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SOE-LRed: a simple and time-efficient method to localize genes with point mutations onto the *Escherichia coli* chromosome

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Abstract

We report a powerful method to replace wild type genes on the chromosome of *Escherichia coli*. Employing a unique form of PCR, we generate easily constructible gene fusions bearing single point mutations. Used in conjunction with homologous recombination, this method eliminates cloning procedures previously used for this purpose.

Keywords

SOE PCR; Transformation; *E. coli*; *dinB*; Lambda Red; Point Mutation

Methods, Results, Discussion

There has been an effort by many labs to develop a quick and easy protocol for gene replacement on the bacterial chromosome (Datsenko and Wanner, 2000; Murphy *et al.*, 2000; Heermann *et al.*, 2008; Link *et al.*, 1997; Gordon *et al.*, 1997). Though a rapid method for chromosomal gene replacement was recently reported by Heermann *et al.* (2008), it includes two major limitations: gene replacement requires two sequential recombination events and must be done in strains bearing the *rpsL150* mutation in the ribosomal protein S12. We present here an efficient and streamlined method, which relies on a single recombination event and could be carried out in any bacterial strain expressing the Lambda red proteins (Datsenko and Wanner, 2000).

We report for the first time the use of Splicing by Overlap Extension (SOE) polymerase chain reaction (PCR) to join noncontiguous DNA sequences with the purpose of constructing a fusion product to replace wild type genes with ones bearing point mutations. We call this method SOE-LRed to indicate the merger of SOE PCR (Horton *et al.*, 1989) and lambda Red (Datsenko and Wanner, 2000) methodologies to localize genes with point mutations onto the chromosome. SOE PCR is a variation of traditional PCR that since its inception has been extremely valuable in joining noncontiguous DNA sequences from different sources (Horton *et al.*, 1989). Notably, it eliminates traditional construction of recombinant DNA molecules, which is not only laborious, but also generally introduces unnecessary DNA sequences to the final product (Murphy *et al.*, 2000).

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Fusion of sequences via SOE PCR relies on two sequential PCR amplification reactions and is directed by primers that introduce a common linker region (CLR). In the first PCR reaction, the CLR is incorporated separately into each gene. Amplification of the full-length fusion sequence draws on the CLR for effective annealing of the two noncontiguous sequences, as well as the forward primer of sequence 1 and the reverse primer of sequence 2 (Figure 1).

In this report we specifically describe the construction of chromosomal *dinB* alleles encoding derivatives of DNA Polymerase IV, an evolutionarily conserved specialized DNA polymerase (Friedberg *et al.*, 2006). *dinB* templates containing the point mutations were available on low-copy number plasmids, which were generated by site-specific mutagenesis of the wild-type sequence using the GeneTailor™ Site-Directed Mutagenesis System (Invitrogen). SOE PCR was utilized to fuse the *dinB* sequences of interest to a selectable marker, the *cat* gene, enabling selection for chloramphenicol resistant (Cm^R) recombinants. Flippase Recognition Target (FRT) sites, which permit the excision of the resistance marker by the site-specific recombinase, flippase (FLP), flank the *cat* gene (Datsenko and Wanner, 2000). Primer design was carried out such that the native *cat* and *dinB* promoters, as well as FRT sites upstream and downstream of *cat*, (Figure 1) were included in the fusion sequences. Replacement of wild-type chromosomal genes with the amplified fusion sequences relied on the lambda Red recombinase system described by Datsenko and Wanner (2000).

Gradient PCR cycle conditions were employed for both amplification reactions (Dieffenbach and Dveksler, 1995) and primer sequences are listed in Table 1. Primers added approximately 50 bp at the 5' end and 80 bp at the 3' end of the fusion sequences. These regions, termed insertion site homology regions (ISHR), are homologous to genes located upstream and downstream of chromosomal *dinB* (*mhbA* and *yafN*, respectively) and direct the replacement of the wild-type chromosomal *dinB* sequence at this locus. Primers also included a 54 bp region of *dinB*'s native promoter, which served as the CLR. Total volume of the PCR reactions was 25 μ L, using GoTaq Master Mix (Promega), and 1 μ L each of a forward and reverse primer at 10 μ M. Template for the first round of amplification was diluted plasmid DNA (~2 μ g/mL). After the first amplification reaction, PCR products were gel purified to a final concentration of about 20 ng/ μ L. Two microliters of each amplicon were used as template for the second round of SOE PCR, and the obtained full-length size product was verified by agarose gel electrophoresis. Examples of all PCR products obtained from the first and second amplification reactions are shown in Figure 2.

The obtained fusion sequences were introduced into Δ *dinB*::Kan^R JW0221 cells (Baba *et al.*, 2006) by electroporation at 1.8 mV for 5 msec following standard protocols (Ausubel *et al.*, 2001). Cells contained the plasmid expressing lambda Red proteins (pKD46) and were made electrocompetent as previously described (Datsenko and Wanner, 2000). After incubation for 1 to 1.5 hours at 37°C to permit *cat* gene expression, cells were plated on LB agar containing 20 μ g/mL of chloramphenicol (Cm), and incubated for 20 hours at 37°C. Cm resistant (Cm^R) colonies were purified on the same plates and screened for sensitivity to kanamycin (Kan^S), the marker replacing *dinB* in JW0221. Since the *dinB-cat* fusion products contained sequences homologous to regions upstream of the *dinB* promoter and downstream of the *dinB* open reading frame (ORF), recombination into the chromosome replaces the Kan^R sequence with the *dinB* fusion products containing the desired point mutations (Figure 3). Therefore, colonies that were both Cm^R and Kan^S were examined for unique junction sequences by standard PCR to confirm that the replacement had taken place at the desired location on the chromosome. DNA sequencing of a PCR amplicon spanning the *dinB* gene and promoter region confirmed the presence of the desired *dinB* allele on the chromosome. All three Cm^R colonies analyzed were also Kan^S because the Kan^R gene was

replaced with the *dinB(D103N)* allele in JW0221 (Wagner *et al.*, 1999). However, just 2 of 11 Cm^R colonies analyzed were also Kan^S when introducing the *dinB(Y79A)* allele (Jarosz *et al.*, 2009) on the chromosome. It is possible that the frequency of obtaining the desired point mutation on the chromosome depends on the efficiency of the SOE PCR reactions. Nevertheless, the desired alleles were recombined into the chromosome in the time frame of a week. The *dinB-cat* fusion was transduced (Miller, 1972) from the JW0221 strain into other *E. coli* strains with P1 virulent phage.

Here we have shown, through use of SOE-LRed, that chromosomal genes may be replaced with sequences encoding variant gene products of interest. Although we have used chloramphenicol, any selectable marker may be chosen. Furthermore, use of the readily available Kan^R Keio collection (Baba *et al.*, 2006), allows for positive selection throughout the process. Gene replacement permits the removal of the Kan^R marker sequence and insertion of the selectable marker linked to the gene of interest. Although we used *dinB*, the same experiment can be carried out with essential genes. Depending on the efficiency of the SOE PCR reactions, it may be necessary to screen a larger number of recombinant colonies for appropriate junctions with PCR.

This technique is very powerful because of its simplicity and time efficiency. It permits the *in vivo* study of strains encoding genes with a variety of point mutations, and allows for their expression at physiologically relevant concentrations. The expression of genes of interest at chromosomal levels facilitates genetic analyses such as structure-function, gene dominance, alteration of activity, and studies of gene expression.

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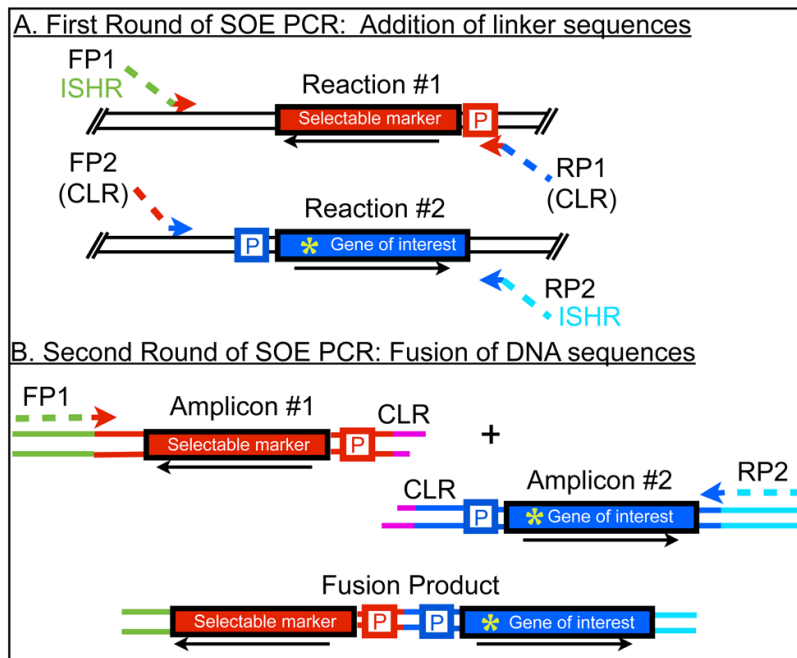


Figure 1. Schematic of products generated by SOE PCR

A. First round of SOE PCR: addition of linker sequences. In these reactions (#1 and #2), primers add a common linker region (CLR) to the genes to be fused. An insertion site homology region (ISHR) is also added to the amplicons of the first reaction; this sequence will later direct the insertion of the gene fusion at a specific locus on the bacterial chromosome. **B. Second round of SOE PCR: fusion of DNA sequences.** In this reaction, the fusion product is generated, relying on the CLR to anneal the two originally noncontiguous sequences. Genes are depicted as red or blue rectangles and flanking sequences as lines. The CLR is shown in pink lines and is common to both amplicons to be fused. P indicates the promoters of each gene and the arrow the orientation of transcription. FP stands for forward primer and RP stands for reverse primer. Green asterisk indicates a point mutation in the gene of interest.

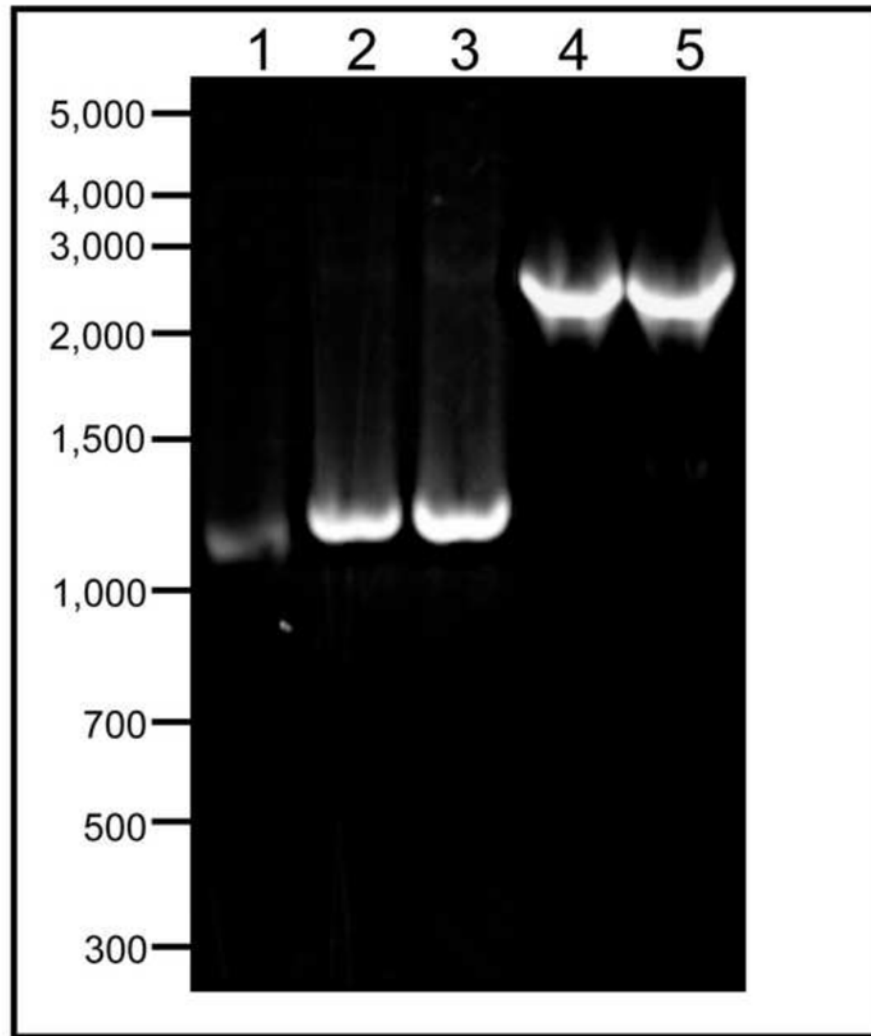


Figure 2. Amplification of DNA sequences using gradient cycling conditions
Amplification of *cat*-CLR, lane 1; *dinB(D103N)*-CLR, lane 2; *dinB(Y79A)*-CLR, lane 3; *dinB(D103N)*-*cat* fusion sequence, lane 4; and *dinB(Y79A)*-*cat* gene fusion, lane 5. A gradient cycle was used in both phases of SOE PCR. Products were separated in a 1% agarose gel in Tris Acetate Buffer (Ausubel *et al.*, 2001). Lanes 1–5 contain 3 μ L of DNA.

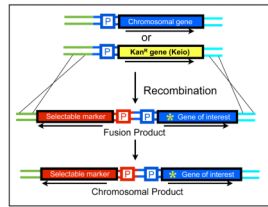


Figure 3. Schematic of the recombination event that replaces the chromosomal gene with the fusion construct

The reaction relies on homologous sequences in the fusion constructs and chromosomal regions flanking the gene to be deleted, depicted as a blue rectangle. Gene deletions of the Keio collection may also be used (shown as a yellow rectangle), since the flanking regions of deleted genes are still present and can be used as the ISHR. Green asterisk indicates a point mutation in the fusion construct. Cells in which the recombination has occurred are resistant to the selectable marker and bear the point mutation of interest. If the Keio strain is used, recombinants would also be resistant to kanamycin.

Table 1

SOE PCR Primer Sequences

Details	Sequence
FP1 Adds <i>mhaB</i> ISHR to 5' end; amplifies <i>cat</i> gene	5'CTGGTGAAAAGCTGGATAAGCAGCAG GTGCTTTCGCAGCGAACGCGTTAATGAGC GATTGTGTAGGCTGG 3'
RP1 Adds <i>dinB</i> sequence as common linker; amplifies <i>cat</i> gene	5'CTGGTAAAGTATACAGTGATTTACAGGGT TTGAGAAATGCGTAAAGATTCAGCATGCC ATGGTCCATATGAATATCCTCC 3'
FP2 Amplifies <i>dinB</i> with promoter; contains linker sequence	5'ATGCTGAATCTTTACGCATTTCTCAAACC CTGAAATCACTGTATACTTTACCAGTGT TGAGAGGTGAGCAATGC 3'
RP2 Amplifies <i>dinB</i> ; adds <i>yafN</i> ISHR at to 3' end	5'GACCGATTTTTACGCGAGAATTCGATGC ATACAGTGATACCCTCATAATAATGCACAC CAGAAATACATAATAGTATAC 3'