Error-suppression mechanism of PCR by blocker strands

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The polymerase chain reaction (PCR) is a central technique in biotechnology. Its ability to amplify a specific target region of a DNA sequence has led to prominent applications, including virus tests, DNA sequencing, genotyping, and genome cloning. These applications rely on the specificity of the primer hybridization, and therefore require effective suppression of hybridization errors. A simple and effective method to achieve that is to add blocker strands, also called as clamp, to the PCR mixture. These strands bind to the unwanted target sequence, thereby blocking the primer mishybridization. Because of its simplicity, this method is applicable to a broad nucleic-acid-based biotechnology. However, the precise mechanism by which blocker strands suppress PCR error remains to be understood, limiting the applicability of this technique. Here, we combine experiments and theoretical modeling to reveal this mechanism. We find that the blocker strands both energetically destabilize the mishybridized complex and sculpt a kinetic barrier to suppress mishybridization. This combination of energetic and kinetic biasing extends the viable range of annealing temperatures, which reduces design constraint of the primer sequence and extends the applicability of PCR.

Introduction

PCR is used in a broad, ever-expanding range of biotechnological applications\textsuperscript{1}. Fidelity of PCR is determined by the specificity of the primer hybridization. In applications, mishybridization leads to unwanted consequences, such as false positives in virus tests and sequencing errors. Given the importance and widespread nature of these applications, methods for suppressing hybridization errors are crucial. A simple and effective method to suppress mishybridization is to add blocker strands, also called as clamp, to the reaction mixture\textsuperscript{2–5}. The blockers bind to the target mutants and suppress mishybridization to the mutant sequence. The hybridization of nucleic acids is essential in many biotechnology such as genome editing\textsuperscript{6}, DNA nanotechnologies\textsuperscript{7}, and RNA interference\textsuperscript{8}. The blocker method may be applicable to these systems.

However, despite its effectiveness, the detailed mechanism by which blockers suppress errors remains unclear. This limits the rational design of the blocker sequence\textsuperscript{4}. One hypothesis is that the blocker strands energetically destabilize the mishybridized state. An alternative hypothesis is that the blocker strands effectively create a kinetic barrier to kinetically suppress the mishybridization. Discerning between these hypotheses would improve our understanding of how PCR works, and suggest design principles for optimized protocols. In this paper, we combine experiments and theory to show that PCR blockers suppress errors in PCR by a combination of energetic and kinetic discrimination. A non-trivial consequence of our theory is that, in the presence of blockers, the range of viable annealing temperatures for the PCR reactions is considerably broadened.

To illustrate the factors that determine the hybridization error, we consider the example of a PCR mixture containing a right template R and a contaminated wrong template W with a mutation in the primer binding region (Fig. 1a). During a PCR cycle, the temperature is lowered from a high denaturing temperature to the annealing temperature \( T_a \). Then, a primer strand P hybridizes to either R or W. Since the primer hybridization is reversible, P repeatedly hybridizes to or dissociates from the template. Eventually, a polymerase binds to the hybridized complex P:R or P:W and elongates P to produce a complementary copy \( \overline{R} \) or \( \overline{W} \), respectively. Important quantities characterizing the PCR are the growth rates \( \alpha_R \) and \( \alpha_W \).
and error rate $\eta$, defined by

$$\alpha_R = \frac{r}{[R]}, \quad \alpha_W = \frac{w}{[W]}, \quad \eta = \frac{\alpha_W}{\alpha_R + \alpha_W}.$$  \hspace{1cm} (1)

Here, $[\cdot]$ denotes a concentration and $r$ and $w$ are the increases in concentrations of $R$ and $W$ in a cycle, respectively. Hence, $\alpha_R$ and $\alpha_W$ represent the fractions of copied strands per template in a cycle. Ideally, one wants to maximize the growth rate $\alpha_R$, also called the PCR efficiency, and at the same time minimize the error rate $\eta$.

The accuracy of conventional PCR relies on primer hybridization to $R$, being energetically more stable than hybridization to $W$, as quantified by the free energy difference $\Delta G$ between $P:R$ and $P:W$ (Fig. 1b). This energetic bias can be increased to reduce $\eta$ by carefully designing the primer sequence and increasing the annealing temperature $T_a$.

However, this approach has an inherent limitation. To see that, we consider the hybridization kinetics (Fig. 1b). The DNA binding rate is usually diffusion-limited and thus does not significantly depend on the sequence\(^9,10\). Hence, we assume that $P$ hybridizes to $R$ and $W$ at the same rate. On the other hand, the dissociation rates depend on $\Delta G$. Repeated hybridization and dissociation of $P$ eventually bring the system to thermodynamic equilibrium, where the amount ratio of the wrong hybridization to the right one is $e^{-\Delta G/k_B T_a}$, where $\Delta G$ is the energetic bias and $T_a$ is the annealing temperature. In the figure, $\Delta G^\ddagger$ is the kinetic bias and $\tau$ is the relaxation time of the binding dynamics.

Equation (2) gives the error rate in this equilibrium limit.

$$\eta_{eq} = \frac{1}{1 + (k_{r_{pol}}^R/k_{r_{pol}}^W)e^{\Delta G/k_B T_a}}$$  \hspace{1cm} (2)

(SI Section S4). $k_{r_{pol}}^R$ and $k_{r_{pol}}^W$ are the elongation rates on $R$ and $W$ by the polymerase. In this case, one can show that the short time error rate is always larger than $\eta_{eq}$\(^11\), see Fig. 1c. In fact, one problem with this approach is that the enzymatic reaction is usually quite efficient and starts elongation before the binding equilibrates. This means that the error rate is usually not as small as one would expect from Eq. (2). Slowing down of the reaction by, for example, reducing polymerase concentration would lower the error rate by allowing sufficient time for equilibration, but at the cost of efficiency.
An alternative strategy is to sculpt a kinetic bias by building asymmetric barriers characterized by a difference $\Delta G^\ddagger$, so that $P$ preferably binds to $R$ (Fig. 1c, right). Theory predicts that such kinetic bias can reduce $\eta$ without sacrificing efficiency$^{11}$. Brief methods

To quantify the error rate, we perform PCR with only a single side of the primer set (Fig. S1). Hence, the product concentration increases linearly, rather than exponentially as in standard PCR (Fig. S2). We mix a primer strand $P$, two variants of 72-nt template DNA strands $R$ and $W$, indicated concentrations of thermostable DNA polymerase, and necessary chemicals for the reactions. We also add blocker strands depending on the experiments. The $R$ and $W$ templates are mixed at the same concentrations $([R] = [W] = 2.5 \text{nM})$, much smaller than the primer concentration $([P] = 100 \text{nM})$. The $P$ strand binds to $R$ without mismatches and to $W$ with a single-base mismatch. In these conditions, the hybridization error is $T_a(T)$ without significantly impairing efficiency, at least for large $m$. 

In contrast, $B_W$ drastically reduced the errors at all the annealing temperatures we tested (Fig. 2c) without significantly impairing efficiency, at least for large $T_a$ (Fig. 2a). At $T_a \approx T_m^{B_W,R} = 56.3^\circ C$, $\alpha_R$ was reduced due to the hybridization of $B_W$ to $R$. We found that the errors are still effectively suppressed at $T_a$ much lower than $T_m^{P,W}$, meaning that fine tuning of $T_a$ is not needed in the presence of blockers.

In contrast, $B_R$ drastically reduced $\alpha_R$ and increased $\eta$. At $T_a < 60^\circ C$, $\eta$ was larger than 50%, meaning that $B_R$ inverted the preference of $P$ hybridization. This setup could be used to amplify rare sequences that would be otherwise difficult to sample.

We used chimeric DNA strands containing LNA bases for the blocker strands, which enhance the specificity of hybridization. The blocker strands with only DNA bases had a limited effect when used with the same concentration and strand length as the chimeric strands (Fig. S7).
Mathematical model. We quantified the PCR kinetics and in particular the role of blockers using a mathematical model (see SI section S4 for details). The model includes reversible hybridization rate of P to the template strands. In contrast, we assume that the blockers are always at chemical equilibrium as their high concentrations make their hybridization and dissociation dynamics very fast. For simplicity, polymerization is modeled as a single rate without explicitly including polymerase binding and dissociation.

Introducing blockers creates an effective kinetic bias, and at the same time enhances the effective energetic bias between right and wrong targets (Fig. 3a). These effects are quantified by

\[
\Delta G^\dagger = k_B T_a \ln \left( 1 + \frac{[B_W]}{K_d^{BW:W}} \right),
\]

\[
\Delta G - \Delta G_0 = k_B T_a \ln \left( 1 + \frac{1}{1 + [P]/K_d^{PW}} \frac{[B_W]}{K_d^{BW:W}} \right).
\]

Here, \(\Delta G_0\) is the energetic bias in the absence of the blocker, and \(K_d\) is the dissociation constant of the specified hybridization.

We measured the hybridization melting curves and estimated the temperature dependence of \(K_d\) (Fig. S4) to evaluate the dependence of \(\Delta G\) and \(\Delta G^\dagger\) on \(T_a\) (Fig. 3b). We found that \(\Delta G \gg \Delta G^\dagger\) at high \(T_a\) and \(\Delta G^\dagger \gg \Delta G\) at low \(T_a\). This result shows that the error suppression by blocker in a broad \(T_a\) range stems from the combined effects of the energetic and kinetic biasing pronounced at high and low \(T_a\), respectively.

Kinetics of error suppression. To analyze the detailed kinetics of error suppression by the blocker strands, we varied the polymerase concentration by more than two orders of magnitude while fixing \(T_a\) to 60 °C, which is an appropriate temperature for our primer sequence (Fig. 4). Since elongation by polymerase...
Figure 3. Effective energetic and kinetic bias in the presence of blockers. (a) The free energy landscape in the presence of B_W. (b) Dependence of energetic ($\Delta G, \Delta G_0$) and kinetic ($\Delta G^\dagger$) bias on $T_a$ calculated by the model (SI section S4).

quenches the hybridization dynamics, a change in the polymerase concentration tunes the time scale available for the hybridization dynamics to relax.

In the absence of blocker strands, $\eta$ decreased as the polymerase concentration decreased (Fig. 4c). In the low polymerase concentration limit, the limiting error rate $\eta_{\text{eq}}$, see Eq. (2), was lower in the presence of blockers B_W. Since the error is determined by the energetic bias in this limit, this result confirms our prediction that the blocker addition increases $\Delta G$.

The blocker addition lowered $\eta$ at high polymerase concentration as well (Fig. 4c), consistently with our prediction that the blockers also create an effective kinetic barrier for the wrong primer strands. The theory approximates the limiting error rate as

$$\eta_{\text{fast}} \simeq \frac{1}{1 + (k_{\text{on}}[P]t_a)^{-1}\exp(\Delta G^\dagger/k_B T_a)}$$

(SI Section S4), depending on the barrier difference $\Delta G^\dagger$. Here, $k_{\text{on}}$ is the binding rate of the primer, and $t_a = 30\text{s}$ is the duration of the annealing step. These results support that the blocker strands suppress errors both energetically and kinetically, as illustrated in Fig. 3a.

The model successfully reproduced the experimental results (dashed lines in Fig. 4). The values of the five fitting parameters were comparable with estimates based on previous work (see SI section S5). Even without blocker strands, a slight reduction of polymerase concentration is effective at suppressing errors without affecting much the efficiency (black arrow in Fig. 4d). However, this strategy requires fine-tuning of the polymerase concentration to maintain the efficiency. On the other hand, the addition of blocker strands is more effective at reducing errors and without reducing the efficiency (blue arrow in Fig. 4d).

**Kinetic regime.** Our theoretical model suggests the presence of a novel error correction regime. This fact is related with the relative magnitude of $\Delta G$ and $\Delta G^\dagger$, which changes at a temperature $T_a \approx 58 \degree C$ where the $\Delta G$ and $\Delta G^\dagger$ curves intersect each other in Fig. 3b. According to a theory\textsuperscript{11}, the relative magnitude of these two quantities determines the behavior of the primer-binding dynamics.
Figure 4. Improved performance of PCR reaction in the presence of blockers is consistent with model predictions. Dependence of the efficiencies (a, b) and error rate (c) on the polymerase concentration. (d) The error rate is plotted against the growth rate $\alpha_R$ (efficiency). Symbols denote the experimental data, and dashed lines correspond to the model fitting. Stars indicate the limiting values at high (open) or low (closed) polymerase concentrations, which correspond to $\eta_{eq}$ given by (2) and $\eta_{fast}$ given by (4), respectively. The polymerase concentration of the standard PCR protocol is 25 units/mL. $T_a = 60^\circ$C. We excluded two points with negative averages due to the statistical errors from (b) and (c) (+B_W with 0.17 and 0.25 units/mL polymerase). The error bars indicate the standard deviations. See Fig. S5 for the plots in the linear scale.

To illustrate this theoretical prediction, we ran a simulation of our model but without the elongation by polymerase. On the one hand, in the “energetic regime” characterized by $\Delta G > \Delta G^\ddagger$ (60 °C), the fraction $\gamma \equiv [P:W]/([P:R] + [P:W])$ decreases with the annealing time $t_a$ and converges to an equilibrium value $\gamma_{eq} = 1/(1 + e^{\Delta G/k_B T_a})$ (Fig. 5a). On the other hand, in the “kinetic regime” characterized by $\Delta G < \Delta G^\ddagger$ (57.5 °C), the kinetic barrier blocks the primer binding to the wrong template (Fig. 1c), and $\gamma < \gamma_{eq}$ is observed. At the limit of vanishing $t_a$, $\gamma$ converges to $\gamma_{short} = 1/(1 + e^{\Delta G^\ddagger/k_B T_a})$ (SI Section S4).

These binding dynamics imply that, in the presence of the polymerase, $\eta$ may increase with $t_a$ and be smaller than $\eta_{eq}$ in the kinetic regime. That is, we may reduce errors and cycle time simultaneously and also achieve significant error suppression. This prediction was confirmed in the case of dilute polymerase
Figure 5. Dynamics of binding and error calculated by simulation. (a) Binding dynamics in the absence of polymerase. \( \gamma \) is the fraction of the complex of the primer and wrong template. \( \eta_{eq} \) is its equilibrium value. See the main text. \( \eta_{eq} = 0.038 \) for 60 °C and 0.057 for 57.5 °C. (b) Error rate dynamics in the presence of polymerase normalized by \( \eta_{eq} \) given by (2). The polymerase concentration is 0.1 units/mL (dashed) or 25 units/mL (dotted). \( \eta_{eq} \approx 0.0033 \) for 60 °C and 0.0045 for 57.5 °C. (c) The growth rates of the right product. For the simulation, the dissociation constants are obtained experimentally. \( k_{off} / k_{on} \) and \( k_{W} \) are obtained by the global fitting of the experimental results at each temperature. See SI Section S5 for details.

(0.1 units/mL) and relatively short \( t_a \) (Fig. 5b). However, at \( t_a \) larger than \( \sim 100 \) s, \( \eta \) increased with \( t_a \) in both regimes. This increase is pronounced with a typical polymerase concentration for PCR (25 units/mL). The increase in \( \eta \) at large \( t_a \) is caused by the saturation of the growth; \( \alpha_W \) increases with \( t_a \) while \( \alpha_R \) is already saturated (Fig. 5c). Nevertheless, we still obtained \( \eta < \eta_{eq} \) at small \( t_a \). At the limit of vanishing \( t_a \), \( \eta \) converges to

\[
\eta_{short} = \frac{1}{1 + \left( k_{on} / k_{off} \right) e^{\Delta G^2 / R T_a}}
\]

which has the same mathematical form as \( \eta_{eq} \) given by (2) but with \( \Delta G^2 \) appearing instead of \( \Delta G \) (SI Section S4).

It remains unclear whether the regime \( \eta < \eta_{eq} \) is experimentally accessible. The expected time scale \( t_a \approx 1 \) s is significantly shorter than the typical time scale (\( \sim 10 \) s) necessary for cooling from \( \sim 95 \) °C at the denaturing step to \( \sim 60 \) °C at the annealing step. Our model assumes an instantaneous temperature jump for simplicity and may no longer be applicable for such a short \( t_a \).

The short \( t_a \) results in a significant reduction of the product amount (Fig. 5c). However, the approach implementing the kinetic regime would be beneficial in the applications such as genome editing, where error suppression is critical.

**Multiple wrong sequences.** In real-world applications of PCR, samples may contain multiple types of unwanted sequences. We study by numerical simulations of our model whether blockers could suppress replication errors in this case. For simplicity, we focus on a scenario in which the sample contains \( N \) types of wrong sequences, and we add \( N \) blocker sequences, each of which perfectly hybridizes to the corresponding wrong sequence. We fix the total concentration of the wrong sequences and the blocker sequences, so that the concentration of each wrong sequence and blocker sequence is proportional to \( 1/N \).
We note that the concentration of $B_W : W$ is roughly proportional to $[W][B_W]$. Since $[W]$ and $[B_W]$ decrease with $N$, blocking may become less effective with $N$. The error rate $\eta$ is defined similarly to (1), but where $\alpha_W$ is the total amount of wrong products (see SI section S4). We find that, although the error increases with $N$, the blocker strands suppress $\eta$ even in the large $N$ limit (Fig. 6). Moreover, our model predicts that the addition of blockers should not significantly affect $\alpha_R$ unless they strongly hybridize to R.

Figure 6. Error rate in the presence of multiple error sequences and blockers predicted by numerical simulation. The blue dotted line and black dashed line correspond to the result for a large number of wrong sequences ($N = 10^4$) and without blocker strands, respectively.

Discussion

Kinetic modeling of PCR reaction has contributed to quantitatively characterize the reaction performance$^{14–16}$ and other aspects such as amplification heterogeneity$^{17}$. However, modeling has been scarcely used to develop new guiding principles. The physics of information processing can provide such principles, thanks to its progress in characterizing general biochemical reactions$^{11, 18–30}$.

We demonstrated that adding blocker strands discriminates the right and wrong sequences by combining energetic and kinetic biasing. The kinetic biasing is effective in decreasing the error rate without affecting the efficiency. An alternative setup we studied is the use of blocker strands targeting the right template. In this case, we could increase the error rate up to a value larger than 80%. This inverted error control can not be achieved without kinetic biasing and may be helpful for sampling rare sequences.

Importantly, we found that error suppression is still effective at $T_a$ much lower than $T_m$ of the primer binding. This implies that we can suppress the hybridization errors in systems with limited temperature controllability, such as the reverse transcription, in which the optimal reaction temperature is not high, and the hybridization inside biological cells. This might also lead to a reduced cost of applications such as virus tests by using a low-cost cycler since we do not require accurate temperature control.

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Methods

Linear PCR experiment. DNA strands were synthesized by Eurofins Genomics, and Integrated DNA Technologies (see SI section S1). DNA/LNA chimeric strands were synthesized by Aji Bio-Pharma. The reaction mixture for polymerization contained Hot-start Taq DNA polymerase (New England Biolabs), Taq standard reaction buffer, $R$, $W$, $P$, and blocker strands. We performed initial heating for 30 s at 95 °C and, then, 10 or 40 cycles of 15 s at 95 °C, 30 s at 60 °C, and 5 s at 68 °C using a PCR cycler. Immediately
after the cycles, the mixture was cooled down on the ice to stop the enzyme reaction and used for the quantification. The number of cycles were 40 when the polymerase concentration was 0.17, 0.25 or 0.5 units/mL and 10 otherwise.

**Quantification of P<sub>R</sub> and P<sub>W</sub>**. Additional quantitative PCR was performed on a real-time PCR cycler after the linear PCR experiment for quantifying r and w (see SI section S2). The reaction mixture contains Luna Universal qPCR Master Mix (New England Biolabs), 200 nM each of the primers, and the diluted sample. The dilution rate is 1/250 in the final concentration. The thermal cycle consists of initial heating for 60 s at 95 °C, 40 cycles of 15 s at 95 °C, 30 s at 66 °C, and 5 s at 72 °C.

**Melt curve analysis.** We measured the melting curves for the hybridization of P, B<sub>R</sub>, and B<sub>W</sub> to R and W from 95 °C and 20 °C and calculated their K<sub>d</sub> (SI section S3). We mixed 100 nM each of DNA and double-strand-specific fluorescent molecule EvaGreen (Biotium). The fluorescent profile was analyzed based on the exponential background method<sup>31</sup> to obtain the melting curve.

**Data availability**

The data that support the findings of this study are available from the corresponding author upon reasonable request.

**Author Contributions**

HA, SP, and ST designed the research, developed the theory, and wrote the paper. HA and SO did experiments.

**Competing Interests**

The authors declare no conflict of interest.

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