An efficient and economic enhancer mix for PCR

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Abstract

Polymerase chain reaction (PCR) has become a fundamental technique in molecular biology. Nonetheless, further improvements of the existing protocols are required to broaden the applicability of PCR for routine diagnostic purposes, to enhance the specificity and the yield of PCRs as well as to reduce the costs for high-throughput applications. One known problem typically reported in PCR experiments is the poor amplification of GC-rich DNA sequences. Here we designed and tested a novel effective and low-cost PCR enhancer, a concentration-dependent combination of betaine, dithiothreitol, and dimethyl sulfoxide that broadly enhanced the quantitative and/or qualitative output of PCRs. Additionally, we showed that the performances of this enhancer mix are comparable to those of commercially available PCR additives and highly effective with different DNA polymerases. Thus, we propose the routine application of this PCR enhancer mix for low- and high-throughput experiments.

Keywords: Polymerase chain reaction; GC-rich sequence; Enhancer; Additive; Promoter PCR; Genomic PCR; PCR template; Taq DNA polymerase

The polymerase chain reaction (PCR) was developed in the 1980s by Kary Mullis and Fred Faloona [1,2]. Starting with the biotechnological application of thermostable DNA polymerases [3], PCR has become a fundamental technique in molecular biology. There are ever-increasing needs for further improvements of PCR protocols for low-cost and efficient high-throughput approaches in a wide range of applications ranging from quantitative analysis at the genome- or transcriptome levels to routine diagnostic purposes [4,5]. Large-scale PCR experiments require broadly applicable and reliable reaction conditions for establishing cost-effective production pipelines. Taq DNA polymerase, originally purified from the thermophilic bacterium Thermus aquaticus [6], is widely used, since it can be produced in every standard laboratory at low-cost [7–9]. One major factor limiting the output of PCR routines is that a number of DNA sequences are poorly or not amplifiable under standard reaction conditions, either because of their intrinsic properties to form secondary structures, and/or because of their high GC-content. Improvements of the PCR conditions can be achieved by modifying the classical reaction conditions, for example, by performing “touchdown” PCR, consisting of a stepwise reduction of the annealing temperature for each cycle [10], or by the use of modified DNA polymerases for carrying out “hot-start” reactions [11]. Typically, to overcome amplification problems of GC-rich DNA, the addition of substances that enhance the specificity and/or the yield of the PCR is necessary. The most prominent PCR enhancing additives that are currently used are either betaine [12], small sulfoxides like dimethyl sulfoxide (DMSO, [13]), small amides like formamide [14] or reducing compounds like β-mercaptoethanol or dithiothreitol (DTT, [10]). However, their capacity to significantly improve PCR yields mainly for
high-throughput experiments is marginal. Commercial enhancers have led to better results but with two major drawbacks, their cost and the fact that their chemical composition is unknown.

Mammalian promoter sequences often contain highly GC-rich regions, which are difficult to amplify under standard reaction conditions [15]. In this study, we tested the efficacy of concentration-dependent combinations of different PCR additives for a reliable amplification of genomic DNA corresponding to a set of human promoter sequences and generated a novel, cheap, and flexible PCR enhancer.

Materials and methods

Primer design. PCR primers for the amplification of ~1000–1600 bp sized DNA fragments from human genomic DNA were designed using the “Primer 3.0” online service [16] on the basis of the human genome annotation build 35.1 (NCBI). The primer sequences, locus information, the overall GC-content, and the size of the expected amplicons are given in the supplementary material Table 1.

Purification of human genomic DNA. Human genomic DNA was prepared from oral mucosa. The mucosal smear was washed with water and dissolved in 400 µl lysis buffer (50 mM Tris–HCl, 10 mM EDTA, and 2% SDS, pH 8.8). The suspension was incubated for 5 min at 65 °C, then supplemented with 250 µl 4.5 M NaCl and cleared by centrifugation. Genomic DNA was recovered from the supernatant by isopropanol precipitation.

PCR conditions. PCRs were performed in a 30 µl volume in 96-well microtiter plates. Reaction buffer contained 65 mM Tris–HCl, 16.6 mM (NH₄)₂SO₄, 3.1 mM MgCl₂, and 0.01% (v/v) Tween 20, pH 8.0. 2.5 U Taq DNA polymerase purified from Escherichia coli according to the method of Engelke et al. [7], 0.6 µmol of each oligonucleotide, and 25 µmol dATP, dTTP, dCTP, and dGTP were added prior to the cycling reaction. The cycling reactions were performed in a PTC-200 Thermocycler (MJ Research) with an initial denaturation for 5 min at 96 °C, followed by the thermal cycles as follows: denaturation step at 98 °C for 15 s, annealing step at 72 °C for 40 s, and an elongation step at 72 °C for 90 s. The annealing step was started at a temperature of 66 °C and declined in 0.5 °C steps for each cycle until a temperature of 56 °C was reached. Subsequently, 30 additional cycles were performed with a constant annealing temperature of 52 °C. The reaction was completed with a final elongation step at 72 °C for 2 min. PCR products were analyzed with agarose-gel electrophoresis and stained with ethidium bromide (Sigma).

Results and discussion

In the context of a systematic project aiming at the functional analysis of promoter elements, we set out to amplify 110 human promoter sequences from genomic DNA using classic touch-down PCR conditions (as described in Materials and methods). We observed that approx. 30% of the promoter regions could not be correctly amplified, either because the PCR products were unspecific or because of the poor yield of the amplicons. Most of these had an overall GC-content of 50–75% (62% in average).

In order to improve these results, we evaluated different PCR enhancing additives for their capacity to promote the amplification of three different gene promoter regions (SIM2, DIP2A, and SLC19A1, please refer to the supplementary material for detailed information) whose GC-content ranged from 71% to 75%. We designed three primer pairs for these promoters (named A for SIM2, B for DIP2A, and C for SLC19A1) and carried out touch-down PCR supplemented with different concentration ranges of the PCR additives betaine [12], dithiothreitol (DTT) [10], dimethyl sulfoxide (DMSO) [13], or formamide [14] as indicated in Fig. 1. We observed that betaine had the best PCR enhancing properties at a concentration of 0.8 M in all PCR samples for primer pairs A, B, and C, whereas DTT and DMSO were less effective since the PCR output was enhanced only for one gene out of three (3.2 mM DTT for primer pair A, or 3.2% DMSO for primer pair B, respectively) (Fig. 1). No PCR enhancing effects were observed by adding formamide to the respective PCRs at any of the indicated concentrations.

On this basis, we generated a 5-times concentrated preliminary combinatorial enhancer solution (preCES-I) composed of 4 M betaine, 16 mM DTT, and 16% DMSO. We included 83 µg/ml bovine serum albumin (BSA) in the solution, since BSA, which has no direct effect on the enzymatic reaction per se, can stabilize enzymes and neutralize inhibitory contaminants that may be present in the DNA.
template preparation or in the reaction buffers [10,17]. Since high compound concentration could potentially inhibit the activity of the Taq DNA polymerase, we additionally tested two other preCESs containing lower concentrations of the respective additives, preCES-II (4 M betaine, 10 mM DTT, and 10% DMSO) and preCES-III, (2 M betaine, 5 mM DTT, and 5% DMSO). To analyze the efficiency of these preliminary enhancer solutions, we selected 12 (9 additional) primer pairs, of which 10 failed to produce adequate PCR products under standard conditions without additive. These primer pairs produced either non-specific products (H and I), prominent additional bands to the expected product (C, D, and L), very low yield (A, E, and K), or no product at all (B and G) (Fig. 2A). Subsequently, PCRs were repeated with these primers with or without preCESs I, II, or III. As demonstrated in Fig. 2A, the output of 10 out of 12 PCRs analyzed was enhanced by at least one of the three preCESs. For primer pairs B and G, which have not resulted in any detectable PCR product, the addition of the preCESs resulted in the amplification of a specific DNA fragment. For primer pairs C, D, H, L, or I, respectively, the preCESs enhanced the specificity of the PCRs, whereas for primer pairs A, E, and K the presence of at least one preCES resulted in a significantly improved product yield. However, the addition of preCESs did not improve the PCR performed with primer pair F, and in one case the addition of preCESs had a negative effect on the PCR yield (primer pair J).

Thus, these experiments clearly demonstrated that the addition of a preCES enhances the yield and/or the specificity of PCRs in virtually all cases, particularly for the amplification of highly GC-rich sequences up to 75%. Among the three tested enhancers, preCES-II containing the intermediate concentrated enhancer solution appeared to perform best, as visualized in Fig. 2B.

In the next step, we further optimized further the compound concentration of the preCES-II. Initially, the 30 μl PCR mixture was supplemented with incremental quantities of preCES-II in 2 μl steps (ranging from 0% to 40% of the final volume) and PCRs were performed with the various primer pairs as indicated (Fig. 2C). Best results in terms of specificity and yield were obtained with addition of 4 μl preCES-II to the 30 μl reaction volume, corresponding to final concentrations of 0.54 M betaine, 1.34 mM DTT, 1.34% DMSO, and 11 μg/ml BSA. Thus, we generated a 5-times concentrated combinatorial enhancer solution termed CES.

In a third step, we compared the efficiency of our CES with those of three commercial PCR enhancer solutions, namely Q-solution (Qiagen), PCR enhancer solution (Invitrogen), and Hi-Spec PCR additive (Bioline). For this comparative analysis, we selected 32 primer pairs designed for the amplification of DNA fragments with a GC-content ranging from 33% to 75% (Fig. 3A). PCRs were performed using the reaction buffer without additives or supplemented with the Q-solution, PCR enhancer solution, Hi-Spec PCR additive or our CES. As demonstrated in Fig. 3A, commercial PCR enhancers could improve 90% of the PCRs. The CES described in this study led to comparable performances in all tested PCRs. A major advantage of the CES is that it is much more economic for laboratory routine applications and that its composition is well described and can thus be tuned whenever necessary for more specific applications. Furthermore, Qiagen’s Q-solution and Invitrogen’s PCR enhancer mix can only be purchased conjoint with the suppliers Taq AB C

Fig. 2. Generation of a combinatorial enhancer solution (CES). (A) Comparative analysis of PCRs without enhancing additives (--) and PCRs supplied with preCES-I, preCES-II, or preCES-III, respectively. Primer pairs are sorted by ascending GC-content of the expected PCR product. Arrows highlight the specific DNA fragments. (B) Like (A), but sorted by descending product specificity and yield. (C) Concentration-dependent application of preCES-II to PCRs performed with primer pairs for the amplification of DNA sequences with varying GC-content.
polymerase (Qiagen) or with a proprietary reaction buffer (Invitrogen). To exclude that the PCR enhancing effects of CES are limited to PCRs performed with our home-made Taq DNA polymerase, we further analyzed the performance of CES with commercial polymerases. Using DNA polymerases like InviTaq (Invitek), HotStartTaq (Qiagen), TaqPlus (Stratagene) or even with Vent polymerase, originally purified from Thermococcus litoralis ([18], New England BioLabs), we amplified eight different genomic DNA fragments with or without CES (Fig. 3B). The majority of PCRs with inadequate products were enhanced and virtually no negative effects resulting from the addition of the CES were observed in all reactions. Thus, our enhancer solution can be used with any of the tested DNA polymerases as indicated. Interestingly, the extremely GC-rich DNA fragments resulting from primer pair A or C (75% and 74%, respectively) that were poorly amplified with the home-made Taq DNA polymerase even in the presence of CES were satisfactorily amplified using CES in combination with the InviTaq or TaqPlus enzyme, respectively (Fig. 3A and B). Finally, to support the broad applicability of this PCR enhancer, we also tested the CES on other types of template DNA like yeast genomic DNA, plasmids, and even glycerol stocks, and detected PCR enhancing effects of the CES (data not shown). The 5-times concentrated CES containing 2.7 M betaine, 6.7 mM DTT, 6.7% DMSO, and 55 μg/ml BSA was stable at −20 °C for at least 3 months. Since different Taq reaction buffers currently in use show diverse performances, we recommend to use a reaction buffer containing final concentrations of 65 mM Tris–Cl, 16.6 mM (NH₄)₂SO₄, 3.1 mM MgCl₂, and 0.01% (v/v) Tween 20 at a pH of 8.8 as described in the Materials and methods section.

In summary, we have demonstrated that the concentration-dependent combination of the known PCR additives betaine, DMSO, and DTT results in a cost-effective PCR enhancer solution showing equivalent performances compared with commercial enhancers, at least under the chosen experimental conditions. Since the CES is composed of low-cost components, the usage of this PCR enhancer solution is especially advantageous and attractive from the economic perspective for large-scale projects and routine applications requiring reliable PCR results.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.bbrc.2006.06.151.

References