Effect of Stalling after Mismatches on the Error Catastrophe in Nonenzymatic Nucleic Acid Replication

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Abstract: The frequency of errors during genome replication limits the amount of functionally important information that can be passed on from generation to generation. During the origin of life, mutation rates are thought to have been quite high, raising a classic chicken-and-egg paradox: could nonenzymatic replication propagate sequences accurately enough to allow for the emergence of heritable function? Here we show that the theoretical limit on genomic information content may increase substantially as a consequence of dramatically slowed polymerization after mismatches. As a result of postmismatch stalling, accurate copies of a template tend to be completed more rapidly than mutant copies and the accurate copies can therefore begin a second round of replication more quickly. To quantify this effect, we characterized an experimental model of nonenzymatic, template-directed nucleic acid polymerization. We found that most mismatches decrease the rate of primer extension by more than 2 orders of magnitude relative to a matched (Watson–Crick) control. A chemical replication system with this property would be able to propagate sequences long enough to have function. Our study suggests that the emergence of functional sequences during the origin of life would be possible even in the face of the high intrinsic error rates of chemical replication.

Introduction

Biological organisms store information in the sequence of their genomes. The information is propagated during genome replication, but each nucleotide incorporation presents an opportunity for error. At a given mutation rate per base (µ), if the genome is too long, the sequence information will be lost as mutants accumulate (an error “catastrophe”). Therefore, the mutation rate limits the total amount of information that can be carried by a genome. In particular, the maximum genome information is inversely proportional to the mutation rate. Experimental data on mutation rates in RNA viruses, which appear to exist near this limit (the error threshold), also support this relationship. Modern organisms have elaborate machinery for error detection and correction, but the first replicators were presumably very simple and had high error rates. Previous work indicates that nonenzymatic, template-directed nucleic acid polymerization has high error rates (close to 20%), corresponding to a genome of roughly 5 bases, but aptamers, ribozymes, and deoxyribozymes are usually at least 30 bases long. This discrepancy raises a paradox for the emergence of functional sequences during the origin of life. Is nonenzymatic replication accurate enough to propagate functional sequences? Previous proposals to address Eigen’s paradox include a mutualistic hypercycle, a spatially structured environment with cooperating sequences, mutational neutrality, or very high fitness differences. However, these approaches either invoke special functions or are relatively limited in magnitude. For example, one analysis of self-cleaving ribozymes found that 25% of bases could be mutated without destroying function, so the physical length of the genome could exceed the informative length by 25%. Here we show that the chemical dynamics inherent in polymerization could offset the error threshold to the extent that sequences long enough to be functional could readily emerge.

The error threshold was first derived by Eigen from the following set of reactions describing replication:

\[
\begin{align*}
2 \text{A} & \rightarrow \text{B} + \text{A} \\
\text{B} & \rightarrow \text{C} + \text{D} \\
\text{C} & \rightarrow \text{E} + \text{F}
\end{align*}
\]

and

\[
\begin{align*}
\text{A} + \text{B} & \rightarrow \text{C} + \text{D} \\
\text{C} + \text{D} & \rightarrow \text{E} + \text{F} \\
\text{E} + \text{F} & \rightarrow \text{G} + \text{H}
\end{align*}
\]

and so on. Note that the replication rates are exponential in the number of nucleotides, leading to a catastrophe for genomes longer than a certain size limit. Eigen showed that this limit could be increased by a factor of 25% by the presence of an error-free self-cleaving ribozyme. This was the origin of the idea that prebiotic evolution could have been driven by copying errors as well as correct copies, and that the presence of error-correcting systems could have increased the genome size limit. If this is true, the total information content that could be passed on from generation to generation would be increased by a factor of 25%. This is the origin of the question of the error catastrophe. The frequency of errors during genome replication limits the amount of functionally important information that can be passed on from generation to generation. During the origin of life, mutation rates are thought to have been quite high, raising a classic chicken-and-egg paradox: could nonenzymatic replication propagate sequences accurately enough to allow for the emergence of heritable function? Here we show that the theoretical limit on genomic information content may increase substantially as a consequence of dramatically slowed polymerization after mismatches. As a result of postmismatch stalling, accurate copies of a template tend to be completed more rapidly than mutant copies and the accurate copies can therefore begin a second round of replication more quickly. To quantify this effect, we characterized an experimental model of nonenzymatic, template-directed nucleic acid polymerization. We found that most mismatches decrease the rate of primer extension by more than 2 orders of magnitude relative to a matched (Watson–Crick) control. A chemical replication system with this property would be able to propagate sequences long enough to have function. Our study suggests that the emergence of functional sequences during the origin of life would be possible even in the face of the high intrinsic error rates of chemical replication.

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In these reactions, X is a “master” sequence that is characterized by higher fitness \((r > 1)\) relative to that of all mutant sequences (Y) and q is the probability of replicating without errors (i.e., \(q = (1 - \mu)^r\), where \(\mu\) is the mutation rate per base and L is the number of functionally informative sites). In this classical model,\(^1\) the master sequence can survive only if \(L < L^*\) (Supporting Information). Because \(L^*\) has a relatively weak logarithmic dependence on \(r\) and the prebiotic fitness is thought to have been relatively small, this equation is often approximated as \(L^* \approx \frac{1}{\mu} (\text{corresponding to } r = e)\) or roughly one error per replication. Beyond this point, the system undergoes a phase transition to a state in which the master sequence disappears and the genomes diffuse randomly through sequence space. \(L^*\) is often thought of as a physical length, although strictly speaking it is the maximum number of informative sites. In essence, full-length mutant sequences, which are mismatched has been observed for several DNA polymerases, which are produced both by the replication of mutants and by mutation from the master sequence, grow in number faster than the master sequence when the genome is too long for a given mutation rate. Thus, the mutants outgrow the master sequences because they all consume resources during replication, and in a finite population, the master sequence would eventually disappear.\(^8\) In this simple model, the existence of complementary strands was ignored but the error threshold is similar for complementary replication.\(^9\)

In the classical model, polymerization was assumed to proceed equally fast regardless of whether an error occurred. However, studies of enzymatic polymerization show that if an incorrect nucleotide is incorporated then primer extension stalls after the mutation, presumably because of a suboptimal conformation at the mismatched terminus.\(^10\) If an incorrect nucleotide is incorporated then primer extension stalls after the mutation, presumably because of a suboptimal conformation at the mismatched terminus.\(^10\) Stalling after base pair mismatches has been observed for several DNA polymerases, with the ratio of extension rates from a matched versus mismatched terminus (the stalling factor, S) ranging from 10 to 100.\(^11\) Intuitively, this effect might slow the production of inaccurate copies of the master sequence, increasing the effective fidelity and the maximum genome information. However, it was previously unknown whether nonenzymatic polymerization would also slow after mutations. We therefore undertook the determination of mutation rates and stalling factors in a model system for template-directed nonenzymatic polymerization. We used 2′-deoxy-5′-phosphorimidazolides (ImpdNs) as the activated monomers, DNA templates, and DNA primers terminated by a complementary replication.

\[
\begin{align*}
X^r &\rightarrow X + X \\
Y^{n(1-r)} &\rightarrow Y + Y \\
Y^{1-r} &\rightarrow Y + Y
\end{align*}
\]

closely. We then calculated the error threshold including the effect of stalling after mutations. Our results indicate that stalling increases the maximum genome information to the extent that functional sequences could have been replicated without enzymes.

**Materials and Methods**

**Preparation of Nucleoside 5′-Phosphorimidazolides.** All chemicals were obtained from Sigma-Aldrich (St. Louis, MO) unless otherwise specified. The protocol used to synthesize the activated nucleotides (ImpdNs) was based on a previously published method.\(^13\) The free acid form of each nucleoside-5′-monophosphate (1.5 mmol) was suspended with imidazole (15 mmol) and 2,2′-dithiodipyridine (4.5 mmol) in 20 mL of a 1:1 mixture of anhydrous dimethyl formamide and anhydrous dimethyl sulfoxide. Subsequently, triethylamine (TEA, 4.5 mmol) and triphenylphosphine (3 mmol) were added and the mixture was stirred at room temperature for \(\sim 4\) h. The reaction progress was monitored by thin layer chromatography using a mobile phase of 50% n-butanol and 20% acetic acid in water. The resulting clear, yellow solution was added dropwise to a flask containing a mixture of anhydrous ether (200 mL)/acetone (125 mL)/TEA (15 mL)/anhydrous sodium perchlorate (0.5 g) and precipitated with gentle stirring for 30 min on ice. The resulting precipitate was filtered, washed with 200 mL of a 1:1 mixture of acetone and ether and with 100 mL of anhydrous ether, and dried overnight in vacuum desiccator over phosphorus pentoxide to give the corresponding nucleoside 5′-monophosphate imidazolide sodium salt. The resulting mixture was analyzed by RP-HPLC (Varian, Inc., Palo Alto, CA) using a C18 column (Varian microsorb, 250 × 41 mm i.d., 5 μm particle size). The conditions for HPLC were the following: solvent A: 0.025 M TEAB, pH 7.3; solvent B: 70% acetonitrile/water; gradient: isocratic 15% B; flow rate: 15 mL/min; and UV detection: 260 nm. The fractions containing the desired ImpdN were collected and frozen. These were then lyophilized to obtain the solid triethylammonium salt of the imidazolide. All ImpdNs were found to be \(\geq 93\%\) pure according to analytical HPLC.

**Oligonucleotides for Nonenzymatic Polymerization.** DNA primers terminated with a 3′-amino-2′,3′-dideoxynucleotide were either radiolabeled or fluorescently tagged for detection and quantification of the reaction products. The primer used to obtain misincorporation rates (AminoG) was synthesized on a dT-CPG column (Glen Research; Sterling, VA). A single 3′-amino-dG residue was added manually using 3′-amino-5′-dMT-dG (RI Chemical Inc.; Orange County, CA) under standard coupling conditions. The remainder of the sequence was synthesized on an Expedite 8900 nucleic acid synthesizer (Millipore; Billerica, MA). After ammonium hydroxide cleavage from the column and deprotection, the oligo was gel purified and then treated with 80% acetic acid overnight to cleave off the terminal phosphoramidite-linked T residue and the hydroxylate was purified by HPLC to isolate the 3′-amino oligo. The correct mass of the oligo was confirmed by matrix-assisted laser desorption ionization–time-of-flight mass spectrometry (MALDI-TOF MS; PerSeptive Biosystems Voyager MALDI-TOF; Framingham, MA). A sample of \(\sim 200\) pmol of oligonucleotide was adsorbed on a C18 ZipTip. Samples were eluted with 1.5 μL of a matrix solution containing a 2:1 mixture of 52.5 mg/mL 3-hydroxypropionic acid in 50% acetonitrile and 0.1 M ammonium citrate in water. Eluates were directly spotted onto a stainless steel MALDI-TOF plate and analyzed in positive mode. The AminoG primer was end-labeled with a T4 polynucleotide kinase (New

England Biolabs; Ipswich, MA) and γ-32-P-ATP (Perkin-Elmer; Waltham, MA) at the 5′-hydroxyl termini of DNA, following an established protocol. This primer was also used for a subset of extension reactions for matched versus mismatched termini.

The three remaining primers (AminoA, AminoT, and AminoC) for these extension reactions were made by reverse synthesis in an established protocol. For these extension reactions were made by reverse synthesis in the W. M. Keck Biotechnology Resource Laboratory at Yale University (New Haven, CT). The synthesis used the following phosphoramidites for the 3′ residue: AminoA: 3′-O-tritylamino-N°-benzoyl-2′,3′-dideoxycytidine-5′-cyanoethyl phosphoramidite; AminoC: 3′-O-tritylamino-N°-benzoyl-2′,3′-dideoxyadenosine-5′-cyanoethyl phosphoramidite; and AminoT: 3′-tritylamino-3′-deoxythymidine-5′-cyanoethyl phosphoramidite. The correct mass of these oligos was verified by MALDI-TOF as described above.

Polymerization. The DNA template sequences were synthesized and PAGE purified by Sigma-Aldrich (St. Louis, MO). Primer and template sequences are given below.

- **primer sequences:**
  - AminoG ("primer G"): 5′ GG GAT TAA TAC GAC TCA CTG-NH2
  - AminoA ("primer A"): 5′ GG GAT TAA TAC GAC TCA CTAG-NH2
  - AminoT ("primer T"): 5′ GG GAT TAA TAC GAC TCA CTAG-NH2
  - AminoC ("primer C"): 5′ GG GAT TAA TAC GAC TCA CTAG-NH2

Template sequences for misincorporation reactions are given below:

- **MisincorpA:** 5′ AGT GAT CTA CAG TGA GTC GTA TTA ATCC CC
- **MisincorpT:** 5′ AGT GAT CTT CAG TGA GTC GTA TTA ATCC CC
- **MisincorpG:** 5′ AGT GAT CTG CAG TGA GTC GTA TTA ATCC CC
- **MisincorpC:** 5′ AGT GAT CTC CAG TGA GTC GTA TTA ATCC CC

Template sequences for mismatch extension reactions are given below:

- **MismatchA:** 5′ AGT GAT CTC AAG TGA GTC GTA TTA ATCC CC
- **MismatchT:** 5′ AGT GAT CTC AAG TGA GTC GTA TTA ATCC CC
- **MismatchG:** 5′ AGT GAT CTC CAG TGA GTC GTA TTA ATCC CC
- **MismatchC:** 5′ AGT GAT CTC CAG TGA GTC GTA TTA ATCC CC

**Primer Extension Reactions and Assay for Nonenzymatic Polymerization.** A primer (0.325 μM) and a template (1.3 μM) (1 μL each) were mixed in water, incubated at 95 °C for 5 min, and annealed by cooling to room temperature on a benchtop for 5–7 min. In a typical reaction of 10 μL volume, 1 μL of 1 M Tris (pH 7) and 0.5 μL of 4 M NaCl were added to final concentrations of 100 mM Tris and 200 mM NaCl. For reactions with ImpdA, ImpdC, or ImpdG, the reaction was initiated by the addition of 1 μL of 100 mM ImpdN to a final concentration of 10 mM ImpdN. For reactions involving ImpdT, 1.38 μL of 289 mM stock solution was added to a final concentration of 40 mM ImpdT. The total volume of the reaction was 10 μL. The reaction mixtures were incubated at room temperature, and aliquots were withdrawn during a certain period of time. Time points were obtained by adding 1 μL of the reaction mixture to 9 μL of the loading buffer with 8 M urea, 100 mM EDTA, and 1.3 μM of a competitor DNA with the sequence 5′ GG GAT TAA TAC GAC TCA CTN 3′ where N = AT/ GC to match the primer employed in the reaction. Time points were heated to 90 °C for 5 min to disrupt primer–template complexes and were run on 20% denaturing PAGE.

The gels were phosphorimagined using a Typhoon TRIO variable-mode imager (Piscataway, NJ), and the scans were analyzed with ImageQuant v5.2 software. The frequency of unreacted primer was calculated by dividing the intensity of the unreacted primer band by the sum of intensities of the unreacted and reacted primer. In some cases, the extended product appeared to be a doublet band that was well separated from the unreacted primer; the doublet intensities were summed. To avoid experimental artifacts late in the reaction, initial rates were estimated by a linear fit to the first several data points.

**Calculation of Mutation Rate and Stalling Factor.** The frequency of incorporation (f_{template base,ImpdN}) of an ImpdN across a particular template base was calculated by dividing its rate of extension by the sum of the rates of extension for all ImpdN’s of the same primer–template complex (i.e., containing the same template base at the position opposite the incoming nucleotide). The stalling factor for template base N (f_{stall}) is the sum of the frequencies of incorrect incorporations (e.g., \( f_{A:ImpdN} = f_{A:ImpdT} + f_{A:C} + f_{A:T} \)). If the fraction of the genome composed of base N is given by \( F_N \), then the average mutation rate of a genome \( (\mu_{ave}) \) is \( \Sigma f_{template base,ImpdN} \). For example, a genome composed of equal parts A, C, G, and T would have \( \mu_{ave} = 0.25(\mu_{A} + \mu_{C} + \mu_{G} + \mu_{T}) \), and a genome composed of equal parts of only G and C would have \( \mu_{ave} = 0.5(\mu_{G} + \mu_{C}) \).

The stalling factor for each mismatch \( (f_{template base,primer terminus}) \) was calculated by dividing the rate of extension from the corresponding matched terminus \( (f_{template base,primer terminus}) \) by the same template sequence, by the rate of extension from the mismatched terminus \( (f_{template base,primer terminus}) \) which has the same template sequence, by the rate of extension from the mismatched terminus \( (f_{template base,primer terminus}) \) which has the same template sequence, by the rate of extension from the mismatched terminus \( (f_{template base,primer terminus}) \) which has the same template sequence.

The stalling factor, \( S_{ave} \), was calculated by weighting each stalling factor by the frequency of incorporation that leads to that stalled complex \( S_{ave} = F_A f_{A:ImpdN} f_{A:ImpdN} / F_A f_{A:ImpdN} f_{A:ImpdN} + F_C f_{C:ImpdN} f_{C:ImpdN} + F_T f_{T:ImpdN} f_{T:ImpdN} + F_G f_{G:ImpdN} f_{G:ImpdN} \). In other words, the most frequent mutations contribute most to the overall stalling factor because they result in the most frequent mismatched termini. Stalling factors are also weighted by the genome composition because mutations across the most common template base (and the corresponding mismatched terminus) would be relatively well represented. In this article, we assume that the genome is equal parts A, C, G, and T for the purpose of the stalling factor calculation \( F_A = F_C = F_G = F_T = 0.25 \). The standard deviation of the overall stalling factor and mutation rate, \( S_{ave} \) and \( \mu_{ave} \), were calculated as the standard deviation of the corresponding values from an initial batch of reactions and a duplicate batch.

**Results and Discussion**

**Mutation Rate of Nonenzymatic Polymerization.** We determined the rates of misincorporation in a series of reactions containing a template sequence, a perfectly complementary primer (either radiolabeled or fluorescently tagged), and one ImpdN (A, C, G, or T). Initial experiments showed that the rate of incorporation of T was particularly slow, causing relatively low fidelity when copying across a template base A, so we increased the concentration of ImpdT to 40 mM in our reactions (compared with 10 mM for the other nucleotides). Adjusting the ratio of monomer concentrations has been used...
previously to improve fidelity in enzymatic reactions.\textsuperscript{17} We followed reactions over time to determine apparent first-order rate constants for all possible correct incorporations (4 reactions) and misincorporations (12 reactions) (Figures 1a,b and 2a and Supporting Information).

We found that the average mutation rate ($\mu_{\text{ave}}$) of a genome composed of equal proportions of A, C, G, and T would be 7.6 ± 1.4% in this system. Misincorporations occurred predominantly when copying A and T, so a GC-rich genome would have a lower mutation rate (e.g., for an entirely GC genome, $\mu \approx 0.8$%; see Methods and Materials for details of the calculation). The absolute rate of incorporation of G and C across their cognate bases was also ~10 times greater than the rate of incorporation of A and T, consistent with trends from previous work\textsuperscript{18} suggesting that hydrogen bonding may also contribute to the reaction rate. Our results differ somewhat from previous


estimates of fidelity in a similar system, probably because of differences in ionic conditions and monomer concentrations.\(^3\) According to the original Eigen model, a mutation rate of 7.6% would be too high to sustain a functional genome at low fitness \((L^* \approx 13)\).

**Stalling Factor of Nonenzymatic Polymerization.** To determine stalling factors for nonenzymatic polymerization, we varied the 3′ terminus of the primer to form either a perfectly matched terminus (4 reactions) or a mismatched terminus (12 reactions) and measured the rate of incorporation of the correct subsequent monomer (Figures 1c,d and 2b and Supporting Information). The overall stalling factor \((S_{\text{ave}})\) was calculated as an average weighted by the misincorporation frequency leading to the terminus. (See Methods and Materials for details of the calculation.) Despite the lack of an enzyme, nonenzymatic polymerization showed substantial stalling, with the extension from any mismatch being slower than its matched counterpart by a factor of 20–300 \((S_{\text{ave}} = 124 \pm 22)\).

**Modified Error Threshold Including the Effect of Stalling after Mutation.** Would the effect of stalling be large enough to permit the nonenzymatic replication of functional sequences? We modified Eigen’s model of replication to include a stalled state after a misincorporation, which progresses to completion at a relatively slow rate. Following Eigen’s model, we assume that the relative fitness of the master sequence is \(r\) and resources for replication are available at constant concentration. In the reactions given below, \(X\) is the fittest sequence, \(Z\) is an incomplete copy in which a mismatched nucleotide was incorporated, and \(Y\) is the finished mutant.

\[
\begin{align*}
\text{no error} & : X \rightarrow X + X \\
\text{error} & : X \rightarrow X + Z \quad \text{(slow)} \rightarrow X + Y
\end{align*}
\]

Mutant sequences undergo an analogous set of reactions when an error occurs. Strand separation is assumed to occur frequently compared with the relatively slow process of chemical replication (e.g., thermal cycling due to day-night changes or in convection cells [Supporting Information]). The error threshold for the corresponding set of differential equations was determined analytically in the limit of large numbers under the condition that the total density of the system is conserved \((\{X\} + \{Y\} + \{Z\} = \text{constant}; \text{see Supporting Information for a full description})\). We obtained a new expression for the maximum genome information corresponding to the condition that \(\{X\} > 0\) in the stationary state, namely that \(L^* = \ln(\{X\} + \mu S\{r - 1\})/\mu\) (Figure 3a). As with the classical model, \(L^*\) is inversely proportional to \(\mu\). As expected, as \(S\) increases (i.e., as stalling becomes more pronounced), \(L^*\) also increases. This effect is weighted by \(\mu\) because the synthesis of new strands is stalled longer if they contain multiple mutations.

Because this limit is always greater than or equal to the original Eigen condition, stalling would be beneficial for a variety of scenarios (i.e., different error rates and stalling factors). We also found that the error threshold was robust to details of the model; a second model in which imperfect copies were more likely to degrade during copying because of their longer copying time (e.g., longer exposure to UV damage or hydrolysis) gave the same error threshold (Supporting Information).

Using our experimentally determined parameters for \(\mu_{\text{ave}}\) and \(S_{\text{ave}}\), we calculated the maximum information of a genome undergoing nonenzymatic replication (Figure 3b). Although the classical Eigen model predicts that the mutation rate is too high to propagate a functional sequence, accounting for stalling after errors in polymerization increases the maximum informative length to 39 (at \(r = e\)). As with the classical threshold, this length increases with higher fitness (Figure 3b). This result demonstrates that an intrinsic feature of nonenzymatic polymerization could circumvent the Eigen paradox, allowing the propagation of functional sequences before enzymes evolved.

**Choice of Experimental Model System.** Our studies were carried out with 3′-amino-2′,3′-dideoxynucleotide-terminated primers. Although DNA was probably a relatively late invention in the course of prebiotic evolution, we use this 3′-amine system as an experimentally tractable model of nonenzymatic polymerization. In preliminary experiments, we had attempted to assay misincorporations in the nonenzymatic polymerization of a 2′,3′-hydroxyl system. However, polymerization in the 2′,3′-hydroxyl system was too slow to measure the rate of misincorporation accurately. There are also other unsolved issues with nonenzymatic RNA replication, such as strand separation, leading many to suggest that a different nucleic acid preceded the RNA world.\(^{18,19}\) Another possible experimentally tractable system would use 2′-amino-2′,3′-dideoxynucleotide-terminated primers.\(^{14}\) Although the 2′-amine system may have superior properties for copying long sequences with the goal of synthesizing a

\[(19)\text{ Eschenmoser, A. Science 1999, 284, 2118–2124.}\]
protocell, our goal here was to estimate the error rates associated with the more biological 3′–5′ linkage. In addition to the fairly efficient 3′-amine polymerization observed by Orgel and colleagues, a different 3′-amine system has also been studied by the Richert group, which exhibited very fast reaction rates with nearly quantitative yield, suggesting that a 3′-amine system has the potential to be efficient enough to copy relatively long sequences. It is possible that the 3′-amine system will have a fidelity differing from that of a 3′-hydroxyl system. Our data may not be representative of mutations in the RNA world itself, but our results do demonstrate that a nonenzymatic system exhibits stalling after mutations and that such a system could be capable of propagating sequences long enough to be functional because of this effect.

Conclusions

We have shown that the error catastrophe could be substantially mitigated through the dynamics of replication in which fidelity should not be considered to be a simple constant. Our experimental model system for nonenzymatic, template-directed nucleic acid polymerization demonstrates that stalling can be important even without enzymes. The presence of a mismatched terminus in the nascent sequence stalls extension and effectively decreases the rate of extension of a mutant sequence by more than 2 orders of magnitude. Interestingly, the same features of the prebiotic world that would reduce the maximum genome information in Eigen’s model—low fitness and high mutation rates—also increase the importance of stalling in offsetting the error catastrophe. Thus, nonenzymatic replication could potentially give rise to sequences long enough to be functional despite a high mutation rate. These dynamic effects could still be important after functional sequences emerged, permitting the genome to encode more sequences or longer sequences with higher activity. Furthermore, stalled primer–template complexes could provide a substrate for the evolution of error-correction machinery. Eventually, these effects would become obsolete as the replication machinery evolved greater accuracy and cooperating networks emerged, but early on they could have served to “kick start” the evolution of functional genomes.

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Supporting Information Available: Supporting text, figures, and a description of the mathematical analysis. This material is available free of charge via the Internet at http://pubs.acs.org.

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Supporting Information

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- Figure S1: Rates of incorporation and mis-incorporation .................. S2
- Figure S2: Rates of extension from matched and mismatched termini ... S3
- Mathematical description of modeling ...................................... S4
- Discussion of modified error threshold model in a prebiotic context .... S9
- Discussion of connection of model parameters to experimental results .. S10
- Discussion of degradation of stalled intermediates (Modeling section 2)..S11
**Figure S1.** Rate of incorporation of different activated nucleotides (C,G,A,T) across each template base. Error bars show standard deviations calculated from duplicate sets of reactions.
**Figure S2.** Rates of extension from matched or mismatched primer-template termini.

Error bars show standard deviations calculated from duplicate sets of reactions.
SUPPLEMENTARY TEXT: MODELS OF THE ERROR THRESHOLD WITH STALLING AND STRAND SEPARATION

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1. Classical error threshold

First, let us briefly discuss the classical case that Eigen solved in 1971. We have self-replicating units (e.g., RNA): master units and mutant units. Master units replicate at rate $r \geq 1$, while mutants replicate at unit rate (which fixes the time scale). The copy is perfect with probability $q$, and there is a mistake in the copy otherwise, with probability $1 - q$. A master unit with a mistake is a mutant, and a mutant with a mistake is also considered to be mutant (i.e., we neglect back mutations). We denote master units as $X$ and mutants as $Y$, so these processes can be depicted as

\[
\begin{align*}
  X & \xrightarrow{rq} X + X \\
  X & \xrightarrow{r(1-q)} X + Y \\
  Y & \xrightarrow{1} Y + Y
\end{align*}
\]

(1)

We keep the population size constant by also removing a random unit at each replication. In the large population size limit, the frequencies $x$ and $y = 1 - x$ of the above units evolve according to

\[
\begin{align*}
  \dot{x} &= r qx - \Phi x \\
  \dot{y} &= r (1 - q) x + y - \Phi y
\end{align*}
\]

(2)

where $\Phi = rx + y$ is chosen to keep the total population $x + y = 1$ constant. It is easy to see that the above two equations are identical, since $y = 1 - x$. In the stationary state $\dot{x} = \dot{y} = 0$, and we can express the frequency of master units as

\[
x = \frac{rq - 1}{r - 1}
\]

(3)

which is positive as long as $q > q_E^*$, where $q_E^* = 1/r$ is the classical error threshold of Eigen. This means that the copy has to be perfect with probability $q \geq q_E^*$ in order to have master units in the population at steady state. This can be understood in simple terms by noting that the master sequence effectively replicates at rate $qr$, which has to be larger than the mutant fitness for the master copy to survive, which immediately leads to $q_E^* = 1/r$. For a sequence of length $L$ and with mutation rate $\mu$ per base pair, the probability of making a perfect copy is

\[
q = (1 - \mu)^L \approx e^{-\mu L}
\]

(4)

where the last expression is valid for large sequences ($L \gg 1$) and small mutation rates ($\mu \ll 1$). Now the classical error threshold can also be formulated for the critical sequence length $L^* = (\ln r) / \mu$.

2. Strand separation and explicit degradation

Now we discuss an extension of the classical model, in which if an error occurs during the copying process, the nascent sequence is completed only with probability $p$. Otherwise, with probability
$1 - p$, it separates from the template before completion (e.g., during thermocycling) and degrades. Graphically

$$X \xrightarrow{rq} X + X$$
$$X \xrightarrow{r(1-q)p} X + Y$$
$$Y \xrightarrow{q} Y + Y$$
$$Y \xrightarrow{(1-q)p} Y + Y$$

The process $Y \rightarrow Y + Y$ happens at rate $q + p(1 - q)$. If we rescale time by $q + p(1 - q)$, $Y$ replicates at rate one, and $X$ replicates at rate $r$. Replication produces a perfect copy with probability

$$\hat{q} = \frac{q}{q + p(1 - q)}$$

and an imperfect copy with probability $1 - \hat{q}$. Hence our model is equivalent to the original model of Eigen but with $\hat{q}$. The critical error threshold is given by $\hat{q}^* = 1/r$, which corresponds to

$$q^* = \frac{p}{r + p - 1}$$

This function is depicted in Figure 1 for several values of $p$.

In the absence of strand separation, $p = 1$, we recover Eigen’s classical result of $q_E^* = 1/r$. As the strand separation becomes stronger, $p \rightarrow 0$, the error threshold tends to zero. This means that the error frequency can be arbitrarily high if mutant copies are not produced at all. We also see from (7) that $q^*$ is a monotonically increasing function of $p$. Hence the degradation of imperfect copies always lowers the error threshold. Note also that one could generalize this model by defining two distinct probabilities of strand separation and degradation: one in the absence of a mistake $p'$, and another one in case of a mistake $p''$. This model, however, would be equivalent to our present model with $p = p''/p'$.

3. STRAND SEPARATION AND RE-ANNEALING

In this section we assume that if a mistake happens during the copying process, the nascent sequence separates from the template (e.g., during thermocycling), and then re-anneals again to either a master or a mutant unit. Completion of the nascent sequence occurs at rate $a$, determined by the stalling factor. We denote a stalled, incomplete mutant unit as $Z$, so the model is

$$X \xrightarrow{rq} X + X$$
$$X \xrightarrow{r(1-q)a} X + Y$$
$$Y \xrightarrow{q} Y + Y$$
$$Y \xrightarrow{(1-q)a} Y + Y$$

In order to keep the total population size constant (counting all $X, Y$ and $Z$ units), we remove a random unit from the system each time a new unit is made. The limit $a \rightarrow \infty$ corresponds to the classical case of Eigen.

In the large population size limit the frequencies of different types of units change according to the differential equations

$$\dot{x} = rqx - \Phi x$$
$$\dot{y} = qy + axz + ayz - \Phi y$$
$$\dot{z} = r(1-q)x - axz + (1-q)y - ayz - \Phi z$$

where $\Phi = rx + y$ is chosen to keep the total population size $x + y + z = 1$ constant.
The stationary state of (9) is the solution of the equations

\begin{align}
0 &= rqx - (rx + y)x \\
0 &= qy + axz + ayz - (rx + y)y \\
0 &= r(1 - q)x - axz + (1 - q)y - ayz - (rx + y)z
\end{align}

At the error threshold, \( x \to 0 \). In order to find the critical surface \( q^*(r,a) \) we assume that \( x \ll 1 \) but still \( x, y, z > 0 \). Now the last two equations of (10) become identical in the leading order in \( x \), which leads to

\begin{equation}
y = \frac{q + a}{1 + a} + O(x)
\end{equation}

Here we also used that \( z = 1 - y + O(x) \). Now inserting (11) into the first equation of (10), and then letting \( x \to 0 \), we arrive at the critical \( q \) value

\begin{equation}
q^* = \frac{a}{r(a + 1) - 1}
\end{equation}

This function is depicted in Figure 1 for several values of \( a \).

In the present model a mistake always leads to strand separation, but a separated copy can re-anneal and be completed at rate \( a \). In the model of Section 2 a mistake leads to strand separation only with probability \( 1 - p \), but the separated copy degrades and never re-anneals. Despite the very different nature of these two models, they lead to equivalent error thresholds. When an imperfect copy re-anneals in the model of this section, it happens at rate \( a(x + y) \), and the copy dies at rate
$\Phi = rx + y$ [see (9)]. Hence the imperfect copy gets completed with probability

$$p = \frac{a(x + y)}{a(x + y) + rx + y}$$

(13)

Close to the error threshold $x \to 0$, the survival probability becomes

$$p = \frac{a}{a + 1}$$

(14)

Indeed, for such parameter values the error threshold $q^*$ is the same for the two models (7) and (12). For example, we recover the classical result of Eigen $q^*_E = 1/r$ for $a \to \infty$ or as $p \to 1$. The optimal case for the master units is the zero error threshold, which is reached for $a \to 0$ or for $p \to 0$. One can see the critical error threshold in Figure 1 for several corresponding values (14) of $p$ and $a$. We see from (7) and (12) that $q^*$ is an increasing function of either $p$ or $a$ for all $r > 1$. Hence stalling and strand separation always lower the error threshold.
Discussion of modified error threshold model in a prebiotic context

Whether stalling affects the error threshold depends on the timescales of replication. In a prebiotic world, if strand separation is infrequent enough that all products, both perfect copies and mutants, would be completed within a single replication cycle (e.g., before the strands melt), then stalling would not affect the products. But if strand separation occurs before mutant copies are completed, then stalling could potentially reduce the effective rate of production of mutants relative to perfect copies, which continue to propagate while mutants are stalled. In experimental models of templated non-enzymatic polymerization of nucleic acids, the half-times range from hours to days per base \(^1\), suggesting that the copying time for a short ribozyme (e.g., 30 bases) would be >18 hours. Prebiotically, the length of time available for replication before strand separation (\(\tau_r\)) might be dictated by thermal cycling. For a diurnal cycle, \(\tau_r\) would be \(~12\) hours assuming a rotational period of 24 hours, or approximately \(~7-10\) hours for the early earth with higher angular velocity \(^2\). For thermocycling in convection cells (e.g., deep sea hydrothermal vent), \(\tau_r\) could be as short as several seconds \(^3\). We therefore modeled polymerization with stalling, assuming that strands separate more quickly than mutant copies are completed.
Discussion of connection of model parameters to experimental results

Close to the error threshold, master sequences constitute a very small proportion of the sequence pool. Therefore, most sequences ($Y$) will be replicated with time $1/L$ per base, with stalled bases requiring additional time $S/L$ per mutation (master sequences replicate faster with an average time of $1/(rL)$ per base). Given $\mu L$ mutations per sequence on average, mutant sequences would take an additional time $\mu S$ to copy through stalled bases. The additional time is related to the completion of $Z$. Therefore, in terms of measurable parameters, $a = 1/(\mu S)$. 
Discussion of degradation of stalled intermediates (Modeling section 2)

To examine the dependence of the modified error threshold on the details of our model, we also solved a model of stalling in which degradation of stalled sequences is explicit during the copying process (Modeling section 2). In this second model, we assume that imperfect copies have a higher probability of degradation during polymerization, because their copying time is longer. This corresponds to longer exposure to chemical damage or simply a higher chance of washing out from the system before copying is complete. The relative probability of degradation of imperfect copies would be the ratio of exposure times. In the terminology of the main model (Modeling section 3), the time taken to complete a perfect copy would be 1, and the time taken to complete an imperfect copy would be 1+1/α. Therefore, the relative probability of survival for an imperfect copy would be 1/(1+1/α), or α/(α+1). Indeed, \( L_a^* \) in the two models agreed when \( p = \alpha/(\alpha+1) \) (Modeling section 3), indicating that the error threshold is robust to differences in the details of modeling.
