# Sequencing by Ligation with Double-Labeled Fluorescent Probes

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**Abstract.** We have synthesized a set of double-labeled fluorescent probes and evaluated the performance of these fluorescent probes in sequencing by ligation (SBL). The results indicated that the ligation efficiency and specificity of double-labeled fluorescent probes was as higher as that of singe-labeled fluorescent probes. Moreover, double-labeled fluorescent probes yielded a r emarkable increase in s ignal i ntensities compared t o s ingle-labeled fluorescent probes. It could also reduce background and shorten exposition time compared to probes labeled with single FAM and FITC dyes. Thus, it could supplant singe-labeled fluorescent probes in SBL to increase sequencing accuracy.

**Keywords:** sequencing by ligation (SBL), double-labeled fluorescent probe, single-labeled fluorescent probe, high-throughput sequencing.

### 1 Introduction

Over the past few years, massively parallel DNA sequencing platforms have become widely available, reducing the cost of DNA sequencing by over two orders of magnitude. Sequencing by ligation (SBL), one of the massively parallel DNA sequencing platforms, uses DNA ligase and either one-base-encoded probes or two-base-encoded probes to determine template sequences [1]. In this method, the specificity and accuracy of probes hybridizing to their complementary sequences adjacent to the primed templates play an important role in the cyclic sequencing cycles. Many researchers have devoted to exploiting new probes for increasing sequencing specificity and accuracy. The SOLiD platform (Applied Biosystems; Foster City, CA, USA) uses specific fluorescent-labeled 8 -mer p robes, whose 4 th and 5 th b ases are encoded by the a ttached fluorescent group [2]. Mir et al. have proposed a sequencing method, named cyclic ligation and cleavage (CycLiC), that uses oligonucleotide probes in which all but one nucleotide is degenerate [3]. Ho et al. have described SBL with deoxyinosinecontaining query oligonucleotides, which could be digested by endonuclease V, leaving a ligatable end extended into the unknown sequence for further SBL cycles [4]. As an al ternative, Li e t a l. have pr esented S BL with a k ind of pr obe i n which deoxynucleoside ph osphorothioates are introduced i nto t he pr obes t o i ncrease t he cleavage accuracy of en donuclease V on double-stranded DNA templates [5]. We

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have also proposed an SBL approach employing oligonucleotide probes with 3'-thiodeoxyinosine and the r ead length c ould r each up to 40 bp with high a ccuracy [6]. However, when this method was ap plied to s equence E. co lig enome, s equencing error rate increased dramatically since the fourth cycle of ligation and the exposition time of FAM dye was almost the sum of the other three dyes. It was unable to improve the ratio of signal to background after the sixth cycle of ligation and cleavage. Herein, a set of double-labeled fluorescent probes for improving the background and exposition t ime was s ynthesized. T he pe rformance o f d ouble-labeled f luorescent probes i n SBL was a lso e valuated. I n a ddition, we exploited whether th e s inglelabeled fluorescent probes could be replaced by double-labeled fluorescent probes for SBL.

# 2 Experimental Setup

#### 2.1 Synthesis of Oligonucleotide Probes

All the ligonucleotide sequences used in this study were shown in Table 1. DNA templates and s equencing p rimers were friendly synthesized by A gene Bioinformative Company (Wuxi, China). T4 DNA ligase and  $5 \times DNA$  ligase reaction buffer (250 mM Tris-HCl [pH 7.6], 50 mM MgCl<sub>2</sub>, 5 mM ATP, 5 mM DTT, 25% [w/v] polyethylene glycol-8000) were purchased from Invitrogen<sup>TM</sup> (Shanghai, China).

Title		oligonucleotide Sequences (5'-3')						
	$T_1$	NH2-CTTTCCTCTCTATGGGCAGTCGGTGATAAGCGTACCCCTA						
Template		GCCCAAATGATCTGCTGTACCGTA						
	<b>T</b> <sub>2</sub>	NH <sub>2</sub> - <u>CTTTCCTCTCTATGGGCAGTCGGTGAT</u> CCTCCTTATCGGGC						
		ATCTCACGCCCTGCTGTACCGTA						
	T <sub>3</sub>	NH2-CTTTCCTCTCTATGGGCAGTCGGTGATGGCCATCTGTGCA						
		CTGCCGAAACACTGCTGTACCGTA						
	$T_4$	NH <sub>2</sub> - <u>CTTTCCTCTCTATGGGCAGTCGGTGAT</u> TACTTGGATCAG						
		GGACGTAGAAGCTGCTGTACCGTA						
S-Primer	$SP_1$	PO4 <sup>3-</sup> -ATCACCGACTGCCCATAGAGAGGAAAG						
S-Probe	<b>P</b> <sub>1</sub>	6-FAM-IIINNNAA	P <sub>5</sub>	(6-FAM) <sub>2</sub> -IIINNNAA				
	P <sub>2</sub>	6-FAM-IIINNNCC	P <sub>6</sub>	(6-FAM) <sub>2</sub> -IIINNNCC				
	P <sub>3</sub>	6-FAM-IIINNNGG	P <sub>7</sub>	(6-FAM)2-IIINNNGG				
	<b>P</b> <sub>4</sub>	6-FAM-IIINNNTT	P <sub>8</sub>	(6-FAM)2-IIINNNTT				

 Table 1. Oligonucleotide sequences used in this study

Underlined bases are areas of sequencing primers hybridization. The bold characters were the queried bases. S-Primer: sequencing primer. S-Probe: sequencing probe. 6-FAM: 6-carboxyfluorescein. (6-FAM)<sub>2</sub>: two 6-FAM dye labels. N: degenerate base. I: deoxyinosine.

#### 2.2 Hybridization of the Sequencing Primers to the ssDNA Templates

The s ingle-stranded DNA (ssDNA) microarrays were f abricated according t ot he previously published literature [7]. Amino-modified oligonucleotides, a s single-stranded DNA templates, were first diluted in sodium c arbonate buffer (0.1 M, pH 9.0) and then transferred to a 384-well plate. The ssDNA templates in the 384-well plate were spotted on the prepared glass slides using a PixSys 5500 microarray (Cartesian Technology). After spotting, the slides were incubated in a humid chamber at room temperature for 4 h and then at 37 °C for 2 h. Finally, the slides were washed by  $2 \times SSC/0.5\%SDS$  and  $0.1 \times SSC/0.5\%SDS$ , and then dried by nitrogen.

To hybridize the sequencing primers to ssDNA templates, a mixture containing sequencing primers (4  $\mu$ M),  $\ltimes$ hybridization buffer (10 mM Tris -HCl, 0.9M NaCl, and 10% sodium dode cyl sulfate) was placed on the arrays by pipette, and then covered with coverslips. Afterwards, the slide was put in a humid chamber. Hybridization was firstly conducted at 80°C for 5 min, then cooled at room temperature, and finally incubated at 37°C for 1 h. After hybridization, the slide was washed and then dried by nitrogen.

#### 2.3 Ligation of Fluorescent-Labeled Probes

After h ybridization, the templates on the microarrays were lig ated with la beled probes. Ligation mixture c ontaining the following was prepared and placed on the arrays:  $1 \times DNA$  ligase r eaction b uffer (50 m M T ris-HCl (pH 7. 5, 25 °C), 10 mM MgCl<sub>2</sub>, 10 mM DTT, 1 mM ATP), 0.025 U/µL T4 DNA ligase and a mixture of 2 µM fluorescent-labeled probes. Afterward, the microarrays with mix reagent were incubated for 30 minutes at 15 °C in a humid chamber. Finally, the slides were washed and dried by n itrogen. Fluorescent s ignals on the microarray were cap tured by using a homemade scanner (Luxscan-10K/A, C apital Biochip C orporation, C hina), and the data were analyzed with Spot Data Pro 3.0 software.

## **3** Results and Discussion

### 3.1 Synthesis of Double-Labeled Fluorescent Probes

The double-labeled fluorescent probes were synthesized as previously described [8, 9]. To obtain double-labeled fluorescent probes (Fig. 1(a)), oligonucleotide sequence was synthesized in a dvance. A symmetric doubler phosphoramidite (Glen Research Corporation, Catalog No. 10-1920-90), which could introduce two hydroxyl radical groups to facilitate the conjunction of fluorescent dyes, was coupled to 5' terminal of oligonucleotide sequence during synthesis (Fig. 1(b)). Subsequently, two fluorephores were linked to the symmetric doubler phosphoramidite. The resultant double-labeled fluorescent probes were characterized by ultraviolet (UV) spectra (Nanophotometer, Implen GmbH) and the results were shown in Fig. 2.



**Fig. 1.** The structure of dou ble-fluorescent l abeled p robe (a) and s ymmetric d oubler phosphoramidite (b).

As shown in Fig. 2, the solid lines represented the normalized absorbance spectra (Groenzin a nd M ullins, 2 000) f or dou ble-labeled f luorescent pr obe  $P_7$  and s ingle-labeled fluorescent probe  $P_3$ , respectively. Compared to single-labeled probes  $P_3$ , the absorbance peak of 6-FAM dye in double-labeled probes  $P_7$  was much higher when their concentrations ar e equal. This confirmed that the label e fficiency of double-labeled fluorescent probes and single-labeled probes were 120% and 89% when the absorbance of oligonucleotide sequences and 6-FAM dyes were  $7.5 \times 104$  (L.mol<sup>-1</sup>.cm<sup>-1</sup>) and  $6.2 \times 104$  (L.mol<sup>-1</sup>.cm<sup>-1</sup>), respectively. That was, the label ratio between single-labeled probes and double-labeled probes were 1:1.35 when their concentration were equivalent. E ach double-labeled f luorescent probe had approximately l abeled 1.2 fluorophore dyes at the 5' terminus on average while each single-labeled probe had labeled about 0.89 fluorophore dyes.



**Fig. 2.** The absorption spectra for double-labeled fluorescent probes (6-FAM)<sub>2</sub>-IIXNNNGG (b) and single-labeled fluorescent probes 6-FAM-IIXNNNGG(a).

#### 3.2 SBL with Double-Labeled Fluorescent Probes

To investigate whether SBL with double-labeled fluorescent probes could be performed successfully, we spotted synthetic templates  $T_1$ ,  $T_2$ ,  $T_3$  and  $T_4$  onto a microarray. As shown in the left subarrays in Fig. 3(a), SBL with double-labeled fluorescent probes could successfully performed and each probe could specifically ligated with

their complementary template. To compare the ligation efficiency of SBL with singlelabeled probes to that of SBL with double-labeled probes, single-labeled probe  $P_4$  (6-FAM-IIXNNNTT) and double-labeled fluorescent probe P8 ((6-FAM)2-IIXNNNTT) were ligated with those templates (Fig. 3). As shown in Fig. 3(a), fluorescence intensity of double-labeled fluorescent probe  $P_8$  was nearly two times stronger than that of the single-labeled fluorescent p robe P4 at the s ame c oncentration. It in dicated that double-labeled fluorescent probes might not quench the fluorephore dyes during ligation reactions, and had nearly no impact on the ligation reactions. The relationship between f luorescence i ntensity a nd l igation c oncentration w as s hown i n F ig. 3 (b), fluorescence intensities of probes P<sub>4</sub> and P<sub>8</sub> increased almost linearly from 0 to 0.5  $\mu$ M, and then reached a plateau with the highest fluorescence intensity at a concentration of 1  $\mu$ M. The highest fluorescence intensity kept stable as the probe concentration continued to increase. It indicated that two fluorophore dyes in probe P<sub>8</sub> might slightly affect the hybridization kinetics, but this unobvious effect might be negligible during the half-hour long reactions. That was, probes  $P_8$  and  $P_4$  had almost the same kinetic during the ligation reactions. In addition, probe  $P_8$  and  $P_4$  had the same ligation specificity during ligation reactions. Thus, single-labeled probes could be supplanted by double-labeled fluorescent probes in SBL.



**Fig. 3.** The fluorescent intensities of probes  $P_4$  and  $P_8$  at the concentration of 0.5, 1.0, 1.5, 2.0, 2.5  $\mu$ M, respectively (a) The microarray contained ten subarrays, and each subarray included 4 spots of the templates  $T_1$ ,  $T_2$ ,  $T_3$  and  $T_4$ . From top to bottom, the left subarrays were ligated with probe  $P_4$  at the concentration of 0.5, 1, 1.5, 2, 2.5  $\mu$ M, respectively while the r ight subarrays were ligated with probe P8 at the concentration of 0.5, 1, 1.5, 2, 2.5  $\mu$ M, respectively. (b) Correlation between the concentration of ligation probe and fluorescence intensity. Single: single-labeled fluorescent probe.

We also evaluated the performance of iterative cycles of ligation and cleavage with four kinds of double 6-FAM labeled fluorescent probes. The results were compared with that of SBL using single 6-FAM labeled fluorescent probes (Table 2). At each given concentration, the fluorescence intensities of SBL with double-labeled fluorescence probes were nearly one times stronger than that of SBL with single-labeled fluorescence probes. That was, the ratio of fluorescence intensities of SBL with double-labeled fluorescence probes.

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ble-labeled to single-labeled fluorescence probes was about 2:1. Therefore, SBL with double-labeled fluorescence probes could obtain stronger fluorescence intensity and could improve sequencing accuracy. In addition, SBL with 6-FAM labeled fluorescent probes could avoid longer exposition time and higher background existing in SBL with one FAM labeled, or FITC dyes labeled probes. Although the method has advantage in the improvement of signal intensity and the extension of exposition time, some disadvantages of this method needed to be emphasized. First, the synthesis of fluorescent dyes might increase the cost of SBL. Second, labeling t wo FAM fluorephore dyes at the 5' terminus might make the ligation more difficult due to the steric hindrance. However, the increase of signal intensity might reduce the sequencing error rate and the exposition time of FAM dye was almost the sum of the other three dyes from the fourth cycle of ligation. Thus, double-labeled fluorescence probes could replace single-labeled probes for SBL. It would be useful for improving sequencing accuracy.

Table 2. Fluorescence intensities of single-labeled and doubled-labeled fluorescence probes

Probes	AFI		Probes	AFI		The ratio of AFI	
	1 μM	1.5 μM		1 µM	1.5 μM	1 µM	1.5 μM
<b>P</b> <sub>1</sub>	10801	12553	P <sub>5</sub>	20845	24359	$P_5/P_1 = 1.93$	P <sub>5</sub> /P <sub>1</sub> =1.94
P <sub>2</sub>	10881	12022	P <sub>6</sub>	21936	23804	P <sub>6</sub> /P <sub>2</sub> =2.02	P <sub>6</sub> /P <sub>2</sub> =1.98
P <sub>3</sub>	11964	15606	P <sub>7</sub>	28613	29729	P <sub>7</sub> /P <sub>3</sub> =2.39	P <sub>7</sub> /P <sub>3</sub> =1.90
P <sub>4</sub>	11663	11883	P <sub>8</sub>	21106	23172	P <sub>8</sub> /P <sub>4</sub> =1.81	P <sub>8</sub> /P <sub>4</sub> =1.95

AFI: the a verage fluorescence in tensities from four i terative experiments. In this table, o ligonucleotide sequences  $P_1 \sim P_4$  were labeled with single fluorescence dyes while oligonucleotide sequences  $P_5 \sim P_8$  were labeled with double fluorescence dyes at the 5' terminus, respectively.

## 4 Summary

In this report, we have synthesized a s et of d ouble-labeled fluorescent probes and have evaluated the p erformance of d ouble-labeled fluorescent probes in S BL. The results de monstrated that the lig ation e fficiency and s pecificity of d ouble-labeled fluorescent probes were as higher as that of singe-labeled fluorescent probes. Moreover, S BL with double-labeled fluorescent probes yielded a r emarkable increase in signal intensities compared to SBL with single-labeled fluorescent probes, making SBL reactions more accurate. It also could reduce background and shortened exposition time. Thus, it could supplant singe-labeled fluorescent probes in SBL to increase sequencing accuracy and specificity.

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