Isothermal Amplification and Multimerization of DNA by \textit{Bst} DNA Polymerase

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ABSTRACT

We have demonstrated the isothermal in vitro amplification and multimerization of several different linear DNA targets using only two primers and the strongly strand-displacing exonuclease-negative Bst DNA polymerase. This reaction has been termed linear target isothermal multimerization and amplification (LIMA). LIMA has been compared with cascade rolling-circle amplification and has been found to be less sensitive but to yield similar variable-length multimeric dsDNA molecules. Products from several different LIMA reactions were characterized by restriction analysis and partial sequence determination. They were found to be multimers of subsets of the target sequence and were not purely primer derived. The sensitivities with respect to target concentration of several different LIMA reactions were determined, and they varied from 0.01 amol to 1 fmol. The sensitivity and specificity of LIMA were further tested using \textit{E. coli} genomic DNA, and the selective amplification of a transposon fragment was demonstrated. A successful strategy for reducing LIMA-dependent background DNA synthesis in CRCA reactions, it is plausible that the ability of displaced strands to act both as primers and as replication substrates results in the reaction being fully self-sustaining.

INTRODUCTION

It is now established that small circular DNA molecules may serve as templates for isothermal in vitro rolling-circle replication reactions catalyzed by strand-displacing DNA polymerases. These reactions have been shown to yield single-stranded linear concatemeric products (1,4,5,11). More recently, there have been reports that, if these reactions are carried out in the presence of two primers, one identical and one complementary to the circular template, then exponential amplification that yields double-stranded variable-length concatemers takes place (12,19). These cascade rolling-circle amplification (CRCA) reactions appear to be limited only by the available dNTPs. From what is currently known about CRCA reactions, it is plausible that the ability of displaced strands to act both as primers and as replication substrates results in the reaction being fully self-sustaining.

Nucleic acid amplification reactions are central to DNA-based diagnostic methods, and there have been several demonstrations of rolling-circle replication-based diagnostic strategies (5,12,18). Probably the most straightforward approach for using CRCA for gene or mutation detection is to make use of padlock oligonucleotides (13). These are designed to hybridize to the target such that they are circularizable with a DNA ligase. In this way, circularization of the padlock is dependent on the presence of target DNA. We have compared the ability of the strand-displacing \textit{Bst} DNA polymerase to amplify and multimerize both circular and linear padlocks. This has led to a novel strategy for the amplification of specific nucleic acid sequences and also a method for reducing the background DNA synthesis in CRCA.

MATERIALS AND METHODS

Bacterial Strains

\textit{E. coli} strain PNG801 is a mutagenized derivative of the wild-type \textit{E. coli} strain W1485 (supplied by N. Kleckner, Harvard University). The \textit{Tn}10-derived mini-transposon 103, which encodes a kanamycin resistance determinant, was inserted into the genome by the method of Kleckner et al. (8). PNG801 is a random isolate from the mutant poolate, and the location of the transposable element in the genome is unknown. \textit{E. coli} strain DH5α™ was supplied by Life Technologies (Melbourne, Victoria, Australia). Genomic DNA was extracted from these strains by the SDS, EDTA, proteinase K method described by Silhavy et al. (17).

Oligonucleotides

All oligonucleotides were purchased from GeneWorks (Adelaide, South Australia, Australia) or Life Technologies and were synthesized using standard phosphoramidite chemistry. Oligonucleotides used as padlocks and/or synthetic amplification targets were gel purified to homogeneity by the suppliers, while all others were supplied as desalted preparations. Padlock and spacer oligonucleotides used in ligation
reactions were either phosphorylated during synthesis or after synthesis using T4 polynucleotide kinase (15).

**Sensitivity Determination of CRCA Using Gel-Purified Circularized Padlock**

It should be noted that oligonucleotides that are designed to anneal to a target such that they may be circularized with a ligase we have termed “padlocks” or “padlock oligonucleotides”. The term padlock probe has been avoided because we have not used the molecules as probes. The padlock B2B (150 pmol) (5'-ACGATCAGCGGTCCACAGTAGAGCAGGTGATCAAAAGAACGAATCTTCTACAGCACGTCATCAATGAGTGACGGTGATAGGACTACCTT) was mixed with 150 pmol target oligonucleotide (5'-CGGGATTCGTTGACCGGTCATCGGGCTCGATCCTTG-3') in a 100-µL reaction containing T4 DNA ligase buffer (Roche Diagnostics Australia, Brisbane, Queensland, Australia). The reaction was heated for 5 min at 95°C and then cooled slowly to room temperature. T4 DNA ligase (5 U; Roche Diagnostics Australia) was then added, and the mixture was incubated at room temperature overnight. The amplification reaction products were electrophoresed through a 18% denaturing polyacrylamide gel, stained with ethidium bromide, and the band representing the circular molecules was excised from the gel. The molecules were eluted from the gel by the "crush and soak" method described by Sambrook et al. (15).

The identity of the isolated molecules was confirmed by digestion with the restriction enzyme *TaqI* and differential electrophoresis mobility in denaturing 6% and 10% polyacrylamide gels. The basis for the latter technique is that circular DNA has different apparent molecular weights when electrophoresed in different concentrations of acrylamide when linear molecular weight standards are used. The concentration of the final circularized product was estimated spectrophotometrically.

The purified circular DNA was amplified in a 60-µL reaction that contained 16 pmol each of primers B4C (5'-TTGATAGGTCATCAGCAT-3') and P5C (5'-AGAGCTTGCCGCTATCAGC-3'), 10 mM KCl, 10 mM (NH₄)₂SO₄, 20 mM Tris-HCl, pH 8.8, 200 µM each dNTP, 2 mM MgSO₄, and 1% Triton® X-100. The mixture was heated to 94°C for 30 s then equilibrated at 60°C for 5 min. At this point 4 U *Bst* DNA polymerase were added, and the reactions were incubated at 60°C for 2 h.

**Monitoring of CRCA in Real Time**

This experiment was carried out using the same batch of gel-purified circularized padlock as was used for the determination of CRCA sensitivity. The amplification conditions were also the same as described, with the exception that 15 µg bovine serum albumin (BSA) and 1 µL SYBR® Green I (1:1000 dilution; Molecular Probes, Eugene, OR, USA) were included in each reaction. The reaction mixtures were denatured, and *Bst* DNA polymerase was added, which point they were placed into cuvettes and then loaded into a LightCycler® (Idaho Technology, Salt Lake City, UT, USA). This was programmed to incubate the samples at 60°C and take fluorescence readings at 1-min intervals.

**Amplification from Other Templates**

Both circular and linear templates were amplified as follows. The reactions contained 16 pmol each primer, 10 mM KCl, 10 mM (NH₄)₂SO₄, 20 mM Tris-HCl, pH 8.8, 200 µM each dNTP, 2 mM MgSO₄, and 1% Triton X-100. The template was added in a volume of 1–2 µL, and the final reaction volume was 60 µL. The mixture was heated to 94°C for 30 s and then equilibrated at 55°C–60°C for 5 min. At this point, 4 U *Bst* DNA polymerase was added, and the reactions were incubated at the equilibration temperature for 1.5–3 h.

**Gel Electrophoresis and Hybridization**

The amplification reaction products were electrophoresed through 2% agarose gels in TAE or TBE buffer (15) and visualized by ethidium bromide staining. When blotting was carried out, the gels were soaked in 0.5 M NaOH, 1.5 M NaCl for 30 min, then in 1 M Tris-HCl, pH 8.0, 1.5 M NaCl for 30 min. The nucleic acids were then capillar transferred to Hybond®-N+ membrane (Amersham Pharmacia Biotech, Sydney, NSW, Australia) according to the manufacturer’s instructions. Oligonucleotide probes were 3'-labeled with digoxigenin (DIG)-dUdUTP using terminal transferase (Roche Diagnostics Australia). Membranes were prehybridized in 5–10 mL Rapid-HybTM buffer (Amersham Pharmacia Biotech) at 42°C for 30 min and were hybridized at 42°C overnight. The membranes were washed and developed using anti-DIG-alkaline phosphatase conjugate and CPDStar® (Roche Diagnostics Australia) according to the manufacturer’s instructions.

**Cloning and Sequence Analysis**

The amplification reaction products were first purified through Wizard® PCR DNA purification columns (Promega, Madison, WI, USA). The purified products were then ligated into pGEM®-T (Promega) at 16°C for 3 h, and the ligated products were introduced into *E. coli* DH5α by electroporation. Inserts were sequenced using dye-labeled terminators and a model 373A sequencer (Applied Biosystems, Foster City, CA, USA).

**Affinity Purification and Amplification of Circularized Padlocks Using Streptavidin-Coated Superparamagnetic Beads**

The annealing and ligation reactions were carried out in a final volume of 50 µL, which contained padlock FV2 (5'-AGGAAATAACGGATATTGGTCCCTTGGCGCGTGAGCTATATGGGAGCTATGAGTTCTTATAGGACTACCTTTCAATCTGTAAGAG-3'), synthetic target FVWT (5'-TTAGTTCAAGGCAAAATACCTGTATTTCCTGCGTGTCAGGATATCAGGCAGATG-3'), synthetic target FVWT (5'-TTAGTTCAAGGCAAAATACCTGTATTTCCTGCGTGTCAGGATATCAGGCAGATG-3'), synthetic target FVWT (5'-TTAGTTCAAGGCAAAATACCTGTATTTCCTGCGTGTCAGGATATCAGGCAGATG-3'), synthetic target FVWT (5'-TTAGTTCAAGGCAAAATACCTGTATTTCCTGCGTGTCAGGATATCAGGCAGATG-3'), synthetic target FVWT (5'-TTAGTTCAAGGCAAAATACCTGTATTTCCTGCGTGTCAGGATATCAGGCAGATG-3') in ligation buffer (20 mM Tris-HCl, pH 8.3, 25 mM KCl, 10 mM MgCl₂, 0.5 mM NAD, and 0.1% (v/v) Triton X-100). The mixture was heated to 94°C, 5 U Ampligase (Epicentre Technologies; supplied by Austral Scientific, Sydney, Australia).

NSW, Australia) were then added, and incubation at 94°C continued for 3 min. The reaction was then cooled to 60°C and incubated at that temperature for 1 h.

The streptavidin-coated superparamagnetic beads were supplied by Dynal Biotech (Melbourne, Victoria, Australia). The supplied bead suspension (125 μL) was transferred to a 1.5-mL microcentrifuge tube and was washed four times in 125 μL 2× BW buffer (10 mM Tris-HCl, pH 7.5, 1 mM EDTA, 2M NaCl). Pelleting of the beads was carried out using a magnetic platform. After the final wash, the beads were resuspended in 250 μL 2× BW buffer. Fifty microliters of the bead suspension were then mixed with the entire 50 μL products of the annealing and ligation reactions. The mixture was incubated at room temperature for 30 min with occasional agitation. The beads were then pelleted using a magnetic platform, the supernatant was removed, and the beads were washed twice in 1× BW buffer (5 mM Tris-HCl, pH 7.5, 0.5 mM EDTA, 1 M NaCl), then three times in 100 μL 0.1 M NaOH to remove any hybridized molecules, then twice in TE buffer (10 mM Tris-HCl, pH 8.0, 1 mM EDTA). For each wash, the beads were suspended in the relevant buffer for 5 min and then pelleted using a magnetic platform. After the washes, the beads were suspended in 50 μL TE buffer. One microliter of the suspension was used as a template in amplification reactions, as described above, using primers PadFV3 (5'-GA-AATTCAATAGTCCCC-3') and PadFV4 (5'-CGCGGTGAGCTATATAT-3').

Affinity Purification and Amplification of Circularized Padlock Using Streptavidin-Coated Microplates

This procedure also made use of the padlock FV2, the biotinylated spacer LigW, and the amplification primers PadFV3 and PadFV4. The targets used were the synthetic oligonucleotide FVWT, human genomic DNA, and E. coli genomic DNA (negative control). These oligonucleotides are based around the Factor V Leiden mutation, and the human DNA used was homozygous normal at this site.

Annealing and ligation reactions containing padlock, biotinylated spacer oligonucleotide, and target DNA were carried out as described above. The entire reaction products were transferred to streptavidin-coated microplates (NEN® Life Science Products, Boston, MA, USA) and incubated at room temperature for 30 min. The wells were then washed twice with 1× BW, twice with 0.15 M NaOH, once with 1× BW, 12 times with phosphate-buffered saline containing 0.05% (v/v) Tween®20 (PBST) (15), and finally once with TE. All washes were 200 μL.
in volume. The amplification reaction components (as described above) were then added to the wells, and the microplates were then heated at 98°C for 3 min, cooled to 60°C, and 4 U Bst DNA polymerase were added to each well. The microplates were then incubated at 60°C for 90 min.

RESULTS

Minimal Circular Template Is Required for the Synthesis of Abundant Linear Amplimer by Bst DNA Polymerase

We are interested in the diagnostic potential of CRCA. It is now clear that if short single-stranded circles of DNA are combined with Bst DNA polymerase, two primers, and dNTPs in a suitable buffer and incubated at approximately 60°C, a self-sustaining exponential amplification reaction takes place (12,19). We have attempted to accurately determine the sensitivity of this reaction. A single-stranded circular
template was constructed through hybridization of a padlock to a synthetic template and ligation of the padlock ends. The circular product was then gel-purified, and the abilities of various dilutions of this circular molecule to serve as rolling-circle templates were determined. It was found that approximately 10 circular molecules were all that was required for amplification to take place (Figure 1). As expected, the reactions yielded large quantities of variable-length concatemers. The self-sustaining nature of the reactions was evident from the lack of any direct relationship between the amounts of circular template DNA and product.

**Relationship between Template Concentration and Reaction Kinetics in CRCA Is Similar to PCR**

In the case of PCR, target quantitation is often done by determining the number of amplification cycles before a predefined quantity of amplimer is synthesized. To determine if CRCA is amenable to a similar quantitation strategy, CRCA reactions containing different dilutions of template were carried out in a LightCycler. This device measures the accumulation of dsDNA in real time by making use of the double-stranded nucleic acid-specific fluorescent dye SYBR Green I. Figure 2 shows that there is a clear relationship between the log of the template concentration and the time taken to reach a threshold level of amplimer.

**Figure 3. Products of LIMA reactions using different primer sets.** LIMA reactions were carried out using 1 pmol target and 16 pmol each primer and 3 h incubation at 60°C. (A) Sequences of the target (B2B padlock) and primers. (B) Ten microliters of the reaction products were electrophoresed in 2% agarose. Lane M, molecular weight markers; lane C, no template; lane 1, primers P1 and B4C; lane 2, primers P5C and B4C; lane 3, primers P5C and B4H; lane 4, primers P5C and B4I; lane 5, primers P5D and B4C; lane 6, primers P5D and B4I; and lane 7, primers P5E and B4I.
In the Presence of Linear Template DNA, CRCA Reactions Are Prone to Background

Padlocks are designed to anneal to the target sequence such that they are circularizable with a conventional DNA ligase (13). As circularization can only occur in the presence of the target, this provides a strategy for CRCA-based gene detection or, if ligation can be made sufficiently specific, a strategy for mutation diagnosis.

In practice, however, it is very difficult to devise reaction conditions in which DNA synthesis does not occur in the absence of target DNA. Electrophoresis of this background material commonly revealed the “ladder” effect of CRCA products that is indicative of variable-length multimers (data not shown).

Linear Molecules May Be Amplified under CRCA Reaction Conditions

The background problems experienced lead us to hypothesize that circular templates are not necessary for exponential DNA amplification under CRCA reaction conditions. This was shown to be the case with seven different synthetic 90-mer templates (data not shown). A series of “linear template”

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**Figure 4. Analysis of LIMA products.** The products of four LIMA reactions (A–D) were analyzed by restriction digestion/electrophoresis and sequence determination. The reactions were carried out using 16 pmol each primer and 1 pmol target (padlock B2B) and were incubated at 60°C for 3 h. The 2% agarose gels are shown in A(i)–D(i). Lane M, molecular weight markers; lane U, undigested reaction products; lane S, *Sac*III-digested amplimer; lane A, *Alu*I-digested amplimer; lane E, *Eco*RI-digested amplimer; lane T, *Taq*I-digested amplimer. The reaction products were also cloned, and partial sequence was obtained from one clone from each LIMA reaction. The relationships between the repeating units in the clone sequences and the target and primer sequences are depicted in A(ii)–D(ii). The boxes show the position of the repeating units. The numbers to the right of the boxes show the number of consecutive repeats in the sequence obtained. The arrows show the directions of the sequence reads. Mutations with respect to the target are indicated within the boxes (B(ii) only). The discontinuous box in D(ii) indicates a deletion in the repeating unit with respect to the target.
were carried out using different (Figure 5). When similar experiments respect to template concentration. It was concluded that, in general, LIMA is sev-
tants. Despite this variability, it may be
function of the sequences of the reac-
tion products. In general, the repeating units were analyzed by electrophoresis. Fig-
ure 6 shows the results.

There was no evidence for amplification when \( E. coli \) DH5\(\alpha \) DNA was used as the template. However, when \( E. coli \) PNG801 DNA was used, amplifi-
cation took place. It was deduced from the single molecular weight species in the \( \text{HindIII} \)-treated samples that, as expected, multiple evenly spaced \( \text{HindIII} \) sites were present in the PNG801-de-
ved amplimers. The derivation of the amplimers from the target was con-
minated by hybridization with a target-

duced probe (data not shown). It was con-
cluded that specific amplification and multimerization of the target se-
quence had taken place.

It is noteworthy that the sizes of the repeating units varied from reaction to reaction, even though the target se-
quence and primers were the same. The molecular basis for this was determined by sequence analysis of three of the re-
action products. Each product contained both primer binding sites and the expected 120-bp sequence between these sites. However, the repeating units from two of the reactions contained additional-
sequences, up to 77 bp in size, clearly
derived from the mini-transposon but from outside the primer binding sites.

### Removal of Linear Padlock before Amplification Reduces LIMA-Derived Background

It is now clear that LIMA is the basis for background reactions in padlock-de-
pendent embodiments of CRCA. Lig-
ation of a padlock to a rare target se-
quence yields a mixture in which the
linear padlock is present in great molar
excess over the circularized padlock, thus offsetting the lower sensitivity of
LIMA as compared with CRCA. In ad-
dition, LIMA will take place in the ab-
ance of target sequence. It may there-
re be predicted that removal of the linear padlock before amplification would reduce the background. This was
tested using a padlock and biotinylated spacer oligonucleotides designed such that the spacer hybridizes to the target
between the padlock arms, thus allow-
ing the affinity-based separation of cir-
cularized padlock from linear padlock. This strategy was tested in two different experimental systems. In the first, a
padlock/spacer set were designed to hy-
bridize to a synthetic oligonucleotide

target, and the effect of incorporating a circle purification protocol on the nature of the amplimer was determined. It may be
seen in Figure 7A that the circle pu-
rification prevented LIMA-based amplification of the linear padlock alone (lanes 1 and 3) and eliminated LIMA derived background in the CRCA reactions (lanes 2 and 4; lane 2 contains a mixture of LIMA and CRCA products). In the second experiment, the ability of CRCA to carry out high-sensitivity gene detection was demonstrated. The protocol was designed to detect the human Factor V-encoding gene and, as expected, gave a positive result from human DNA and a negative result from E. coli genomic DNA (Figure 7B).

**DISCUSSION**

It has been shown by several researchers that strongly strand-displacing DNA polymerases are able to exponentially amplify multimeric DNA (12,19). The role of rolling-circle replication in CRCA is to create a multimeric molecule that is then amplified. CRCA is therefore something of a misnomer because it may be predicted that any reaction that yields linear multimers may be substituted for the rolling-circle reaction. Although the precise mechanism of LIMA is obscure, our current model is that it is a form of in vitro evolution in which a variety of inefficient replication initiation events occurs early in the reaction. This results in Darwinian selection of molecules that are more easily replicated. As would be predicted from the rapid and exponential kinetics of CRCA, such molecules are usually multimeric. It is highly likely that the initial formation of easily amplifiable multimers is the limiting factor for the sensitivity of the reaction and, to a great extent, defines the reaction duration. The precise events that occur early in LIMA reactions will be very difficult to elucidate, but it may be deduced from the nature of the reactants and the fact that LIMA takes place at all that “illegitimate” events such as primer invasion of dsDNA must be involved. The random nature of the early stages of the reaction would account for the variabilities found with respect to both sensitivity and the end points of the multimer repeating units. The production of RNA multimers from linear templates has been previously reported by Krupp (9), and a rolling-circle-like mechanism us-

![Figure 6. Use of LIMA to amplify a specific sequence from an E. coli genomic preparation.](image)
ing a non-covalently closed template was suggested. A similar mechanism cannot be ruled out for LIMA, although the smaller size of the repeating units in LIMA as compared with CRCA would suggest that this is not the case.

The use of LIMA in diagnostic applications is an intriguing possibility. We have demonstrated reproducible and selective amplification of a gene fragment in an E. coli genome sample, thus indicating a useful level of specificity. The variable nature of the relationship between the starting material and the reaction products, and the low sensitivity as compared with CRCA, would suggest that applications would need to be individually optimized and that using LIMA for high sensitivity gene detection (e.g., direct diagnosis of infectious agents in clinical samples) may be problematic. However, these disadvantages may well be offset by the small number of components required and the isothermal nature of the reaction. We have not yet attempted to determine whether LIMA can be made specific to the level of single nucleotide polymorphisms, but if proved to be feasible, then applications in the area of routine genotyping may be envisaged.

It may be inferred from the ability of Bst DNA polymerase to mediate LIMA and the great molar excess of padlock over target DNA in any sensitive CRCA-based gene or mutation detection reaction that the removal of linear padlock before CRCA would reduce background problems. This was tested by affinity purifying circular DNA before amplification. As expected, background DNA synthesis was greatly reduced. The use of a biotinylated spacer would be expected to increase the effective specificity of the ligation reaction and eliminate LIMA-derived background. This is because only in the presence of the target would the spacer be brought into close proximity with the padlock and, as a result, any circular molecules formed by annealing of the padlock to a spurious target will not include the biotin molecule and will not be captured before amplification. These results are consistent with those reported by Zhang et al. (19), who have used a target purification protocol to reduce the background of DNA amplification reactions including those based on rolling-circle replication. When applied to CRCA, their protocol also would be expected to reduce the amount of linear padlock present.

It is possible that LIMA and/or CRCA may be useful in applications outside DNA-based diagnostics. It has already been established that concatemerization of response elements can enhance their affinity for transcription factors (20), and this phenomenon has been exploited for affinity chromatography and functional analysis of transcription factors (2,7,16). Techniques previously used to generate concatemers include DNA ligation (7), direct cloning of repetitive DNA (10,14), or amplification of large oligonucleotides (6). LIMA and/or CRCA could easily be used in place of these reactions. Indeed, Daubendiek and Kool (3) have recently demonstrated the production by rolling circle of self-cleaving concatemers of catalytic RNA molecules.

In summary, we have demonstrated that Bst DNA polymerase readily multi-

Figure 7. CRCA from affinity-purified circular DNA templates. The amplification products were electrophoresed in 2% agarose. The DNA species used are listed in the Materials and Methods section. (A) The effect on background of circular template purification using superparamagnetic beads was determined. Lane M, molecular weight markers; lane 1, no target or ligase added annealing and ligation reaction, no affinity purification; lane 2, standard annealing and ligation reaction, no affinity purification; lane 3, as for lane 1, with affinity purification step; lane 4, as for lane 2, with affinity purification step. (B) Background-free gene detection in a human genomic sample using circular template purification in a streptavidin-coated microplate was demonstrated. Lane M, molecular weight markers; lane 1, 200 ng human genomic DNA, no ligase added annealing and ligation reaction; lane 2, 100 ng E. coli DH5α genomic DNA used as target; lane 3, 200 ng human genomic DNA used as target; lane 4, 20 ng human genomic DNA used as target; lane 5, no target added; and lane 6, 1 pmol synthetic target.
merizes and exponentially amplifies both circular and linear DNA with no need for temperature cycling or additional enzymes. The potential applications of both reactions in DNA-based diagnostics are worth investigating fully. While there is no doubt that CRCA is more sensitive and predictable than LIMA, the necessity of using a padlock or similar device before amplification adds inherent complexity to CRCA-based embodiments. This complexity is accentuated by the background problems caused by LIMA and the measures needed to avoid this. In contrast, LIMA is extremely simple and straightforward to carry out—all that is required are the primers and the polymerase.

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REFERENCES