

Zebularine-DNA:

First nM

APOBEC3A

inhibitor

Design, Synthesis, and Evaluation of a Cross-Linked Oligonucleotide as the First Nanomolar Inhibitor of APOBEC3A

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ABSTRACT: Dru	ug resistance is a major proble	em assoc	iated with anticancer chemo- and	3′	N=N 2 ^{5'} Cross-linked

immunotherapies. Recent advances in the understanding of resistance mechanisms have revealed that enzymes of the APOBEC3 (A3) family contribute to the development of drug resistance in multiple cancers. A3 enzymes are polynucleotide cytidine deaminases that convert cytosine to uracil $(C \rightarrow U)$ in single-stranded DNA (ssDNA) and in this way protect humans against viruses and mobile retroelements. On the other hand, cancer cells use A3s, especially A3A and A3B, to mutate human DNA, and thus by increasing rates of evolution, cancer cells escape adaptive immune responses and resist drugs. However, as

A3A and A3B are non-essential for primary metabolism, their inhibition opens up a strategy to augment existing anticancer therapies and suppress cancer evolution. To test our hypothesis that pre-shaped ssDNA mimicking the *U*-shape observed in ssDNA–A3 complexes can provide a better binder to A3 enzymes, a Cu(I)-catalyzed azide–alkyne cycloaddition was used to cross-link two distant modified nucleobases in ssDNA. The resultant cytosine-containing substrate, where the cytosine sits at the apex of the loop, was deaminated faster by the engineered C-terminal domain of A3B than a standard, linear substrate. The cross-linked ssDNA was converted into an A3 inhibitor by replacing the 2'-deoxycytidine in the preferred TCA substrate motif by 2'-deoxyzebularine, a known inhibitor of single nucleoside cytidine deaminases. This strategy yielded the first nanomolar inhibitor of engineered A3B_{CTD} and wild-type A3A ($K_i = 690 \pm 140$ and 360 ± 120 nM, respectively), providing a platform for further development of powerful A3 inhibitors.

INTRODUCTION

Nucleic acid editing by members of the APOBEC3 (apolipoprotein B mRNA editing enzyme catalytic polypeptide-like 3; A3) family is an important part of the innate immune system involved in combating pathogens but often having detrimental consequences in cancers.¹⁻⁴ Unlike singlenucleoside cytidine deaminases (CDA), which act on individual cytidines, A3 enzymes deaminate cytosine to uracil (C→U) (Figure 1A) only on single-stranded DNA (ssDNA), although some activity has been reported on RNA.^{5,6}

There are seven human A3 enzymes (A3A-A3H, excluding A3E). Although A3 enzymes are a part of the human immune system, they can also mutate human genomic DNA and thus contribute to cancer genesis, cancer mutagenesis, and drug resistance.^{1-4,8-12} Single-domain A3A and double-domain A3B have been identified as the major contributors to cancer mutagenesis; their mutational signatures have been found in different cancers including bladder, breast, cervix, head/neck, and lung cancers.¹³⁻¹⁵ A3B is also associated with poor survival prognosis in some cancers.^{4,8,16} Since A3A and A3B are not essential to primary metabolism and A3B deletion is prevalent in some populations,¹⁷ their inhibition (especially of nucleus-targeted A3B) offers a potent strategy to suppress cancer evolution and thereby make existing anticancer therapies more efficient.^{11,18} As A3 enzymes deaminate

predominantly ssDNA, we have been exploring the possibilities of ssDNA-based inhibitors that can be used as conjuvants to existing cancer therapies, i.e. be used in combination with existing therapies to extend their efficacy.^{19–21}

+1

The enzymes A3 and CDA share a structurally homologous Zn^{2+} -containing active site; consequently, they are proposed to have a similar mechanism of cytosine deamination. We have recently developed the first rationally designed competitive inhibitor of A3 by incorporating a cytidine analogue 2'-deoxyzebularine, dZ, a known inhibitor of CDA (Figure 1B), into ssDNA fragments.¹⁹ We have also demonstrated that dZ as a single nucleoside does not inhibit A3, but once embedded into ssDNA, it becomes a micromolar competitive inhibitor of A3 enzymes.¹⁹ By analogy to the binding of dZ to CDA, this key observation supports a mechanism in which ssDNA delivers dZ to the active site of A3 for inhibition. In addition, we and others^{7,22,23} have shown that ssDNA bound to both A3A and to an engineered mutant of the catalytically active C-

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Figure 1. (A) Deamination of dC to dU in ssDNA by A3 enzymes. C^0 represents target cytosine; nucleobases (thymine T and adenine A) at 5' relative to C^0 are represented as T^{-1} , T^{-2} , and so forth and those at 3' relative to C^0 as A^{+1} , T^{+2} , and so forth. (B) Inhibitors of cytidine deamination used in this work, 2'-deoxyzebularine (**dZ**) and 5-fluoro-2'-deoxyzebularine (**FdZ**), incorporated into ssDNA (left) and their proposed hydrates in the active site of A3 (right), which mimic an intermediate of C-to-U conversion. (C) Crystal structure of an engineered C-terminal domain of A3B, A3B_{CTD} (light brown ribbon), bound to ssDNA, showing C^0 flipped out (yellow) to project into the Zn²⁺-containing active site (gray ball), +1 2'-deoxyadenosine (red), and -2 thymidine (blue) are close together in the U-shaped conformation in ssDNA (PDB: STD5).⁷



Figure 2. Terminal (A) and internal (B) cross-linking strategies by CuAAC to pre-shape a linear ssDNA as a substrate (dC-containing oligo) and as an inhibitor of $A3B_{CTD}$ (dZ-containing oligo). (C) Alkyne (Y) and azide (d R^{N3}) modifications used in the sequence for terminal cross-linking strategy. (D) Alkyne (d U^E) and azide (d A^{N3}) modifications used for internal cross-linking strategy.

terminal domain of A3B, $A3B_{CTD}$ (PDB: 5TD5 and 5SWW), is not linear but adopts a distinctive *U*-shape, projecting the target dC (or **dZ**) into the active site (Figure 1C). We hypothesized that locking DNA sequences into the observed *U*-shape may provide not only better substrates but also, appropriately modified, better inhibitors of A3 enzymes.

METHODOLOGY

To test our hypothesis, we evaluated two different approaches for pre-shaping DNA using covalent cross-links between distant nucleotides in the sequence, that is, terminal and internal cross-linking strategies (Figure 2A,B). For pre-shaping ssDNA (both strategies), we used copper(I)-catalyzed azide– alkyne cycloaddition (CuAAC), which is compatible with DNA synthesis and does not change the chemical structure of native nucleotides.^{24–27} For terminal cross-linking strategy, terminal alkyne and azide moieties were incorporated at 5' and 3'-ends of ssDNA, respectively. Commercially available alkyne Y^{28} (Figure 2A) was used and in-house synthesized azido-sugar dR^{N3} was attached to the solid support (controlled-pore glass, CPG). ssDNAs of different lengths were synthesized by standard DNA phosphoramidite chemistry. The second strategy used detailed structural information from the A3ssDNA complexes. Relative to the central 5'-TCA motif of the ssDNA, the positions of the alkyne and azide for the internal cross-linking strategy were selected based on the observation that 2'-deoxyadenosine at the +1 position immediately next to the target dC (position 0) and thymidine at the -2 position are close together and form hydrogen bond interactions with each other through a water molecule for the bound ssDNA in $A3B_{CTD}$ and A3A crystal structures. This makes dA in the +1 position and dT in the -2 position suitable for cross-linking (Figure 1C).^{7,29-31} We employed an in-house synthesized azide-containing 2'-deoxyadenosine (dA^{N3}) phosphoramidite to place it in position +1 and commercially available 5-ethynyl-2'-deoxyuridine (dU^E , Figure 2D) phosphoramidite³² to mimic dT in position -2 for the internal cross-linking strategy.

After the oligonucleotide synthesis, for terminal and internal cross-links, we employed CuAAC as a post-synthetic step, monitored by analytical reverse-phase high-performance liquid chromatography (HPLC) and mass spectrometry. The crosslinked oligonucleotides (oligos) were then evaluated as A3 substrates and compared to a standard linear substrate by a nuclear magnetic resonance (NMR)-based assay. Real-time NMR assays are advantageous because they are direct, utilizing only A3 enzymes and oligos in a suitable buffer system, unlike many fluorescence-based assays where a secondary enzyme and a fluorescently modified oligo are used.³³ The NMR-based assay yields directly the initial velocity of deamination of various ssDNA substrates, including the modified ones, in the presence of A3 enzymes. Consequently, the Michaelis-Menten kinetic model is used to characterize substrates and inhibitors of A3. Therefore, currently, the NMR-based direct assay is best suited to study modified ssDNA-based substrates and inhibitors of A3.

The engineered A3 enzyme used in our experiments is the well-characterized and very active $A3B_{CTD}$ -QM- $\Delta L3$ -AL1swap,¹⁹ where loop 3 is deleted and loop 1 is replaced with the corresponding loop 1 from A3A. This is the same enzyme used in the crystal structure of the A3-ssDNA complex (except that for the X-ray structure, the active-site Glu255 was mutated to Ala to prevent deamination of the substrate, dC-containing ssDNA).⁷ Our recent work on A3G has shown that inhibition of the C-terminal catalytic domain is mirrored by the inhibition of full-length enzyme, validating our approach to use catalytic C-terminal domains for inhibitor development. ssDNA substrates were converted into A3 inhibitors by incorporating dZ in place of the target dC. The evaluation and optimization strategy is summarized in Figure 3. The inhibitory potential of dZ-containing oligonucleotides was evaluated by the NMR-based assay in the presence of a linear substrate. We also characterized by isothermal titration calorimetry the thermodynamics of binding of our cross-linked oligos with A3 enzymes in comparison with linear ssDNA. The inactive E72A mutant of A3A (the same as in A3A-ssDNA crystal structure, PDB: 5SWW) was chosen to evaluate substrate binding, and the active $A3B_{CTD}\mbox{-}QM\mbox{-}\Delta L3\mbox{-}AL1\mbox{swap}$ was chosen for inhibitor binding. Finally, we demonstrated that



Figure 3. Methodology for the design and evaluation of cross-linked substrates and inhibitors.

the cross-linked DNA inhibitor suppressed the activity of wild-type A3A.

Our strategy has led to the design, synthesis, and evaluation of the first nanomolar inhibitor of A3 enzymes.

RESULTS

In the past 15 years, alkyne- and azide-modified oligos have been widely used for the synthesis of a vast variety of bioconjugates via azide-alkyne "click chemistry". Usually, terminal alkynes can be introduced into oligos using phosphoramidite chemistry, and several reagents are commercially available, such as the Y and dU^E modifications used in the current work. However, there are limited options for the introduction of the azide functionality into oligos, especially in the middle of the DNA sequence. Usually, isolation of azidocontaining phosphoramidite and its use in the DNA synthesis is problematic^{34,35} due to plausible Staudinger reaction between organic azide and a trivalent phosphorus in the phosphoramidite function. Indeed, the Staudinger reaction has been used recently for modification of a phosphate backbone.^{36–41} Interestingly, a Staudinger reaction does not occur on the azide group in the support-bound oligonucleotide chain during subsequent elongation using nucleoside phosphoramidites.⁴² This led to the development of azide-containing solid supports to produce oligos bearing organic azides at the 3'-end.⁴³⁻⁴⁶ In 2014, Fomich et al.⁴⁷ demonstrated that azidecarrying phosphoramidite can be obtained and used in DNA synthesis to produce oligos having up to three organic azides in the middle of the sequence. This is achievable if the azidecarrying phosphoramidite is not concentrated and is used directly in automated DNA synthesis after aqueous work-up and drying of the solution over sodium sulfate. These advancements in the field were used in the current work to produce oligos containing an organic azide at the 3'-end as well as in the middle of the sequence using azide-modified 2deoxyribose and dA described below, respectively.

Synthesis of DNA Precursors Containing Organic Azides. A method reported recently⁴⁸ described the synthesis of compound 2 from Hoffer's chlorosugar 1 (Scheme 1) using a Lewis acid and trimethylsilyl azide, but low β/α selectivity and moderate yields were reported. In contrast, we employed a phase-transfer protocol⁴⁹ with minor changes using NaN₃ and Bu₄NHSO₄ under vigorous stirring in saturated aqueous NaHCO₃ and chloroform followed by addition of Hoffer's chlorosugar 1. The reaction was finished in 20 min, and after

Scheme 1. Synthesis of Azido Sugar dR^{N3} Modified CPG-Support^a



^aReagents and conditions: (i) NaN₃, Bu₄NHSO₄, satd. NaHCO₃/CHCl₃ (1:1), rt, 20 min; (ii) 30% aq. NH₃, MeOH, rt, 3 days; (iii) 4,4'dimethoxytrityl chloride (DMTCl), pyridine, 0°C to rt, overnight; (iv) long-chain aminoalkyl CPG (LCAA-CPG-CO₂H), N-(3dimethylaminopropyl)-N'-ethylcarbodiimide (EDC), Et₃N, 4-(dimethylamino)pyridine (DMAP), pyridine/DMF (1:1), rt, 3 days.

Scheme 2. Synthesis of Modified 2'-Deoxyadenosine Phosphoramidite Containing a Two-Carbon Linker with Azide^a



"Reagents and conditions: (i) acetic anhydride, pyridine, rt overnight then 60° C for 6 h; (ii) compound 5, Cs₂CO₃, CH₃CN, 60° C, overnight; (iii) Et₃N in MeOH/H₂O, rt, 15 min; (iv) 4,4'-dimethoxytrityl chloride, pyridine, 0°C to rt, overnight; (v) *N*,*N*-diisopropylamino-2-cyanoethoxychlorophosphine (CEP-Cl), Et₃N, dry DCM, 0°C.

work-up resulted in almost pure azide 2 with high yield (88%) and β/α ratio of 16:1 (6% of α -anomer). Subsequent recrystallization from EtOH slightly increased the β/α ratio to 19:1 (5% of α -anomer). This protocol was optimized to a multi-gram scale synthesis of azide 2 in our laboratory. The CPG conjugated to azido-sugar 4 was prepared from a 5'dimethoxytrityl (DMT) protected azido-sugar 3 by a slight modification of reported methods^{44,50} (Scheme 1). The load on CPG was found to be 39 μ mol/g by UV absorption of the DMT cation released upon treatment of a sample of the modified support 4 with a 3% solution of dichloroacetic acid in dichloromethane.

The second strategy relied on an internal cross-link created between azide-containing dA^{N3} and commercially available alkyne-modified dU^{E} (Figure 2B,D).

Here, we decided to introduce an azidoalkane linker into the N^6 position of dA, due to close proximity of an exocyclic amine of dA⁺¹ and 5-methyl group of thymidine⁻² in the crystal structure (Figure 1C). In the past, various methods have been employed for N^6 alkylation of dA, which usually relied upon alkylation in the N^1 position followed by subsequent Dimroth rearrangement to the N^6 -alkylated product.^{51,52} In contrast to previous reports, we observed a direct alkylation of the N^6 position in triacetylated dA (7) using 2-azidoethyl-4-tosylate ($\mathbf{5}$)⁵³ and Cs₂CO₃ as the base (Scheme 2). The alkylation at the N^6 position of dA providing compound $\mathbf{8}$ was confirmed by ¹H and ¹³C heteronuclear multiple-bond correlation (HMBC) NMR experiments, in which three-bond correlations were

detected. We observed that proton signals of the CH₂ group of the azidoalkyl linker at 4.31 ppm had a cross-peak not only with C^6 of adenine at 152.58 ppm but also with the carbonyl carbon of the acetyl group at 170.06 ppm (Figure S49, see Supporting Information). This is only possible to observe for N^6 and not for N^1 alkylated dA (three versus five bonds between hydrogen of CH₂ and carbon of CO, respectively). After the selective deprotection of 3'- and 5'-acetyl groups, the 5'-hydroxy group was converted into the 4,4'-dimethoxytrityl derivative 10 under standard conditions. Next, a pilot phosphitylation reaction was performed on compound 10 using previously described conditions⁴⁷ in dry CDCl₃, producing phosphoramidite 11 with a purity of more than 70% as shown by ¹H, ¹³C, and ³¹P NMR spectra (Figures S59-S61, Supporting Information). This suggests that compound 11 can be used for automated DNA synthesis, with a limitation that the freshly prepared phosphoramidite should be used immediately after synthesis because the storage of this solution at room temperature for 24 h results in significant degradation of compound 11 as evidenced by NMR spectroscopy (Figure S62, Supporting Information).

For automated DNA synthesis, the reaction was performed under the same conditions using dry dichloromethane as the solvent. To prevent a Staudinger reaction between the azide and phosphoramidite present in one molecule, we did not purify the product or concentrate it. Instead, it was used directly for the oligonucleotide synthesis. The concentration of reagent **11** in DCM was found to be 0.2 M, determined by UV

DNA sequence, $5' \rightarrow 3'$		name ^a	retention time (min)	ESI-MS [Da] found/calculated
YTTTCATTdR ^{N3}	linear	dC-9-mer	17.5	2552.47/2552.48
	cross-linked	dC-9-mer-X	14.1	2552.47/2552.48
YTTCATdR ^{N3}	linear	dC-7-mer	18.0	1944.37/1944.37
	cross-linked	dC-7-mer-X	13.6	1944.37/1944.37
YTCAdR ^{N3}	linear	dC-5-mer	17.8	1336.29/1336.30
	cross-linked	dC-5-mer-X	13.5	1336.29/1336.30
AdU ^e TTCdA ^{N3} TTT	linear	$dC[U^{E}(-3), A^{N3}(+1)]$	16.7	2757.46/2757.49
	cross-linked	$dC[U^{E}(-3), A^{N3}(+1)]X$	15.6	2757.47/2757.49
AT dU^eTCdA^{N3}TTT	linear	$dC[U^{E}(-2), A^{N3}(+1)]$	16.3	2757.46/2757.49
	cross-linked	$dC[U^{E}(-2), A^{N3}(+1)]X$	15.3	2757.46/2757.49
AdU ^e TTdZdA ^{N3} TTT	linear	$dZ[U^{E}(-3), A^{N3}(+1)]$	16.9	2742.45/2742.48
	cross-linked	$dZ[U^{E}(-3), A^{N3}(+1)]X$	15.7	2742.46/2742.48
AT dU^eTdZdA^{N3}TTT	linear	$dZ[U^{E}(-2), A^{N3}(+1)]$	16.9	2742.45/2742.48
	cross-linked	$dZ[U^{E}(-2), A^{N3}(+1)]X$	15.7	2742.45/2742.48

Table 1. List of Linear Oligos with Azide/Alkyne Modifications as well as Cross-Linked Oligos Having 1,4-Disubstituted 1,2,3-Triazole Modification

^{*a*}d stands for DNA backbone, C or Z denotes substrate (dC) or inhibitor (dZ) of A3 present in position 0 in the DNA sequence. The length of terminally cyclized oligos is described as being 9-, 7-, or 5-mer in relation to the length of the native DNA sequence having the same number of phosphate groups. In other words, modifications Y and dR^{N3} are counted as modified nucleotides. X stands for a cross-linked oligo containing 1,4-disubstituted 1,2,3-triazole, that is, dC-9-mer-X refers to a cross-linked oligo. For internally cross-linked oligos, the abbreviation of alkyne- and azido-containing nucleotides is presented in square brackets, and the position of each modification relative to the position 0 being dC or dZ is shown in brackets.



Figure 4. Initial speed of A3B_{CTD}-QM- Δ L3-AL1swap catalyzed deamination of terminally (A) and internally (B) cross-linked oligos in comparison to the linear substrate. Substrate concentration: 100 μ M in (a) and 800 μ M in (b). Enzyme concentration: 50 nM in (A) and 300 nM in (B). Buffer: 50 mM citrate-phosphate (pH 5.5), 200 mM NaCl, 2 mM β -mercaptoethanol, 200 μ M 4,4-dimethyl-4-silapentane-1-sulfonic acid, and 10% D₂O at 25°C. Error bars are estimated standard deviations from triplicate measurements.

absorption of the 4,4'-dimethoxytrityl cation released upon treatment of an aliquot of this sample with a 3% solution of dichloroacetic acid in dichloromethane.

Synthesis of Cross-Linked Oligonucleotides by CuAAC. Oligodeoxynucleotides (oligos) were synthesized by the phosphoramidite method on a MerMade-4 DNA/RNA synthesizer (BioAutomation) on a 5 μ mol scale using the standard manufacturer's protocol for native nucleotides, whereas increased coupling time from 2 to 10 min was used for modified alkyne and azide phosphoramidites. Detailed procedures are given in Supporting Information.

The purified linear oligos were cross-linked by CuAAC following the protocol implemented in our laboratory (see Supporting Information). Progress of the reaction was monitored by reverse-phase HPLC. Gratifyingly, a new peak appeared in the chromatogram for all cross-linking experiments over a period of 16 h, characterized by shorter retention times than those of the starting materials (Table 1). These results agree with previously published properties of circularized DNA, which were also prepared using CuAAC.⁴⁴ After separation by reverse-phase HPLC, the products were desalted

and analyzed by mass-spectrometry, showing, as expected, the same mass as for the starting materials, confirming the composition of monomeric cross-linked oligos (Table 1). Abbreviation of individual oligos is explained in the caption of Table 1.

Additionally, to prove that our cross-linking strategy works for internal modifications, we performed NMR experiments on linear and cross-linked oligos (see Supporting Information, Section S5.1). Differences in chemical shifts between the starting material and the product in ¹H NMR were observed for all methyl peaks of thymidines located between 1.6 and 2.0 ppm, suggesting the formation of a new product (cross-linked oligo) as the result of CuAAC reaction (Figure S13 in the Supporting Information). In addition, acetylenic proton at 2.44 ppm in the linear oligo that was assigned by HMBC NMR having a cross peak with the <u>CH</u>₂ (18.30 ppm in ¹³C NMR) immediately next to it (Figures S11 and S12 in the Supporting Information) was absent in the cross-linked oligo, which provided additional evidence of the successful cross-linking by CuAAC.

Evaluation of DNA Substrates by NMR-Based Activity **Assay.** Using the 9-mer oligonucleotide 5'-AT₃CAT₃ as the standard substrate, we compared the deamination of both terminally and internally cross-linked oligos using our previously described real-time NMR assay. 19,20 For the terminally cross-linked oligos, we observed decreased rates of deamination from 9-mer to 5-mer. One should note that the number of phosphates in the modified 9-mer sequence is the same as for the 9-mer DNA control (Figure 4A). This means that terminally cross-linked oligos are poor mimics of the DNA shape seen in the complex, although the flexibility associated with the longer oligos allows some deamination to occur. This agrees with earlier observations that longer linear oligos bind A3A better than shorter oligos.³⁰ In marked contrast, a faster deamination rate was observed for the internally cross-linked oligo $dC[U^{E}(-2), A^{N3}(+1)]X$ compared to the standard, linear substrate (Figure 4B). The linked -2 and +1 positions were spatially close in the X-ray structure of the inactive A3B_{CTD}-QM- Δ L3-AL1swap(E255A)/DNA complex. Moreover, an extremely slow deamination rate was detected for $dC[U^{E}(-3)]$, $A^{N3}(+1)$]X oligo, which we attribute to a shape and conformation that together hinder proper binding with the enzyme compared to the dC[U^E(-2), A^{N3}(+1)]X oligo. We did not create a cross-link between dA at position +1 and T at position -1 as the X-ray structure (pdb: 5TD5) shows a clear binding pocket for T in the -1 position, from which we infer that cross-linking in this position will disrupt the DNAprotein interaction.

Evaluation of Inhibitors by NMR Assay. The increased speed of deamination of the $dC[U^{E}(-2), A^{N3}(+1)]X$ oligo prompted us to investigate the inhibition potential of this oligo by changing dC to dZ. We used the NMR assay to compare the internally cross-linked dZ-containing oligos with a linear DNA inhibitor containing dZ (dZ-linear, 5'-AT₃dZAT₃) that was characterized earlier.¹⁹ The NMR assay was performed using unmodified oligo (5'-T₄CAT) as a substrate in the presence of A3B_{CTD}-QM- Δ L3-AL1swap and a known concentration of dZ-containing inhibitors (linear and cross-linked).

The results revealed that the internally cross-linked dZcontaining oligo derived from the faster deaminated substrate, $dZ[U^{E}(-2), A^{N3}(+1)]X$, inhibited the A3-catalyzed deamination by greater than an order of magnitude more potently than our best linear dZ oligo (Figure 5). On the other hand, $dZ[U^{E}(-3), A^{N3}(+1)]X$ was found to be a much weaker inhibitor, in line with the earlier observation that $dC[U^{E}(-3), A^{N3}(+1)]X$ was a very poor substrate. Interestingly, its inhibitory potential is comparable with that of the linear oligo, probably because of dZ interacting strongly with the active site of the enzyme.

For determination of inhibition constants (K_i) of inhibitors, we first evaluated kinetic parameters of A3B_{CTD}-QM- Δ L3-AL1swap on two DNA substrates (Table 2), which were used to detect the residual deamination rate in the presence of inhibitors (Table 3). Fitting the experimental data by nonlinear least-squares using a global fit provided values of kinetic parameters similar to those obtained by the linearized Lineweaver–Burk plot (Table 2). Then, the inhibition constant (K_i) for our cross-linked inhibitor was calculated by several methods assuming a competitive mode of inhibition¹⁹ (Table 3), based on the obtained apparent K_m and experiments in which the concentration of the individual inhibitor was varied, and the residual deamination initial rate was measured.



Figure 5. Inhibition of A3B_{CTD}-QM-ΔL3-AL1swap-catalyzed deamination of S'-T₄CAT by **dZ**-containing linear and cross-linked oligos. Experimental conditions are 400 μM of S'-T₄CAT, 8 μM of **dZ**-containing oligos, 300 nM of A3B_{CTD}-QM-ΔL3-AL1swap in a 50 mM sodium phosphate buffer (pH 6.0) containing 100 mM NaCl, 2.5 mM β-mercaptoethanol, 0.1 M EDTA, and 50 μM 3-(trimethylsilyl)-2,2,3,3-tetradeuteropropionic acid at 25°C. K_i value of linear **dZ**-oligo on A3B_{CTD}-QM-ΔL3-AL1swap has been established previously by our group.¹⁹

The K_i for dZ[U^E(-2), A^{N3}(+1)]X is 0.69 \pm 0.14 μ M, a value around 10 times lower than that for the linear dZ-oligo (7.0 \pm 1.3 μ M).¹⁹ The nanomolar inhibitory potential exhibited by our internally cross-linked oligo shows that properly pre-shaped, cross-linked DNA can become a better inhibitor than the linear one and that the better substrate becomes the better inhibitor, which supports our original hypothesis.

Binding of Cross-Linked Oligos to A3 Enzymes. To understand if the improved inhibition and quality of substrate is due to enthalpic or entropic contributions, we used ITC to determine the thermodynamics of binding. We tested how the cross-linked oligos bind to A3 enzymes and compared this binding to that of linear oligos. For oligos containing dZ, catalytically active $A3B_{CTD}$ -QM- Δ L3-AL1swap was used, as it has the active-site glutamic acid required for protonation of N3 in dZ. This protonation activates C4 in dZ to accept nucleophilic $^{-}OH/H_2O$ coordinated to the Zn²⁺, which converts dZ into a transition-state analogue of cytidine deamination (Figure 1B).⁵⁴ In contrast, A3A-E72A, in which the active-site glutamic acid is replaced with alanine, was used particularly for oligos containing dC so that the complications of deamination can be avoided. This mutation has no change to the active-site structure but does cause a small but significant diminution in ligand binding¹⁹ through loss of hydrogen bonding to the ligand. We expect that the binding of inactivated A3B_{CTD}-QM- Δ L3-AL1swap (as the E255A mutant) to the substrate will be close to or even worse than $K_{\rm m}$ of 200 μ M, which places substrate binding outside ITC applicability. Results of ITC experiments are presented in Table 4. The raw titration data and binding curves are available in Supporting Information (Figures S23-S32). One should emphasize that the compositions of buffers used for NMR kinetic and ITC binding experiments are different, mainly because proteins in ITC experiments need to be at much higher concentration than for kinetic assays (10–100 μ M for ITC versus 50-300 nM in kinetic experiments). This required a search for buffer components that prevent protein precipitation during ITC titrations. Our optimized ITC buffers

oligo	parameters	linear analysis (Lineweaver–Burk plot)	non-linear regression analysis
5'-AT ₃ CAT ₃	$K_{\rm m}$, $\mu { m M}$	200 ± 30	150 ± 30
	$V_{\rm max} \ \mu { m M/s}$	0.014 ± 0.002	0.0126 ± 0.0008
	$k_{\rm cat}$ s ⁻¹	0.29 ± 0.04	0.26 ± 0.02
	$k_{\rm cat}/K_{\rm m}$, s ⁻¹ $\mu{ m M}^{-1}$	0.0014	0.0018
5′-T ₄ CAT	$K_{\rm m}$, $\mu { m M}$	350 ± 70	290 ± 70
	$V_{\rm max} \ \mu { m M/s}$	0.056 ± 0.008	0.051 ± 0.005
	$k_{\rm cat}$ s ⁻¹	0.19 ± 0.03	0.171 ± 0.015
	$k_{\rm cat}/K_{\rm m}$, s ⁻¹ $\mu {\rm M}^{-1}$	0.00054	0.00059

Table 2. Kinetic Parameters of $A3B_{CTD}$ -QM- Δ L3-AL1swap on Two DNA Substrates Used in This Work

Table 3. Inhibition Constants (K_i) for dZ-Containing Oligo Inhibitors of A3B_{CTD}-QM- Δ L3-AL1swap Obtained by Various Calculation Methods^{*a*}

oligo name	parameters	Lineweaver–Burk plot for substrate/Dixon plot for inhibitor	non-linear regression analysis for both substrate and inhibitor
dZ-linear	K_{i} , μ M	7.5 ± 1.7	7.0 ± 1.3
	$V_{ m max}$ $\mu{ m M/s}$		0.0126 ± 0.0008
$dZ[U^{E}(-2), A^{N3}(+1)]X$	K_{i} , μM	0.66 ± 0.14	0.69 ± 0.14
	$V_{ m max}~\mu{ m M/s}$		0.052 ± 0.006

 a dZ-linear inhibitor was evaluated against 5'-AT₃CAT₃ substrate using 50 nM of A3B_{CTD}-QM- Δ L3-AL1swap; dZ[U^E(-2), A^{N3}(+1)]X was evaluated against 5'-T₄CAT substrate using 200 nM of A3B_{CTD}-QM- Δ L3-AL1swap.

Table 4.	Thermody	vnamic I	Parameters	of	Oligos	with .	A3	Enzymes	Obtained	bv	ITC ^a
		/						/		- /	

oligo	protein	ITC buffer ^b	$K_{\rm d}$ (μ M)	$\Delta H(ext{kcal/mol})$	$T\Delta S(\text{kcal/mol})$
5'-AT ₃ CAT ₃	A3A-E72A	1	0.20 ± 0.04	-22.9 ± 1.2	-13.4 ± 1.2
5′-T ₄ CAT	A3A-E72A	1	0.27 ± 0.04	-19.8 ± 0.9	-13.1 ± 0.9
dC-9-mer-X	A3A-E72A	1	0.63 ± 0.13	-18.1 ± 0.9	-9.7 ± 0.9
dC-7-mer-X	A3A-E72A	1	0.48 ± 0.10	-14.1 ± 1.4	-5.4 ± 1.4
dC-5-mer-X	A3A-E72A	1	4.5 ± 0.9	-10.9 ± 1.2	-3.8 ± 1.1
5'-AT ₃ CAT ₃	A3A-E72A	2	0.15 ± 0.03	-53 ± 3	-44 ± 3
5′-T ₄ CAT	A3A-E72A	2	0.75 ± 0.15	-45 ± 2	-36 ± 2
$dC[U^{E}(-2), A^{N3}(+1)]X$	A3A-E72A	2	0.030 ± 0.006	-67 ± 3	-57 ± 4
dZ -linear	$A3B_{CTD}$ -QM- Δ L3-AL1swap	3	11.4 ± 2.3	-13.0 ± 0.6	-6.2 ± 0.6
$dZ[U^{E}(-2), A^{N3}(+1)]X$	$A3B_{CTD}$ -QM- Δ L3-AL1swap	3	2.5 ± 0.5	-24.6 ± 1.2	-16.7 ± 1.7

 ${}^{a}K_{d}$ is the dissociation constant for the ligand from A3; ΔH and $T\Delta S$ are for association of the ligand to A3; the temperature is 25°C. b ITC buffer 1: 50 mM MES, pH 6.0, 100 mM NaCl, 200 μ M EDTA, and 1 mM β -mercaptoethanol; ITC buffer 2: 50 mM Na⁺/K⁺ phosphate, pH 6.0, 50 mM NaCl, 50 mM choline acetate, 2.5 mM TCEP, and 200 μ M EDTA with 30 mg/mL bovine serum albumin; ITC buffer 3: 50 mM Na⁺/K⁺ phosphate, pH 6.0, 200 mM trimethylamine *N*-oxide dihydrate, and 2.5 mM TCEP, 200 μ M EDTA.

were also different for A3A(E72A) and A3B_{CTD}-QM- Δ L3-AL1swap, which means that K_d and K_i values cannot be directly compared. However, our aim was to compare the binding of different oligos to A3 to ascertain if general trends can be established.

The dissociation constants (K_d) obtained by ITC showed no gain in affinity of A3A-E72A for terminally cross-linked oligos in comparison with the standard linear oligos (5'-AT₃CAT₃ and 5'-T₄CAT). The dC-5-mer-X had noticeably weaker binding to A3A-E72A and lower deamination rate as a substrate to A3B_{CTD}-QM- Δ L3-AL1swap compared to its 7mer and 9-mer analogues; this is partly consistent with previous observations on linear substrates of varying lengths^{19,30} but also with not enough flexibility to achieve the right conformation, when a short oligo is cyclized. A reduced entropic component to the free energy of binding is not sufficient to overcome a markedly depressed enthalpic component resulting in relatively weak interactions of ligand with protein.

The internally cross-linked oligo dC[U^E(-2), A^{N3}(+1)]X, which gave higher substrate activity for A3B_{CTD}-QM- Δ L3-AL1swap, has bound five times more strongly ($K_d = 0.030 \pm$

0.006 μ M) in comparison to the linear substrate oligo (5'-AT₃CAT₃, $K_d = 0.15 \pm 0.03 \,\mu$ M) in the same buffer for A3A-E72A. Both unfavorable entropic and favorable enthalpic contributions increased in magnitude, indicating that affinity was improved by the better fitting of active site (enthalpic contribution). The entropic contribution to the Gibbs free energy of binding became more negative (more unfavorable), despite the pre-shaped DNA. This somewhat unexpected observation is attributed to the decrease in protein conformational flexibility upon protein—oligo complex formation as the main source of the entropic contribution, with this decrease being greater for the cross-linked oligo. In contrast, the terminally linked oligos had less favorable enthalpic and less unfavorable entropic contributions upon binding to A3A-E72A.

Next, we evaluated the binding data of **dZ**-containing oligos toward catalytically competent A3B_{CTD}-QM- Δ L3-AL1swap. The dissociation constant of dZ[$U^{E}(-2)$, A^{N3}(+1)]X (K_{d} = 2.5 \pm 0.5 μ M) was around 4.5 times smaller (i.e., stronger binding) than that of **dZ**-containing linear oligo (K_{d} = 11.4 \pm 2.3 μ M), similar to the difference seen in substrates. This trend is also consistent with the trend observed in inhibition constants (K_i) determined by our real-time NMR assay. This effect is smaller in ITC measurements, most likely due to the differences in experimental conditions, as the ITC experiments were conducted in a different buffer and in the presence of trimethylamine *N*-oxide (TMAO) to stabilize the enzyme.

Inhibition of Wild-Type A3A. To evaluate inhibitory potential of cross-linked hairpin oligo against wild-type A3, we expressed in *E. coli* and purified wild-type A3A possessing a (his)₆ tag at the C-terminal end. Recently, we have shown that linear DNA containing 5-fluoro-2'-deoxyzebularine (FdZ-linear)²¹ is superior to the corresponding DNA containing dZ in the inhibition of A3B_{CTD} and A3A.²¹ The best FdZ-linear and cross-linked oligo dZ[U^E(-2), A^{N3}(+1)]X were tested in the NMR assay of A3A activity, where the hairpin DNA 5'-T(GC)₂TTC(GC)₂T (dC-hairpin, deaminated C is bold) was used as a substrate. (Table 5).⁵⁵ The kinetic data

Table 5. K_i Values of Inhibitors of Wild-Type A3A^{*a*}

inhibitor	$K_{\rm i}$ (nM)
F dZ -linear, ²¹ 5'-AT ₃ F dZ AT ₃	2400 ± 940
cross-link, $dZ[U^{E}(-2), A^{N3}(+1)]X$	360 ± 120

^{*a*}Conditions: 50 mM Na⁺/K⁺ phosphate buffer, pH 7.4 supplemented with 100 mM NaCl, 1 mM TCEP, 100 μ M DSS, and 10% D₂O; enzyme concentration: 140 nM; substrate concentration (dC-hairpin): 500 μ M. K_m for dC-hairpin against A3A is 21 μ M, k_{cat} is 0.13 s⁻¹, and k_{cat}/K_m is 6.2 s⁻¹ mM⁻¹.

were measured over time until exhaustion of the substrate and analyzed by global regression using Lambert's W function, which provides superior estimates for $K_{\rm m}$ and $V_{\rm max}$ than nonlinear regression analysis of initial rate data or any of the known linearized transformations of the Michaelis–Menten equation, such as Lineweaver–Burk, Hanes–Woolf, and the Eadie–Hofstee transformations.⁵⁶

Obtained K_i values show that **dZ**-containing cross-linked oligo is a superior inhibitor of the wild-type A3A in comparison with the best F**dZ**-linear oligo. This is consistent with the results obtained for A3B_{CTD}-QM- Δ L3-AL1swap.

DISCUSSION

Recently, we have developed the first selective A3 inhibitors $^{19-21}$ based on incorporation of cytidine-like 2'-deoxyzebularine 57,58 and 5-fluoro-2'-deoxyzebularine (dZ and FdZ on Figure 1B) into short, linear ssDNA, achieving low micromolar inhibition constants (K_i) . Recent structural studies^{7,22,23,59} have revealed that flexible ssDNA adopts a more rigid U-shaped loop upon binding to A3, projecting cytosine into the active-site pocket (Figure 1C). One should note, however, that these studies were conducted on A3A, chimeric $A3B_{CTD}$ (as $A3B_{CTD}$ -QM- Δ L3-AL1swap), and modified $A3G_{CTD}$. Chimeric $A3B_{CTD}$ had four mutations distant from the active site, loop 3 was truncated, and loop 1 of A3B_{CTD} was swapped with loop 1 of A3A; this loop 1, constitutes the main differences between the amino acid sequences of A3A and A3B_{CTD}. This A3B construct has the substrate specificity of A3A, at least in the context of the CCC motif,⁶⁰ which has been used by us recently to develop A3B-specific ssDNA-based inhibitors.²⁰ Strictly speaking, there is no A3B_{CTD} crystal or NMR structure with ssDNA bound that would support a notion that DNA adopts a U-shape in complex with wild-type A3B_{CTD} as well. However, A3A and, to a lesser extent, A3B_{CTD} have been reported recently to

deaminate cytosine in short 3-4 nucleotide loops of DNA hairpins.⁵⁵ This reassures that both A3A and A3B_{CTD} accept Ushaped loops. Interestingly, A3H haplotype I (A3H-I) does not deaminate cytosine in DNA hairpins at all,⁵⁵ suggesting that Ushaped DNA may have some specificity toward A3 Z1 sequence signatures present in A3A, A3B, and A3G.⁶¹ Here, we demonstrate that by cross-linking dA^{+1} and T^{-2} , mimicking the close proximity observed in the crystal structure,⁷ a faster deaminated substrate and a better inhibitor can be obtained in comparison with linear ssDNA of the same length and nucleotide content. These results obtained in our NMRbased activity assay have been corroborated in ITC binding experiments. It is interesting that creating a covalent cross-link between the same dA^{+1} but with the T in position -3completely abolished substrate activity. The cytosine-containing circularized (terminally cyclized) oligos showed lower deamination rates compared to the linear DNA substrate and for that reason were not converted to inhibitors.

The overall inhibition effect was improved from 25-fold (**dZ**linear) to nearly 200-fold ($dZ[U^{E}(-2), A^{N3}(+1)]X$) when the apparent inhibition constants (K_i) of oligos are compared with the K_m of the linear ssDNA substrate 5'-ATTTCATTT ($K_m =$ 150 μ M) on A3B_{CTD}-QM- Δ L3-AL1swap. The same trend was observed for inhibition of wild-type A3A. As K_m (and K_i) depends strongly on the surrounding environment (e.g., buffer composition, pH, and ionic strength—see Table 4), inhibition with even lower K_i may be observed in the cellular environment, because K_m and K_d values in the low $\mu M^{30,31}$ and nM⁶² range have been reported. During our study, two articles have been published on drug discovery efforts to obtain small molecule inhibitors of A3.^{63,64} None approaches the potency of our inhibitors. This highlights the interest and need for powerful A3 inhibitors.

In future, we plan to test if selectivity can be achieved, analogous to that observed with linear ssDNA inhibitors,²⁰ by varying the nucleobases adjacent to the dZ or FdZ in the context of the cross-linked oligos. The pre-shaping of oligos, using our "click coupling" of alkyne and azide, may become a useful general synthetic strategy for oligonucleotide-based inhibitors, as U-shaped nucleic acids feature in complexes with several cellular proteins. So far, the U-shape has been observed in APOBEC-related tRNA adenosine deaminase TadA (PDB 2B3J)⁶⁵ as well as in functionally and structurally unrelated proteins, such as telomere protein Pot1pc (PDB 4HIO)⁶⁶⁻⁶⁸ and bacterial cold-shock protein Bs-CspB (PDB 2ES2 and 3PF4).^{69,70} More generally, the structural information can be harvested to design pre-shaped DNA/RNA inhibitors for any DNA/RNA-interacting protein, if a significant change in polynucleotide shape is found upon complex formation.

CONCLUSIONS

Inhibition of APOBEC enzymes, especially A3A and A3B, is clinically relevant as these enzymes are involved in cancer mutagenesis and development of drug resistance. In this study, we developed pre-shaped ssDNAs as potentially better substrates for A3 enzymes using Cu(I)-catalyzed "click chemistry" to cross-link alkyne and azide moieties embedded in a ssDNA fragment. This has led to the first nM inhibitor of $A3B_{CTD}$ -QM- Δ L3-AL1swap and A3A by incorporating **dZ** in place of the target dC in the cross-linked DNA fragment. This work provides the stepping stone for further development of modified ssDNAs as potential A3 inhibitors.⁷¹

Article

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.biochem.2c00449.

Experimental details about the synthesis of azido-2deoxyribose CPG **4**, adenosine **11**, modified oligos, cross-linking protocol, and NMR analysis of linear versus cross-linked oligo; protein expression and purification; ITC and NMR-based kinetic assays; examples of the calculation of inhibition of A3 enzyme by dZ-containing cross-linked oligos; sequence alignment of proteins used in this study; ¹H, ¹³C, ³¹P NMR, IR, and HRMS (ESI) spectra of new compounds synthesized; and RP-HPLC profiles and HRMS (ESI) spectra of linear and crosslinked oligos (PDF).

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Notes

The authors declare the following competing financial interest(s): The authors have filed a patent application on cross-linked ssDNA as enzyme inhibitors (WO 2022162536).

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