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Application of 4'-C- α -aminoethoxy-2'-O-methyl-5-propynyl-uridine for antisense therapeutics†

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Owing to the increased public interest and advances in chemical modifications, the approval of antisense therapeutics, a class of mRNA-targeting DNA-based oligonucleotide therapeutics, has accelerated in recent years. It was previously reported that siRNAs with several 4'-C- α -aminoethoxy-2'-O-methyl-uridine (4AEoU) analogs could maintain moderate thermal stability similar to the native ones while showing robust nuclease stability. In this study, we further expanded the application of 4AEo modification to antisense therapeutics and achieved superior thermal stability by adding the uracil 5-propynyl modification. Antisense oligonucleotides containing 4'-C- α -aminoethoxy-2'-O-methyl-5-propynyl-uridine (4AEo^{PU}) could efficiently activate RNase H-mediated antisense *in vitro* in the presence of native DNA gaps. These results encourage future studies of 4AEo^{PU}-containing antisense therapeutics.

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Introduction

The COVID-19 pandemic has brought public attention to mRNA vaccines that use long-stranded RNAs encoding antigens as drug molecules. As a novel and attractive modality, nucleotide-based therapeutics are considered useful in treating diseases related to viral infections and genetic mutations. Compared to long mRNA vaccines, antisense oligonucleotides (ASOs) composed of short single-stranded DNA-like oligonucleotides are representative and constitute the majority of approved oligonucleotide therapeutics.^{1–3} Since they were first approved by the Food and Drug Administration (FDA) in 1998, 12 types of antisense therapeutics have been approved till 2024.^{4–15} ASOs hybridize with their target mRNA strands in the cytoplasm or nucleus and modulate the gene expression by the RNase H-mediated mRNA-selective cleavage or steric hindrance during pre-mRNA splicing or protein translation.^{16–18} Because native oligonucleotides are too unstable to survive nuclease-mediated degradation and exhibit poor cellular uptake due to their negatively charged phosphate linkages, various chemically

modified ASOs have been developed and utilized instead of native oligonucleotides.^{3,19–21} There is a special platform for RNase H-dependent ASOs called a “gapmer,” which has been employed in several approved antisense therapies.^{5,8,9,13,14} The antisense gapmer consists of a central DNA gap region to maintain the ability to activate the RNase H-mediated antisense mechanism and two chemically modified wing regions on either side to improve thermal and enzymatic stabilities.²² Generally, phosphorothioate (PS) linkages are utilized instead of native phosphate (PO) linkages to enhance cellular uptake and protect ASOs from enzymatic degradation.^{23–27}

In our previous studies, siRNAs, another typical oligonucleotide therapeutic consisting of double-strand RNAs, containing 4'-C- α -aminoalkyl-2'-O-methyl-modified uridine analogs showed superior nuclease resistance and tended to be more stable against nucleases with relatively longer side chains.²⁸ Nevertheless, incorporation of the aminoalkyl side chain into C4' led to a C2'-*endo* preference and the significant thermal destabilization of siRNAs. Through inserting an oxygen atom to form a 4'-C- α -aminoalkyl-2'-O-methyl (4AEo) modification, the modified siRNAs showed moderate thermal stability as the native ones and were more stable than those with a 4'-C- α -aminopropyl-2'-O-methyl (4AP) modification.²⁹ This improvement prompted further investigations into the application of 4AEo modification in antisense therapeutics. Furthermore, C2'-O-modifications such as 2'-O-methyl (2'-OMe) and 2'-O-methoxyethyl (2'-MOE) are well known to inhibit RNase H recruitment, resulting in poor RNase H-mediated cleavage.^{30,31} Another study showed that gapmers containing 4AEo-modified deoxythymidine without any 2'-OMe modification in the gap region reduced the antisense activity, indicating an unfavorable effect of the aminoethoxy side chain on the RNase H

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mechanism.³² Therefore, we hypothesized that it would be better for the 4AEo-modified analogs to be located in the wing regions of antisense gapmers. Based on this hypothesis, the thermal stability of the 4AEo-modified analogs must be further improved to obtain sufficient target RNA binding affinity for the antisense gapmer.

In this study, we designed two novel 4AEo-modified uridine analogs: 4'-C- α -aminoethoxy-2'-O-methyl-5-methyl-uridine (4AEo^mU) and 4'-C- α -aminoethoxy-2'-O-methyl-5-propynyl-uridine (4AEo^PU) (Fig. 1). Both 5-methyl and 5-propynyl modifications have been used to stabilize the siRNAs formation.³³ The same thermal stabilization was also confirmed in ASOs, and no inhibition towards RNase H was observed, as the 5-propynyl moiety was located in the major groove of the DNA/RNA duplexes.³⁴ The synthesis and physical and biochemical evaluations of the 16-mer thymidine-cytidine(TC)-repeat DNA oligomers containing these novel uridine analogs are described. A series of 4AEo-modified antisense gapmers based on a previously well-studied 16-mer KRAS-targeting sequence was prepared and used for cellular evaluation.

Synthesis of novel nucleoside analogs

The novel synthesis of the phosphoramidite corresponding to 4'-C- α -aminoethoxy-2'-O-methyl-5-methyl-uridine (4AEo^mU) is shown in Scheme 1. Commercially available 2'-O-methyl-5-methyl-uridine (**1**) was subjected to iodination *via* the Appel reaction, followed immediately by dehalogenation using 1,8-diazabicyclo[5.4.0]-7-undecene (DBU) to obtain the olefin derivative. Subsequently, 3'-O-silyl protection was performed to obtain compound **4**. After epoxidation with Oxone® in acetone, the aminoethoxy side chain was incorporated into the C4' position using *N*-trifluoroacetyl aminoethanol in the presence of a Lewis acid. Two kinds of products were obtained from this reaction with a ratio of 2 : 1. According to an earlier study in 1993, epoxidation of the 4',5'-double bonds could yield diastereomeric epoxide derivatives, which would be attacked by the alcohol reactant and transform into the corresponding diastereomeric 4'-C-substituted ones.³⁵ Furthermore, the 4'-C- α -aminoethoxy derivatives would be the major products in the presence of the 3'-O-silyl protecting group.^{29,32,36,37} Therefore, we hypothesized that the desired compound **6** was also predominantly obtained in this study. The configurations of the C4' position in the major product were determined *via* nuclear Overhauser effect spectroscopy (NOESY) to confirm our

hypothesis. As shown in ESI,† consistent with the previous studies, the aminoethoxy side chain was proven to be inserted at the 4'-C- α -site by the correlations observed between H1' and H6', H3' and H5', and H3' and OH5'.^{29,32,38} Thus, compound **6** was the major product, as expected, and was used for the following process.

Next, the 5'-hydroxy moiety was protected using a 4,4'-dimethoxytrityl (DMTr) group, followed by deprotection of the 3'-O-silyl group with 0.1 M tetrabutylammonium fluoride (TBAF) solution in tetrahydrofuran (THF). Phosphoramidite **9** was synthesized using a standard phosphitylation procedure with *N,N*-diisopropylethylamine (DIPEA) and 2-cyanoethyl-*N,N*-diisopropylchlorophosphoramidite (CEPCI) as reagents.

The phosphoramidite corresponding to 4'-C- α -aminoethoxy-2'-O-methyl-5-propynyl-uridine (4AEo^PU) was prepared from a previously reported compound **10** (Scheme 2).²⁹ Initially, the 3'-O-silyl protecting group was removed due to its low stability in the subsequent steps. Selective 5-iodination was accomplished using diammonium cerium(IV) nitrate (CeAN) as the catalyst, and the propynyl group was introduced by Sonogashira-Hagihara cross-coupling. Finally, after 5'-O-DMTr protection, the corresponding phosphoramidite **15** was prepared using the standard phosphitylation procedure.

Synthesis of oligonucleotides

Novel nucleoside analogs were incorporated into the DNA-based oligonucleotides utilizing the solid-phase phosphoramidite method with a DNA/RNA synthesizer. After oligosynthesis, to prevent the reaction of acrylonitrile with the amino groups at the end of the 4'-C- α -aminoethoxy side chain, controlled-pore glass (CPG) beads were treated with 10% dimethylamine in acetonitrile (MeCN), followed by rinsing with MeCN to selectively remove the cyanoethyl groups. The oligonucleotides were then treated with an aqueous ammonia solution for cleavage from the CPG beads and deprotection and finally purified by 20% denaturing polyacrylamide gel electrophoresis (PAGE) containing 7 M urea. The sequences of the oligonucleotides used in this study are shown in Tables 1 and S1.†

Thermal stability

Because it is important for RNase H-dependent antisense oligonucleotides to hybridize to their target mRNAs with excellent affinity, the 50% melting temperature (T_m) values of

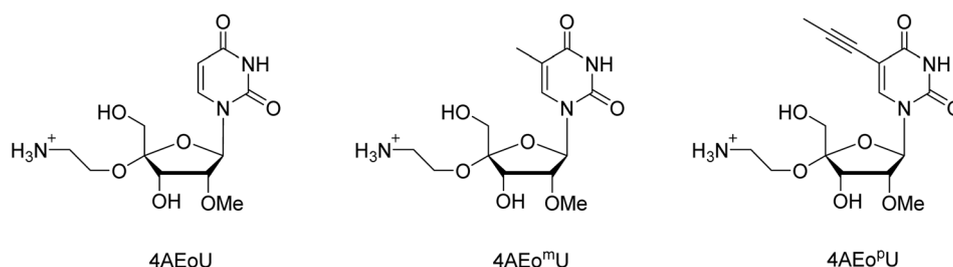
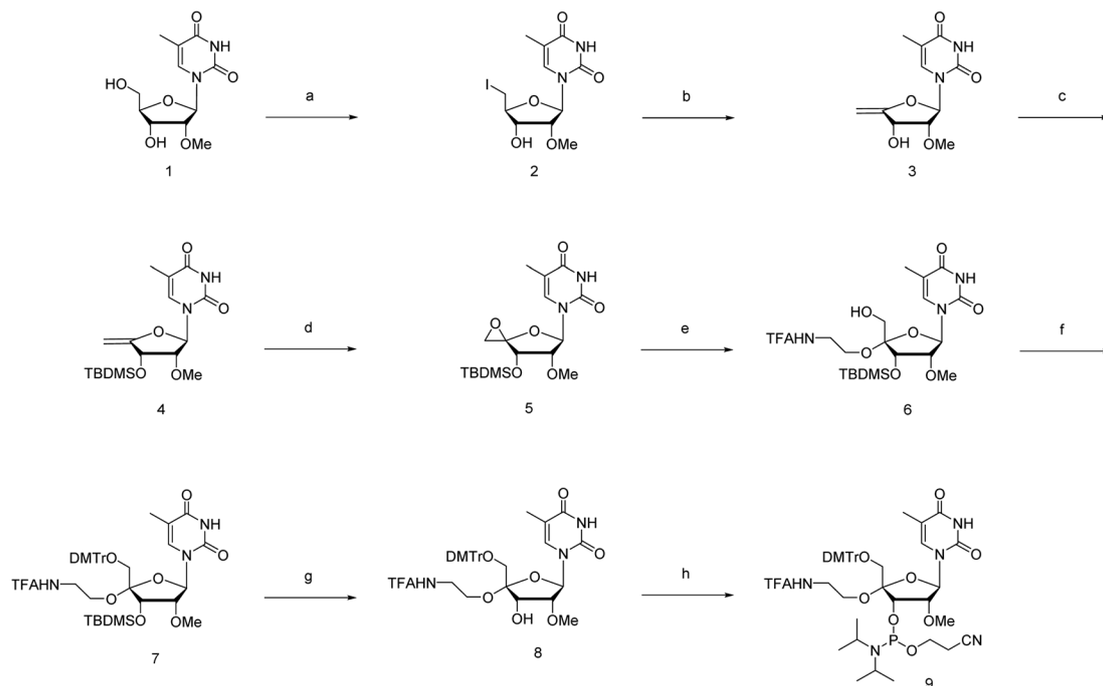


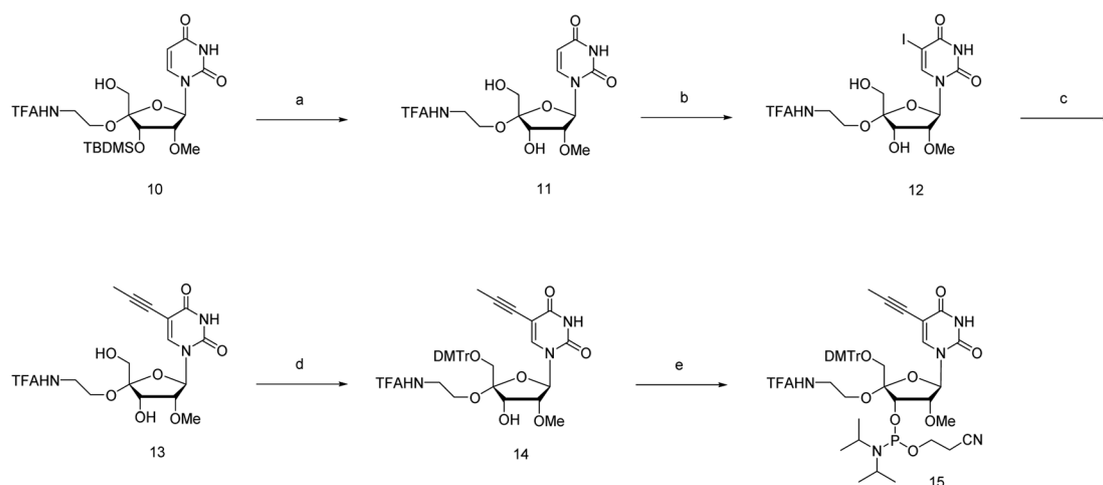
Fig. 1 The 4AEo-modified uridine analogs.



Scheme 1 Synthesis of 4'-C- α -aminoethoxy-2'-O-methyl-5-methyl-uridine phosphoramidite **9**.^a Reagents and conditions: (a) PPh₃, imidazole, I₂, THF, r. t., 4 h; (b) DBU, 24 h; (c) TBDMSCl, imidazole, DCM, 3 h; **4**: 74% in 3 steps; (d) Oxone®, sat. NaHCO₃ aq., DCM, acetone, r. t., 2 h; (e) *N*-trifluoroacetylaminoethanol, ZnCl₂, THF, -60 °C; then, r. t., 24 h; **6**: 29% in 2 steps; (f) DMTrCl, DIPEA, pyridine, r. t., overnight; **7**: 79%; (g) TBAF, THF, r. t., 1 h; **8**: 86%; (h) CEPCI, DIPEA, THF, r. t., 2 h; **9**: 55%.

DNA/RNA duplexes containing modified oligonucleotides with their complementary RNAs strand should be evaluated. Here, we employed a TC-repeat modeling sequence tested in our previous studies.^{32,39–42} For the T_m measurement, each unmodified or 25% modified 16-mer DNA was hybridized with a 16-mer complementary RNA strand, and 3 μ M of these DNA/RNA duplexes were utilized to obtain the T_m values. The results in Table 1 show that, compared with the unmodified TC-Nat., the DNA/RNA duplex containing four 4AEo^mU analogs resulted in

a slight thermal destabilization ($\Delta T_m -1.2$ °C; -0.3 °C per modification). Conversely, incorporating 4AEo^pU led to a significant increase in the T_m value ($\Delta T_m +4.2$ °C; $+1.4$ °C per modification), suggesting thermal stabilization by the 5-propynyl moiety. Furthermore, thermodynamic parameters were calculated using 1, 3, 6, 12, 15, 30, and 60 μ M of these DNA/RNA duplexes, providing more insight into the differences in T_m values. As shown in Table 1, the free energy of duplex formation (expressed as ΔG°) of the unmodified DNA/RNA duplexes was



Scheme 2 Synthesis of 4'-C- α -aminoethoxy-2'-O-methyl-5-propynyl-uridine phosphoramidite **15**.^a Reagents and conditions: (a) TBAF, THF, r. t., 1 h, **11**: 95%; (b) CeAN, I₂, 80 °C (reflux), 1 h; (c) Pd(PPh₃)₂Cl₂, CuI, TEA, propyne, DMF, 35 °C, overnight; **13**: 79% in 2 steps; (d) DMTrCl, DIPEA, pyridine, 50 °C, 24 h, **14**: 73%; (e) CEPCI, DIPEA, THF, r. t., 2 h, **15**: 72%.



Table 1 Sequence of TC-repeat modeling oligomers, T_m values, and thermodynamic parameters of the modified duplexes

Oligo	Sequence ^a	T_m^b (°C)	ΔT_m^c (°C)	ΔH°^d (kcal mol ⁻¹)	ΔS°^d (cal mol ⁻¹ K ⁻¹)	ΔG°^d (kcal mol ⁻¹)
TC-Nat.	5'-TCT TTC TCT TTC CCT T-3'	59.3	—	-148.4	-417.7	-23.9 (25 °C) -18.9 (37 °C)
TC-m	5'-TCT T _m C TCT mTC CC _m T-3'	58.1	-1.2	-129.9	-364.1	-21.4 (25 °C) -17.1 (37 °C)
TC-p	5'-TCT T _p C TCT pTC CC _p T-3'	63.5	+4.2	-156.5	-436.2	-26.5 (25 °C) -21.3 (37 °C)

^a T and C denote native thymidine and deoxycytidine, respectively. m and p in red denote the 4AEo^mU and 4AEo^pU analogs, respectively. ^b The T_m values were measured in 10 mM sodium phosphate buffer (pH 7.0) containing 100 mM NaCl. The sequence of the complementary RNA strand was 5'-r(a agg gaa aga gaa aga)-3', where a and g denote native adenosine and guanosine, respectively. The concentrations of the duplexes were 3 μM. All measurements were performed in triplicate, and the data are presented as average values. ^c ΔT_m represents [T_m (modified) - T_m (unmodified)]. ^d Thermodynamic parameters were calculated based on the slope of the $1/T_m$ vs. $\ln(C_T/4)$ plot. All measurements were performed in triplicate, and the data are presented as average values.

-23.9 kcal mol⁻¹ at 25 °C and -18.9 kcal mol⁻¹ at 37 °C. In contrast, the ΔG° values of the 4AEo^mU-containing duplex were slightly lower (-21.4 kcal mol⁻¹ at 25 °C and -17.1 kcal mol⁻¹ at 37 °C). This result was consistent with the T_m measurements. Similarly, the decrease in ΔG° for duplexes containing 4AEo^pU (-26.5 kcal mol⁻¹ at 25 °C and -21.3 kcal mol⁻¹ at 37 °C) indicated that it could more readily bind with the target RNA strand.

Circular dichroism (CD) spectra showed that the 4AEo-modified DNA/RNA duplexes formed the A-type-like helix, similar to those of the unmodified ones (Fig. S3†), which is consistent with a previous study on the secondary structure of duplexes consisting of homopyrimidine DNA strands and homopurine RNA strands.⁴³ No significant change was observed in the negative maximum around 210 nm, but either the wavelength or the intensity of the positive maximum shifted slightly due to the incorporation of the 4AEo^mU analogs and significantly shifted in the case of the 4AEo^pU analogs.

Since all nucleotides adopt the C3'-endo sugar pucker when forming a typical A-type helix, a more favorable pre-orientation and a resulting helix more akin to the A-type could be predicted if nucleoside analogs in the DNA strand prefer a C3'-endo sugar pucker. In a previous study, we investigated whether RNA/RNA duplexes containing several 4AEoU analogs could achieve thermal stability equivalent to the unmodified uridines.²⁹ It was possible that a favorable C3'-endo sugar pucker resulted from the $n_{O4'} \rightarrow \sigma_{C4'eO}^*$ anomeric and $\sigma_{C3'eH} \rightarrow \sigma_{C4'eO}^*$ gauche effects of the 4'-C-α-aminoethoxy side chain, as well as the 2'-O-methyl modification.⁴⁴ It could also explain the favorable entropy observed in the 4AEo^mU-containing DNA/RNA duplex (ΔS° -364.1 cal mol⁻¹ K⁻¹) compared with the unmodified one (ΔS° -417.7 cal mol⁻¹ K⁻¹). However, regardless of the favorable entropy, the 4AEo^mU-containing duplex exhibited slightly lower thermal stability, according to the T_m measurements. Several factors affect the thermal stability of the nucleotide duplexes, including hydrogen bonding, stacking interactions, conformational entropy, counterion condensation, and hydration.⁴⁵ Since there

was favorable counterion condensation due to the aminoethoxy moiety and no change in thymine, its thermal destabilization might be due to unfavorable hydration resulting from steric repulsion between the 4'-C-α-aminoethoxy side chain and the 2'-O-methyl group. Conversely, the ΔS° value for the 4AEo^pU-modified duplex (-436.2 kcal mol⁻¹) was unfavorable, while the ΔH° value (-156.5 kcal mol⁻¹) was more favorable than the other duplexes. It has been previously reported that incorporating a propynyl group at the C5 position of thymine can result in an enhanced stacking effect.⁴⁶ Therefore, we hypothesized that 4AEo^pU analogs also prefer a C3'-endo sugar pucker similar to 4AEo^mU analogs, and the improved stacking interaction seems to play a major role in this case, showing more favorable enthalpy.

Molecular dynamics simulation

As described above, the incorporation of the 4'-C-α-aminoethoxy side chain was expected to further induce the sugar pucker into C3'-endo because of the $n_{O4'} \rightarrow \sigma_{C4'eO}^*$ anomeric and $\sigma_{C3'eH} \rightarrow \sigma_{C4'eO}^*$ gauche effects. Based on molecular dynamics (MD) simulations, we performed *in silico* calculations of these 4AEo-modified nucleoside analogs to investigate the changes in nucleoside sugar pucker and molecular fluctuations caused by chemical modifications. The results of the MD simulations are presented in Table 2, Fig. S4 and S5.† Native thymidine prefers the C2'-endo sugar pucker due to the absence of the C2'-substituent. Consistent with that, the percentage of C3'-

Table 2 Features of thymidine, 4AEo^mU, and 4AEo^pU obtained from the MD simulation

	Thymidine	4AEo ^m U	4AEo ^p U
C3'-endo ^a (%)	5	76	67
RMSD ^b (Å)	0.52 (±0.15)	1.18 (±0.23)	1.04 (±0.33)

^a The percentages of C3'-endo sugar pucker were calculated based on a criterion that considers the sugar pucker as C3'-endo if $0^\circ < P < 90^\circ$ and $270^\circ < P < 360^\circ$. ^b The mean values and standard deviations of the RMSD (Å). The values are shown as average (±standard deviation).



endo sugar puckering in the case of thymidine was only 5% from the MD simulation. In contrast, the incorporation of the 4AEo modification resulted in a significant shift towards the *C3'-endo* sugar puckering, as expected, with 76% for the 4AEo^mU analog and 67% for the 4AEo^pU analog. The 4AEo^pU analog was slightly less favorable for *C3'-endo* sugar puckering than the 4AEo^mU analog, suggesting steric hindrance of the 5-propynyl moiety. The root mean square deviations (RMSDs) were calculated relative to the reference structure obtained from the energy minimization calculation. The distribution range of the RMSD, shown as the standard deviation (SD), is considered an indicator of fluctuations in nucleoside molecules. As shown in Table 2, the SD of the RMSDs of thymidine, 4AEo^mU analog, and 4AEo^pU analog were ± 0.15 , ± 0.23 and ± 0.33 , respectively. Therefore, the fluctuation of the 4AEo^mU analog was more pronounced than that of native thymidine because of the flexibility of the 4'-C- α -aminoethoxy side chain. Furthermore, the nucleoside molecule vibrated more violently when the 5-methyl group was changed to a propynyl group, indicating a negative effect of the 5-propynyl modification on the nucleoside structural stability.

Consequently, our hypothesis proved that either the 4AEo^mU or 4AEo^pU analog that prefers *C3'-endo* sugar puckering would result in favorable entropy for the DNA/RNA duplex formation. Notably, the 4AEo^pU analog was more structurally unstable than the 4AEo^mU analog, resulting in unfavorable entropy due to the extra energy required to strain the structure for duplex formation. However, the enhanced stacking effect of the 5-propynyl modification was strong enough to recover the structural energy loss, resulting in favorable enthalpy and higher thermal stability than the 4AEo^mU analog.

Base-discrimination ability

To investigate the base-discrimination ability of the 4AEo-modified analogs, the binding affinities of these modified DNAs towards one-base mismatched RNA strands were evaluated according to their T_m values. As the results show in Table 3 and Fig. S1,† the T_m values of the unmodified duplexes with

a mismatched base pair with uridine (u), guanosine (g), and cytosine (c) were 49.1, 55.9, and 49.2 °C, respectively. Compared with the full-matched duplex (T_m (T:a) 59.3 °C), pyrimidine:pyrimidine mismatching decreased the T_m value significantly by approximately 10 °C, whereas a mismatched base pair consisting of thymidine and guanosine led to a decrease of approximately 3 °C. The incorporation of one 4AEo^mU analog yielded results similar to those of the unmodified thymidine. The T_m values were 59.3 °C for the full-matched duplex and 49.3 and 49.2 °C for the pyrimidine:pyrimidine mismatched ones. However, a highly beneficial decrease of 5 °C was observed for 4AEo^mU:guanosine mismatching. The same trend was observed for the 4AEo^pU analog. One thymidine substituted with a 4AEo^pU analog increased the T_m value of the full-matched duplex by 2 °C (T_m (p:a) 61.5 °C). The T_m values of the p:u and p:c mismatching were 51.0 and 51.2 °C, respectively. These T_m values were 2 °C higher than those of the unmodified and 4AEo^mU-containing ones, while the ΔT_m values were the same as the others, approximately -10 °C. Conversely, the T_m (p:g) was 54.2 °C, which was not significantly different from T_m (m:g) (54.0 °C), but 7 °C lower than the full-matched T_m (p:a).

Thymidine not only hybridizes with adenosine to form the Watson-Crick base pairs but can also hybridize with other nucleobases to form the so-called “wobble” base pairs, which have been studied to be as stable as the Watson-Crick base pairs.⁴⁷ While the formation of wobble base pairs is important for some biological mechanisms, such as translation, it is unsuitable for antisense therapeutics because of the associated risk of off-target effects. In this study, the thymidine:pyrimidine mismatched duplexes were severely destabilized, suggesting that no wobble base pairs were formed. However, a 2 °C increase was still observed in the case of T_m (p:u) and T_m (p:c). This indicates that thermal stabilization from improved stacking interactions by the 5-propynyl modification was still present without base pairing. Conversely, because the ΔT_m value of thymidine: guanosine mismatching was -3 °C, there is no doubt that thymidine formed a wobble base pair with guanosine. The ΔT_m values were also moderate in the cases of 4AEo^mU:guanosine and 4AEo^pU:guanosine mismatching, suggesting the formation of wobble base pairs. However, their T_m values were comparable (T_m (m:g) 54.0 °C vs. T_m (p:g) 54.2 °C), indicating that the formation of wobble base pairs altered the torsion of nucleobases and changed the relative position of the 5-propynyl moiety, resulting in no enhanced stacking effect. Overall, the 4'-C- α -aminoethoxy-2'-O-methyl-5-propynyl modification exhibited higher thermal stability and base discrimination ability.

Nuclease resistance

The enzymatic stability of the modified DNAs was investigated in OPTI-MEM supplemented with 3% (v/v) bovine serum (BS) containing various exonucleases and endonucleases. In this study, the 5'-fluorescein-labeled TC-Nat-F, TC-m-F, and TC-p-F, corresponding to TC-Nat., TC-m, and TC-p, respectively, were used. Their sequences are shown in Table S1.† Samples aliquoted at each timepoint were analyzed using 20% denaturing

Table 3 T_m values of full-matched and one-mismatched DNA/RNA duplexes

T_m^c (°C) (ΔT_m^d (°C))		X^b			
		a	u	g	c
T^a	Thymidine (T)	59.3(-)	49.1(-10.2)	55.9(-3.4)	49.2(-10.1)
	4AEo ^m U (m)	59.3(-)	49.3(-10.0)	54.0(-5.3)	49.2(-10.1)
	4AEo ^p U (p)	61.5(-)	51.0(-10.5)	54.2(-7.3)	51.2(-10.3)

^a The sequence of the DNA strand: 5'-TCT TTC ICT TTC CCT T-3'. T and C denote native thymidine and deoxycytidine, respectively. T denotes native thymidine, 4AEo^mU, or 4AEo^pU analog, as shown in the table.

^b The sequence of the RNA strand: 3'-aga aag Xga aag gga a-5'. a, u, g, and c denote native adenosine, uridine, guanosine and cytosine, respectively. X denotes a mismatched site. ^c The T_m values were measured in 10 mM sodium phosphate buffer (pH 7.0) containing 100 mM NaCl. The concentrations of the duplexes were 3 μ M. All measurements were performed in triplicate and the data are presented as average values. ^d ΔT_m represents [T_m (mismatched) - T_m (full-matched)].



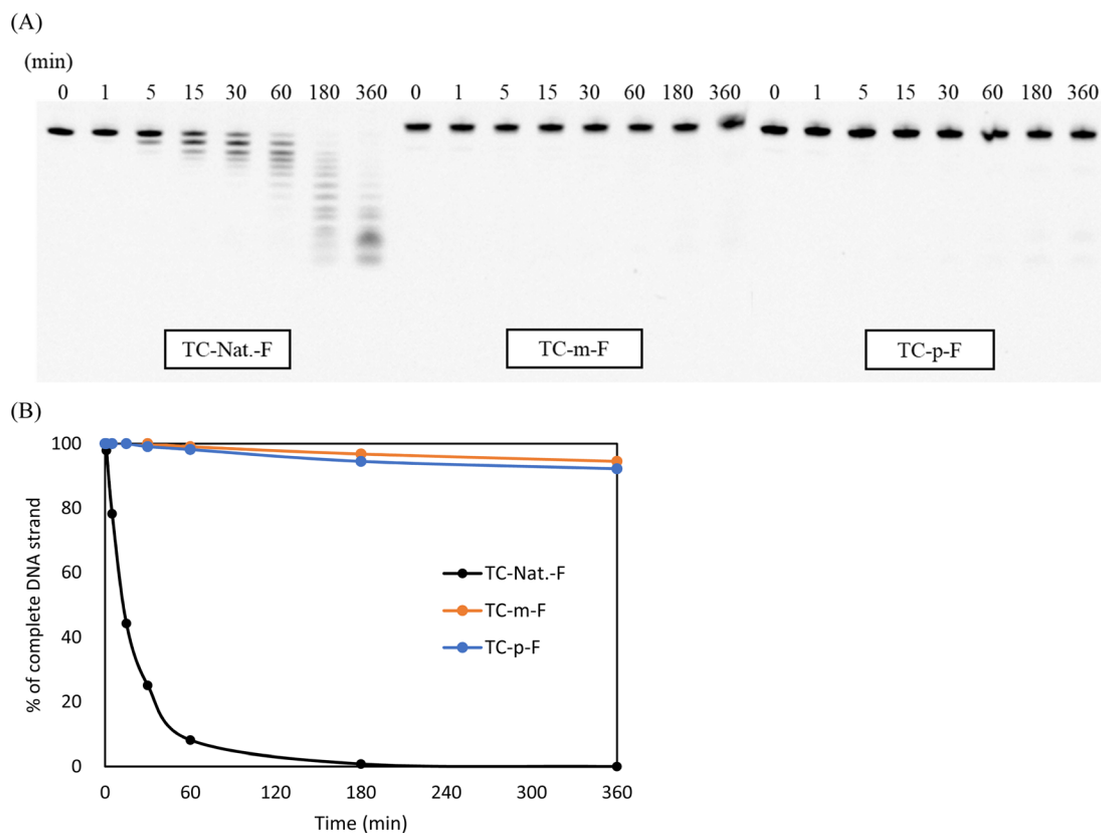


Fig. 2 PAGE analysis of single-stranded oligomers treated with 3% bovine serum^a. ^aThe 5'-fluorescein-labeled TC-Nat.-F, TC-m-F, and TC-p-F, corresponding to TC-Nat., TC-m, and TC-p, respectively, were incubated in OPTI-MEM containing 3% bovine serum, and samples from the reaction mixtures at various incubation time points (0, 1, 5, 15, 30, 60, 180, and 360 min) were analyzed via 20% PAGE with 7 M urea. (A) PAGE images obtained from luminescent analysis; (B) plot of the percentage of complete DNA strands remaining at each time point analyzed via ImageJ.

PAGE. The resulting images were acquired using LAS-4000 (Fujifilm) and quantified using ImageJ software, as shown in Fig. 2. Unmodified TC-Nat.-F was degraded to half its amount within 15 min, with no complete strand remaining after 180 min. In contrast, more than 90% of the full-length strands of 4AEo-modified TC-m-F and TC-p-F remained even after 360 min of the BS treatment. We further tested the nuclease resistances of TC-m-F and TC-p-F under 10%, 25% and 50% BS conditions (Fig. S6[†]). No significant differences were observed between the 4AEo-modified DNAs. With 360 min of incubation, approximately 20% of the full-length strands of the 4AEo-modified DNAs survived in a 50% BS-containing environment, indicating robust nuclease resistance due to the 4AEo modification. Substitution of the 2'-hydroxy group with a methoxy group might avoid nuclease-mediated hydrolysis of the 3'-side phosphate diester linkage. Additionally, the aminopropyl modification at the C2' site was shown to be highly nuclease-resistant and contributed to an electrostatic effect of the positively charged amino group occupying the metal ion binding site of exonucleases instead of a magnesium ion, as well as the steric effect of removing the phosphate group from a zinc ion.⁴⁸ Altering the aminoalkyl side chain into the C4' or C5' site could also significantly increase stability against nucleases. Furthermore, because the variety of cleavage fragments from 4AEo-

modified DNAs was reduced to 3 or 4, the 4AEo modification appeared to prevent not only the 3'-phosphate but also at least two phosphate diester bonds on the 5'-side.

E. coli RNase H1 activation

To confirm the ability of the modified DNAs to activate the RNase H-mediated antisense mechanism, we utilized RNase H1 purified from *Escherichia coli* (*E. coli*) and the TC-repeat modeling sequence containing several native 4-mer DNA gaps that are sufficient for recognition and cleavage by *E. coli* RNase H1.⁴⁹ To evaluate the turnover, unmodified TC-Nat., modified TC-m, and TC-p were annealed to their complementary 5'-fluorescein-labeled RNA strand in a ratio of 1:5. Then, these duplexes treated with 3.5 U of *E. coli* RNase H1 at 37 °C and samples aliquoted at each time point were analyzed by 20% denaturing PAGE (Fig. 3). Consequently, the *E. coli* RNase H1-mediated cleavage was efficiently activated by the unmodified DNA/RNA duplex, leaving no full-length RNA strands within 5 min. However, when comparing each result at the 5 min timepoint, the cleavage rate of duplexes containing the 4AEo^mU analogs was slower than that of the unmodified ones and even slower in the case of 4AEo^pU analogs. It was reported that chemical modifications inside the minor groove of DNA/RNA duplexes, such as 2'-O-methyl modification, influence RNase H1 recognition, which



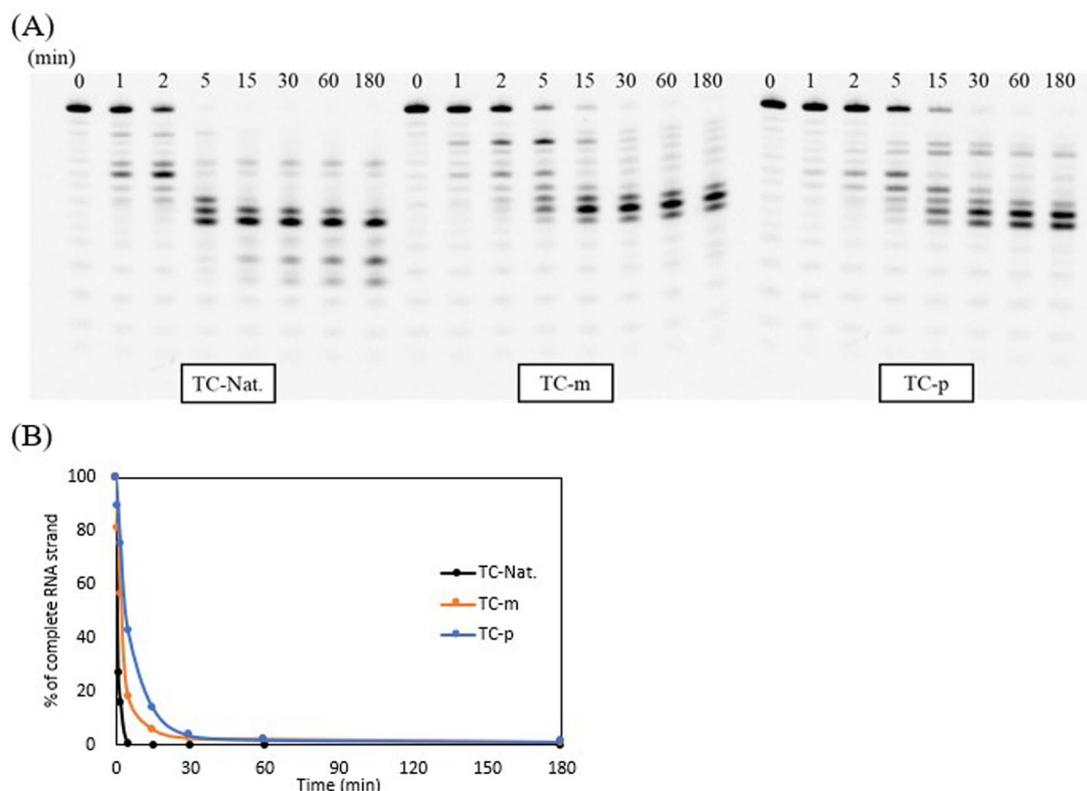


Fig. 3 PAGE analysis of native and modified duplexes treated with *E. coli* RNase H¹. ^aThe duplexes containing TC-Nat., TC-m, and TC-p within the 5'-fluorescein-labeled complementary RNA strand were incubated in a reaction buffer with a diluted RNase H1 solution (50 units per L in water). Samples from the reaction mixtures at various incubation time points (0, 1, 2, 5, 15, 30, 60, and 180 min) were analyzed via 20% PAGE with 7 M urea. (A) PAGE images obtained from luminescent analysis; (B) plot of the percentage of complete RNA strands remaining at each time point analyzed via ImageJ.

was not cleaved when fully modified with the 2'-O-methyl group. In addition, metal ions are required for the *E. coli* RNase H1-mediated cleavage.^{49–51} There was concern that the 4'-C-aminoethoxy side chain would electronically and sterically interfere with RNase H1 recognition. Conversely, 5-propynyl modification, a chemical modification located in the major groove, efficiently knocked down the target mRNA in cells.³⁴ However, it has also been reported that the 5-propynyl moiety incorporated into siRNAs can interfere with interactions at the major groove of the RISC complex, adversely affecting RNAi activity.³³ Similar disruptions might also appear in DNA/RNA duplexes and RNase H1. Therefore, considering the change in the variety of cleavage products, it seems that the binding, recognition, and migration of *E. coli* RNase H1 were affected by the 4AEo modification, resulting in a shift in the initial and final cleavage sites and a decrease in the cleavage rate.

Nevertheless, the 4AEo-modified duplexes also activated *E. coli* RNase H1 for RNA-selective cleavage if native DNA sections were long enough for activation, indicating their potential for application as antisense gapmers.

Antisense activity in mammal cells

As aforementioned, we demonstrated that the 4'-C- α -aminoethoxy-2'-O-methyl-5-propynyl modification exhibited superior thermal

and biochemical stability in the TC-repeat modeling sequence and was able to activate the RNase H-mediated RNA-selective cleavage with several DNA gaps. We further investigated its effect on antisense activity when incorporated into antisense gapmers. Here, a cancer-related gene, *KRAS*, was chosen as the endogenous target in NCI-H460 cells, and a 3-10-3 LNA-wing antisense gapmer served as a positive control. Based on this platform, a series of *KRAS*-targeting antisense gapmers ("KRASs" in short) were prepared by replacing an LNA-T in the wing region as well as a thymidine in the gap region with a 4AEo^mU or 4AEo^pU analog, leaving at least an 8-mer DNA-gap for Human RNase H activation.⁵²

Before operating the cell assays, duplexes composed of the KRASs and their complementary RNA strands were checked to determine if they were sufficiently stable under biological conditions (37 °C). As shown in Table 4, substituting an LNA-T with a 4AEo^mU analog caused a significant decrease in T_m values, -4.6 °C for KRAS-m-1 and -3.6 °C for KRAS-m-4, whereas substituting a native thymidine showed comparable or even higher T_m values, -0.2 °C for KRAS-m-2 and $+1.3$ °C for KRAS-m-3. Notably, when thymidine was replaced, the 4AEo^mU analog caused a slight decrease in the TC-repeat modeling sequence but an increase in KRAS-m-3, indicating the possibility of being affected by the position inside the oligonucleotide and the variety of surrounding nucleobases. It was considered that it would be better to add the 4AEo modification



Table 4 Sequence of KRAS gapmers and T_m values of duplexes containing these gapmers

No. of gapmer	Sequence ^a	T_m ^b (°C)	ΔT_m ^c (°C)
KRAS-pos	5'- <u>G</u> · <u>C</u> · <u>T</u> ·A·T·T·A·G·G·A·G·T·C· <u>T</u> · <u>T</u> · <u>T</u> -3'	62.5	—
KRAS-m-1	5'- <u>G</u> · <u>C</u> · <u>m</u> ·A·T·T·A·G·G·A·G·T·C· <u>T</u> · <u>T</u> · <u>T</u> -3'	57.9	-4.6
KRAS-m-2	5'- <u>G</u> · <u>C</u> · <u>T</u> ·A· <u>m</u> ·T·A·G·G·A·G·T·C· <u>T</u> · <u>T</u> · <u>T</u> -3'	62.2	-0.2
KRAS-m-3	5'- <u>G</u> · <u>C</u> · <u>T</u> ·A·T·T·A·G·G·A·G· <u>m</u> ·C· <u>T</u> · <u>T</u> · <u>T</u> -3'	63.8	+1.3
KRAS-m-4	5'- <u>G</u> · <u>C</u> · <u>T</u> ·A·T·T·A·G·G·A·G·T·C· <u>m</u> · <u>T</u> · <u>T</u> · <u>T</u> -3'	58.9	-3.6
KRAS-p-1	5'- <u>G</u> · <u>C</u> · <u>p</u> ·A·T·T·A·G·G·A·G·T·C· <u>T</u> · <u>T</u> · <u>T</u> -3'	59.4	-3.0
KRAS-p-2	5'- <u>G</u> · <u>C</u> · <u>T</u> ·A· <u>p</u> ·T·A·G·G·A·G·T·C· <u>T</u> · <u>T</u> · <u>T</u> -3'	63.8	+1.4
KRAS-p-3	5'- <u>G</u> · <u>C</u> · <u>T</u> ·A·T·T·A·G·G·A·G· <u>p</u> ·C· <u>T</u> · <u>T</u> · <u>T</u> -3'	66.4	+4.0
KRAS-p-4	5'- <u>G</u> · <u>C</u> · <u>T</u> ·A·T·T·A·G·G·A·G·T·C· <u>p</u> · <u>T</u> · <u>T</u> · <u>T</u> -3'	62.3	-0.2

^a Underlined letters denote LNAs. m and p in red denote the 4AEo^mU and 4AEo^pU analogs, respectively. Other letters denote native DNAs. The dots denote PS linkages. ^b The T_m values were measured in 10 mM sodium phosphate buffer (pH 7.0) containing 100 mM NaCl. The sequence of the complementary RNA strand was 5'-r(aaa gac ucc uaa uag c)-3'. The concentrations of the duplexes were 3 μ M. All measurements were performed in triplicate, and the data are shown as average values. ^c ΔT_m represents [T_m (KRAS-m/p-1-4) - T_m (KRAS-pos)].

into the 3'-side of the gapmer rather than the 5'-side in terms of thermal stability. The same trend was also confirmed in the case of the 4AEo^pU analog since the T_m (ΔT_m) values of KRAS-p-1, 2, 3 and 4 were 59.4 (-3.0), 63.8 (+1.4), 66.4 (+4.0) and 62.3 (-0.2) °

C, respectively. Meanwhile, despite the dispersion of T_m values, each gapmer exhibited a T_m value much higher than 37 °C, suggesting that it could form stable duplexes with the target mRNA.

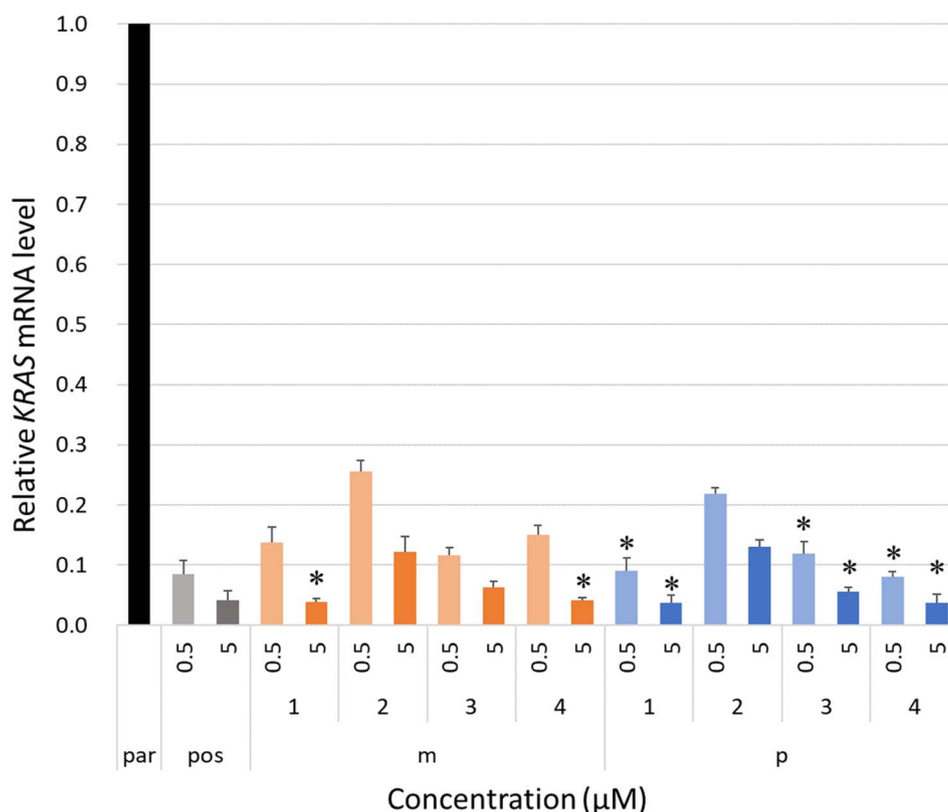


Fig. 4 Relative *KRAS* mRNA levels of each KRAS gapmer.^a NCI-H460 cells were treated with the KRAS gapmers at various final concentrations (0.5 or 5.0 μ M). Subsequently, the cells were incubated with the gapmers at 37 °C for 24 h and further incubated for 24 h after replacing the medium. Treatment only with PBS was used as a control ("par"). Total cellular mRNA was extracted, and the target *KRAS* mRNA was reverse transcribed. Quantitative real-time polymerase chain reaction (qRT-PCR) was performed in triplicate, and relative *KRAS* mRNA levels were calculated using the $\Delta\Delta C_T$ method. Results are expressed as the mean \pm SE ($n = 3$ or 4). Asterisks indicate no significant difference compared with the KRAS-pos-treated group at the corresponding concentration (* $P \geq 0.05$).



The intracellular antisense activity of KRASs was then evaluated in NCI-H460 cells. The cells were treated with KRASs at final concentrations of 0.5 and 5.0 μM without lipofection for 48 h at 37 $^{\circ}\text{C}$ with 5% carbon dioxide. The relative KRAS mRNA levels corresponding to each KRAS-treated group were calculated by qRT-PCR using the $\Delta\Delta\text{CT}$ method with beta-actin (ACTB) as a reference. A group treated only with PBS buffer was designated as the negative control, "par." As shown in Fig. 4, the 4AEo^mU-containing gapmers showed significantly lower antisense activity than KRAS-pos, except for KRAS-m-1 and KRAS-m-4 at a concentration of 5.0 μM . Conversely, KRASs containing one 4AEo^pU analog, except for KRAS-p-2, efficiently knocked down the target mRNA without any significant difference from the corresponding positive control. Incorporating these 4AEo-modified analogs at the gap-2 position negatively influenced the intracellular antisense activity, suggesting a disadvantage to Human RNase H recruitment, which is consistent with the results for *E. coli* RNase H1. Nevertheless, compared with 4AEo^mU and 4AEo^pU, the combination with 5-propynyl modification obviously improved the antisense performance. As the thermal stability and antisense activity of KRAS-p-4 and its LNA counterpart were comparable, LNA might be replaced with 4AEo^pU to mitigate LNA-associated hepatotoxicity.^{26,53}

Overall, the 4'-C- α -aminoethoxy-2'-O-methyl-5-propynyl modification exhibited excellent thermal and biochemical stability and could be tolerated while maintaining sufficient antisense activity in the 3'-wing region of the LNA gapmer, indicating its potential clinical use as an alternative to LNA.

Conclusion

To summarize, two novel uridine analogs, 4'-C- α -aminoethoxy-2'-O-methyl-5-methyl-uridine (4AEo^mU) and 4'-C- α -aminoethoxy-2'-O-methyl-5-propynyl-uridine (4AEo^pU), were designed and synthesized to expand the application of the 4AEo modification in antisense therapeutics. In the case of 4AEo^pU, the 5-propynyl moiety was introduced after the addition of the 4'-aminoethoxy side chain, providing an opportunity to combine the 4AEo modification with other base modifications in future studies. These uridine analogs could be smoothly incorporated into the DNA-based oligonucleotides using an automated DNA/RNA synthesizer in a manner similar to native uridine. In the TC-repeat modeling sequence, the presence of three 4AEo^mU analogs in the complementary DNA/RNA duplex resulted in a slight thermal destabilization, whereas the incorporation of the 4AEo^pU analogs thermally stabilized the duplex significantly. According to the results obtained from calculations of the thermodynamic parameters, CD spectra and molecular dynamics simulations, both 4AEo^mU and 4AEo^pU analogs preferred the C3'-endo sugar pucker, which is entropically favorable for the thermal stability of the DNA/RNA duplexes. Furthermore, these analog molecules exhibited slight structural fluctuations in *in silico* calculations. However, the enhanced stacking effect of the 5-propynyl moiety in the 4AEo^pU analog could overcome this structural disadvantage, resulting in favorable enthalpy and much higher thermal stability. The

difference between these novel uridine analogs in terms of thermal stability was attributed to the 5-propynyl modification. Combination with the 5-propynyl modification also improved the base discrimination ability by reducing the affinity with the complementary RNA strand containing a single guanosine mismatch. From a biochemical perspective, the novel uridine analogs used in this study showed either robust nuclease resistance or efficient *E. coli* RNase H activation. These results encouraged us to prepare LNA-wing antisense gapmers containing one 4AEo^mU or 4AEo^pU analog and to confirm their effects on cellular antisense activity. Incorporating one 4AEo^mU analog in the wing region resulted in a significant decrease in thermal stability and antisense activity, whereas in the gap region, the changes were modest. In contrast, the 4AEo^pU analog was as thermally stable as its LNA counterpart in the 3'-wing region, and no negative effects were observed in antisense activity except for KRAS-p-2.

Owing to its excellent thermal stability, the novel 4AEo^pU analog could be an alternative to LNA, providing insight into the design of antisense gapmers containing the 4AEo^pU wings for further evaluation. This analog was also expected to reduce the number of phosphorothioate linkages surrounding the 4AEo^pU analog to avoid additional interactions with undesirable proteins while maintaining robust nuclease resistance. This approach might be suitable for splicing-switching ASOs. Therefore, the 4'-C- α -aminoethoxy-2'-O-methyl-5-propynyl-modified nucleoside analogs are potential candidates as antisense therapeutics.

Experimental section

General remark

All chemicals and dry solvents (DCM, DMF, MeCN, THF, and pyridine) were obtained from commercial sources and used without any further purification. Thin layer chromatography (TLC) was performed on silica gel plates pre-coated with a fluorescent indicator and visualization by UV light or by dipping into a solution of 5% (v/v) concentrated H₂SO₄ in a mixture of *p*-anisaldehyde and methanol, followed by heating. Silica gel (63–210 mesh) was used for column chromatography. ¹H NMR (400 or 500 MHz), ¹³C {¹H} NMR (101 MHz), and ³¹P NMR (162 MHz) were recorded on 400 or 500 MHz NMR equipment. CDCl₃ or DMSO-*d*₆ was used as a solvent to obtain the NMR spectra. Chemical shifts (δ) are given in parts per million (ppm) from CDCl₃ (7.26 ppm) and DMSO-*d*₆ (2.50 ppm) for ¹H NMR spectra, and CDCl₃ (77.2 ppm) and DMSO-*d*₆ (39.52 ppm) for ¹³C NMR spectra. The abbreviations s, d, t, q, and m signify singlet, doublet, triplet, quadruplet, and multiplet, respectively. High-resolution mass spectra (HRMS) were obtained in the positive ion electrospray ionization (ESI-TOF) mode. The purity of each oligonucleotide synthesized in this study was confirmed by MALDI-TOF/MS.

3'-O-(*tert*-Butyldimethylsilyl)-4',5'-didehydro-5'-deoxy-2'-O-methyl-5-methyl-uridine (4)

Under an argon atmosphere, triphenylphosphine (2.31 g, 8.81 mmol), imidazole (21.00 g, 14.69 mmol) and iodine (2.23 g, 8.81



mmol) were added to the solution of compound 1 (2.01 g, 7.34 mmol) in THF (38.20 mL). After stirring at room temperature for 4 hours, DBU (4.40 mL, 29.38 mmol) was added dropwise to the reaction solution and stirred at room temperature for 24 hours. The solution was then evaporated, and the resulting residue was dried overnight *in vacuo*. After that, the crude residue was dissolved in DCM (20.00 mL) under the argon atmosphere. Imidazole (1.51 g, 22.02 mmol) and *tert*-butyldimethylsilyl chloride (3.33 g, 22.09 mmol) were added to the reaction solution and stirred at room temperature for 3 hours. The reactant was then quenched by saturated NaHCO₃ aq. and extracted with ethyl acetate and water. The organic layer was washed with brine, dried over Na₂SO₄, filtered, and then concentrated *in vacuo*. The resulting residue was purified by column chromatography (33% ethyl acetate in hexane) to afford the desired product 4 (2.01 g, 5.44 mmol, 74%). ¹H NMR (400 M Hz, CDCl₃) δ: 9.02 (s, 1H), 6.98 (d, *J* = 1.4 Hz, 1H), 6.07 (d, *J* = 3.7 Hz, 1H), 4.60–4.59 (m, 2H), 4.33 (q, *J* = 1.2 Hz, 1H), 3.83 (t, *J* = 4.4 Hz, 1H), 3.49 (s, 3H), 1.94 (d, *J* = 1.4 Hz, 3H), 0.93 (s, 9H), 0.14 (d, *J* = 1.8 Hz, 6H); ¹³C {¹H} NMR (101 M Hz, CDCl₃) δ: 163.7, 160.7, 150.1, 134.9, 111.8, 88.8, 86.7, 82.2, 69.9, 58.4, 25.8, 18.4, 12.8, −4.5, −4.6; HRMS (ESI-TOF) *m/z* calcd for C₁₇H₂₈N₂NaO₅Si [M + Na]⁺, 391.1665, found 391.1682.

3'-O-(*tert*-Butyldimethylsilyl)-2'-O-methyl-5-methyl-4'-C-α-trifluoroacetyl aminoethoxy-uridine (6)

Compound 4 (2.01 g, 5.44 mmol) was dissolved in a mixture of DCM (30.00 mL), acetone (30.00 mL) and saturated NaHCO₃ aq. (30.00 mL), followed by cooling in an ice bath. Oxone® (2.48 g, 16.32 mmol) in water (20.00 mL) was added to the reaction solution. After stirring at room temperature for 2 hours, the reactant was extracted with chloroform and water. The organic layer was washed with brine, dried by Na₂SO₄, filtered, and then concentrated *in vacuo*. The crude residue was dissolved in THF (20.00 mL) under the argon atmosphere. *N*-Trifluoroacetyl aminoethanol (1.83 mL, 16.32 mmol) was then added and immediately stirred at −60 °C for at least 10 minutes. After adding zinc chloride (1.0 M solution in THF; 5.40 mL, 5.40 mmol), the reaction solution was slowly brought to room temperature and stirred for 24 hours. The reactant was then quenched by saturated NaHCO₃ aq. and filtered by Celite®. The resulting filtrate was extracted with ethyl acetate and water. The organic layer was washed with brine, dried over Na₂SO₄, filtered, and then concentrated *in vacuo*. The resulting residue was purified by column chromatography (50% ethyl acetate in hexane) to afford the desired product 6 (0.86 g, 1.60 mmol, 29%). ¹H NMR (400 M Hz, DMSO-D₆) δ: 11.41 (s, 1H), 9.22 (bs, 1H), 7.67 (s, 1H), 6.01 (d, *J* = 5.5 Hz, 1H), 5.40 (t, *J* = 5.5 Hz, 1H), 4.44 (d, *J* = 6.0 Hz, 1H), 3.93 (t, *J* = 5.7 Hz, 1H), 3.76–3.66 (m, 2H), 3.49 (dq, *J* = 31.1, 5.8 Hz, 2H), 3.30 (s, 5H), 1.78 (s, 3H), 0.88 (s, 9H), 0.08 (d, *J* = 2.3 Hz, 6H); ¹³C {¹H} NMR (101 M Hz, DMSO-D₆) δ: 163.7, 150.5, 136.0, 109.9, 106.7, 86.3, 81.5, 71.1, 62.0, 60.2, 58.2, 40.1, 40.0, 25.7, 18.1, 12.3, −4.7, −5.0; HRMS (ESI-TOF) *m/z* calcd for C₂₁H₃₄F₃N₃NaO₈Si [M + Na]⁺, 564.1965, found 564.1963.

3'-O-(*tert*-Butyldimethylsilyl)-5'-O-(4,4'-dimethoxytrityl)-2'-O-methyl-5-methyl-4'-C-α-trifluoroacetyl aminoethoxy-uridine (7)

Under an argon atmosphere, *N,N*-diisopropylethylamine (0.70 mL, 4.00 mmol) and 4,4'-dimethoxytrityl chloride (0.67 g, 1.98 mmol) were added to the solution of compound 6 (0.87 g, 1.60 mmol) in pyridine (8.70 mL). After stirring overnight at room temperature, the reactant was quenched by saturated NaHCO₃ aq. and extracted with ethyl acetate and water. The organic layer was washed with brine, dried over Na₂SO₄, filtered, and then concentrated *in vacuo*. The resulting residue was purified by column chromatography (50% ethyl acetate in hexane) to afford the desired product 7 (1.06 g, 1.26 mmol, 79%). ¹H NMR (500 M Hz, CDCl₃) δ: 8.19 (s, 1H), 7.64 (s, 1H), 7.43 (s, 1H), 7.35 (d, *J* = 6.9 Hz, 2H), 7.31–7.22 (m, 7H), 6.83 (q, *J* = 4.6 Hz, 4H), 6.00 (s, 1H), 4.60 (d, *J* = 6.3 Hz, 1H), 3.79 (d, *J* = 1.1 Hz, 6H), 3.75–3.71 (m, 2H), 3.63–3.57 (m, 2H), 3.55 (s, 3H), 3.54–3.50 (m, 1H), 3.42–3.36 (m, 1H), 3.23 (d, *J* = 9.7 Hz, 1H), 1.26 (s, 3H), 0.80 (s, 9H), 0.05 (s, 3H), −0.04 (s, 3H); ¹³C {¹H} NMR (101 M Hz, CDCl₃) δ: 163.8, 159.1, 150.0, 143.8, 135.2, 134.8, 130.4, 130.3, 128.4, 128.2, 127.6, 113.4, 113.4, 111.6, 106.7, 90.1, 87.5, 84.1, 71.4, 61.7, 60.6, 60.0, 55.4, 39.9, 25.7, 18.2, 11.6, −4.4, −4.8; HRMS (ESI-TOF) *m/z* calcd for C₄₂H₅₂F₃N₃NaO₁₀Si [M + Na]⁺, 866.3272, found 866.3275.

5'-O-(4,4'-Dimethoxytrityl)-2'-O-methyl-5-methyl-4'-C-α-trifluoroacetyl aminoethoxy-uridine (8)

Under an argon atmosphere, tetrabutylammonium fluoride (1.0 M solution in THF, 0.48 mL, 0.48 mmol) was added to a solution of compound 7 (0.20 g, 0.24 mmol) in THF (4.30 mL). After stirring at room temperature for 1 hour, the reactant was quenched with saturated NaHCO₃ aq. and extracted with ethyl acetate and water. The organic layer was washed with brine, dried over Na₂SO₄, filtered and concentrated *in vacuo*. The resulting residue was purified by column chromatography (67% ethyl acetate in hexane) to afford the desired product 8 (0.15 g, 0.21 mmol, 86%). ¹H NMR (400 M Hz, CDCl₃) δ: 8.78 (s, 1H), 7.52 (d, *J* = 0.9 Hz, 1H), 7.39–7.35 (m, 2H), 7.32–7.9 (m, 10H), 6.84 (d, *J* = 8.7, 4H), 5.89 (s, 1H), 4.70 (dd, *J* = 11.2, 6.6 Hz, 1H), 3.83–3.79 (m, 7H), 3.75–3.70 (m, 2H), 3.65 (s, 3H), 3.50–3.45 (m, 2H), 3.37 (d, *J* = 10.1 Hz, 1H), 3.26 (dd, *J* = 14.2, 10.5 Hz, 1H), 2.94 (d, *J* = 11.4 Hz, 1H), 1.34 (d, *J* = 0.9 Hz, 3H); ¹³C {¹H} NMR (101 M Hz, CDCl₃) δ: 163.7, 159.0, 149.9, 144.0, 135.0, 134.9, 130.2, 128.3, 128.2, 127.5, 113.5, 111.6, 106.2, 89.9, 87.5, 84.2, 70.4, 61.4, 61.2, 60.3, 55.4, 39.4, 11.8, 1.2; HRMS (ESI-TOF) *m/z* calcd for C₃₆H₃₈F₃N₃NaO₁₀ [M + Na]⁺, 752.2407, found 752.2430.

3'-O-[2-Cyanoethoxy(diisopropylamino)phosphino]-5'-O-(4,4'-dimethoxytrityl)-2'-O-methyl-5-methyl-4'-C-α-trifluoroacetyl aminoethoxy-uridine (9)

Under an argon atmosphere, *N,N*-diisopropylethylamine (0.88 mL, 5.04 mmol) and 2-cyanoethyl-*N,N*-diisopropylchlorophosphoroamidite (0.45 mL, 2.00 mmol) were added to a solution of compound 8 (0.73 g, 1.00 mmol) in THF (7.30 mL). After stirring at room temperature for 2 hours, the



reactant was quenched by saturated NaHCO₃ aq. and extracted with ethyl acetate and water. The organic layer was washed with brine, dried over Na₂SO₄, filtered, and then concentrated *in vacuo*. The resulting residue was purified by column chromatography (50% ethyl acetate in hexane) to afford the desired product **9** (0.51 g, 0.55 mmol, 55%). ³¹P NMR (162 M Hz, CDCl₃) δ: 151.8, 151.3; HRMS (ESI-TOF) *m/z* calcd for C₄₅H₅₅F₃N₅NaO₁₁P [M + Na]⁺, 952.3486, found 952.3458.

2'-O-Methyl-4'-C-α-trifluoroacetylaminooxy-uridine (11)

Under an argon atmosphere, tetrabutylammonium fluoride (1.0 M solution in THF, 9.80 mL, 9.80 mmol) was added to a solution of compound **10** (2.57 g, 4.87 mmol) in THF (88.00 mL). After stirring at room temperature for 1 hour, the reactant was quenched with saturated NaHCO₃ aq. and extracted with ethyl acetate and water. The organic layer was washed with brine, dried over Na₂SO₄, filtered and concentrated *in vacuo*. The resulting residue was purified by column chromatography (9% methanol in chloroform) to afford the desired product **11** (1.92 g, 4.64 mmol, 95%). ¹H NMR (400 MHz, DMSO-D₆) δ: 11.39 (s, 1H), 9.25 (s, 1H), 7.79 (d, *J* = 8.2 Hz, 1H), 5.96 (d, *J* = 4.1 Hz, 1H), 5.67 (d, *J* = 8.2 Hz, 1H), 5.34 (t, *J* = 5.5 Hz, 1H), 4.67 (d, *J* = 9.2 Hz, 1H), 4.29–4.25 (m, 1H), 3.83–3.80 (m, 1H), 3.70–3.54 (m, 3H), 3.43 (q, *J* = 5.8 Hz, 1H), 3.35 (s, 4H); ¹³C{¹H} NMR (101 M Hz, CDCl₃) δ: 163.0, 150.3, 140.4, 106.3, 102.2, 87.0, 81.9, 79.2, 69.6, 60.8, 60.1, 58.1; HRMS (ESI-TOF) *m/z* calcd for C₁₄H₁₈F₃N₃NaO₈ [M + Na]⁺, 436.0944, found 436.0956.

2'-O-Methyl-5-(1-propynyl)-4'-C-α-trifluoroacetylaminooxy-uridine (13)

Under an argon atmosphere, diammonium cerium(IV) nitrate (1.14 g, 2.08 mmol) and iodine (0.64 g, 2.52 mmol) were added to a solution of compound **11** (2.18 g, 5.28 mmol) in MeCN (40.00 mL). After stirring at reflux at 80 °C for 1 hour, the reactant was quenched with methanol, and the solvent was evaporated. The resulting residue was then dried *in vacuo* for 2 hours. After that, the crude residue was dissolved in DMF (10 mL) under argon atmosphere. Bis(triphenylphosphine) palladium(II) dichloride (0.34 g, 0.49 mmol), copper(I) iodide (0.19 g, 0.98 mmol), triethylamine (1.40 mL, 9.80 mmol) and propyne (1.0 M solution in DMF; 15.00 mL, 15.00 mmol) were added to the reaction solution. After stirring overnight at 35 °C, the reactant was quenched with methanol, and the solvent was evaporated. The resulting residue was purified by column chromatography (9% methanol in chloroform) to afford the desired product **13** (1.89 g, 4.18 mmol, 79%).

¹H NMR (400 MHz, DMSO-D₆) δ: 11.65 (s, 1H), 9.24 (s, 1H), 8.08 (s, 1H), 5.92 (d, *J* = 3.7 Hz, 1H), 5.45 (t, *J* = 5.5 Hz, 1H), 4.65 (d, *J* = 9.2 Hz, 1H), 4.28–4.24 (m, 1H), 3.87–3.84 (m, 1H), 3.68 (d, *J* = 10.1 Hz, 1H), 3.59 (q, *J* = 5.2 Hz, 2H), 3.44 (q, *J* = 5.5 Hz, 1H), 3.35 (s, 4H), 1.98 (s, 3H); ¹³C{¹H} NMR (101 M Hz, DMSO-D₆) δ: 161.7, 149.3, 142.4, 117.4, 106.5, 99.5, 89.4, 87.4, 82.0, 71.9, 69.4, 60.6, 60.1, 58.2, 4.1; HRMS (ESI-TOF) *m/z* calcd for C₁₇H₂₀F₃N₃NaO₈ [M + Na]⁺, 474.1100, found 474.1103.

5'-O-(4,4'-Dimethoxytrityl)-2'-O-methyl-5-(1-propynyl)-4'-C-α-trifluoroacetylaminooxy-uridine (14)

Under an argon atmosphere, *N,N*-diisopropylethylamine (1.00 mL, 5.73 mmol) and 4,4'-dimethoxytrityl chloride (1.36 g, 4.01 mmol) were added to a solution of compound **13** (0.80 g, 2.00 mmol) in pyridine (9.00 mL). After stirring at 50 °C for 24 hours, the reactant was quenched by saturated NaHCO₃ aq. and extracted with ethyl acetate and water. The organic layer was washed with brine, dried over Na₂SO₄, filtered, and then concentrated *in vacuo*. The resulting residue was purified by column chromatography (60% ethyl acetate in hexane) to afford the desired product **14** (1.10 g, 1.46 mmol, 73%). ¹H NMR (400 MHz, CDCl₃) δ: 9.23 (s, 1H), 7.87 (s, 1H), 7.42 (d, *J* = 6.9 Hz, 2H), 7.34–7.20 (m, 8H), 6.84 (d, *J* = 8.2 Hz, 4H), 5.81 (s, 1H), 4.70 (q, *J* = 6.0 Hz, 1H), 3.86 (d, *J* = 6.4 Hz, 1H), 3.79 (s, 6H), 3.75–3.67 (m, 2H), 3.65 (s, 3H), 3.47–3.43 (m, 1H), 3.42 (s, 2H), 3.29–3.22 (m, 1H), 3.02 (d, *J* = 11.4 Hz, 1H), 1.81 (s, 1H), 1.60 (s, 3H); ¹³C{¹H} NMR (101 M Hz, CDCl₃) δ: 162.0, 158.8, 149.1, 144.3, 140.9, 135.3, 135.1, 130.1, 130.0, 128.2, 127.9, 127.1, 113.5, 106.4, 101.4, 91.7, 90.1, 87.4, 84.0, 70.2, 69.7, 61.4, 61.2, 60.3, 55.4, 39.3, 4.5; HRMS (ESI-TOF) *m/z* calcd for C₃₈H₃₈F₃N₃NaO₁₀ [M + Na]⁺, 776.2407, found 776.2398.

3'-O-[2-Cyanoethoxy(diisopropylamino)phosphino]-5'-O-(4,4'-dimethoxytrityl)-2'-O-methyl-5-(1-propynyl)-4'-C-α-trifluoroacetylaminooxy-uridine (15)

Under an argon atmosphere, *N,N*-diisopropylethylamine (2.00 mL, 11.45 mmol) and 2-cyanoethyl-*N,N*-diisopropylchlorophosphoramidite (1.00 mL, 4.48 mmol) were added to a solution of compound **14** (1.88 g, 2.50 mmol) in THF (19.00 mL). After stirring at room temperature for 2 hours, the reactant was quenched by saturated NaHCO₃ aq. and extracted with ethyl acetate and water. The organic layer was washed with brine, dried over Na₂SO₄, filtered, and then concentrated *in vacuo*. The resulting residue was purified by column chromatography (60% ethyl acetate in hexane) to afford the desired product **15** (1.72 g, 1.80 mmol, 72%). ³¹P NMR (162 M Hz, CDCl₃) δ: 151.2, 151.1; HRMS (ESI-TOF) *m/z* calcd for C₄₇H₅₅F₃N₅NaO₁₁P [M + Na]⁺, 976.3486, found 976.3514.

Solid-phase oligonucleotide synthesis

The synthesis was carried out with an automated DNA/RNA synthesizer using the solid-phase phosphoramidite method. After the synthesis, oligonucleotides containing 4AEO modification were treated with a mixture of dimethylamine/MeCN (1 : 9, v/v) for 5 minutes, followed by rinsing with MeCN to selectively remove the cyanoethyl groups. After that, the DNA oligonucleotides were treated with 28% aqueous ammonium solution for 12–14 hours at 55 °C. In contrast, the RNA oligonucleotides were cleaved from CPG beads and deprotected by treatment with 28% aqueous ammonium solution/40% methylamine (1 : 1, v/v) for 10 minutes at 65 °C. Then, 2'-O-TBDMS groups in RNA oligonucleotides were removed by Et₃N·3HF (125 μL) in DMSO (100 μL) for 1.5 hours at 65 °C. The reaction was quenched with 0.1 M TEAA buffer (pH 7.0), and the



mixture was desalted using a Sep-Pak C18 cartridge. All oligonucleotides were purified by 20% PAGE containing 7 M urea to give highly purified ones. The analysis of each synthesized oligonucleotide in this study was carried out with MALDI-TOF/MS, and the results are shown in Table S1.†

T_m measurement

The solution containing 3.0 μM DNA/RNA duplexes was prepared by mixing each DNA with an equal amount of cRNA1 in a buffer of 10 mM sodium phosphate (pH 7.0) containing 100 mM NaCl. The solution was heated at 90–100 °C and allowed to stand for at least 1.5 hours at room temperature to cool down gradually. Thermally induced transitions were monitored at 260 nm with a UV/vis spectrometer fitted with a temperature controller in quartz cuvettes with a path length of 1.0 cm. The sample temperature was increased by 0.5 °C min^{-1} . The thermodynamic parameters of the duplexes on duplex formation were determined by calculations based on the slope of a $1/T_m$ vs. $\ln(C_T/4)$ plot, where C_T (1, 3, 6, 12, 15, 21, 30 and 60 μM) is the total concentration of the single strands.

CD spectroscopy

All CD spectra were recorded at 25 °C. The following instrument settings were used: resolution, 1 nm; response, 1.0 s; speed, 50 nm min^{-1} ; accumulation, 10.

Molecular dynamics simulation

The initial geometries of the native thymidine, 4'-C- α -aminoethoxy-2'-O-methyl-5-methyl-uridine and 4'-C- α -aminoethoxy-2'-O-methyl-5-propynyl-uridine were modeled by a molecular modeling software, GaussView. The geometry of each nucleoside was optimized, and the electrostatic potential (ESP) of each nucleoside was calculated using Gaussain16 at the HF/6-31G* level of theory.⁵⁴ The restrained ESP (RESP) charges were derived using the Antechamber RESP fitting procedure.⁵⁵ The generalized AMBER force field (gaff) was used for the native thymidine and modified nucleosides.⁵⁶ Each of the nucleosides was solvated in a cubic box with the TIP3P water model, and the size of the cubic box was determined to be 42.0 Å (in length).⁵⁷ Chloride ions were added to the solution to neutralize the charge of the systems. The energy minimization calculation was performed using the SANDER module of AMBER18. The solute molecule was optimized by 2500 steps of the steepest descent method, followed by 2500 steps of the conjugate gradient method by applying a restraint force constant of 2.0 $\text{kcal mol}^{-1} \text{Å}^{-2}$ to the nucleosides. Then, the calculation of the entire system was carried out using 5000 steps of the conjugate gradient method without any restraints. The minimized structure assigned to the atom type and calculated RESP charges for each nucleoside are shown in Fig. S7.† After that, the temperature of the system was heated gradually from 0 K to 300 K over a period of 100 ps of *NVT* dynamics with a restraint force constant of 2.0 $\text{kcal mol}^{-1} \text{Å}^{-2}$ to the nucleoside, followed by 100 ps of *NPT* equilibration at 300 K and 1 bar pressure with the same restraint force. After the equilibration phase, 500 ns MD simulations were performed in an *NPT* ensemble at 300 K and 1

bar pressure without any restraints. During the MD simulations, a time step of 2 fs was used. The cut-off distance towards the van der Waals force was 10 Å, the SHAKE algorithm was used to constrain all bonds involving hydrogen atoms, and the Particle Mesh Ewald (PME) method was used for long-range electrostatic interaction.^{58,59} The temperature was regulated by Langevin dynamics with a collision frequency of 2.0 ps^{-1} .⁶⁰ All the MD simulations were performed using the PMEMD module of AMBER18. Trajectory analysis was done for 50 000 snapshots, every 10 ps, sampled during the full 500 ns simulation.

Nuclease resistance of single-strand DNAs

Fluorescein-labeled DNAs (300 pmol) were dissolved in OPTI-MEM (37 μL) and used for the serum stability test. 1.0 μL of the oligomer solution was diluted in 10 μL stop solution (10% formamide in 10 mM EDTA) as the control sample (0 min). Then, bovine serum was added to achieve various final concentrations (3%, 10%, 25% or 50%; v/v), and subsequently, the mixture was incubated at 37 °C for the required time. Aliquots of 2.0 μL were diluted with a 10 μL stop solution. Samples were subjected to electrophoresis in 20% PAGE containing 7 M urea and quantified by Luminescent Image analyzer LAS-4000 (Fujifilm).

E. coli RNase H activation

The solution of DNA/RNA duplexes used for the RNase H assay was prepared by mixing the native or modified DNAs (600 pmol) with fluorescein-labeled cRNA-2 (3000 pmol) in 75 μL of 50 mM Tris-HCl (pH 8.0) containing 75 mM KCl, 3 mM MgCl_2 and 10 mM dithiothreitol, followed by heating at 90–100 °C for 5 min and cooling gradually to room temperature. Then, 70 μL diluted RNase H solution (60 units per L in H_2O) was added, and subsequently, the mixture was incubated at 37 °C for the required time. Aliquots of 5 μL were diluted with 100% formamide (10 μL). Samples were subjected to electrophoresis in 20% PAGE containing 7 M urea and quantified by a Luminescent Image analyzer LAS-4000 (Fujifilm).

In vitro antisense activity of KRAS-gapmers

NCI-H460 cells, transfected with *KRAS* and *ACTB* genes, were used for the gene expression suppression assays, and the *KRAS* & *ACTB* primer probes for qPCR were provided by Professor Tokuhirro Chano (Shiga University of Medical Science). Cells were cultured in D-MEM containing 10% fetal bovine serum (FBS), penicillin (50 units per mL) and streptomycin (50 mg mL^{-1}). Cells were incubated at 37 °C in a humidified chamber supplemented with 5% carbon dioxide. NCI-H460 cells were plated into 96-well plates at 400 cells per well in 50 μL D-MEM containing 10% FBS. Then, the *KRAS*-targeting antisense gapmers were added into specified wells at the final concentration of 0.5 or 5.0 μM . After a 24 hours incubation at 37 °C with 5% carbon dioxide, an additional 50 μL medium was added, followed by a further 24 hours incubation. Reverse transcription was directly performed in each of the 96-well plates using a TaqMan™ Fast Advanced Cells to CT™ Kit (Thermo Fisher Scientific). Treatment without gapmers was



used as a control, named “par”. Quantitative real-time polymerase chain reaction (qRT-PCR) of the targeted *KRAS* mRNA was performed using a TaqMan® Fast Advanced master mix and StepOnePlus™ Real-Time PCR system (Thermo). Each PCR reaction was performed in duplicate, and the relative *KRAS* mRNA levels were calculated by the $\Delta\Delta CT$ method using beta-actin (ACTB) as a reference.

Data availability

The authors confirm that the data supporting the findings of this study are available within the article and/or its ESI.†

Author contributions

Yujun Zhou: investigation, data curation, writing – original draft. Hitotaka Sato: data curation. Miwa Kawade: data curation. Kenji Yamagishi: data curation. Yoshihito Ueno: writing – review & editing, funding acquisition, conceptualization.

Conflicts of interest

There are no conflicts to declare.

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