Synthesis, Affinity for Complementary RNA and DNA, and Enzymatic Stability of Triazole-Linked Locked Nucleic Acids (t-LNAs)

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Supporting Information

ABSTRACT: Dinucleoside phosphoramidites containing a triazole internucleotide linkage flanked by locked nucleic acid (LNA) were synthesized and incorporated into oligonucleotides (ONs). ONs bearing both LNA and triazole at multiple sites were obtained and their biophysical properties including enzymatic stability and binding affinity for RNA and DNA targets were studied. t-LNAs with four incorporations of a dinucleoside monomer having LNA on either side of the triazole linkage bind to their RNA target with significantly higher affinity and greater specificity than unmodified oligonucleotides, and are remarkably stable to nuclease degradation. A similar but reduced effect on enzymatic stability and binding affinity was noted for LNA only on the 3′-side of the triazole linkage. Thus, by combining unnatural triazole linkages and LNA in one unit (t-LNA), we produced a promising class of ONs with reduced anionic charge and potential for antisense applications.

INTRODUCTION

Antisense oligonucleotides (ASO) are short single-stranded nucleic acids (typically 15–25 nucleotides in length) that bind to their RNA target in a sequence-specific manner and modulate translation (protein synthesis) or RNA splicing.1 Unmodified natural DNA or RNA strands are not optimum antisense candidates because of their poor serum stability and modest binding affinity for RNA. Chemically modified nucleic acids2−4 offer a promising alternative to their unmodified counterparts and have been extensively studied since the ASO concept was first introduced.5 Some of these modifications possess favorable biophysical properties, which is evident from the recent success in the clinic.6−8 Despite this, ideal antisense candidates, with the desired characteristics of efficient unaided cellular uptake, long serum half-life and high RNA binding affinity and target specificity, remain elusive and there is a need to explore new chemically modified nucleic acids.

Recently, we have been interested in backbone-modified oligonucleotides (ONs) with a special focus on triazole-linked nucleic acids.9 A series of studies over the past decade led us to a highly versatile and biocompatible six-atom triazole linkage (monomer W, Figure 1).9−12 This and other triazole linkages13,14 have been shown to improve the stability of modified ONs to degradation by DNase enzymes14−17. Unfortunately, the thermal stability of duplexes carrying triazole internucleotide linkages is compromised by the modified backbone.13,15,17−23 This imposes a restriction on the use of triazole-linked oligonucleotides in antisense applications, even though they possess the favorable properties of resistance to enzymatic degradation and the lack of anionic charge that can hinder cell uptake of natural DNA. Building on the seminal work by Wengel,24 we have previously shown16 that introducing a conformationally restricted locked nucleic acid (LNA)25−27 next to the triazole linkage (monomers X and Z, Figure 1) counterbalances the reduction in binding affinity caused by the triazole, resulting in modified ONs that bind to their RNA target with affinity similar to unmodified ONs. Interestingly, Watts et al. found in contrast that LNA does not stabilize duplexes containing a more rigid four-atom triazole linkage internally.15 Thus, the biocompatible six-atom triazole linkage and LNA (monomers X and Z, Figure 1) is a promising combination and warrants further investigation.

In our previous study, we incorporated monomers W−Z into oligonucleotides by ligating together two short ONs, one with a 5′-azide group and the other with a 3′-propargyl group.16 This approach was chosen for ease of ON synthesis. However, multiple incorporations of the triazole linkage into short ONs and the introduction of the linkage at the ends of ONs is not practical using this strategy. Here, we report the synthesis of dinucleoside phosphoramidites containing the triazole linkage flanked by LNA as suitable building blocks for the efficient and flexible incorporation of monomers W−Z into ONs using standard solid phase phosphoramidite DNA synthesis methodology. Using this approach, 13-mer ONs with 1, 2, or 4 incorporations of the modified monomers W−Z were readily obtained, enabling us to study their enzymatic stability and
hybridization (duplex-forming) properties. Monomer Z stands out among those tested; ONs carrying four incorporations of Z are strikingly resistant toward nucleolytic degradation, and show enhanced binding to RNA targets with improved affinity and specificity relative to unmodified ONs. Thus, the combination of LNA and the triazole linkage provides a new class of potentially significant antisense oligonucleotide candidates.

**RESULTS AND DISCUSSION**

**Chemical Synthesis.** First, nucleosides with a 3′-O-propargyl group and 5′-azido group were prepared (Scheme 1). Accordingly, 5′O-(4,4′-dimethoxytrityl)-3′-O-propargyl-LNA 5-methylcytosine (1) and 5′O-(4,4′-dimethoxytrityl)-3′-O-propargyl-5-methylcytosine (2) were reacted with benzoic anhydride to obtain protected nucleosides 3 and 4 in yields of 86 and 92%, respectively. 5′-Azido-LNA thymidine (6) was obtained from protected nucleoside 5 by removing the 3′-O-benzyl group followed by nucleophilic displacement of 5′-O-mesylate with azide. 5′-Azidothymidine (7, Scheme 2) was prepared following a literature procedure.

Dinucleosides 8–11, which contain the triazole linkage with and without LNA, were prepared in good yields (79–87%) by Cu(I) catalyzed alkyne azide cycloaddition reaction between nucleosides 3, 4, 6, and 7 (Scheme 2). 3′-O-Phosphitylation of 8–11 gave phosphoramidites 12–15 as suitable building blocks for incorporating monomers W–Z, respectively, into DNA via solid-phase DNA synthesis. For introducing monomers W–Z at the 3′-end of ONs, dinucleosides 8–11 were loaded onto an activated amino-SynBase resin using N-ethyl-N′-(3-dimethylaminopropyl)carbodiimide hydrochloride (EDC) as the coupling reagent.

**Hybridization Studies.** Hybridization to Complementary RNA Targets. Having dinucleoside phosphoramidites in hand allowed multiple incorporations of the modified monomers internally and at the termini of ONs. Hence, phosphoramidites 12–15 were used to introduce monomers W–Z, respectively, into chosen ON sequences (Table 1). Six sequences were prepared: ON1–ON2 with modified monomers at 3′- or 5′-end, ON3–ON4 bearing modified monomers in the middle region, ON5 containing two modifications in the middle, and ON6 incorporating four modified monomers (at the ends and in the middle). The modified ONs were then mixed with the complementary RNA sequences in a 10 mM phosphate buffer containing 200 mM NaCl at pH 7.0 and the melting temperatures (T_m) of the resulting duplexes were determined from the UV melting curves (Figure 2).

In accordance with the literature, duplexes formed between triazole-linked DNA and fully complementary RNA strands were less stable than the unmodified DNA/RNA duplex (compare monomer W with U). However, the drop in T_m caused by the triazole linkage was fully compensated for by either the introduction of LNA on the 3′-side (monomer X) or
on both sides of the linkage (monomer Z). This observation is in full agreement with our previous study.\textsuperscript{16} Furthermore, the binding affinity of ONs carrying the modified monomers (W–Z) at the 3′-end (ON1W–ON1Z) was comparable to that of the unmodified ON (ON1U), suggesting that all the monomers are well tolerated at this position. ONs bearing monomers W or Y at the 5′-end bind to their RNA targets with a slightly lower affinity than the unmodified DNA strand (compare ON2U with ON2W and ON2Y, \(\Delta T_m\) of \(-2.5\) and \(-3.3\) °C, respectively). In contrast, monomers X and Z with \(\Delta T_m\) of \(+1.7\) and \(+0.8\) °C (compare ON2U with ON2X and ON2Z), respectively, induced a slight increase in binding affinity. A larger drop in thermal stability was observed with incorporation of either W or Y at the center of modified duplexes (\(\Delta T_m\) in the range of

\[\text{Scheme 2. Synthesis of Triazole-Linked Dinucleoside Building Blocks}\]

\[\text{Table 1. Thermal Melting (}\text{\(T_m\)}\text{) Data for the DNA/RNA Duplexes}\]

| ON   | ON sequence (5′–3′) | \(B = \text{^31}C\text{pT (U)}\) | \(\text{^31}C\text{pT}^\text{a} (V)\) | \(\text{^31}C\text{rT (W)}\) | \(\text{^31}C\text{rT}^\text{a} (Y)\) | \(\text{^31}C\text{rT}^\text{a} (Z)\) |
|------|--------------------|----------------|----------------|----------------|----------------|----------------|
| ON1  | 5′-CTC ACT ATC TGB | 54.8           | nd             | 53.7 (−1.1)    | 54.6 (−0.2)    | 54.7 (−0.1)    |
| ON2  | 5′-BCA CTA TCT GCT | 54.0           | nd             | 51.5 (−2.5)    | 55.7 (1.7)     | 50.7 (−3.3)    |
| ON3  | 5′-CTC ABA TCT GCT | 55.1           | 66.0 (+10.9)   | 49.6 (−5.5)    | 57.1 (2.0)     | 49.8 (−5.3)    |
| ON4  | 5′-CTC ACT ATB GCT | 54.4           | nd             | 49.4 (−5.0)    | 55.0 (0.6)     | 49.2 (−5.2)    |
| ON5  | 5′-CTC ABA TGB CT  | 55.7           | nd             | 43.7 (−12.0)   | 57.0 (1.3)     | 44.1 (−11.6)   |
| ON6  | 5′-BCA BAT BGB     | 56.7           | >75            | 39.4 (−17.3)   | 57.8 (1.1)     | 38.3 (−18.4)   |

\(\text{a}\)Melting temperatures (\(T_m\)) were obtained from the maxima of the first derivatives of the melting curves (\(A_{260}\) vs temperature) recorded in a buffer containing 10 mM phosphate and 200 mM NaCl at pH 7.0 using 3.0 \(\mu\)M concentrations of each strand. \(\Delta T_m = \text{change in } T_m\) for a modified duplex relative to the unmodified duplex. RNA target: 5′-AGC AGA UAG UGA G. nd = not determined. p = phosphodiester backbone. \(^*\) = LNA nucleotide. t = triazole backbone.
-5.0 to −5.5 °C, ON3W, ON4W, ON3Y, and ON4Y). In contrast, one incorporation of monomers X or Z increased the thermal stability (ΔT_m of +0.6 to +3.0 °C, ON3X, ON4X, ON3Z, and ON4Z), with monomer Z being more effective than monomer X (compare ON3X with ON3Z and ON4X with ON4Z). Indeed, monomer Z with LNA sugars on both sides of the triazole linkage induced an increase of 3.0 °C (ON3Z) in T_m which is the largest increase in this series for a single modification. Importantly, the thermal stability of a duplex with four incorporations of monomer Z was significantly higher than that of the unmodified duplex (compare ON6U with ON6Z, ΔT_m = +5.6 °C), and for monomer X (with LNA on the 3′-side of the triazole), a small increase in thermal stability was observed (compare ON6U with ON6X, ΔT_m = +1.1 °C). In contrast, the modified duplexes carrying four incorporations of monomer W (triazole linkage without LNA) or Y (LNA on 5′-side of the triazole) were strikingly less stable than the unmodified duplex (compare ON6W or ON6Y with ON6U, ΔT_m = −17.3 and −18.4 °C, respectively).

**Hybridization to Mismatched RNA Targets.** To avoid “off target” effects that can lead to the inhibition of a wrong mRNA, ASOs must be able to differentiate between matched and mismatched RNA targets. ON3 with a backbone modification at the center was chosen to assess the binding specificity of the modified monomers. The nucleotides opposite to MeC or T on either side of the triazole linkage were changed, and the T_m differences between matched and mismatched duplexes were recorded (Table 2). In general, monomers W−Z discriminated between a matched and a single-mismatch target with the efficiency similar to that of the unmodified base (U). Monomer W (triazole linkage without LNA) was least effective in sensing a mismatched nucleotide. Monomer V (LNA without a triazole linkage) gave a good mismatch discrimination in all the cases (e.g., compare ON3U with ON3V), which is in agreement with the literature.23 Monomer X, with LNA on the 3′-side of the triazole linkage showed the greatest discrimination for a mismatched nucleotide on the 3′-side of the triazole linkage. However, marginally inferior discrimination was observed for monomer Y, which carries LNA on the 5′-side of the linkage. Monomer Z, which has LNA on both sides of the triazole linkage showed improved discrimination for the mismatch nucleotide on the 3′-side of the triazole linkage (T−C and T−G mismatch). Monomer Z discriminated against a mismatch guanine base (T-G mismatch) with greater efficiency than that of the unmodified ON (compare ON3U with ON3Z). The T−G mismatch discrimination for Z was also noticeably superior to that observed for monomers X or V (compare ON3Z with ON3X or ON3V). The greater relative destabilization of the T−G mismatch is significant, as this mismatch is generally quite stable. The ability to discriminate between mispaired nucleotides was then studied for ONs containing four incorporations of monomers X and Z (Table S2) and similar results were observed. Again, significant destabilization of duplexes containing a T−G mismatch was observed (compare ON6Z with ON6U).

In summary, the combined hybridization data (Tables 1 and 2) indicate that LNA on the 3′-side of the triazole linkage maintains its beneficial effects on RNA binding affinity and specificity, but this is not true for LNA on the 5′-side of the triazole linkage. Our explanation for this behavior is that LNA

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**Table 2. Mismatch Data for ONs Containing Modified Monomer in the Center (ON3U–ON3Z)**

| ON code | ON sequence (5′−3′) | mismatch data against MeC (3′-side of the linkage) | mismatch data against T (3′-side of the linkage) |
|---------|---------------------|---------------------------------------------------|--------------------------------------------------|
|         |                     | ΔT_m<sup>a</sup> | MeC−G | MeC−U | MeC−A | T_m<sup>a</sup> | ΔT_m<sup>a</sup> | T−A | T−C | T−G |
| ON3U    | S′−CTC A<sub>MeC</sub>T A TCT GCT | 55.1 | −17.7 | −16.1 | 55.1 | −12.3 | −3.4 |
| ON3W    | S′−CTC A<sub>MeC</sub>T<sub>4</sub>A TCT GCT | 66.0 | −18.1 | −17.2 | 66.0 | −12.9 | −4.6 |
| ON3Y    | S′−CTC A<sub>MeC</sub>T<sub>4</sub>T A TCT GCT | 49.6 | −17.0 | −15.2 | 49.6 | −10.7 | −2.1 |
| ON3Z    | S′−CTC A<sub>MeC</sub>T<sub>4</sub>C A TCT GCT | 57.1 | −16.9 | −13.5 | 57.1 | −13.8 | −5.6 |

<sup>a</sup>Melting temperatures (T_m<sub>a</sub>) were obtained from the maxima of the first derivatives of the melting curves (∆A<sub>260</sub> vs temperature) recorded in a buffer containing 10 mM phosphate and 200 mM NaCl at pH 7.0 using 3.0 μM concentrations of each strand. ΔT_m<sup>a</sup> = change in T_m<sub>a</sub> for a mismatch duplex relative to the corresponding matched duplex. p = phosphodiester backbone. t = triazole backbone. Matched RNA target: S′-AGCAGAUAGUGAG. Mismatch RNA targets: S′-AGCAGAUUUGAG (MeC−U mismatch), S′-AGCAGAUAUGAG (MeC−A mismatch), S′-AGCAGAUCUGAG (T−C mismatch), S′-AGCAGAUGUGAG (T−G mismatch). DOI: 10.1021/acsomega.8b01086

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Table 3. Thermal Melting ($T_m$) Data for the DNA/DNA Duplexes

| ON code | ON sequence ($5’-3’$) | $B = ^{16}$CpT (U) | $^{30}$CpT (W) | $^{30}$CpTl (X) | $^{30}$CpT (Y) | $^{30}$CpTl (Z) |
|---------|--------------------------|---------------------|---------------|---------------|---------------|---------------|
| ON1     | 5′-CTC ACT ATC TGB        | 5.4                 | 53.7 (−0.8)   | 52.8 (−1.7)   | 53.1 (−1.4)   | 53.5 (−1.0)   |
| ON2     | 5′-BCA CTA TCT GCT        | 53.8                | 50.9 (−2.9)   | 49.1 (−4.7)   | 49.4 (−4.4)   | 49.6 (−4.2)   |
| ON3     | 5′-CTC ABT TCT GCT        | 54.9                | 47.5 (−7.4)   | 51.4 (−3.5)   | 44.4 (−10.5)  | 53.4 (−1.5)   |
| ON4     | 5′-CTC ACT ATB GCT        | 54.4                | 49.7 (−4.7)   | 51.0 (−3.4)   | 45.0 (−9.4)   | 52.1 (−2.3)   |
| ON5     | 5′-CTC ABT BGB CT         | 55.9                | 41.0 (−14.9)  | 48.6 (−7.3)   | 34.2 (−21.7)  | 51.4 (−4.5)   |
| ON6     | 5′-BCA BAT BGB            | 58.0                | 37.6 (−20.4)  | 45.8 (−12.2)  | <−30.0        | 48.3 (−9.7)   |

“Melting temperatures ($T_m$) were obtained from the maxima of the first derivatives of the melting curves ($A_{lim}$ vs temperature) recorded in a buffer containing 10 mM phosphate and 200 mM NaCl at pH 7.0 using 3.0 μM concentrations of each strand. $\Delta T_m$ = change in $T_m$ for a modified duplex relative to the unmodified duplex ($5′$-CTC ACT ATC TG$^{30}$CpT. $p$ = phosphodiester backbone. $l$ = LNA nucleotide. $t$ = triazole backbone.

Figure 3. CD spectra of duplexes formed between ON6U–ON6Z and complementary RNA: ON6: $5′$-BCA BAT BGB (ON6U, $B = ^{16}$CpT; ON6V, $B = ^{30}$CpTl; ON6W, $B = ^{30}$CpT; ON6X, $B = ^{30}$CpTl; ON6Y, $B = ^{30}$CpTl; ON6Z, $B = ^{30}$CpTl).

on the $3′$-side of the triazole is attached to a natural phosphodiester group, but when present on the $5′$-side of the triazole, the LNA sugar is attached to a more flexible triazole linkage. LNA stabilizes DNA/RNA duplexes by reducing phosphate backbone flexibility,24,33 which may not be possible when the LNA sugar is attached directly to the more flexible modified linkage and/or is close to an unstable mismatched base pair.

Hybridization to DNA Targets. Next, modified ONs were mixed with their DNA targets in a buffer containing 10 mM phosphate and 200 mM NaCl at pH 7.0 to give DNA/DNA duplexes and $T_{m,8}$ were recorded. These were then compared with the $T_m$ of the unmodified DNA/DNA duplex and differences ($\Delta T_{m,8}$) were determined (Table 3). In all the cases, the thermal stabilities of the modified duplexes were lower than those of the unmodified duplexes (negative $\Delta T_{m,8}$). However, as observed for DNA/RNA duplexes, modifications were better tolerated at the $3′$-end relative to other positions ($\Delta T_{m,8}$ only in the range of $−0.8$ to $−1.7$ °C, compare ON1W–ON1Z with ON1U). A large drop in $T_m$ was observed for monomer $W$ at internal positions, which is consistent with the literature (compare ON3W and ON4W with ON3U).19,20

Surprisingly, monomers $X$ and $Z$ were better accommodated at the center of the duplex compared to the $5′$-end, and the thermal stability of the modified duplex incorporating monomer $Z$ in the center was similar to that of unmodified duplex (compare ON3Z with ON3U, drop of only 1.5 °C). As with the DNA/RNA duplexes, monomer $Z$ gave the highest melting temperatures for all DNA/DNA duplexes. It is not surprising that the triazole–LNA combination stabilizes the DNA/RNA duplexes to a greater degree than it does the DNA/DNA duplexes. This is because the LNA sugar exists in the C3′-endo conformation, which occurs in A-type helices (DNA/RNA and RNA/RNA) but is disfavored in DNA/DNA duplexes in solution.24,34

Circular Dichroism (CD) Spectroscopy. CD spectra of the modified duplexes obtained by mixing ON6 (ON6U–ON6Z, Table 1) with RNA targets were recorded in a buffer containing 10 mM phosphate and 200 mM NaCl at pH 7.0 and the results are shown in Figure 3. Neither LNA nor the triazole linkage significantly alters the global conformation of the DNA/RNA duplexes (compare ON6V–ON6Z with ON6U). However, subtle shifts in CD maxima near 272 nm were observed. For instance, the CD spectrum of the duplex containing LNA but no triazole linkage (monomer $V$) displayed a maximum at 263 nm (a hypsochromic shift of 9 nm). For duplexes containing LNA on the $3′$-side or both sides of the triazole linkage (monomers $X$ and $Z$), the CD maxima were observed at around 267 nm (a hypsochromic shift of 5 nm), whereas monomer $W$ (triazole linkage but no LNA) or monomer $Y$ (LNA on the $5′$-side of the linkage) did not induce a noticeable shift in their CD maxima. Interpretation of these shifts in terms of DNA conformation is not straightforward, but the changes are modest, and overall geometry of the duplexes...
carrying LNA–triazole modifications is likely to be similar to the unmodified DNA/RNA duplex. Nevertheless, an interesting and significant trend emerges. The hypsochromic shifts observed for LNA-containing duplexes in Figure 3 are in line with previously reported data and could be attributed to enhanced A-form-geometry due to the restricted conformational mobility of LNA. Assuming that flexible triazole linkages partially (triazole–3′-LNA) or almost totally (5′-LNA–triazole) eliminate beneficial LNA-driven A-form conformational changes, the extent of the hypsochromic shift should decrease in the following order: ON6V (uncompensated LNA effects) > ON6X and ON6Z (partially compensated LNA effects) > ON6Y (almost totally compensated LNA effect) > ON6W and ON6U (no LNA, zero shift). The relative magnitudes of these hypsochromic shifts are also proportional to differences in melting temperatures, precisely as would be expected.

Enzymatic Stability of t-LNAs. Oligonucleotides ON6U–ON6Z (Table 1) were incubated with phosphodiesterase 1 from Crotalus adamanteus, snake venom, in 50 mM Tris buffer containing 10 mM MgCl₂ at pH = 9.0. (A) Comparison of nuclease stability of ON6U, ON6V, and ON6Z. (B) Comparison of nuclease stability of ON6W, ON6X, and ON6Z.

Figure 4. Denaturing polyacrylamide gel electrophoresis (PAGE) analysis of ON6U–ON6Z after digestion with phosphodiesterase 1 from C. adamanteus, snake venom. This shows that the introduction of LNA next to the triazole improves stability against nucleases. Digestions were performed using snake venom in 50 mM Tris buffer containing 10 mM MgCl₂ at pH = 9.0. (A) Comparison of nuclease stability of ON6U, ON6V, and ON6Z. (B) Comparison of nuclease stability of ON6W, ON6X, and ON6Z.

Figure 5. Denaturing PAGE analysis of ON6U–ON6Z after digestion by FBS in DPBS supplemented with 50% FBS. (A) Comparison of enzymatic stability of ON6U, ON6V, and ON6Z. (B) Comparison of enzymatic stability of ON6V, ON6X, and ON6Z.
being more stable than ON6X, which, in turn, is more stable than ON6W. It is surprising that this 3′-exonuclease cleaves ON6W so readily, as this must require the removal of an entire triazole-linked dinucleotide from the 3′-end of the oligonucleotide (by digestion of the closest available unmodified phosphodiester linkage). In contrast, the extreme stability of ON6Z to snake venom phosphodiesterase shows that there is a synergistic effect between LNA and triazole, justifying the modified DNA linkage design on which this study is based.

Next, the stability of ON6V–ON6Z in fetal bovine serum (FBS) was tested and compared with that of the unmodified ON6U (Figures 5 and S2). FBS contains predominantly endonucleases, so it has different enzymatic DNA degrading activity to snake venom, which is a 3′-exonuclease. FBS was chosen for this study because it will simulate the extracellular conditions to which a therapeutic oligonucleotide would be exposed. ONs were incubated with FBS (50%) in Dulbecco’s phosphate-buffered saline (DPBS). Unmodified DNA (ON6U) was completely digested within 4 h of incubation, whereas both ON6Z and ON6V were stable for up to 8 h (Figure 5A). The observation that LNA enhances the resistance of triazole-linked ONs against nucleolytic degradation in serum was further supported by the fact that ON6Z (LNA on both sides of the triazole linkage) is more stable (visible up to 24 h) than the ON6X (LNA only on 3′-side of the linkage), and the observation that ON6W containing triazole linkages without LNA is less stable (Figure S2). Interestingly, ON6V (eight LNAs) was more stable in serum than ON6X (four triazole linkages and four LNAs), whereas ON6V and ON6Z with eight LNA sugars have similar stability (both stable up to 24 h). This confirms that LNA is a significant contributor to enzymatic stability in serum. The triazole linkage (monomer W) is a mimic of phosphodiester linkage and is biocompatible, so it might be able to bind to serum endonucleases in a fashion similar to unmodified linkage and allow adjacent phosphodiester bonds to be cleaved (ON6X has eight unmodified phosphodiester linkages). In contrast, LNA has a locked sugar (3′-endo), which could inhibit the binding of nucleases to the single-stranded ON. To summarize, t-LNAs with LNA on either side of the triazole linkage have the highest stability against nucleolytic degradation, in addition to the strongest affinity for RNA targets.

## CONCLUSIONS

The efficient synthesis of dinucleoside phosphoramidites containing a triazole linkage and LNA sugars has been achieved. Access to these dinucleoside phosphoramidites allowed the incorporation of the triazole linkage flanked by LNA at multiple positions in oligonucleotides, including the central region and the 3′- and 5′-ends. This was not feasible with our earlier ligation-based strategy where short ONs were joined together using click chemistry. There are several examples of a short triazole linkage being introduced into oligonucleotides via phosphoramidite monomers, but in all the cases, the duplexes were greatly destabilized.19–21 There is also one report of this short triazole being combined with LNA and added to oligonucleotides by phosphoramidite chemistry, but unlike our triazole, this particular linkage was strongly duplex-stabilizing despite the presence of LNA.15 Another study described the synthesis of triazoles-linked DNA by phosphoramidite chemistry using an inverted version of the triazole unit employed in this study, and this linkage slightly destabilized double-stranded DNA.17 In contrast, in this study, we have shown that oligonucleotides (t-LNAs) that contain the triazole linkage in Figure 1 flanked by LNA on the 3′- or both sides (monomers X and Z, respectively) bind to their RNA targets with higher affinity and specificity than the equivalent unmodified ONs. In addition, t-LNAs containing multiple incorporations of monomers X and Z are highly resistant to nuclease degradation, and there is a synergistic effect on enzymatic stability from the combined triazole–LNA linkages.

Further biochemical and biological studies are required, but so far the indications are that t-LNAs constitute a promising class of oligonucleotides with potential antisense and splice-modulating applications.36 They display extreme stability to enzymatic degradation, high RNA-binding affinity and RNA target specificity, and reduced anionic charge. The effects on cellular uptake of reducing the overall anionic nature of oligonucleotides by incorporation of LNA–triazole linkages will be interesting to study, given that oligonucleotide delivery is a major issue in therapeutic applications. The dinucleotide phosphoramidite strategy described in this article also allows for the synthesis of LNA triazoles, which incorporate phosphorothioate linkages, which might further enhance cell uptake. It is also possible that such oligonucleotide analogues will have different toxicological properties than LNA because it is not possible for any of the LNA–triazole dimers to be degraded into LNA nucleotides, so they cannot enter the mononucleotide triphosphate pool and therefore they cannot be incorporated into genomic DNA or RNA. The results presented here and previously16 suggest that, in addition to potential therapeutic applications, modified t-LNAs could find uses in imaging, diagnostics, and nontherapeutic siRNA applications.

## EXPERIMENTAL SECTION

All the reagents were purchased from Sigma-Aldrich, Alfa Aesar, Fisher Scientific, or Link Technologies and used without further purification. Pyridine (from KOH) and POCl₃ were freshly distilled before use, and THF was obtained using the MBrän SPS Bench Top solvent purification system (SPS). All air/moisture-sensitive reactions were carried out under inert atmosphere (argon) in an oven-dried glassware. Reactions were monitored by thin-layer chromatography using Merck Kieselgel 60 F24 silica gel plates (0.22 mm thickness, aluminum backed). The compounds were visualized by UV irradiation at 254/265 nm and by staining in p-anisaldehyde solution. Column chromatography was carried out under pressure using Biotage SNAP Ultra columns. The columns were primed with CH₂Cl₂ containing 1% pyridine before use for all DMTr-containing compounds.1H and 13C spectra were measured on a Bruker AVII 500 spectrometer at 500 and 126 MHz, respectively. Chemical shifts are given in parts per million and were internally referenced to the appropriate residual solvent signal, all coupling constants (J) are quoted in hertz (Hz). Assignment of compounds was aided by correlated spectroscopy, heteronuclear single quantum coherence, heteronuclear multiple bond correlation, and distortionless enhancement by polarization transfer-135 experiments. The high-resolution mass spectra were measured on a Bruker 9.4 FT-ICR-MS mass spectrometer, and the samples were run in MeOH.

**N**₆-Benzoyl-5′-O-(4,4′-dimethoxytrityl)-3′-O-propargyl-5-methyl-LNA Cytidine (3). To a solution of nucleoside 1 (0.74 g, 1.21 mmol) in DMF (5 mL) was added benzoic anhydride (0.41 g, 1.81 mmol). The reaction mixture was stirred at room temperature for 18 h. The solvent was removed.
and the residue was taken up in EtOAc (100 mL), washed with sat. aqueous NaHCO₃ (50 mL), brine (2 × 50 mL), dried (Na₂SO₄), and concentrated. The crude mixture was purified using column chromatography (0–50% EtOAc in hexane) to obtain 3 (0.75 g, 86%) as a white foam. Rf (0.4, 40% EtOAc in hexane). ESI HRMS m/z 312.2663 ([M – H]⁻, C₁₁H₈O₄N₄⁻) calc. 312.2664. ¹H NMR (500 MHz, DMSO) δ 13.12 (bs, 1H, NH), 8.18 (m, 2H, Ph), 7.85 (s, 1H, H6), 7.61–7.58 (m, 1H, Ph), 7.51–7.47 (m, 4H, DMTr, Ph), 7.37–7.33 (m, 6H, DMTr), 7.28–7.25 (m, 1H, DMTr), 6.95–6.93 (m, 4H, DMTr), 5.59 (s, 1H, H1′), 4.68 (s, 1H, H3′), 4.39 (s, 1H, H3′), 4.35 (dd, J = 16.0, 2.4 Hz, 1H, CH₃C(=CH)CH), 4.29 (dd, J = 16.0, 2.4 Hz, 1H, CHH₂C(=CH)CH), 3.75–3.71 (m, 7H, 2 × OCH₂H₂S₃), 3.74–3.72 (dd, J = 8.0 Hz, 1H, H5′s), 3.55 (t, J = 2.4 Hz, 1H, C(=CH)CH), 3.44 (d, J = 11.1 Hz, 1H, H5′), 3.38 (dd, J = 11.1 Hz, 1H, H5′), 1.81 (s, 3H, CH₃). ¹³C NMR (126 MHz, DMSO) δ 158.2, 158.2 (C4/DMTr), 144.5 (C2), 136.7 (C6), 135.2, 134.9, 132.5, 129.8, 129.3, 128.3, 128.0, 127.7, 126.9, 113.4 (DMTrPh), 109.2 (C5), 87.1 (C4), 86.9 (C1′), 85.9 (DMTr), 79.8 (C=CCH), 78.0 (C=CCH), 75.9 (C2′), 75.5 (C3′), 71.7 (C5′), 57.9 (CH₂), 57.1 (C5′), 55.1 (2 × OCH₂H₂S₃), 13.5 (CH₃). A small impurity of ethyl acetate and CH₂Cl₂ as residual solvent was found. The compound was used as such in the next step.

N₅-Benzoyl-5′-O-(4,4′-dimethoxytrityl)-3′-O-propargyl-5-methyl-2′-deoxyctydine (4). To a solution of nucleoside 2 (0.12 g, 2.07 mmol) in DMF (5 mL) was added benzoic anhydride (0.93 g, 4.11 mmol). The reaction mixture was stirred at room temperature for 20 h. The solvent was removed and residue was taken up in EtOAc (100 mL), washed with sat. aqueous NaHCO₃ (50 mL), brine (2 × 50 mL), dried (Na₂SO₄), and concentrated. The crude mixture was purified using column chromatography (0–50% EtOAc in hexane) to obtain 4 (1.30 g, 92%) as a white foam. Rf (0.5, 50% EtOAc in hexane). ESI HRMS m/z 686.2861 ([M + H]⁺, C₁₁H₈O₄N₄⁺) calc. 686.2861. ¹H NMR (400 MHz, DMSO) δ 12.95 (bs, 1H, NH), 8.17 (m, 2H, Ph), 7.80 (s, 1H, H6), 7.62–7.58 (m, 1H, Ph), 7.52–7.48 (m, 2H, Ph), 7.43–7.40 (m, 2H, DMTr), 7.35 (t, J = 7.6 Hz, 2H, DMTr), 7.30–7.24 (m, 5H, DMTr), 6.93 (d, J = 8.8 Hz, 4H, DMTr), 6.15 (t, J = 6.8 Hz, 1H, H1′), 4.25–4.50 (m, 1H, H3′), 4.25 (apparent t, J = 2.4 Hz, 2H, CH₂C(=CH)CH), 4.13–4.12 (m, 1H, H4′), 3.74 (s, 6H, 2 × OCH₃), 3.53 (t, J = 2.4 Hz, 1H, C(=CH)CH), 3.32–3.29 (m, 1H, H3′), 3.23 (dd, J = 10.4, 2.8 Hz, 1H, H5′), 2.46–2.39 (m, 2H, H2′), 1.81 (s, 3H, CH₃). ¹³C NMR (100 MHz, DMSO) δ 159.2 (C4), 158.7 (DMTr), 145.0 (C2), 138.5 (C6), 137.0, 135.8, 135.6, 133.0, 130.2, 129.8, 128.8, 128.1, 128.7, 113.8 (DMTrPh, C5), 110.7 (C5), 86.7 (DMTr), 85.6 (C1′), 83.4 (C4′), 80.4 (C=CCH), 78.2 (C=CCH), 78.0 (H₃′), 63.9 (C5′), 56.5 (CH₂), 55.5 (2 × OCH₂H₂S₃), 37.2 (C2′), 13.2 (CH₃).
Synthesis of LNA/DNA Triazole Nucleoside (10). Nucleosides 3 (0.50 g, 0.73 mmol) and 7 (170 mg, 0.66 mmol) were dissolved in THF/H2O/t-BuOH (10 mL, 3:1:1, v/v/v). To this solution was added pyridine (2 μL, 0.01 mmol) and CuSO4 (1.5 mL, 7.5% aqueous, w/v), and sodium ascorbate (1.7 mL, 1 M aqueous). The reaction mixture was degassed with argon and stirred at room temperature for 2 h. The reaction was diluted with EtOAc (100 mL) and washed with H2O (50 mL) and sat. aqueous solution of EDTA (3 × 50 mL). The combined aqueous phase was back extracted with EtOAc (50 mL) and the combined organic phase was dried (Na2SO4) and concentrated under reduced pressure. The residue was purified using column chromatography (0–6% MeOH in CH2Cl2) to obtain 10 (0.54 g, 87%) as a white foam. Rf (0.5, 7% MeOH in CH2Cl2). ESI HRMS m/z 979.3621 ([M – H]+), C61H70O13N10P+ calc. 1181.4856 [M + H]+. 1H NMR (500 MHz, DMSO) δ 13.16 (s, 1H, NH (MeC)), 11.31 (s, 1H, NH (T)), 8.23 (m, 2H, Ph), 8.03 (s, 1H, triazole-H), 7.85 (s, 1H, H6 (MeC)), 7.61 (m, 1H, Ph), 7.52 (m, 2H, Ph), 7.42–7.42 (m, 2H, DMTr), 7.34–7.23 (m, 8H, DMTr, H6 (T)), 6.93–6.90 (m, 4H, DMTr), 6.16 (t, J = 7.0 Hz, 1H, H1′ (T)), 5.60 (s, 1H, H1′ (MeC)), 5.50 (d, J = 4.3 Hz, 1H, 3′-OH (T)), 4.72 (d, J = 12.0 Hz, 1H, CH2), 4.64–4.64 (m, 3H, H2′ (MeC), CH2, H5′ (T)), 4.58 (dd, J = 14.3, 7.6 Hz, 1H, H5′ (T)), 4.41 (s, 1H, H3′ (MeC)), 4.29–4.25 (m, 1H, H3′ (T)), 4.07–4.04 (m, 1H, H4′ (T)), 3.75–3.71 (m, 8H, 2 × OCH2 × 2 × H5′ (MeC)), 3.41 (d, J = 11.2 Hz, 1H, H5′ (MeC)), 3.36–3.31 (m, 1H, H5′ (MeC), merged with H2O signal from DMSO), 2.21–2.15 (m, 1H, H2′ (T)), 2.12–2.07 (m, 1H, H2′ (T)), 1.86 (s, 3H, CH3 (MeC)), 1.76 (d, J = 1.2 Hz, 3H, CH3 (T)). 13C NMR (126 MHz, DMSO) δ 178.7, 167.4, 159.7, 158.6, 150.9, 145.8, 138.9, 137.3, 137.1, 136.5, 135.7, 135.6, 134.4, 130.6, 121.1, 126.9, 126.6, 125.6, 123.7, 119.7, 118.3, 110.3, 109.9, 87.6, 87.4, 86.3, 84.5, 84.4, 76.6, 76.5, 72.2, 71.2, 63.1, 58.4, 55.5, 51.7, 38.4, 14.0, 12.5.

Synthesis of LNA/LNA Triazole Nucleoside (11). Nucleosides 3 (120 mg, 0.40 mmol) and 6 (0.36 g, 0.50 mmol) were dissolved in THF/H2O/t-BuOH (5 mL, 3:1:1, v/v/v). To this solution was added pyridine (2–3 drops), CuSO4 (0.9 mL, 7.5% aqueous, w/v), and sodium ascorbate (1.0 mL, 1 M aqueous). The reaction mixture was degassed with argon and stirred at room temperature for 2 h. The reaction was diluted with EtOAc (50 mL) and washed with H2O (30 mL) and a sat. aqueous solution of EDTA (3 × 30 mL). The combined aqueous phase was back extracted with EtOAc (20 mL) and the combined organic phase was dried (Na2SO4) and concentrated under reduced pressure. The residue was purified using column chromatography (0–6% MeOH in CH2Cl2) to obtain 11 (0.32 g, 79%) as a white foam with a small amount of EtOAc as an impurity. Rf (0.4, 6% MeOH in CH2Cl2). ESI HRMS m/z 1009.3721 ([M + H]+), C62H72O13N10P+ calc. 1009.3726. 1H NMR (500 MHz, DMSO) δ 13.16 (bs, 1H, NH (MeC)), 11.33 (s, 1H, NH (T)), 8.19 (2H, Ph), 8.12 (s, 1H, triazole-H), 7.86 (s, 1H, H6 (MeC)), 7.60 (m, 1H, Ph), 7.50 (2H, Ph), 7.43–7.41 (m, 2H, DMTr), 7.34–7.22 (m, 7H, DMTr), 6.92–6.88 (4H, DMTr), 6.62 (s, 1H, H6 (T)), 6.05 (d, J = 4.2 Hz, 1H, 3′-OH (T)), 5.60 (s, 1H, H1′ (MeC)), 5.40 (s, 1H, H1′ (T)), 4.97 (d = 15.2 Hz, 1H, H5′ (T)), 4.82 (d, J = 15.2 Hz, 1H, H5′ (T)), 4.77 (d = 11.2 Hz, 1H, CH2), 4.70–4.66 (m, 2H, CH2, H3′ (MeC)), 4.43 (s, 1H, H2′ (MeC)), 4.16 (s, 1H, H2′ (T)), 3.98 (d = 8.0 Hz, 1H, H5′ (T)), 3.80 (d = 4.2 Hz, 1H, H3′ (T)), 3.75–3.72 (m, 8H, 2 × OCH2 × 2 × H5′ (MeC)), 3.50 (d = 8.0 Hz, 1H, H5′ (T)), 3.41–3.36 (m, 2H, 2 × H5′ (MeC)), 1.85 (s, 3H, CH3 (MeC)), 1.61 (s, 3H, CH3 (T)). 13C NMR (126 MHz, DMSO) δ 164.1, 158.7, 150.2, 145.1, 144.0, 143.7, 136.5, 134.2, 133.0, 130.8, 128.8, 128.5, 128.1, 127.3, 126.3, 113.8, 110.9, 87.1, 86.6, 83.9, 79.5, 79.0, 71.8, 70.4, 64.0, 62.4, 55.5, 46.5, 13.1, 12.5.

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0.58 mmol) were added and the reaction mixture was stirred at room temperature for 2 h. The reaction was diluted with CH₂Cl₂ (30 mL) and washed with sat. aqueous KCl (30 mL). The organic phase was dried (Na₂SO₄) and concentrated under reduced pressure. The residue was purified using column chromatography (0–3% MeOH in CH₂Cl₂) to obtain 15 (245 mg, 73%) as a white foam. Rₛ (0.5, 4% MeOH in CH₂Cl₂) ESI HRMS m/z 1209.4813 ([M + H])⁺, C₅₀H₆₀O₁₄N₁₉P₁ calc. 1209.4805. 3¹P NMR (162 MHz, CD₃CN) δ 149.27, 148.87.

**Synthesis of Resin Loaded with DNA/DNA Triazole Nucleoside (17).** Activated resin 16⁻²⁸ (500 mg) was soaked in anhydrous pyridine (1 mL) for 10 min. EDC (170 mg, 1.09 mmol), DMAP (36 mg, 0.29 mmol), and triethylamine (44 μL, 0.32 mmol), and dinucleoside 8 (56 mg, 59 μmol) were added to the resin. The reaction vessel was rotated for 20 h at room temperature, after which pentachlorophenol (26 mg, 98 μmol) was added and the vessel was rotated for an additional 3 h. The solvents were removed by filtration, and the support was washed with pyridine, CH₂Cl₂, and diethyl ether. Piperidine (10% in DMF, 2 mL) was added and the vessel was rotated for 5 min at room temperature. The solvent was removed by filtration and the support was washed with CH₂Cl₂ and diethyl ether. Capping reagent (oligonucleotide synthesis grade, acetic anhydride/pyridine/THF:N-methylimidazole in THF, 1:1, 2 mL) was added and the vessel was rotated at room temperature for 1 h. The solvent was removed by filtration, and the resin was washed with pyridine, CH₂Cl₂, and diethyl ether and dried under high vacuum overnight to obtain 17. The loading of nucleoside 8 on the support was determined by cleaving the DMTr group and was found to be 28 μmol/g.

**Synthesis of Resin Loaded with DNA/DNA Triazole Nucleoside (18).** Activated resin 16⁻²⁸ (800 mg) was soaked in anhydrous pyridine (1 mL) for 10 min. EDC (0.329 g, 2.12 mmol), DMAP (14 mg, 0.11 mmol), triethylamine (85 μL, 0.61 mmol), and compound 9 (100 mg, 0.102 mmol) were added to the resin. The reaction vessel was rotated for 20 h at room temperature, after which pentachlorophenol (49 mg, 0.18 mmol) was added and the vessel was rotated for an additional 3 h. The solvents were removed by filtration, and the support was washed with pyridine, CH₂Cl₂, and diethyl ether. Piperidine (10% in DMF, 2 mL) was added and the vessel was rotated for 5 min at room temperature. The solvent was removed by filtration and the support was washed with CH₂Cl₂ and diethyl ether. Capping reagent (oligonucleotide synthesis grade, acetic anhydride/pyridine/THF:N-methylimidazole in THF, 1:1, 2 mL) was added and the vessel was rotated at room temperature for 1 h. The solvent was removed by filtration, and the resin was washed with pyridine, CH₂Cl₂, and diethyl ether and dried under high vacuum overnight to obtain 18. The loading of nucleoside 9 on the support was determined by cleaving the DMTr group and was found to be 26 μmol/g.

**Synthesis of Resin Loaded with LNA/DNA Triazole Nucleoside (19).** Activated resin 16⁻²⁸ (300 mg) was soaked in 1 mL of anhydrous pyridine for 10 min. EDC (0.132 g, 0.850 mmol), DMAP (6 mg, 49 μmol), triethylamine (34 μL, 0.24 mmol), and compound 10 (40 mg, 41 μmol) were added to the resin. The reaction vessel was rotated for 20 h at room temperature, after which pentachlorophenol (20 mg, 75 μmol) was added and the vessel was rotated for an additional 3 h. The solvents were removed by filtration, and the support was washed with pyridine, CH₂Cl₂, and diethyl ether. Piperidine (10% in DMF, 2 mL) was added and the vessel was rotated for 5 min at room temperature. The solvent was removed by filtration and the support was washed with CH₂Cl₂ and diethyl ether. Capping reagent (oligonucleotide synthesis grade, acetic anhydride/pyridine/THF:N-methylimidazole in THF, 1:1, 2 mL) was added and the vessel was rotated at room temperature for 1 h. The solvent was removed by filtration and the resin was washed with pyridine, CH₂Cl₂, and diethyl ether and dried under high vacuum overnight. The loading of nucleoside 10 on the support was determined by cleaving the DMTr group and was found to be 29 μmol/g.

**Synthesis of Resin Loaded with LNA/LNA Triazole Nucleoside (20).** Activated resin 16⁻²⁸ (250 mg) was soaked in 1 mL of anhydrous pyridine for 10 min. EDC (0.106 g, 0.683 mmol), DMAP (5 mg, 41 μmol), triethylamine (28 μL, 0.20 mmol), and compound 11 (33 mg, 33 μmol) were added to the resin. The reaction vessel was rotated for 20 h at room temperature, after which pentachlorophenol (16 mg, 60 μmol) was added and the vessel was rotated for an additional 3 h. The solvents were removed by filtration, and the support was washed with pyridine, CH₂Cl₂, and diethyl ether. Capping reagent (oligonucleotide synthesis grade, acetic anhydride/pyridine/THF:N-methylimidazole in THF, 1:1, 2 mL) was added and the vessel was rotated at room temperature for 1 h. The solvent was removed by filtration and the resin was washed with pyridine, CH₂Cl₂, and diethyl ether and dried under high vacuum overnight. The loading of nucleoside 11 on the support was determined by cleaving the DMTr group and was found to be 28 μmol/g.

**Synthesis of Oligonucleotides.** Standard DNA phosphoramidites, solid supports, and reagents were purchased from Link Technologies and Applied Biosystems. LNA phosphoramidites were obtained from Exiqon. Automated solid-phase synthesis of oligonucleotides (trityl off) was performed on an Applied Biosystems 394 synthesizer. The synthesis was performed on 1.0 μm scale involving cycles of acid-catalyzed detritylation, coupling, capping, and iodine oxidation. Standard DNA phosphoramidites were coupled for 60 s, whereas extended coupling time of 10 min was used for the modified phosphoramidites including LNA phosphoramidites. Modified phosphoramidites 12, 13, 14, and 15 were used to prepare ONs with monomers W, X, Y, and Z respectively. Coupling efficiencies and overall synthesis yields were determined by the inbuilt automated trityl cation conductivity monitoring facility and were ≥98.0% in all the cases. The oligonucleotides were then cleaved from the solid support and protecting groups from the nucleobase and backbone were removed by exposure to concentrated aqueous ammonium hydroxide for 60 min at room temperature followed by heating in a sealed tube for 5 h at 55 °C. RNA was purchased from Integrated DNA Technologies BVBA, Leuven, Belgium.

**Purification of Oligonucleotides.** The fully deprotected oligonucleotides were then purified by reverse-phase high-performance liquid chromatography (HPLC) on a Gilson system using a Luna 10 μm C8(2) 100 Å pore Phenomenex column (250 × 10 mm²) with a gradient of acetonitrile in triethylammonium bicarbonate (TEAB) over 20 min at a flow rate of 4 mL/min. Buffer A: 0.1 M TEAB, pH 7.5; buffer B: 0.1 M TEAB, pH 7.5, with 50% acetonitrile were used. Elution was monitored by UV absorption between 260 and 295 nm. A gradient of acetonitrile in hexylammonium acetate (HAA) over 20 min at a flow rate of 4 mL/min was used to purify
oligonucleotides, which were not separable using a gradient of acetonitrile in TEAB. Buffer A: 0.1 M HAA, pH 7.5; buffer B: 0.1 M TEAB, pH 7.5, with 50% acetonitrile were used. The elution was monitored by UV absorption between 260 and 298 nm.

Thermal Denaturation Experiments. UV DNA melting curves were recorded in a Cary 4000 Scan UV−vis spectrophotometer using 3 μM of each oligonucleotide in a 10 mM phosphate buffer containing 200 mM NaCl at pH 7.0. The samples were annealed by heating to 85 °C (10 °C/min) and then slowly cooling to 20 °C (1 °C/min). Six successive cycles (heating and cooling) were performed at a gradient of 1 °C/min and the change in UV absorbance at 260 nm was recorded. The melting temperature was calculated from the first derivative of the melting curve using an in-built software.

Snake Venom Phosphodiesterase Stability. Five nanomole of oligonucleotide was dissolved in 100 μL buffer (50 mM Tris–HCl, 10 mM MgCl₂, pH = 9.0). Twenty microliter of this solution was removed as a control (0 min). To the remaining solution was added 2 μL aqueous solution of phosphodiesterase 1 from C. adamanteus venom (from Sigma-Aldrich, catalogue number P3243, 0.45 units, dissolved in 7 mL H₂O). The reaction was incubated at 37 °C and aliquots (20 μL) were taken at different time intervals, mixed with formamide (20 μL), and stored at −20 °C. The samples were then analyzed by denaturing 20% polyacrylamide gel electrophoresis.

Serum Stability. Five nanomole of oligonucleotide was dissolved in 50 μL Dulbecco’s PBS and 50 μL of FBS (Gibco, standard (sterile-filtered)) was added. The sample was thoroughly mixed by vortexing before 20 μL of this solution was removed, mixed with formamide (20 μL), and stored at −80 °C as a control (0 h). The remaining reaction mixtures were incubated at 37 °C and aliquots (20 μL) were taken at different time intervals, mixed with formamide (20 μL), and stored at −80 °C. The samples were then analyzed by denaturing 20% polyacrylamide gel.

CD Spectroscopy. CD spectra (200−340 nm) were recorded on a Chirascan Plus spectropolarimeter using a quartz optical cells with a path length of 5.0 mm. The scans were performed at 10 °C using a step size of 0.5 nm, a time per point of 1.0 s, and a bandwidth of 2 nm; the average of four scans is recorded. The melting temperature was calculated from the first derivative of the melting curve using an in-built software.

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