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Synthesis and properties of DNA oligomers containing stereopure phosphorothioate linkages and C-5 modified deoxyuridine derivatives†

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Phosphorothioate (PS) modification, where a non-bridging oxygen atom in a phosphodiester linkage is replaced by a sulfur atom, is widely used to improve the properties of nucleic acid drugs. Each PS linkage can be found in two stereoisomers, Rp and Sp. Since one non-bridging oxygen or sulfur atom in Sp-PS or Rp-PS, respectively, is located close to the C-5 substituent of uracil in a DNA/RNA hybrid duplex, the combination of the stereochemistry of the PS linkages and the type of the C-5 modification of uracil bases is expected to affect the properties of the hybrid duplexes. Herein, DNA oligomers containing both stereopure phosphorohioate linkages and C-5 modified deoxyuridine derivatives were synthesized. The thermodynamic stability of the DNA/RNA and DNA/DNA duplexes and RNase H activity of the DNA/RNA duplexes were evaluated. The combination of 5-propynyluracil and (Rp)-PS linkages in a DNA strand could significantly increase the thermal stability of a DNA/RNA hybrid duplex without reducing its RNase H activity.

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Introduction

Nucleic acid therapeutics, namely drugs based on nucleic acid molecules, such as DNA, RNA, and their derivatives, have been an attractive field for drug development for several decades. Nucleic acid drugs are classified into various categories, including antisense oligonucleotides (ASOs), small interfering RNAs (siRNAs), and aptamers. Particularly, the recent progress in ASOs has been so impressive that an increasing number of ASOs have been approved as therapeutics since 2010.¹

ASOs are classified into two types, the ribonuclease H (RNase H)-dependent and the RNase H-independent ASOs, according to their mechanism of action after the duplex formation.² Although both types of ASOs bind to a complementary region of the target RNA, *e.g.*, mRNA, to form an ASO/RNA hybrid duplex, the RNase H-dependent ASOs induce the cleavage of the target RNAs by RNase H after the formation of the hybrid duplex. The RNase H-dependent ASOs contain several consecutive DNA regions and RNase H can recognize the DNA/RNA hybrid duplex structures in the ASO/RNA hybrid, thus cleaving the RNA strand.²⁻⁵ In contrast, the RNase H-independent ASOs bind to

Since nucleic acid molecules are susceptible to cleavage by endogenous nucleases *in vivo*, their chemical modification is essential for therapeutic application. Among them, phosphorothioate (PS) modification, where a non-bridging oxygen atom in a phosphodiester linkage is replaced by a sulfur atom, is the most employed modification for P-atoms. PS modification improves the properties of ASOs, including nuclease stability and pharmacokinetics, ^{2,6} and has already been adopted for the development of several ASO drugs. Although the PS modification seems to be established for the chemical modification of ASOs, it should be noted that the P-atom in PS linkages is chiral different from that in phosphodiester (PO)-linkages, and the (*R*p)-PS and (*S*p)-PS diastereomers exhibit completely different physicochemical and biological properties (Fig. 1).⁷⁻¹²

[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/d0ra06970a

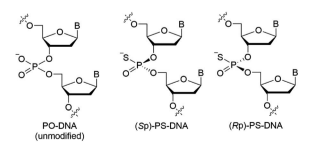


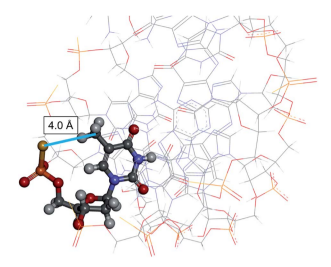
Fig. 1 Structures of PO-DNA and the two stereoisomers of PS-DNA.

the target RNA without cleaving the target RNA, but they sterically block some biological processes by the duplex formation.

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In an ASO/RNA duplex, which is expected to form an A-type duplex structure like DNA/RNA and RNA/RNA duplexes, a non-bridging oxygen atom of a phosphodiester linkage in the DNA strand is close to the 5-H atom/5-CH₃ group of pyrimidine nucleobases or to the 8-H atom of purine nucleobases in the DNA strand. Accordingly, in (*R*p)-PS modified DNA, the sulfur atom is close to the hydrogen atom or to the methyl group (Fig. 2),¹³ whereas in (*S*p)-PS modified DNA, the non-bridging oxygen atom is close to the hydrogen atom or the methyl group. Therefore, the different location of the kind of non-bridging atoms is expected to affect the interactions between the non-bridging atoms and the nucleobase-derived substituents, as well as the properties of the nucleic acid duplexes.

Regarding the interactions between the non-bridging sulfur or oxygen atoms and nucleobases, C5-modified pyrimidine nucleobases have possibilities to exhibit discriminating features when combined with stereoregulated PS-DNAs. C5-Modified deoxyuridine derivatives have been studied to improve the properties of nucleic acids such as the affinity to their complementary RNAs and the nuclease resistance. 14-29 C5-Modified deoxycytidine derivatives have also been explored for similar purposes, 21,24,26 while 5-methyl cytosine has been used to inhibit immune responses derived from CpG sequences. 30,31 In this study, we focused on the combination of stereoregulated PS linkages and C-5 modified uracil nucleobases using 5-methyl

Fig. 3 5'-Modified (and unmodified) uracil bases used in this study.

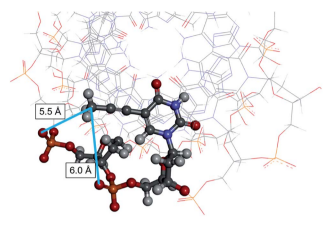


Fig. 4 3D view of the ^{pr}U-contained-DNA/RNA hybrid duplex, dG^{pr}C^{pr}U^{pr}C^{pr}U pr CPrU pr UPr C/rGAAGAGAAGC²⁸ (PDB 1007, NMR solution structure). The underlined ^{pr}U residue and two phosphate groups close to the methyl group of the ^{pr}U are displayed in ball-and-stick style. The distance between the carbon atom of the methyl group of ^{pr}U and the sulfur atom in a pro-*R*p oxygen atom in a phosphodiester linkage was 5.5 Å, and the distance between the same carbon atom and another pro-*R*p oxygen atom was 6.0 Å.

deoxyuridine, known as thymidine, 5-bromo deoxyuridine ($d^{Br}U$), 5-iodo deoxyuridine ($d^{I}U$), and 5-propynyl deoxyuridine ($d^{Pr}U$) units (Fig. 3). While the methyl, bromo and iodo substituents were expected to be close to one of the non-bridging atoms of a PS linkage in the duplex structure, the propynyl group in $d^{Pr}U$ was expected to be located close to two non-bridging atoms of two different internucleotide linkages, as shown in Fig. $4.^{28}$

In this study, DNA oligomers containing consecutive base sequences of these deoxyuridine derivatives, dCG(^xU_{PS})₈CG (d^xU represents 5-modified or unmodified deoxyuridine), were designed and synthesized in a stereoregulated manner. The properties of these DNA oligomers including duplex forming ability and RNase H activity were also evaluated.

Results and discussion

Synthesis of stereopure PO/PS chimeric DNA oligomers containing C-5 modified deoxyuridine derivatives

Oxazaphospholidine monomer units were initially prepared for the synthesis of DNA oligomers with stereoregulated PS linkages by an oxazaphospholidine approach.³²⁻³⁴ The thymidine monomers (*Rp*)- and (*Sp*)-3b were synthesized according to our previous report.³³ 5'-DMTr-protected dU (1a),³⁵ d^{Br}U (1c),³⁶ and d^IU (1d)³⁶ were prepared following already reported procedures, while 5'-DMTr-protected d^{Pr}U (1e) was synthesized from 1d *via* the Sonogashira coupling.³⁷ 1a, 1c, 1d, and 1e were then used to prepare the corresponding oxazaphospholidine monomers (3a, 3c, 3d, and 3e) by the conventional method.^{33,38} All the *Sp*- and *Rp* monomers were synthesized in moderate yields with high stereopurities (Table 1).

Subsequently, ten DNA 12mers containing eight consecutive U or modified-U sequences (dCG(^xU_{PS})₈CG) were synthesized based on the automated stereocontrolled solid-phase synthesis

Table 1 Synthesis of the oxazaphospholidine monomer units

Entry	Compound	^x U	Yield (%)	$\mathrm{dr}^a\left(R\mathrm{p}:S\mathrm{p}\right)$	Purity ^a (%)
1	(Rp)-3a	U	45	>99:1	89
2	(<i>R</i> p)-3b	T	60	>99:1	98
3	(Rp)-3c	$^{\mathrm{Br}}\mathrm{U}$	41	>99:1	97
4	(<i>R</i> p)-3d	$^{\mathrm{I}}\mathrm{U}$	42	>99:1	97
5	(<i>R</i> p)-3e	$^{ m pr}$ U	42	>99:1	>99
6	(Sp)-3a	U	57	>1:99	95
7	(Sp)-3b	T	44	>1:99	>99
8	(Sp)-3c	$^{\mathrm{Br}}\mathrm{U}$	49	>1:99	99
9	(Sp)-3d	$^{\mathrm{I}}\mathrm{U}$	50	>1:99	96
10	(Sp)-3e	$^{\mathrm{pr}}\mathrm{U}$	53	>1:99	98

of (PO/PS) chimeric DNA oligomers (Fig. 5).³⁴ Eight PS modification sites were designed to all the *R*p or *S*p configurations in the respective DNA oligomers. For the synthesis of DNA containing halouracil derivatives (d^{Br}U and d^IU), dC^{ac}-, and dG^{ipr-pac}-phosphoramidites and CPG-dG^{ipr-pac} were used instead of dC^{bz}- and dG^{ibu}-phosphoramidites and CPG-dG^{ibu}, for deprotection under milder conditions in order to avoid 5-amination by ammonia.³⁹ All the DNA oligomers (*Rp*)- or (*Sp*)-4a-4e were successfully synthesized as shown in Table 2.

PO/PS Chimeric DNA 4. deprotection & release conc Nity-ECIDH (St., v/v) 25° C(one Nity-ECIDH

Fig. 5 Automated solid-phase synthesis of PO/PS chimeric DNA oligomers in a stereoregulated manner.

UV melting analysis

UV melting analysis was conducted to evaluate the duplex forming ability of the synthesized DNA oligomers. The UV melting curves of the DNA oligomers and their complementary 12mer RNA, rCGA₈CG, are depicted in Fig. 6, and the melting temperatures $(T_{\rm m})$ are listed in Table 3. In all cases, the DNA/ RNA hybrid duplexes of the (Rp)-PO/PS-DNA oligomers exhibited higher T_m values than those of the (Sp)-PS/PO-DNA oligomers. These results were consistent with previous reports and the differences in the properties between the two diastereomers. 7,8,11,34,40 Moreover, the $\Delta T_{\rm m}$ values between the PO/(Rp)-PS-DNAs and the PO/(Sp)-PS-DNAs were different depending on the substituents on the C-5 position. In the case of dCG(U_{PS})₈CG (4a) and $dCG(T_{PS})_8CG$ (4b), the ΔT_m values between the PO/(Rp)-PS-DNAs and PO/(Sp)-PS-DNAs were relatively small, 2.8 °C and 2.9 °C, respectively. The halogen modification also increased the $\Delta T_{\rm m}$ values, with (Rp)- dCG(BrU_{PS})₈CG/RNA and (Rp) $dCG(^{I}U_{PS})_{8}CG/RNA$ duplexes showing higher ΔT_{m} than their corresponding (Sp)-DNA/RNA duplexes by 5.1 and 5.7 °C, respectively. In the case of PrU, the (Rp)-dCG(PrUPS)8CG/RNA duplex showed a quite high $T_{\rm m}$ value (61.8 °C, (Rp)-4e) compared to the (Sp)-dCG(PrUPS)8CG/RNA duplex (44.1 °C, (Sp)-**4e**), and its $\Delta T_{\rm m}$ value was 17.7 °C. The $T_{\rm m}$ value was even higher than that of the corresponding PO-DNA/RNA duplex (60.3 °C), while in all other DNA oligomers, the $T_{\rm m}$ values of both (Sp)- and (Rp)-DNA/RNA duplexes were by more than 10 °C lower than

Table 2 Synthesis of the PO/PS chimeric DNA oligomers

Entry	Compound	Sequence and modification	CPG	Amidite monomer	Yield/%
1	(Rp)-4a	(Rp) -d $(CG(U_{PS})_8CG)$	$\mathrm{d}G^{\mathrm{ibu}}$	dG^{ibu}, dC^{bz}	20
2	(Sp)-4a	(Sp)-d(CG(U _{PS}) ₈ CG)		,	28
3	(Rp)-4b	(Rp) -d $(CG(T_{PS})_8CG)$			19
4	(Sp)-4b	(Sp) -d $(CG(T_{ps})_8CG)$			24
5	(Rp)-4c	(Rp) -d $(CG(^{Br}U_{PS})_{8}CG)$	$dG^{ipr-pac}$	dG ^{ipr-pac} , dC ^{ac}	10
6	(Sp)-4c	(Sp)-d(CG(BrU _{PS}) ₈ CG)			2
7	(<i>R</i> p)-4d	(Rp) -d $(CG(^{I}U_{PS})_{8}CG)$			2
8	(Sp)-4d	(Sp) - $d(CG(^{I}U_{PS})_{8}CG)$			6
9	(Rp)-4e	(Rp) -d(CG($^{pr}U_{PS})_8$ CG)	$\mathrm{dG}^{\mathrm{ibu}}$	dG^{ibu} , dC^{bz}	29
10	(Sp)-4e	(Sp) - $d(CG(^{pr}U_{PS})_8CG)$		•	5

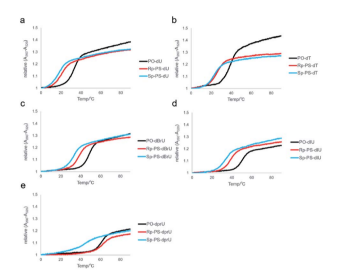


Fig. 6 UV melting curves of the hybrid duplex of $dCG(^{x}U_{PS})_{8}CG$ or $dCG(^{x}U_{PO})_{8}CG$ and $rCGA_{8}CG$. (a) $^{x}U=U$, (b) $^{x}U=T$, (c) $^{x}U=^{Br}U$, (d) $^{x}U=^{I}U$, and (e) $^{x}U=^{Pr}U$.

those of each PO-DNA/RNA duplex. These results suggested the existence of specific stabilization effects deriving from the combination of the (*R*p)-PS linkage and the propynyl group of (*R*p)-4e. In the (*R*p)-4e/RNA duplex, each propynyl group is expected to be located close to two sulfur atoms of two consecutive (*R*p)-PS-nucleotides (Fig. 4). The manners of interaction of the sulfur atom in a phosphorothioate linkage are different from those of the oxygen atom in many points;^{41,42} electrostatic interactions, hydrogen bonding, hydrophobic interactions, and hydration. Since propynyl groups have no ability to form hydrogen bonds and electrostatically interact with other moieties, the stabilization in the case of (*R*p)-4e might be derived from the hydrophobic interactions between the sulfur atoms in the PS linkages and the propynyl group.

UV melting analysis was also performed for the duplexes of the synthesized DNAs and their complementary DNA. The corresponding curves and $T_{\rm m}$ values of ${\rm dCG}(^{\rm x}{\rm U}_{\rm PS})_{\rm 8}{\rm CG}/{\rm dCGA}_{\rm 8}{\rm CG}$ are presented in Fig. 7 and Table 4. In the case of ${\rm dCG}({\rm U}_{\rm PS})_{\rm 8}{\rm CG}$ and ${\rm dCG}({\rm T}_{\rm PS})_{\rm 8}{\rm CG}$, the $T_{\rm m}$ values of ${\rm PO}/(R{\rm p})$ -PS-DNA/DNAs and ${\rm PO}/(S{\rm p})$ -PS-DNA/DNAs were slightly different than those of DNA/RNAs. Even for ${\rm dCG}(^{\rm Bt}{\rm U}_{\rm PS})_{\rm 8}{\rm CG}$ and ${\rm dCG}(^{\rm I}{\rm U}_{\rm PS})_{\rm 8}{\rm CG}$ (4c and

Table 3 $T_{\rm m}$ values of the hybrid duplex of dCG(^XU_{PS})_8CG or dCG(^XU_{PO})_8CG and rCGA_8CG

Entry	×U	PO or PS	Compound	$T_{ m m}/^{\circ}{ m C}$	$\Delta T_{ m m}/^{\circ}{ m C}$ (PS $-$ PO)	$\Delta T_{\rm m}/^{\circ}{ m C}$ $(R{ m p}-S{ m p})$
1	U	PO	_	32.7	_	2.8
2	U	(Rp)-PS	(Rp)-4a	20.6	-12.1	
3	U	(Sp)-PS	(Sp)-4a	17.8	-14.9	
4	T	PO		38.7	_	2.9
5	T	(Rp)-PS	(Rp)-4b	26.2	-12.5	
6	T	(Sp)-PS	(Sp)-4b	23.3	-15.4	
7	$^{\mathrm{Br}}\mathrm{U}$	PO		47.9	_	5.1
8	$^{\mathrm{Br}}\mathrm{U}$	(Rp)-PS	(Rp)-4c	37.8	-10.1	
9	$^{\mathrm{Br}}\mathrm{U}$	(Sp)-PS	(Sp)-4c	32.7	-15.2	
10	$^{\mathrm{I}}\mathrm{U}$	PO	_	49.9	_	5.7
11	$^{\mathrm{I}}\mathrm{U}$	(Rp)-PS	(Rp)-4d	37.9	-12.0	
12	$^{\mathrm{I}}\mathrm{U}$	(Sp)-PS	(Sp)-4d	32.2	-17.7	
13	$^{\mathrm{pr}}\mathrm{U}$	PO		60.3	_	17.7
14	$^{\mathrm{pr}}\mathrm{U}$	(Rp)-PS	(Rp)-4e	61.8	1.5	
15	^{pr} U	(Sp)-PS	(Sp)-4e	44.1	-16.2	

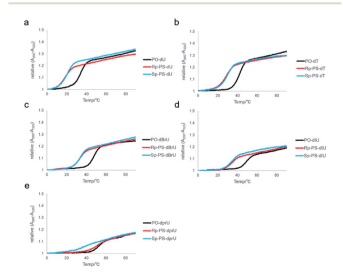


Fig. 7 UV melting curves of the duplex of $dCG(^{x}U_{PS})_{8}CG$ or $dCG(^{x}U_{PO})_{8}CG$ and $dCGA_{8}CG$. (a) $^{x}U=U$, (b) $^{x}U=T$, (c) $^{x}U=^{Br}U$, (d) $^{x}U=^{I}U$, and (e) $^{x}U=^{Pr}U$.

Table 4 $T_{\rm m}$ values of the duplex of dCG(^XU_{PS})_8CG or dCG(^XU_{PO})_8CG and dCGA $_{\rm 8}$ CG

Entry	^x U	PO or PS	Compound	$T_{ m m}/^{\circ}{ m C}$	$\Delta T_{ m m}/^{\circ}{ m C}$ (PO – PS)	$\Delta T_{\rm m}/^{\circ}{ m C} \ \left(R{ m p}-S{ m p}\right)$
1	U	РО	_	33.5	_	-1.8
2	U	(Rp)-PS	(Rp)-4a	18.0	-15.5	
3	U	(Sp)-PS	(Sp)-4a	19.8	-13.7	
4	T	PO	_	41.4	_	-0.9
5	T	(Rp)-PS	(Rp)-4b	28.4	-13.0	
6	T	(Sp)-PS	(Sp)-4b	29.3	-12.1	
7	$^{\mathrm{Br}}\mathrm{U}$	PO	_	47.3	_	-0.4
8	$^{\mathrm{Br}}\mathrm{U}$	(Rp)-PS	(Rp)-4c	33.0	-14.3	
9	$^{\mathrm{Br}}\mathrm{U}$	(Sp)-PS	(Sp)-4c	33.4	-13.9	
10	$^{\mathrm{I}}\mathrm{U}$	PO		48.7	_	0.0
11	$^{\mathrm{I}}\mathrm{U}$	(Rp)-PS	(Rp)-4d	33.2	-15.5	
12	$^{\mathrm{I}}\mathrm{U}$	(Sp)-PS	(Sp)-4d	33.2	-15.5	
13	$^{\mathrm{pr}}\mathrm{U}$	PO		51.1	_	9.7
14	$^{\mathrm{pr}}\mathrm{U}$	(Rp)-PS	(Rp)-4e	50.1	-1.0	
15	$^{\mathrm{pr}}\mathrm{U}$	(Sp)-PS	(Sp)-4e	40.4	-10.7	

4d), almost no differences were observed in the $T_{\rm m}$ values of the two isomers. Only in the case of dCG($^{\rm pr}U_{\rm PS}$) $_{\rm 8}$ CG (**4e**), the $T_{\rm m}$ value of (Rp)-dCG($^{\rm pr}U_{\rm PS}$) $_{\rm 8}$ CG/DNA was similar to that of the corresponding PO-DNA/DNA, and higher than that of the PO/(Sp)-PS-DNA/DNA duplex by 9.7 °C. The observed tendency was similar to that indicated for the DNA/RNA hybrid duplexes, suggesting also a slightly lower stabilization effect deriving from the combination between (Rp)-PS linkages and 5-propynyl groups in the DNA/DNA duplexes.

The difference in the $T_{\rm m}$ values between the two stereoisomers was lower in the DNA/DNA duplexes than in the DNA/RNA hybrid duplexes in all cases. It is possible to be due to their different duplex structures. The DNA/DNA duplexes adopt a B-type duplex structure, whereas the DNA/RNA duplexes basically form an Atype duplex. Additional variations have also been observed in several structural parameters, 43 which can be visually identified, as the A-type duplexes have a narrower and deeper major groove and a wider and shallower minor groove than B-type duplexes. These structural differences possibly affect the distance between atoms and substituents. Regarding the hydrophobic or other interactions between the substituents at the 5-position and the sulfur atom of the (Rp)-PS linkage, the distance between the sulfur atom and the carbon, bromine, or iodine atom at the 5position is expected to be longer in the DNA/DNA duplexes than in the DNA/RNA hybrids by about 1 Å (Fig. S38†),44 possibly leading to the different duplex forming abilities.

Evaluation of RNase H activity

The RNase H activity of the stereoregulated PO/PS-chimeric DNA 12mers containing $^{\rm pr}$ U bases was explored. For the evaluation experiments, **4b**, **4e**, and the corresponding PO-DNA oligomers were used. Each DNA and an excess amount of the 12mer complementary RNA were treated with *E. coli* RNase H at 20 $^{\circ}$ C for 30 min. Fig. 8b–g illustrate the HPLC profiles of each mixture after treatment with RNase H. It was previously reported that (*R*p)-PS rich DNAs show higher RNase H activity than (*S*p)-PS rich DNAs with respect to *E. coli* RNase H. 10 As shown in Fig. 8b and c,

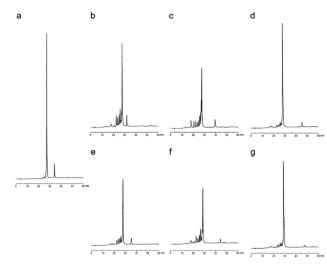


Fig. 8 (a) RP-HPLC of the mixture of dCG(Pr U)₈CG and the complementary RNA (rCGA₈CG) without RNase H treatment; (b)–(g) RP-HPLC profiles of the mixture of PO- or PO/PS chimeric-DNAs and their complementary RNA (rCGA₈CG) after treatment with 4 U/100 μ L RNase H at 20 °C for 30 min: (b) PO-dCGT₈CG, (c) (Rp)-dCG(T_{PS})₈CG (Rp-(4b)), (d) (Sp)-dCG(T_{PS})₈CG (Sp-(4b)), (e) PO-dCG(Pr U)₈CG, (f) (Rp)-dCG(Pr U)_{PS})₈CG (Rp-(4e)), (g) (Sp)-dCG(Pr U)_{PS})₈CG (Sp-(4e)). The RP-HPLC analyses (UV detection at 260 nm) were performed with a linear gradient of 0–30% CH₃CN in 0.1 M triethyl ammonium acetate (TEAA) buffer (pH 7.0) at 20 °C for 60 min with a flow rate of 0.5 mL min⁻¹.

(Rp)-4b had similar RNase H activity as PO-dCG(T_{PS})₈CG. In contrast, only small peaks of cleaved RNA fragments were observed in the presence of (Sp)-4b (Fig. 8d), complying with the previous results. Furthermore, in the case of ^{PT}U , (Rp)-4e showed high RNase H activity (Fig. 8f), similar to that of (Rp)-4b. (Sp)-4e also exhibited similar profiles to (Sp)-4b, and only small amounts of RNA was cleaved (Fig. 8g). Additional reaction conditions with higher RNase H concentrations (Fig. S39†) were also applied, but no significant differences were detected in the RNase H activity by base modifications. Therefore, it was indicated that the propynyl modification combined with the (Rp)-PS modification significantly increased the T_m values of the DNA/RNA hybrid duplexes without altering the RNase H activity.

Conclusions

This study revealed that the combination of the stereochemistry of PS linkages with C5-modifications of uracil bases on a DNA strand could strongly affect the thermal stability of DNA/RNA duplexes. In particular, the C5-propyny modification significantly increased the $T_{\rm m}$ value of a DNA/RNA hybrid duplex when combined with (Rp)-PS linkages while maintaining the RNase H activity. These results would be useful for the design of ASOs and other nucleic acid drugs with stereopure PS linkages.

Experimental section

General information

All reactions were conducted under argon atmosphere. The ¹H NMR spectra were obtained at 400 MHz on a JEOL instrument

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with tetramethylsilane as an internal standard (δ 0.0 ppm). The ¹³C NMR spectra were obtained at 100 MHz with CDCl₃ as an internal standard (δ 77.0 ppm). The ³¹P NMR spectra were obtained at 162 MHz with 85% H_3PO_4 as an external standard (δ 0.0 ppm). The PO-DNA and RNA oligomers were purchased from Japan Bio Services Co., LTD. Silica gel column chromatography was performed using silica gel 60 N (neutral silica) or Chromatorex NH-DM1020 (amino silica). Reverse phase (RP)-HPLC was performed for the analysis and purification of the synthesized analogs using a μBondasphere column (5 μm, C18, 100 Å, 3.9×150 mm) (Waters) or Source 5 RPC ST 4.6/150 (GE Healthcare). The organic solvents were dried before use. The mass spectra were recorded on a 910-MS FTMS system (Varian, ESI-MS) or a JMS-700 instrument (JEOL, FAB-MS). The nucleoside phosphoramidites (B^{pro} = C^{bz}, C^{ac}, G^{ibu}, and G^{ipr-pac}) and solid supports were purchased from Glen Research Co., Ltd.

Automated solid-phase synthesis

The DNA oligomers were synthesized using an NTS M2 DNA/ RNA synthesizer (Nihon Techno Service Co., Ltd) based on the following procedure.

		Cycle for PO		Cycle for PS		
Step	Operation	Reagents	Time	Reagents	Time	
1	Detritylation	3% (w/v) TCA in CH ₂ Cl ₂	12 s	3% (w/v) TCA in CH ₂ Cl ₂	12 s	
2	Washing	Dry CH ₃ CN	_	Dry CH ₃ CN	_	
3	Condensation	0.1 M phosphoramidite monomer and 0.25 M ETT in dry CH ₃ CN	30 s	0.1 M oxazaphospholidine monomer and 0.5 M CMPT in dry $\mathrm{CH_3CN}$		
4	Washing	Dry CH ₃ CN	_	Dry CH ₃ CN	_	
5	Capping	Ac_2O or Pac_2O and 6% (v/v) NMI in dry THF	40 s	$0.5~M~CF_3COIm~and$ $16\%~(v/v)~NMI~in~dry$ THF		
6	Washing	Dry CH ₃ CN	_	Dry CH ₃ CN	_	
7 8	Oxidation/ sulfurization Washing	1 M <i>t</i> -BuOOH in dry toluene Dry CH ₃ CN	30 s	0.3 M POS in dry CH ₃ CN Dry CH ₃ CN	8 min —	

After synthesis, each solid support was treated with a mixture of conc. NH₃(aq)-EtOH (5:1, v/v) at 50 °C for 8-12 h. In the case of oligomers containing ^{Br}U or ^IU bases (4c and 4d), each solid support was treated with conc. NH₃(aq)-EtOH (5:1, v/v) at 25 °C for 13-24 h. Then, the mixture was filtered and the filtrate was concentrated. The crude product was purified by RP-HPLC to afford pure stereoregulated PS/PO-chimeric DNA oligomers. The synthesized DNA oligomers were quantified by their UV absorbance and predicted extinction coefficients at 260 nm, 19,45 except for 4e. The quantification of 4e was performed by the high temperature UV absorbance and predicted extinction coefficients at 280 nm.23

(Rp)-4a ((Rp)-dCG $(U_{PS})_8$ CG). The automated solid-phase synthesis procedure was applied using 5'-O-DMTr-dGibu-CPG (0.5)μmol) as the solid support, (Sp)-3a as the oxazaphospholidine monomer, and dGibu and dCBz phosphoramidite monomers.

ESI-HRMS: m/z calcd for $C_{110}H_{132}N_{32}O_{70}P_{11}S_8^{5-}[M-5H]^{5-}$; 723.4532. Found; 723.4550.

(Sp)-4a ((Sp)-dCG $(U_{PS})_8$ CG). The automated solid-phase synthesis procedure was applied using 5'-O-DMTr-dGibu-CPG (0.5 µmol) as the solid support, (Rp)-3a as the oxazaphospholidine monomer, and dGibu and dCBz phosphoramidite monomers.

ESI-HRMS: m/z calcd for $C_{110}H_{132}N_{32}O_{70}P_{11}S_8^{5-}[M-5H]^{5-}$; 723.4532. Found; 723.4547.

(Rp)-4b ((Rp)-dCG $(T_{PS})_8$ CG). The automated solid-phase synthesis procedure was applied using 5'-O-DMTr-dGibu-CPG (0.5 μmol) as the solid support, (Sp)-3b as the oxazaphospholidine monomer, and dG^{ibu} and dC^{Bz} phosphoramidite monomers.

ESI-HRMS: m/z calcd for $C_{118}H_{148}N_{32}O_{70}P_{11}S_8^{5-}[M-5H]^{5-}$; 745.8782. Found; 745.8804.

(Sp)-4b $((Sp)-dCG(T_{PS})_8CG)$. The automated solid-phase synthesis procedure was applied using 5'-O-DMTr-dG^{ibu}-CPG (0.5 µmol) as the solid support, (Rp)-3b as the oxazaphospholidine monomer, and dGibu and dCbz phosphoramidite monomers.

ESI-HRMS: m/z calcd for $C_{118}H_{148}N_{32}O_{70}P_{11}S_8^{5-}$ for [M -5H]⁵⁻; 745.8782. Found; 745.8792.

(Rp)-4c ((Rp)-dCG(BrUPS)8CG). The automated solid-phase synthesis procedure was applied using 5'-O-DMTr-dGipr-pac-CPG (0.5 μ mol) as the solid support, (Sp)-3c as the oxazaphospholidine monomer, and dGipr-pac and dCac phosphoramidite monomers.

ESI-HRMS: m/z calcd for $C_{110}H_{124}Br_8N_{32}O_{70}P_{11}S_8^{5-}$ [M -5H]⁵⁻; 850.1097. Found; 850.1099.

(Sp)-4c ((Sp)-dCG(BrU_{PS})₈CG). The automated solid-phase synthesis procedure was applied using 5'-O-DMTr-dGipr-pac-CPG (0.5 μ mol) as the solid support, (Rp)-3c as the oxazaphospholidine monomer, and dGipr-pac and dCac phosphoramidite monomers.

ESI-HRMS: m/z calcd for $C_{110}H_{124}Br_8N_{32}O_{70}P_{11}S_8^{5-}$ [M -5H]⁵⁻; 850.1097. Found; 850.1089.

(Rp)-4d ((Rp)-dCG($^{I}U_{PS})_{8}$ CG). The automated solid-phase synthesis procedure was applied using 5'-O-DMTr-dGipr-pac-CPG (0.5 µmol) as the solid support, (Sp)-3d as the oxazaphospholidine monomer, and dGipr-pac and dCac phosphoramidite monomers.

ESI-HRMS: m/z calcd for $C_{110}H_{124}I_8N_{32}O_{70}P_{11}S_8^{5-}$ [M -5H]⁵⁻; 924.8878. Found; 924.8895.

(Sp)-4d ((Sp)-dCG($^{I}U_{PS})_{8}$ CG). The automated solid-phase synthesis procedure was applied using 5'-O-DMTr-dGipr-pac-CPG (0.5 µmol) as the solid support, (Rp)-3d as the oxazaphospholidine monomer, and dGipr-pac and dCac phosphoramidite monomers.

ESI-HRMS: m/z calcd for $C_{110}H_{124}I_8N_{32}O_{70}P_{11}S_8^{5-}$ [M -5H]⁵⁻; 924.8878. Found; 924.8882.

(Rp)-4e ((Rp)-dCG($^{pr}U_{PS})_8$ CG). The automated solid-phase synthesis procedure was applied using 5'-O-DMTr-dG^{ibu}-CPG μmol) as the solid support, (Sp)-3e as the

oxazaphospholidine monomer, and dG^{ibu} and dC^{bz} phosphoramidite monomers.

ESI-HRMS: m/z calcd for $C_{134}H_{148}N_{32}O_{70}P_{11}S_8^{\ 5-}[M-5H]^{5-}$; 784.2782. Found; 784.2794.

(*Sp*)-4e ((*Sp*)-dCG($^{pr}U_{PS})_8CG$). The automated solid-phase synthesis procedure was applied using 5'-O-DMTr-dG^{ibu}-CPG (0.5 µmol) as the solid support, (*Rp*)-3e as the oxazaphospholidine monomer, and dG^{ibu} and dC^{bz} phosphoramidite monomers.

ESI-HRMS: m/z calcd for $C_{134}H_{148}N_{32}O_{70}P_{11}S_8^{5-}[M-5H]^{5-}$; 784.2782. Found; 784.2797.

UV melting analyses

The UV absorbance profiles over temperature were measured using an eight-sample cell changer in quartz cells of 1 cm path length. All the experiments were conducted in a 10 mM phosphate buffer containing 100 mM NaCl (pH 7.0), and the UV absorbance at 260 nm was monitored over temperature. Each sample containing 12mer DNA (2.5 μ M) and its complementary oligonucleotide (DNA or RNA) was first rapidly heated to 95 °C, and, after 10 min, cooled to 0 °C at a rate of 0.5 °C min $^{-1}$. The dissociation process was recorded by heating to 95 °C at a rate of 0.5 °C min $^{-1}$.

RNase H assay

 $0.5~\mu\text{M}$ of 12mer DNA and 5 μM of the complementary RNA were dissolved in 10 mM Tris–HCl buffer containing 100 mM NaCl and 0.5 mM MgCl $_2$ (pH 7.5 at 20 °C). The solution was rapidly heated to 95 °C, and, after 10 min, cooled to 0 °C at a rate of 0.5 °C min $^{-1}$. 4 U or 20 U of RNase H was added to the solution (100 $\mu\text{L})$ and then the mixture was warmed to 20 °C. After 30 min, the mixture was rapidly heated to 95 °C, left for 1 min, and then rapidly cooled to 4 °C. The mixture was analyzed by RP-HPLC.

Conflicts of interest

The authors have no conflicts of interest to declare.

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