A New 1,5-Disubstituted Triazole DNA Backbone Mimic with Enhanced Polymerase Compatibility

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Abstract: Triazole linkages (TLs) are mimics of the phosphodiester bond in oligonucleotides with applications in synthetic biology and biotechnology. Here we report the RuAAC-catalyzed synthesis of a novel 1,5-disubstituted triazole (TL2) dinucleoside phosphoramidite as well as its incorporation into oligonucleotides and compare its DNA polymerase replication competency with other TL analogues. We demonstrate that TL2 has superior replication kinetics to these analogues and is accurately replicated by polymerases. Derived structure–biocompatibility relationships show that linker length and the orientation of a hydrogen bond acceptor are critical and provide further guidance for the rational design of artificial biocompatible nucleic acid backbones.

Introduction

The replacement of natural phosphodiester (PO) bonds in DNA or RNA by artificial internucleoside linkages can generate remarkable biomimetic oligonucleotides (ONs) with applications in synthetic biology and biotechnology. For example, the CuAAC click ligation of alkynyl–azido ONs with 5′-alkyne adaptors forms TL1 efficiently by several polymerases through TL1–dependent slowdown in PCR replications. Many applications of such triazole linkages (TLs) are mimics of the phosphodiester bond in oligonucleotides with applications in synthetic biology and biotechnology. Here we report the RuAAC-catalyzed synthesis of a novel 1,5-disubstituted triazole (TL2) dinucleoside phosphoramidite as well as its incorporation into oligonucleotides and compare its DNA polymerase replication competency with other TL analogues. We demonstrate that TL2 has superior replication kinetics to these analogues and is accurately replicated by polymerases. Derived structure–biocompatibility relationships show that linker length and the orientation of a hydrogen bond acceptor are critical and provide further guidance for the rational design of artificial biocompatible nucleic acid backbones.

Further modification can be achieved by alkylation of the triazole as exemplified by cationic TL7, which is made by methylation of TL6. Many applications of such triazole backbones exist. For example, the CuAAC click ligation of 3′-azido ONs with 5′-alkyne adaptors forms TL1 which was recently described for the preparation of next-generation sequencing libraries. However, the authors reported low replication efficiencies by several polymerases through TL1. Furthermore, click ligation has been used to assemble long DNA templates with isolated TL6 modifications which can be replicated or transcribed by polymerases in bacterial or mammalian cells while retaining high fidelity read-through. However, TL6 suffers from reduced binding affinity to a complementary DNA or RNA target and induces a TL6-dependent slowdown in PCR replications. In search of the ideal nucleic acid triazole linkage, we recently developed a 1,5-disubstituted 1,2,3-triazole internucleoside linkage which was prepared by RuAAC-catalyzed azide–alkyne cycloaddition (CuAAC) (TL3; Figure 1). Altogether, a diverse toolbox of TLs has been reported over the past two decades, all having distinct triazole orientations and linker lengths as indicated in Figure 1. However, whether the more recent triazole isoforms, such as the 1,5-disubstituted TL3,
silyl-protected thymidine further biomimetic nucleic acid linkage designs. 

Evaluation of the molecular characteristics of the TLs with respect to conferred biocompatibility provides guidance for enzymatic read-through studies (Figure 2). For the 1,5-disubstituted 1,2,3-triazole (TL 2), TBAF-mediated deprotection of 4 gave alcohol 5, which was phosphorylated to dimer phosphoramidite 6 for ON synthesis.

In parallel, CuAAC of alkyne 2 and azide 3 resulted in the formation of 1,4-disubstituted dimer 7 (Scheme 1B). Desilylation gave alcohol 8, and phosphorylation using standard conditions yielded phosphoramidite 9. The 1,4-substitution of dimer 7—as opposed to the 1,5-substitution of dimer 4—was confirmed by 2D NMR characterization (Figure S3).

**Hybridization Studies. Duplex Stability.** Phosphoramidites 6 and 9 were coupled during standard ON synthesis to introduce TL2 and TL5 into ON2 and ON5, respectively (Table 1). The ON sequence and position of the TLs are identical to previously reported ONs with TL1, TL3, TL4, and TL6 (ON1, ON3, ON4, and ON6)38 to facilitate accurate comparisons among the TLs in UV melting experiments (Table 1). In comparison to unmodified ON7, introduction of TL2 results in destabilization of the duplex formed with DNA or RNA by −5.1 °C and −3.4 °C, respectively (Table 1). TL5 in ON5 destabilizes the duplexes with DNA and RNA by −8.6 °C, confirming previously reported destabilization by the TL5 modification in a different sequence context.32 Destabilizing effects have also been reported for TL1, TL3, TL4, and TL6 (Table 1).38 All 6-bond TLs (TLs 2–4) generally show more favorable duplex stability properties than their shorter 5-bond (TL1) or longer 7-bond (TL5 and TL6) counterparts. Rigid TL1 is the most destabilizing among the TLs. It is noteworthy that TL4 introduces minimal duplex destabilization (only −0.8 °C) when hybridized to an RNA target. To summarize, the relative duplex stabilities of TL-modified ONs with a DNA target are TL3 > TL4 > TL2 > TL6 > TL5 > TL1, and with an RNA target, TL4 > TL3 ≥ TL2 > TL6 > TL5 > TL1.

**Duplex Structure.** Despite some destabilizing effects, TL1, TL3, TL4, and TL6 are known to retain global B- or A-form duplex structures with the DNA or RNA target, respectively. In the current study, duplexes formed by TL2- and TL5-modified ON2 and ON5 with complementary DNA or RNA were analyzed by circular dichroism (CD) spectroscopy (Figure 2). The duplexes of ON2 with DNA and RNA show almost identical spectra to the equivalent duplexes formed by unmodified ON7. Duplexes of ON5 with DNA or RNA also retain global B- or A-form helical structures, respectively.

**Enzymatic Read-through Studies.** The ability of TL-modified ONs (TLs 1–6) to form duplexes with DNA and RNA targets while maintaining global structural integrity suggests that these TLs could be promising biocompatible DNA backbone mimics. Indeed, TL1, TL6, and methylated TL7 have been extensively studied as modifications in templates for replication by DNA and RNA polymerases.9,13,16,19,28,42 Moreover, a previous study reported encouraging results for TL4 within modified primers in qPCR.28 However, no read-through compatibilities of 1,4-disubstituted TL5 or 1,5-disubstituted TLs (TL2 and TL3) have been reported. Thus, we compared TLs 1–7 in primer extension studies (Figure 3 and Figure 4). We synthesized unmodified templates and TL-modified templates, each containing a single TL (TLs 1–7), using phosphoramidite 6 (TL2), 9 (TL5), or previously reported phosphoramidites (TL1,19 TL3,38 TL4,27 TL6,43 and TL7).53 We then assessed the polymerase compatibilities of TLs 1–7 by enzymatic extension of fluorescein (FL)-labeled primers followed by analysis of the FL-labeled products after gel electrophoresis.

![Figure 1. Summary of TLs connecting the 5′- and the 3′-ribose rings through multiple bonds as indicated by the boxed numbers. Trigonal planar orbital geometries (sp2) along the linkages are highlighted with gray circles. The structure of the natural PO is shown for reference. The TL linkage (TL2) reported here is highlighted. Abbreviations: b, base.](https://doi.org/10.1021/jacs.1c08057)
The sequences of the templates and primers can be found in Table S1.

Klenow Fragment. Extension of an FL-labeled primer by the large fragment of DNA polymerase I (Klenow fragment) using unmodified Template 0 (PO) resulted in formation of the expected full-length product, which was confirmed by mass spectrometry (MS) analysis (FL-replicon 0**; Figure 3A,B). Primer extension through the TL1-modified Template 1 gave a band that migrated slightly faster than FL-replicon 0 and a mass corresponding to a replicon with an A-deletion (FL-replicon 1**; Figure 3A,B). The fact that read-through of TL1 using unmodified Template 0 (PO) resulted in formation of the expected full-length product, which was confirmed by mass spectrometry (MS) analysis (FL-replicon 0**; Figure 3A,B). Primer extension through the TL1-modified Template 1 gave a band that migrated slightly faster than FL-replicon 0 and a mass corresponding to a replicon with an A-deletion (FL-replicon 1**; Figure 3A,B). The fact that read-through of TL1

"Abbreviations: DIPEA, N,N-diisopropylethylamine; DMTr, 4,4′-dimethoxytrityl; T, thymine; TBAF, tetrabutylammonium fluoride; TBDPS, tert-butyldiphenylsilyl.

Table 1. Melting Temperatures (Tm) of TL-Modified ONs with a DNA† and RNA‡ Target

| ON   | Sequence (5′ → 3′) | DNA† target Tm (ΔTm)/°C | RNA‡ target Tm (ΔTm)/°C |
|------|-------------------|-------------------------|-------------------------|
| ON1  | CGACGTTLGCAGC     | (-11.2)58 (-13.6)58     |                         |
| ON2  | CGACGTTLGCAGC     | 57.0 (-5.1) 55.5 (-3.4)  |                         |
| ON3  | CGACGTTLGCAGC     | (-2.9)56 (-3.3)56       |                         |
| ON4  | CGACGTTLGCAGC     | (-3.2)56 (-0.8)56       |                         |
| ON5  | CGACGTTLGCAGC     | 53.5 (-8.6) 50.3 (-8.6)  |                         |
| ON6  | CGACGTTLGCAGC     | (-7.4)56 (-3.3)56       |                         |
| ON7  | CGACGTTLGCAGC     | 62.1 58.9               |                         |

"Values were obtained from the maxima dA260/dT vs T for 3 μM of each ON in 10 mM phosphate buffer, 200 mM NaCl, pH 7.0. The graphs of dA260/dT vs T are shown in Figure S4. ΔTm are relative to the unmodified control ON7. "Sequence of DNA target = 5′-GCTGCAAACGTCG-3′. "Sequence of RNA target = 5′-GCUGCAACGUCG-3′. "Indicates the site where a PO is replaced by TLs 1–6.

Figure 2. CD spectra of ON2, ON5 and ON7 in duplex with (A) DNA and (B) RNA containing 3 μM of ON and 3 μM of the target in 10 mM phosphate buffer, 200 mM NaCl, pH 7.0. Data points were taken as an average of four scans at rt. DNA target = 5′-GCTGCAAACGTCG-3′, RNA target = 5′-GCUGCAACGUCG-3′.
by Klenow fragment results in an A-deletion mutation which is situated next to the TL1 was previously shown by sequencing of the replicons. Templates bearing 6-bond TLs resulted in the formation of the expected full-length products as confirmed by MS (FL-replicons 2**, 3**, and 4**; Figure 3A,B). However, replication of the TL3 template also produced a truncated product, shown by MS analysis to be a consequence of premature extension termination directly before the triazole (FL-replicon 3*; Figure 3A,B). The longer 7-bond TL5 in Template 5 was replicated but resulted in a product with an A-deletion (FL-replicon 5**; Figure 3A,B), and additional faint truncation bands were visible on the gel. Extension through TL6 produced the expected full-length product (FL-replicon 6**; Figure 3A,B) and truncated products, one of which results from termination directly before TL6 (FL-replicon 6*; Figure 3A,B). The presence of
Replication of the TL or after the triazole (FL-replicon 5 resulted in truncated products from termination directly before primer extension experiment with Klenow fragment, replication around the artificial linkage with a Taq polymerase. 19

Hot-Start Taq Polymerase. It is known that replication by Taq polymerase is slowed down at high DNA concentrations such as those used in the primer extension assays. 44 Moreover, a previous study showed that the replication of an unmodified template can be significantly slowed down under these conditions (up to 120 min). 19 Thus, primer extensions along the TL-modified templates were performed with hot-start Taq polymerase at 60 °C for an extended time of 160 min (Figure 4). Replication of unmodified Template 0 (PO) resulted in the exclusive formation of the expected full-length product containing a Taq-characteristic 3′-A overhang as determined by MS (FL-replicon 0*; Figure 4A,B). Replication of the TL1-modified template resulted in minor amounts of a product for which MS suggested deletion of an A-nucleotide (FL-replicon 1**; Figure 4A,B). The major products from TL1 were truncated ONs where replication stopped directly before or after the triazole (FL-replicons 1*; Figure 4A,B). This supports the mutagenic effect of TL1 in causing deletions around the artificial linkage with a Taq polymerase. 19

Remarkably, replication through TL2 yielded a clean full-length product including the expected Taq-characteristic 3′-A overhang (FL-replicon 2*; Figure 4A,B). No truncated products were detected on the gel or by MS. In contrast, a significant amount of truncated product was observed for the TL3 template where extension terminated directly before the triazole (FL-replicon 3*; Figure 4A,B). Extension through TL4 resulted in the expected full-length product without truncation (FL-replicon 4*; Figure 4A,B). In contrast to the previous primer extension experiment with Klenow fragment, replication through TL5 using hot-start Taq polymerase mainly resulted in truncated products from termination directly before or after the triazole (FL-replicons 5*; Figure 4A,B). Replication of the TL6 template produced the correct full-length construct (FL-replicon 6**; Figure 4A,B) and a truncated product (FL-replicon 6*; Figure 4A,B). As seen for Klenow fragment, methylated TL7 completely blocks extension (FL-replicon 7*; Figure 4A,B). Across the Klenow and hot-start Taq primer extension assays, TL2 and TL4 stand out as being the only TLs resulting in the clean formation of the expected full-length product without any detectable mutations or truncated fragments observed on gel or by MS. The excellent read-through compatibility of TL2 by Klenow fragment and hot-start Taq polymerase was confirmed with an alternative sequence (Template 9; Figures S11, S12 and Table S4).

qPCR Kinetics. To assess the time dependencies of TLs 1–7, the modified templates were used in qPCR experiments with various extension times. Impaired read-through of a given template results in an increase of the threshold cycle (Ct) —the number of qPCR cycles after which a signal overcomes a set threshold. If impairment can be overcome with time, the resulting Ct value will decrease with longer extension times if within the range of the time dependency. Amplification of unmodified templates can be considered as being time independent under the conditions of the study, as all internucleoside linkages are natural POs which are formed rapidly. Ct values of qPCR reactions with extension times ranging from 15–240 s were determined for templates with TLs 1–7 using hot-start Taq polymerase (Figures S13–S18). The differences between the determined Ct values of the modified templates to the Ct values of the respective isosequential unmodified PO templates (ΔCt) are plotted for each extension time in Figure 5A. Except for TL2 and TL7, all TLs show time dependency for the read-through of the artificial linkage. The apparent time independency of TL7 has been previously explained by a thermodynamic barrier created by TL7 that can only be overcome when transitioning from extension (60 °C) to denaturation (95 °C) in each cycle. 19 Most notably, the shorter 5- and 6-bond TLs 1-4 are less time dependent than the longer 7-bond TL5 and TL6. Indeed, ΔCt’s of TL5 and TL6 continued to decrease up to 240 s, the highest extension time tested in this study. The ΔCt’s of TL1, TL3, and TL4 decrease with longer extension times, but the

Figure 5. qPCR kinetics and PCR product sequencing. (A) Differences of threshold cycles (ΔCt’s) of TL-modified templates to their respective isosequential unmodified control template from qPCRs with hot-start Taq polymerase as a function of extension time. Unmodified templates: Template 0 and Templates 10–12. TL-modified templates: Template 1 (TL1), Template 2 (TL2), Template 3 (TL3), Template 4 (TL4), Template 5 (TL5), Template 6 (TL6), and Template 7 (TL7). ΔCt = Ct(TL-modified) − Ct(unmodified) for each extension time. Data are presented as the average from triplicates ± standard error of the mean. For C’s see Figure S22. (B) Representative Sanger sequencing results for hot-start Taq PCR amplicons cloned into a vector. Amplicons result from PCR amplifications of Template 2 (TL2), Template 3 (TL3), Template 4 (TL4), and Template 5 (TL5) using hot-start Taq polymerase with an extension time of 30 s each cycle. Corresponding positions of the TLs in the templates are indicated as pentagons. For full template sequences see Table S1, Figure S23, and Figure S24.
fitted curves flatten toward an extension time of 240 s. In contrast, the ΔCₜ of TL2 showed no time dependency within the tested conditions. Together with the superior results from the primer extension experiments, this suggests that the observed time independency of TL2 does not originate from transitions between the cycles as seen for TL7. Remarkably, there is no significant kinetic barrier imposed by TL2 for read-through of hot-start Taq polymerase under the tested conditions.

**DNA Sequencing.** To verify that read-through of the TL-modified templates results in correct replication, PCR products from hot-start Taq polymerase-mediated amplification were sequenced. While TL1, TL6, and TL7 have been previously studied for replication fidelity, the question whether TLs 2–5 can be replicated correctly in PCR remains unanswered. Thus, PCR amplicons generated using hot-start Taq polymerase and templates containing TL2, TL3, TL4, or TL5 were cloned into a vector and transformed into *Escherichia coli*. Several colonies were randomly picked, and the recovered vectors were analyzed by Sanger sequencing (Figure 5B). The 6-bond triazole backbones (TLs 2–4) resulted in the correct read-through producing the expected sequence for all colonies picked (n = 4). In contrast, the longer 7-bond TL5 resulted in a deletion mutation around the artificial linkage for which the T̃TₗT̃ motif has been read as a single T in four out of five colonies picked.

**Structure–Biocompatibility Relationship.** Analysis of the molecular properties of TLs 1–7 provides insight into the structure–biocompatibility relationship among the TLs for replication (Table 2). Most noticeable are the differences in TL backbone lengths. For 5-bond TL1, whose number of bonds is closest to the natural PO, inefficient read-through by hot-start Taq was observed confirming previous reports. Moreover, TL1 is mutagenic and results in single point mutations and other areas. A profound understanding of the structure–biocompatibility relationship among different TLs is crucial for their successful application in biochemical and biological systems. Here we report the synthesis of novel 1,5-disubstituted TL2 by RuAAC and directly compare this to its 1,4-disubstituted equivalent TL5, which is formed by CuAAC. In duplex melting experiments, TL2-modified ON2 forms more stable duplexes with DNA and RNA targets than TL5-modified ON5. Moreover, the read-through kinetics and amplification of TL2 by Klenow fragment and Taq DNA polymerase exceed all other TLs tested. To the best of our knowledge, TL2 has the fastest read-through kinetics and the highest efficiency of full-length product formation by Taq polymerase among the TLs reported to date. Further studies will be necessary to determine if the excellent read-through compatibility of TL2 by Taq can be translated into a cellular environment and be accommodated by RNA polymerases as

| TL   | linker length 1−7 | No. sp³ centers | kinetic barrier (Taq) | Correct read-through (Taq) |
|------|------------------|-----------------|----------------------|-----------------------------|
| PO   | 5                | 0               | –                    | –                           |
| TL1  | 5                | 3               | medium/high          | no, deletion                |
| TL2  | 6                | 2               | not significant      | yes                         |
| TL3  | 6                | 2               | medium               | yes                         |
| TL4  | 6                | 3               | medium               | yes                         |
| TL5  | 7                | 3               | High                 | no, deletion                |
| TL6  | 7                | 3               | High                 | yes                         |
| TL7  | 7                | 3               | very high            | no, deletion                |

*Defined as the minimal number of bonds connecting the 5’- and the 3’-furanose rings. *Along the internucleoside linkage. *Defined as having the correct sequence after read-through has occurred.

In conclusion, triazole linkages are interesting phosphodiester mimics for applications in synthetic biology, biotechnology, and other areas. A profound understanding of the structure–biocompatibility relationship among different TLs is crucial for their successful application in biochemical and biological systems. Here we report the synthesis of novel 1,5-disubstituted TL2 by RuAAC and directly compare this to its 1,4-disubstituted equivalent TL5, which is formed by CuAAC. In duplex melting experiments, TL2-modified ON2 forms more stable duplexes with DNA and RNA targets than TL5-modified ON5. Moreover, the read-through kinetics and amplification of TL2 by Klenow fragment and Taq DNA polymerase exceed all other TLs tested. To the best of our knowledge, TL2 has the fastest read-through kinetics and the highest efficiency of full-length product formation by Taq polymerase among the TLs reported to date. Further studies will be necessary to determine if the excellent read-through compatibility of TL2 by Taq can be translated into a cellular environment and be accommodated by RNA polymerases as

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previously demonstrated for TL5.\(^8,9,29\) Moreover, direct comparison between TL2 and TL5 revealed a remarkable difference in read-through accuracy. While TL2 was read correctly by hot-start Taq polymerase, TL5 resulted in a deletion mutation around the triazole linkage. We have discussed the structure–biocompatibility relationship of several TLs, and we provide a rationale to assist future designs of artificial backbone mimics in replication templates. Importantly, the structure–biocompatibility relationship presented in this study is derived from ONs with TLs between two thymidines, providing a rather challenging sequence context for polymerase read-through. Efficiencies can vary with the nucleobase sequence and generally improve with greater duplex stability (a higher GC-content) around the triazole. Indeed, enhanced read-through compatibility was previously observed for a C\(^{\text{TLC}}\)C motif.\(^8,19,29\) Hence, we anticipate that further investigations using different sequences will provide additional validation of the proposed model. The application of RuAAC to form TL2 exemplifies a powerful strategy to increase TL flexibility through formation of 1,5-disubstituted 1,2,3-triazoles. In contrast to the widely adopted CuAAC reaction, the feasibility of the RuAAC reaction to efficiently ligate azide- and alkyne-modified ONs in high yield, and without inducing DNA damage, remains elusive. However, in light of recent advances to obtain 1,5-disubstituted 1,2,3-triazoles via water and air compatible Ni-catalyzed click chemistry,\(^40\) TL2 represents a promising backbone for use in a plethora of applications across the life sciences.

**ASSOCIATED CONTENT**

1. Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.1c08057.

Experimental section, characterization data and NMRs of new compounds, additional primer extension experiments, \(T_m\) curves, oligonucleotide HPLC traces, oligonucleotide mass spectra, qPCR and sequencing data (PDF)

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**Notes**

The authors declare no competing financial interest.

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**ABBREVIATIONS**

Cas9, CRISPR associated protein 9; CRISPR, clustered regularly interspaced short palindromic repeats; DIPEA, N,N-diisopropylamine; DMTr, 4,4-dimethoxy trityl; TL, triazole linkage; qPCR, quantitative polymerase chain reaction

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