Solid-Phase PCR in Microwells: Effects of Linker Length and Composition on Tethering, Hybridization, and Extension

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ABSTRACT

During the solid-phase PCR (SP-PCR), DNA oligonucleotides complementary to a soluble template and immobilized on a surface are extended in situ. Although primarily used for pathogen detection, SP-PCR has the potential for much broader application, including disease diagnostics, genotyping, and expression studies. Current protocols for SP-PCR in microwells are suitable for enzymatic detection of immobilized products, but yields are generally insufficient for direct detection of products using conventional fluorescent probes. Here, we quantitatively measure the outcome of tethering, hybridization, and solid-phase extension, and examine the effect of composition and length of the spacer at the 5' end of tethered oligonucleotides. Our results indicate that steric hindrance primarily affects polymerase activity rather than the efficiency of hybridization between the template and the tethered oligonucleotide. SP-PCR yields are significantly higher for a five-unit hexaethyleneglycol (HEG) spacer than for the more commonly used 10-residue deoxythymidine (dT) spacer. The optimal 5' HEG spacer resulted in a 60-fold increase in extension efficiency relative to a previously reported value for SP-PCR on a glass surface. Thus, optimized spacers should allow direct quantification of SP-PCR products, providing a simple, quantitative, and cost-effective means of sample analysis for a variety of applications.

INTRODUCTION

The solid-phase PCR (SP-PCR) (9) is analogous to standard PCR, except that one of the oligonucleotide primers, normally free to bind template DNA in solution, is attached to a surface (usually glass slides, beads, or plastic microwells). The potential uses of SP-PCR include genotyping, disease diagnostics, pathogen detection, and gene expression studies. During SP-PCR, a complementary single-stranded template anneals to the tethered oligonucleotide, DNA polymerase extends the molecule, the strands are denatured, and the cycle repeats as the liquid-phase template concentration increases. After amplification is complete, untethered molecules are washed away, and the single- or double-stranded tethered extension products may be detected by a variety of means (2,10,13).

Microplate-based solid-phase extension products are usually detected by enzymatic assays (8–11). For some applications, however, it would be preferable to employ direct detection of fluorescent products, which would allow quantitative estimation of yield over a wide dynamic range, as well as having the advantages of simplicity, flexibility, and cost. Thus, it would facilitate the use of microplate-based SP-PCR in high-throughput, automated applications. In our laboratory, however, SP-PCR yields have not been sufficient for direct fluorescence detection with standard plate readers. We require approximately 100 fmol product in a 50 µL volume/well for reliable quantification.

Previous studies have demonstrated that steric hindrance inhibits the hybridization of DNA in solution to immobilized oligonucleotides (6,12). Steric hindrance might also affect solid-phase polymerization by impeding the attachment of Taq DNA polymerase to tethered oligonucleotides that directly abut the supporting surface. It has been demonstrated that SP-PCR efficiency is enhanced when a polydeoxythymidine (dT) spacer is included at the 5' end of the solid-phase primer (1,10,13). Solid-phase oligonucleotides containing 5' (dT)n spacers are desirable because they are inexpensive and easy to synthesize. However, we tend to observe high background signals when using these primers to amplify AT-rich plant DNA templates.

In this study, we quantitatively measured the outcome of tethering, hybridization, and solid-phase extension in microwells using common fluorescent labels and a standard plate reader. We also evaluated the effect of 5' hexaethyleneglycol (HEG) spacers of varying lengths in mitigating steric hindrance to hybridization versus polymerization, and compared the performance of solid-phase primers with 5-unit HEG and (dT)10 spacers (both type spacers were coupled to a 5' amino modifier C6 residue). The goals were to demonstrate that SP-PCR was occurring in microwells and to develop a SP-PCR protocol suitable for use in quantitative assays that employ direct fluorescence detection in plate format.

MATERIALS AND METHODS

Oligonucleotides

Primers and probes (Table 1) were derived from the first exon of the Ara-
Table 1. Oligonucleotides Used for Evaluation of Solid-Phase Extension and PCR

<table>
<thead>
<tr>
<th>Oligonucleotide</th>
<th>5’ Modification</th>
<th>DNA Sequence (5’→3)(^a)</th>
<th>Experimental Role</th>
</tr>
</thead>
<tbody>
<tr>
<td>(F^{(HEG)})(_n)</td>
<td>aminoC6(^b) ((HEG))(^c)</td>
<td>GCCTTTTTATGCGATTCTGC</td>
<td>Tethered in microwells</td>
</tr>
<tr>
<td>(F^{(dT)})(_{10})</td>
<td>aminoC6</td>
<td>TTTTTTTTTTGGCTTTTTGATTCTGC</td>
<td>Tethered in microwells</td>
</tr>
<tr>
<td>80(^f)</td>
<td>fluorescein</td>
<td>CAGGACCCTCATCGAAGACTACAGATCCAATCTA</td>
<td>Hybridized to tethered primers (solid-phase extension assay)</td>
</tr>
<tr>
<td>F</td>
<td>none</td>
<td>GCCTTTTTATGCGATTCTGC</td>
<td>Liquid-phase primer (SP-PCR)</td>
</tr>
<tr>
<td>R(^f)</td>
<td>fluorescein</td>
<td>CGGTTAGAGTGACCTTGAAT</td>
<td>Liquid-phase primer (SP-PCR)</td>
</tr>
<tr>
<td>R(^r)</td>
<td>Texas Red</td>
<td>CGGGTAGAGTAGACTTGAAT</td>
<td>Probe (SP-PCR)</td>
</tr>
</tbody>
</table>

\(^a\)80\(^f\) contains a \(Hpa\) II site (underlined) not found in the \textit{Arabidopsis} genomic sequences.
\(^b\)Amino modifier C6 (Glen Research, Sterling, VA, USA).
\(^c\)0–20 molecules of hexaethyleneglycol (Spacer Phosphoramidite 18, Glen Research)

Optimization of Spacer Length

Optimal spacer length was determined as outlined in Figure 2. Initially the \(F^{(HEG)}\)\(_n\) oligonucleotides with spacer lengths of 0, 5, 10, and 20 units were evaluated. Four trials (repetitions) were performed with eight 8-well strips/trial, and two wells/treatment/strip. Placement of treatments within strips was randomized. After tethering, the amount of covalently bound primer/well was determined for one strip/trial using YOYO-1 iodide (Molecular Probes, Eugene, OR, USA), a fluorescent dye that has a strong affinity for ss-DNA. Following the published protocol (7), fluorescence was measured with a SPECTRAFluor Plus plate reader (TECAN, Research Triangle Park, NC, USA). After initial results were obtained, a second experiment was performed in which spacers containing from 1–8 HEG residues were evaluated. Here, each spacer length was assigned to one well/strip, but as before, eight strips were used (seven experimental strips and one strip for quantification of tethered oligonucleotide).

The 80\(^f\) oligonucleotide (5 pmol) was hybridized to tethered oligonucleotides in 100 \(\mu\)L 5× SSC (1.25 M NaCl, 0.125 M sodium citrate, pH 7.0) for 16 h at 50°C. Wells were washed three times with 1× SSC at room temperature to remove unhybridized 80-mer, 100 \(\mu\)L 1× SSC were added to

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**bidopsis thaliana** phytochrome C (PhyC) gene (4). Figure 1 shows the relative positions of oligonucleotides within PhyC.

**Covalent Binding of Oligonucleotides to Wells**

In all experiments, 5’ amino-modified oligonucleotides were tethered in NucleoLink™ (Nalge Nunc International, Rochester, NY, USA) strips (eight wells/strip) by standard carbodiimide-mediated condensation chemistry (10). Oligonucleotide tethering and blocking of unreacted primer binding sites followed the microwell manufacturer’s protocol (http://nunc.nalgenunc.com/resource/technical/nag/DP0063.htm).

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**Figure 1.** The PhyC gene (3572 transcribed bp) and the molecules used in this study derived from Exon I: the 251-bp PCR product, the synthetic 80-mer (80\(^f\)), and the R\(^r\) probe. The \(Hpa\) II restriction site in 80\(^f\) was introduced by modifying one nucleotide of the genomic sequence, while the \(Hpa\) II site in the PCR product is native to the Columbia allele. Fluorescent labels are shown by colored circles: green, fluorescein; red, Texas Red.
each well, and the amount of fluorescein/well was determined using the plate reader. Tethered oligonucleotides (F\(^{(HEG)}_n\)) were extended in 50-µL reaction volumes containing 2.5 mM MgCl\(_2\), 0.2 mM each dNTP, and 2.5 U Taq DNA polymerase in 1× PCR buffer (Promega, Madison, WI, USA). Reactions were incubated for 1 h at 50°C, and wells were washed three times with 1× SSC. Restriction digests were done in 50-µL volumes with 1× One-Phor-All Buffer PLUS (Amersham Biosciences, Piscataway, NJ, USA), 0.10 mg/mL BSA (New England Biolabs, Beverly, MA, USA), and 1 U HpaII (Invitrogen, Carlsbad, CA, USA). Reactions were incubated for 1 h at 37°C. The reaction mixture (40 µL) was then transferred to 96-well black plates (Corning Costar, Cambridge, MA, USA), a 60-µL aliquot of TE (10 mM Tris-HCl, pH 8.0, 1 mM EDTA) was added, and fluorescence was measured.

Fluorescein-labeled restriction fragments were purified with Centri-Sep™ spin columns (Princeton Separations, Adelphia, NJ, USA). Samples were concentrated, and approximately one-fourth of the original reaction volume was assayed on a model 377 automated DNA fragment analyzer using established protocols (GeneScan® Reference Guide; both from Applied Biosystems, Foster City, CA, USA).

Quantification of SP-PCR Products

Experiments for confirmation and quantification of SP-PCR products are diagrammed in Figure 3. Three trials were performed with four 8-well strips/trial and one treatment/strip. The presence of Taq DNA polymerase and tethered oligonucleotides were varied in each treatment, and one well/strip was reserved for quantification of tethered oligonucleotides with YOYO-1.

Total genomic DNA was extracted from A. thaliana cv. Columbia seedlings using a standard method (5). F\(^{(HEG)}_5\) oligonucleotides (5-unit spacers) were tethered as described. SP-PCR buffers were as above, except that they contained 1 pmol F (unlabeled), 8 pmol R\(^{11}\) (5'-fluorescein) primers, and 25 ng Arabidopsis genomic DNA. PCR was performed using a Primus 96-plus thermal cycler (MWG Biotech, Ebers-
berg, Germany) with the following temperature profile: 95°C for 5 min, 35 cycles of 95°C for 1 min, 55°C for 1 min, 72°C for 2 min, followed by a 90-min incubation at 50°C. To confirm that liquid-phase PCRs were successful, 10 µL of the amplification reactions were run on a 1% agarose gel stained with ethidium bromide. For selected wells, HpaII digests were performed, and products were sized on a DNA fragment analyzer, as before, to confirm the presence of the expected 161-bp fragment (Figures 1 and 3A).

Quantitative estimates of SP-PCR yield were made as follows. After completion of SP-PCR, wells were washed three times with 1× SSC, and the amount of bound fluorescein signal was determined. Then, the dsDNA was denatured by heating to 95°C for 5 min, the solution was aspirated, the wells were washed three times with 1× SSC, and fluorescein readings were obtained to measure the residual fluorescein-labeled complementary template. The tethered DNA strands were then probed with R<sup>tr</sup> (Texas Red<sup>®</sup>-labeled) oligonucleotide (5 pmol in 50 µL 5× SSC) for 16 h at 50°C. Wells were washed, as before, and measurements of both hybridized R<sup>tr</sup> and residual fluorescein

Figure 3. Experiments for verification and quantification of SP-PCR. Experiment consisted of three trials with four 8-well strips/trial and one treatment/strip. The presence of Taq DNA polymerase and tethered oligonucleotides were varied in each treatment, and the amount of covalently bound primer was determined for one well/strip using YOYO-1. Fluorescein-labeled double-stranded products were generated by inclusion of liquid-phase R<sup>tr</sup> primer. (A) Solid-phase extension was confirmed in selected wells by HpaII digestion of tethered dsDNA and visualization of the resulting 161-bp fragment on a denaturing polyacrylamide gel. (B) Solid-phase extension was quantified by denaturation of tethered double-stranded products, washing, and hybridization of a fluorescent probe (R<sup>tr</sup>) complementary to the 3’ end of the single-stranded product. Wells were washed, and fluorescence was quantified by comparison to a standard. The amount of fluorescein-labeled complementary strand (and/or R<sup>tr</sup> liquid-phase primer) was also determined after completion of SP-PCR, denaturation, probing, and additional washing.
were taken. Fluorescence was determined one final time for both dyes after three additional washes with 1× SSC.

Comparison of 5′ HEG and dT10 Spacers

F(HEG)5 and F(dT)10 were each tethered to all eight wells of three NucleoLink strips apiece, and SP-PCRs were then performed as described above. In addition, there were three control strips that contained all reaction components except tethered oligonucleotides. Wells were washed, probed with Rr, and fluorescence was measured as described above. The quantity of tethered oligonucleotide was determined by YOYO-1 assay for one well/strip.

Statistical Analyses

ANOVA was performed on fluorescence data using the JMP statistical software package (version 3.0; SAS Institute, Cary, NC, USA). Box-Cox transformations were used to obtain normally distributed residuals (3). Linear contrast tests (14) were used for planned comparisons among specific treatments. Variance components were estimated by equating observed to expected mean squares (14).

RESULTS AND DISCUSSION

Optimization of 5′ HEG Spacer Length

We first sought to determine the optimum length of 5′ HEG spacers on tethered primers for hybridization and extension. This experiment (Figure 2) was designed so that hybridization could be measured independent of solid-phase extension.

Previous reports based on enzymatic assays indicated that well-to-well variability was low for the NucleoLink surface (10). However, our ANOVA results indicated that most of the variation in hybridization and extension experiments (65% and 83%, respectively, of the experiment-wide variance) was due to inherent differences between wells of the same strip (likely due to variability in manufacture). By contrast, there was little variability among strips (4% for hybridization and 8% for extension experiments, based on average values for eight wells/strip) and trials (31% and 9%, respectively).

The hybridization and extension results for each spacer length are shown in Figure 4. To establish a range for more detailed study, spacers with 0, 5, 10, and 20 HEG residues were initially evaluated. While the amount of hybridized 80-mer decreased as a function of spacer length, solid-phase primer extension increased from 0 to 5-unit spacers, was roughly equivalent for 5- and 10-unit spacers, and decreased at 20-unit spacers (Figure 4, A and B). Thus, the comparatively inefficient extension of tethered primers without a 5′ spacer appears to be due to steric hindrance of Taq DNA polymerase rather than to lowered hybridization efficiency. Conversely, the decline in primer extension with 20-unit spacers might be related to decreased hybridization of the 80-mer (Figure 4A). The optimal spacer length for efficient solid-phase extension by Taq DNA polymerase appears to be 5–10 HEG units. When we evaluated spacers between one and eight HEG units in length, the results indicated a slight decline in hybridization with increasing spacer length and an increase in the efficiency of solid-phase extension, with little difference in overall yield between five to eight linkers (Figure 4, C and D). Since shorter spacers are easier and less expensive to synthesize, subsequent SP-PCR experiments were performed using tethered primers with 5 units of HEG spacer.

We should note that the decrease in hybridization yield observed with increasing spacer length was surprising. It could be argued that this result was due to fluorescence quenching, as increased concentrations of labeled oligomers were hybridized to tethered primers. Because we do not know the spatial distribution of hybridized fluorescein-labeled oligomers on the well surface, we cannot disregard the possibility that quenching might occur. However, our standard concentration curves for fluorescein in solution were linear (R² = 0.99) at 1 nmol dye/50 µL, about 2000-fold greater than the quantities of fluorescein detected in the SP-PCR experiments. We should also mention that other researchers have observed decreased hybridization yields for long spacers relative to their shorter counterparts using radioactive hybridization assays. For three different

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<table>
<thead>
<tr>
<th>Treatment</th>
<th>After PCR</th>
<th>After Denaturation</th>
<th>After Rf Hybridization</th>
<th>After Further Washes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Taq</td>
<td>Tether</td>
<td>Fluorescein</td>
<td>Fluorescein</td>
<td>Texas Red</td>
</tr>
<tr>
<td>- -</td>
<td>25 (6)*</td>
<td>21 (6)</td>
<td>6 (8)</td>
<td>91 (10)</td>
</tr>
<tr>
<td>- +</td>
<td>17 (5)</td>
<td>24 (5)</td>
<td>4 (5)</td>
<td>73 (5)</td>
</tr>
<tr>
<td>+ -</td>
<td>33 (8)</td>
<td>26 (5)</td>
<td>2 (5)</td>
<td>53 (5)</td>
</tr>
<tr>
<td>+ +</td>
<td>124 (10)</td>
<td>29 (7)</td>
<td>21 (6)</td>
<td>263 (25)</td>
</tr>
</tbody>
</table>

* Experiment consisted of three trials with four 8-well strips per trial and one treatment per strip.

†F(HEG)5 was tethered in microwells.

‡From Rf-labeled PCR products and/or Rf primer alone.

§From Rf probe.

*2 standard error is shown in parentheses.

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glycol spacers (propanediol, diethyleneglycol, and triethyleneglycol), a steady increase in duplex yield was reported with increasing spacer length up to 8–10 units, but hybridization declined with further length increases until, at 25–30 units, duplex yield equaled that found with no spacer at all (12).

After hybridization of the 5′ fluorescein-labeled 80-mer and extension of tethered primers (Figure 2), fragments detected after digestion with HpaII should represent double-stranded extension products, since HpaII does not cut ssDNA. To verify that the observed fluorescence was associated with the appropriate restriction fragment and not residual, uncut 80-mer, aliquots from selected wells were loaded on a DNA fragment analyzer. There were intense fluorescent signals around 54 bp, the size of the expected HpaII restriction fragment, and no fluorescence in the 80-bp region (data not shown). Therefore, HpaII activity was either not affected by steric hindrance or the re-

<table>
<thead>
<tr>
<th>Spacer</th>
<th>Tethered Oligonucleotide</th>
<th>Product Extended</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>fmol</td>
<td>YOYO-1</td>
</tr>
<tr>
<td>(HEG)₅</td>
<td>744 (81)b</td>
<td>228 (8)</td>
</tr>
<tr>
<td>(dT)₁₀</td>
<td>713 (23)</td>
<td>147 (3)</td>
</tr>
<tr>
<td>Control</td>
<td>55 (7)</td>
<td>52 (2)</td>
</tr>
</tbody>
</table>

*Experiment consisted of three 8-well strips per spacer with one treatment per strip. YOYO-1 was quantified in one well per strip.

*b2× standard error is shown in parentheses.

*cControl wells contained all reactants except tethered primers.

Figure 4. Hybridization and solid-phase extension of tethered oligonucleotides with 5′ HEG spacers of various lengths. Error bars represent 2× standard errors. (A) Quantity of fluorescent label (in fmol) after hybridization of 80 to tethered primers (solid line) and after primer extension (dashed line), the latter measured as the quantity of labeled liquid-phase restriction fragment. Results are shown for spacers with 0, 5, 10, and 20 HEG residues. (B) Percent efficiency of extension (extension × 100/hybridization) shown in panel A. (C) Quantity of probe hybridized (solid line) and extended (dashed line) for spacers with 1–8 HEG residues, as in panel A. (D) Efficiency of extension for reactions shown in panel C.
striction enzyme excess (approximately 140-fold) compensated for possible steric constraints.

**SP-PCR Verification and Quantification**

The previous experiments demonstrated solid-phase extension in the presence of abundant template. To test for solid-phase extension coupled with template amplification, two experiments were performed (Figure 3). SP-PCRs were done using primers that amplify a 251-bp fragment from exon I of the *A. thaliana* Phyc gene (Figure 1). The 5′-aminated F[HEG]5 oligonucleotides were tethered, and PCRs were performed using a liquid-phase primer ratio of 1:8 (F:R) to produce an excess of template strands complementary to the tethered oligonucleotide (10,11).

Visualization of appropriately sized (161 bp) fluorescent restriction fragments confirmed the presence of SP-PCR products in wells containing both tethered primers and *Taq* DNA polymerase (Figure 5A). Some but not all experimental wells (those with tethered primers and *Taq*) and some controls (those with *Taq* but without tethered primers) also contained a small amount of full-length, presumably residual single-stranded liquid-phase product (251 bp) that was not removed from wells by washing (Figure 5B).

Table 2 presents fluorescence data from the experiment outlined in Figure 3B. Statistical analyses indicated that there was a highly significant interaction term between the presence of *Taq* DNA polymerase and the presence of tethered primers (*P* = 6 × 10⁻⁴), and a linear contrast between the treatment having all SP-PCR components and the other three treatments revealed that the difference in fluorescence was highly significant (*P* = 3 × 10⁻⁴).

In this experiment (Figure 3B), fluorescein was quantified after completion of SP-PCR and after subsequent washings and hybridizations. The fluorescein signal represented either specific binding of unincorporated R° liquid-phase primers and/or fluorescein-labeled complementary PCR products to extended primers or nonspecific background. After completion of SP-PCR, approximately 100 fmol fluorescein were detected in wells containing all reaction components (Table 2). Fluorescein signal dropped to background after heat denaturation, indicating that the fluorescein-labeled complements/primers were removed from wells. The quantity of solid-phase oligonucleotides extended during PCR was estimated by hybridization to R° (Texas Red-labeled probe). Although background fluorescence was relatively high in the controls, the Texas Red signal from wells containing all SP-PCR components was approximately 3-fold greater. After correction for background fluorescence in the Texas Red data (Table 2), we estimate that approximately 180 fmol tethered primers were extended by SP-PCR. This is consistent with the values obtained in earlier hybridization/extension experiments where approximately 160 fmol product were detected for tethered oligonucleotides with 5'-unit HEG spacers (Figure 4A).

The Texas Red background remained after three additional washes with 1× SSC at room temperature (Table 2). Among the controls, the highest background readings were in wells containing neither *Taq* DNA polymerase nor tethered oligonucleotides. However, the fluorescein readings in these wells were low, even immediately following completion of temperature cycling. Since the R° probe appears to be interacting directly with the well surface, additional blocking steps, different blocking solutions, shorter hybridizations, or an alternative dye may improve the signal-to-noise ratio.

Based on combined YOYO-1 assays from all experiments, we estimate that the amount of tethered primer was 780 ± 30 fmol/well (*n* = 72), and the density of primers on the well surface was 14 fmol (or 8 × 10⁹ molecules)/mm². Coupled with the SP-PCR yields obtained above, these data show that 20%, or 1 in 5, of the covalently bound primers was extended during SP-PCR. This result is a substantial improvement over estimates of SP-PCR efficiency using other approaches (e.g., 1 in 300 primers extended at an equivalent density on a glass surface in Reference 1).

**5′ HEG versus (dT)₁₀ Spacers**

We compared the SP-PCR yields from wells that had been tethered with

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Figure 5. Electropherogram results showing the *HpaII* restriction fragment from SP-PCR and residual full-length product from a liquid-phase PCR “control”. Top scale is size in base pairs. Color coding denotes internal size standards (red peaks) and fluorescein-labeled products (green peaks). (A) Experimental well containing tethered F[HEG]₅ oligonucleotide, *Taq* DNA polymerase, and all other PCR components. Arrow denotes the 161-bp *HpaII* restriction fragment from cleavage of double-stranded SP-PCR products. (B) Liquid-phase PCR control well containing all reactants except tethered oligonucleotides. Arrow indicates a weak signal at 251 bp representing residual, full-length, liquid-phase PCR product (not cleaved by *HpaII*).
CONCLUSION

In general, the surface density of tethered oligonucleotides, the abundance and accessibility of complementary template molecules in solution, and the accessibility of tethered primers to Taq DNA polymerase will affect the efficiency of SP-PCR (1.6). Since SP-PCR products are usually detected by enzymatic reactions that are sensitive to less than 1 amol product/20-μL volume (9), enzyme-based assays succeed if only a small proportion of tethered oligonucleotides are extended during SP-PCR, or if residual liquid-phase products remain in reaction wells. Although enzymatic detection can be done in high-throughput format, it is not quantitative and requires multiple handling steps, chemically modified probes, and expensive substrates. Here, we have demonstrated that direct fluorescent detection of SP-PCR products is feasible in NucleoLink strips. Similar results are to be expected using other commercial strips or plates as long as the tethering chemistry results in immobilization of sufficient quantities of oligonucleotides on the well surface. SP-PCR yields from tethered oligonucleotides with 5′ (HEG)₅ spacers are significantly higher than yields from oligonucleotides with 5′ (dT)₁₀ spacers. Our protocol results in a 60-fold increase in extension of tethered oligonucleotides relative to reported values (1). Thus, direct detection of solid-phase amplification products should now provide a simple, quantitative, and cost-effective means of sample analysis in a variety of molecular applications.

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