nucleosides **2t**-**2g** (Scheme 1).9 Each synthetic cycle of monomer addition consists of the following steps: detritylation, phosphitylation of the 3′-hydroxyl, hydrolysis, and oxidation followed by coupling as previously described by Gryaznov et al. (Scheme 2) (see Supporting Information for detailed procedures).^{7,8} Controlled-pore glass (CPG) resins premodified with thymidine were used in the synthesis (Scheme 2), which led to an extra *Scheme 1.* Synthesis of 2′-Amino Glycerol Nucleoside Monomers*^a*

Scheme 2. Synthesis of npGNA Using Solid-Phase, Oxidative Amination Coupling*^a*

^a Conditions: (a) 3% dichloroacetic acid in CH₂Cl₂; (b) 0.4 M 2-cyanoethyl-*N,N*′*-*diisopropylchlorophosphite/0.4 M diisopropylethylamine in CH2Cl2; (c) 0.4 M tetrazole in 9:1 acetonitrile/water; (d) 0.2 M **1t**, **1c**, **1a**, or **1g** and 0.2 M triethylamine in 1:1 CCl₄/acetonitrile. Functional groups involved in each transformation are highlighted in red.

thymidine at the 2′-terminus in all npGNA sequences synthesized in this study (Table S1).

This method was validated by the synthesis of dinucleotide model compounds (**5t**-**5g**, Figure S1). Only one diastereomer was observed for all 4 dinucleotides in ¹ H and 31P NMR spectra, indicating a clean S_{N2} attack by azide leading to $4t-4g$. In addition, the dinucleotides had $31P$ chemical shifts of ∼7 to 8 ppm (Supporting Information, Figure S1) and underwent acid-catalyzed decomposition, both characteristic of a phosphoramidate linkage. The coupling efficiency of each cycle is [∼]40% for **1a** and 70-80% for **1t**, **1c**, and **1g**. The decreased coupling efficiency compared with phosphoramidate DNA synthesis is thought to be related to the decreased nucleophilicity of the $2'$ -NH₂ due to the presence of

Figure 2. Thermal denaturation curve of a 1:1 mixture of A9 and T10 npGNA compared with A10:T10 duplexes of DNA, GNA, and GNA: npGNA.

Table 1. Summary of Thermal Denaturation Studies

sequence	top:bottom strand	T_m (°C)
$3'$ -aaa aaa aaa a-2'	$npGNA: npGNA^a$	$34.5(32.5^{b})$
$2'$ -TTT TT T TT TT T $-3'$	GNA:npGNA	$37.6(37.5^{b})$
	GNA:GNA	49.5
	DNA:DNA	20.1
$3'-CGT$ ACG ACA $T-2'$	npGNA:npGNA	$50.0(52.5^{b})$
$2'$ -GCA TGC TGT A-3'	GNA:npGNA	64.1
	GNA:GNA	74.0
	DNA·DNA	45.0
$3'$ -(CGA ATT CG) ₂ -2'	npGNA:npGNA	32.0
	GNA:GNA	40 ^c
	$TNA: TNA^d$	29.8^{d}
	DNA:DNA	26 ^c

^a For A9/T10. *^b* Measured in CD experiments. *^c* Reference 1. *^d* TNA: α -L-threose nucleic acid, ref 11.

the 1'-nucleobase substitution.¹⁰ Nevertheless, we were able to isolate several npGNA oligomers (up to 10-mer) in amounts sufficient for base-pairing studies. The unreacted monomers $(1t-1g)$ were quantitatively recovered after each coupling step. Crude npGNA products were further purified by denaturing polyacrylamide gel electrophoresis or by anion-exchange HPLC (Figure S2) and characterized by MALDI-TOF MS (Table S1). All subsequent UV and CD experiments were performed on gel- or HPLC-purified npGNA oligomers.

UV thermal denaturation studies on A9:T10 npGNA showed that npGNA formed a stable duplex (Figure 2). The 1:1 stoichiometry of each strand was established by a Job plot (Figure S3). Stable duplex formation was also observed in several other npGNA sequences (Table 1). The T_m values suggested that npGNA duplexes are more stable than DNA duplexes, but weaker than GNA duplexes of the same sequence (Table 1). Formation of duplex npGNA was confirmed by circular dichroism (CD) spectroscopy, which revealed a temperature-dependent conformational change upon denaturation of npGNA duplexes (Figures 3 and S4). The transition temperatures measured by CD experiments were consistent with those obtained in UV thermal denaturation studies (Table 1).

Thermal denaturation (Figure 2) and CD (Figure $S5-S7$) studies also revealed stable heteroduplex formation between GNA and npGNA. Comparison of the T_m values in the same sequence context suggests that thermostability increases in the order of npGNA, npGNA:GNA, and GNA duplexes (Table 1). Interestingly, the reverse trend of thermostability was reported for N3^{'-P5'} phosphoramidate DNA (npDNA), which increases in the order of DNA, npDNA:DNA, and npDNA duplexes.7 Previous X-ray structural studies and molecular dynamics simulations revealed that npDNA adopts an A-form conformation and its backbone is more rigid than

Figure 3. Temperature-dependent CD studies on a 1:1 A9:T10 npGNA complex. The temperature range is between 10 and 55 at 5 °C intervals. The inset shows temperature-dependent CD signal change monitored at 273 nm.

Scheme 3. Synthesis of Imidazole-Activated Nucleotides for Templated Polymerization*^a*

^a CDI, carbonyl diimidazole; Im, imidazole; Nvoc, 6-nitroveratryl carbamate.

that of DNA.¹² We found that GNA, GNA:npGNA, and npGNA duplexes share similar CD features (Figure S5), suggesting that all three share a similar conformation in solution.

We then investigated the template-dependent synthesis of npGNA oligomers using 3′-imidazole-activated precursors (Figure 1B). A photolytic protecting group (6-nitroveratryl carbamate or $Nvoc$)¹³ was used to protect the 2'-NH₂ before activation of the 3'-phosphate group with carbonyl diimidazole (**10a**, Scheme 3A). After photolysis, however, we found that the imidazolide **11a** underwent rapid intramolecular cyclization at pH < 13 to yield **12a** (Scheme 3A). The rate of cyclization was too fast to be accurately measured by either NMR or HPLC analysis under the conditions for templated polymerization (at 4 °C, pH 8.4). However, we found that an activated dinucleotide (**14a**), which can be generated by photodeprotection of its precursor (**13a**, Scheme 3B), was much more stable than **12a** with a half-life of 53.7 h at 4 °C and pH 8.4 as measured by HPLC (Figure S8). The breakdown products of **14a** were cyclic and linear dinucleotides with a ratio of ∼3.5:1 (Figure S8).

We envisioned that **13a** could be used directly in templated polymerization reactions by generating **14a** in situ via photolysis. Because GNA forms a stable duplex with npGNA and can be conveniently synthesized in a large quantity, $GNA-(T)₁₀$ was used as the template for studying polymerization of **14a**. As shown in Figure 4A, we observed polymerization of **14a** by MALDI-TOF MS analysis. The full-length product (10-mer) was detected together with shorter oligomers after photolysis (Figure 4A). In the absence of the GNA template, only the 4-mer and a small amount of 6-mer products were observed with no full-length or 8-mer product (Figure 4B), suggesting that the template is necessary for efficient polymerization of **14a**. We also investigated incorporation of **14a** into a primer-template complex. Instead of using a GNA or npGNA

Figure 4. Template-dependent polymerization of the activated dinucleotide **14a** on T-3′-TTT TTT TTT T-2′-T analyzed by MALDI-TOF MS. (The underlined denotes the GNA 10-mer sequence): (A) with the template; (B) without the template. In B, the GNA template was cospotted with the sample as an internal control. The polymerization products were identified by comparing the signals to their expected masses. $[M + H]$ ⁺ for 4-mer: calcd, 1101.733; obsd, 1100.01. For 6-mer: calcd, 1642.349; obsd, 1640.67. For 8-mer: calcd, 2184.443; obsd, 2182.59. For 10-mer: calcd, 2725.798; obsd, 2723.68. For the template: calcd, 3168.994; obsd, 3167.08. (* denotes the $n - 1$ synthetic byproduct of the template).

primer-template complex, we used a well-characterized, 5′-[32P] labeled, 3'-NH₂-terminated DNA primer^{5,14} and a 3'-DNA-GNA hybrid template (Figure 5) for the preliminary primer extension studies on **14a**. As shown in Figure 5, we observed multiple extension products after photolysis (lane 3, arrows). Extension products were significantly reduced without the template (lane 2) or with a mismatched template (lane 4), suggesting that polymerization of **14a** was template dependent. The relatively low primer utilization (∼35%) compared with previous chemical extension studies (up to 100%)^{5,15} suggests that the B-form conformation in the primer-template pairing region is not favorable for primerextension with **14a** on the GNA template, which would lead to a duplex-GNA-like conformation.¹ In fact, primer-independent formation of 4, 6, and 8-mer was observed by MALDI-TOF MS analysis of the primer extension reactions. Future studies will be focused on the kinetics and fidelity of polymerization of short npGNA oligomers in an all-GNA or npGNA system to avoid unfavorable conformational transitions.

In summary, we have demonstrated that npGNA can form stable homo- and heteroduplexes with a conformation similar to that of

Figure 5. Template-dependent polymerization of **14a** analyzed by primer extension: (lane 1) primer only as control; (lane 2) without template; (lane 3) with template; (lane 4) with control template. The reaction mixture of 10 μ L contained 1 μ M 5'-^{[32}P]-primer/template, 2 mM **13a**, 20 mM MgCl₂, 0.2 M NaCl, 0.1 M HEPBS, pH 8.4. The reaction was initiated by photolysis at 4 °C for 6 h and, after another 12 h incubation at 4 °C, was analyzed by 20% denaturing polyacrylamide gel electrophoresis.

GNA. In addition, $N2' \rightarrow P3'$ linkages can be formed by templatedependent polymerization, which suggests that sequence information transfer can be achieved nonenzymatically in a genetic system based on GNA oligonucleotides.

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Supporting Information Available: Procedures for synthesis and characterization of **1t**-**1g**, **5t**-**5g**, and **6a**-**14a**, and solid-phase synthesis of npGNA, HPLC profile of the npGNA- $(T)_{10}$ crude product, MALDI-TOF analysis of GNA and npGNA oligonucleotides in this study, UV melting curves, and additional CD spectra. This material is available free of charge via the Internet at http://pubs.acs.org.

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