

Synthesis and Properties of Peptide Nucleic Acids Containing a Psoralen Unit

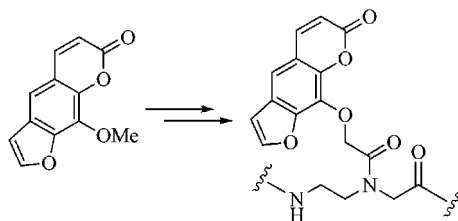
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ABSTRACT



We prepared the psoralen PNA unit from 8-methoxypsoralen and synthesized various PNAs containing psoralen by a typical 'Boc method. PNAs containing psoralen (P-PNA) at strand end formed a stable duplex with complementary DNA. The hybridization of P-PNA with complementary DNA resulted in a considerable decrease of the psoralen fluorescence.

Studies on the molecular basis for photosensitization induced by psoralen derivatives have attracted considerable attention in recent years.¹ These include investigations of the luminescence properties of their excited states,² the formation of psoralen–pyrimidine base cross-link via [2 + 2] photocycloaddition,³ cell killing,⁴ and the cure of skin diseases such as psoriasis.⁵ Psoralens exhibit strong fluorescence by UVA irradiation, and the labeling of DNA and RNA by psoralen is been a promising tool for monitoring interactions with other biomolecules.⁶

Peptide nucleic acid (PNA) is a completely artificial nucleic acid consisting of a peptide backbone. PNA can

recognize a complementary DNA sequence in a highly sequence-specific manner.⁷ The PNA–DNA hybrid is thermodynamically more stable than DNA duplex,⁸ and it is fairly independent of ionic strength in solutions.⁹ Moreover, the incorporation of modified nucleobases into PNA is relatively easy. Therefore, incorporation of a psoralen unit into PNA seems to be very attractive. The sequence-specific DNA recognition by psoralen-containing PNA would potentially be used for the readout of a DNA sequence by monitoring characteristic psoralen fluorescence.

Herein we report the synthesis and properties of PNAs containing psoralen (P-PNA). We synthesized the psoralen PNA unit and then incorporated it into PNA by the 'BOC method. The fluorescence of P-PNA considerably decreased upon forming a PNA–DNA hybrid.

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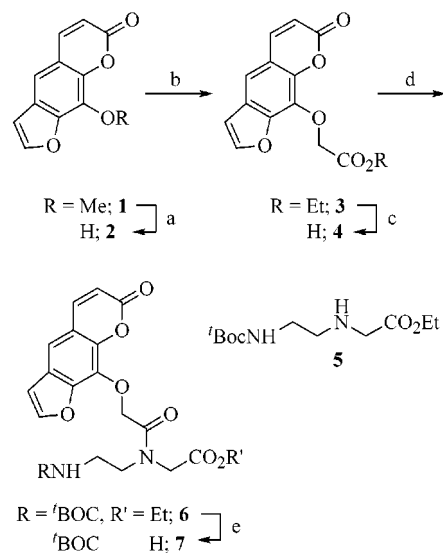
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The synthetic route to the psoralen PNA unit is shown in Scheme 1. The methyl group of 8-methoxypsoralen (**1**) was

Scheme 1. Synthesis of Psoralen PNA Unit^a



^a Reagents and conditions: (a) boron tribromide, dichloromethane, 0 °C, 2 h, 71%; (b) ethyl bromoacetate, potassium carbonate, DMF, rt, 4 h, 81%; (c) lithium hydroxide, ethanol–water (2:1), 0 °C, 10 min, 99%; (d) EDCI, HOBt, DMF, rt, 1 h, and then **5**, rt, 4 h, 87%; (e) lithium hydroxide, ethanol–water (4:3), rt, 4 h, 86%.

removed (71%), and then the hydroxy group of **2** was coupled with ethyl bromoacetate to obtain **3** (81%). Ester **3** was hydrolyzed to **4** (99%), which was coupled with *N*-[2-(*tert*-butoxycarbonylamino)ethyl]glycine ethyl ester (**5**) (87%). Ester **6** was converted to free carboxylic acid to give the psoralen PNA unit **7** (86%). PNA oligomers were synthesized according to a typical ^tBoc solid-phase peptide synthesis using psoralen PNA unit **7**.¹⁰ The crude P-PNA was purified by reversed phase HPLC. A typical example of the HPLC profile is shown in Figure 1. The composition of purified P-PNA oligomers was confirmed by MALDI-TOF mass spectrometry as shown in Table 1.

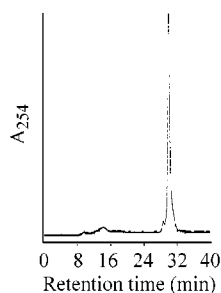


Figure 1. C₁₈ reverse-phase HPLC profile of the crude reaction mixture of **P-PNA 5** obtained by solid-phase peptide synthesis (29.9 min; 0.05% TFA–water, 0–20% acetonitrile over 40 min).

Table 1. Mass Spectral Data of PNAs and *T_m* of Their Hybrids with Complementary DNA Strand

	PNA ^a		<i>T_m</i> (°C) ^b
	MALDI-TOF (M+H) ⁺		
	calcd	found	
PNA 1	H-GTTCCGC-NH ₂		38.0
	1886.83	1886.18	
P-PNA 2	H-PGTTCCGC-NH ₂		46.0
	2229.13	2229.62	
P-PNA 3	H-GTTPCGC-NH ₂		<i>c</i>
	1977.89	1977.91	
P-PNA 4	H-GTTCCPC-NH ₂		27.5
	1937.87	1938.69	
P-PNA 5	H-GTTCCGCP-NH ₂		45.0
	2229.13	2229.61	(20.7) ^d

^a “H” denotes a free amine end, “NH₂” denotes carboxamide end, and “P” denotes psoralin unit. ^b *T_m* value of hybrid with complementary DNA 5′-d(CGCCGAACC)-3′. ^c The sigmoidal melting curve was not observed. ^d *T_m* value of hybrid with 5′-d(CGCCGAACC)-3′ which forms a mismatched base pair.

P-PNA was hybridized with the corresponding complementary DNA strand, and the stability of the P-PNA–DNA hybrid (2.5 μM duplex concentration) was examined by monitoring the melting temperature in 10 mM sodium cacodylate buffer (pH 7.0). The results are shown in Table 1. The P-PNA containing the psoralen unit at the strand end (**P-PNA 2** and **P-PNA 5**) showed 7–8 °C higher *T_m* values than that of the corresponding psoralen-free PNA oligomer (**PNA 1**). Stabilization of the duplex by psoralen at the strand end is probably due to the π-stacking of the hydrophobic aromatic ring of psoralen with the flanking base. However, the *T_m* values of the P-PNA–DNA hybrids containing the psoralen unit in the interior of the PNA strand (**P-PNA 3** and **P-PNA 4**) significantly decreased. It was known that a single mismatched base pair considerably destabilizes the PNA–DNA hybrid.¹¹ Actually, **P-PNA 5**-DNA duplex containing a single-mismatched base pair showed ca. a 24 °C *T_m* decrease. The reason for the destabilization of the duplexes of **P-PNA 3** and **P-PNA 4** is that the psoralen in the interior of the PNA strand cannot form base pairs.

The fluorescence spectra of P-PNAs were measured before and after addition of an equimolar amount of the complementary or mismatched DNA strand. The emission maximum of P-PNAs was 485 nm at 330 nm excitation. Single-stranded **P-PNA 4** and **P-PNA 5** exhibited strong fluorescence, but the fluorescence of **P-PNA 2** was very weak. The hybridization of **P-PNA 4** and **P-PNA 5** with a complementary DNA strand remarkably decreased their fluorescence intensities as compared with that of single-stranded P-PNA without a shift of the emission maximum. It is noteworthy that the fluorescence intensity of **P-PNA 5** decreased 43% by hybridization with complementary DNA (Figure 2). On the other hand,

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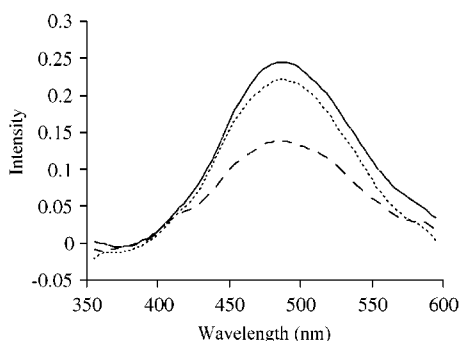


Figure 2. Fluorescence spectral changes caused by **P-PNA 5**–DNA hybrid formation. A solution of 20 μM **P-PNA 5** or **P-PNA 5**–DNA duplex in 10 mM sodium cacodylate (pH 7.0) was used. The fluorescence spectra were measured at 330 nm excitation at 21 $^{\circ}\text{C}$. Solid line, single-stranded **P-PNA 5**; dashed line, **P-PNA 5**–5'-d(CGCGGAACC)-3' (matched duplex); dotted line, **P-PNA 5**–5'-d(CGCCGAACC)-3' (mismatched duplex).

the fluorescence intensity of the **P-PNA 5**–DNA hybrid containing a single-mismatched base pair decreased by only 7% as compared with that of single-stranded **P-PNA 5**. Thus, the full-matched P-PNA–DNA duplex is clearly distinguishable from the mismatched duplex or single-stranded P-PNA by monitoring their fluorescence intensities.

To understand the effect of the P-PNA sequence on the fluorescence intensity of P-PNA, we measured the fluorescence intensities of P-PNAs **H-PT-NH₂**, **H-PC-NH₂**, **H-PA-NH₂**, and **H-PG-NH₂**. The fluorescence intensities of **H-PT-NH₂**, **H-PC-NH₂**, and **H-PA-NH₂** were close to that of **H-P-NH₂**, whereas almost no fluorescence of **H-PG-NH₂** was observed (Figure 3). This result suggests that the low fluorescence intensity of **P-PNA 2** and the remarkable decrease of the fluorescence intensity by the hybridization of **P-PNA 5** and the complementary DNA

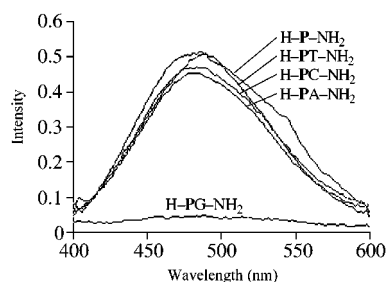


Figure 3. Fluorescence spectra of P-PNAs **H-PN-NH₂** ($N = \text{T, C, A}$ or G) and **H-P-NH₂**. A solution of 20 μM P-PNA in 10 mM sodium cacodylate (pH 7.0) was used. The fluorescence spectra were measured at 330 nm excitation at 19 $^{\circ}\text{C}$.

are due to the quenching of the fluorescence by the flanking G base.

In summary, we have disclosed a facile synthetic route to P-PNA from 8-methoxypsoralen. PNA containing a psoralen unit at strand end forms a stable duplex with complementary DNA. The fluorescence intensity of P-PNA considerably decreased by P-PNA–DNA hybrid formation because of the quenching of the fluorescence by a flanking G base. The fluorescence intensity change of P-PNA on hybridization makes it possible to use for monitoring PNA–DNA hybrid formation. P-PNA would also be useful as a tool for molecular biology and as a potent phototherapeutic drug for PUVA (psoralen plus UVA irradiation) therapy.^{5b}

Supporting Information Available: Experimental procedures and spectral data. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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