

REVIEW ARTICLE

Antimutator variants of DNA polymerases

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Abstract

Evolution balances DNA replication speed and accuracy to optimize replicative fitness and genetic stability. There is no selective pressure to improve DNA replication fidelity beyond the background mutation rate from other sources, such as DNA damage. However, DNA polymerases remain amenable to amino acid substitutions that lower intrinsic error rates. Here, we review these 'antimutagenic' changes in DNA polymerases and discuss what they reveal about mechanisms of replication fidelity. Pioneering studies with bacteriophage T4 DNA polymerase (T4 Pol) established the paradigm that antimutator amino acid substitutions reduce replication errors by increasing proofreading efficiency at the expense of polymerase processivity. The discoveries of antimutator substitutions in proofreading-deficient 'mutator' derivatives of bacterial Pals I and III and yeast Pol δ suggest there must be additional antimutagenic mechanisms. Remarkably, many of the affected amino acid positions from Pol I, Pol III, and Pol δ are similar to the original T4 Pol substitutions. The locations of antimutator substitutions within DNA polymerase structures suggest that they may increase nucleotide selectivity and/or promote dissociation of primer termini from polymerases poised for misincorporation, leading to expulsion of incorrect nucleotides. If misincorporation occurs, enhanced primer dissociation from polymerase domains may improve proofreading *in cis* by an intrinsic exonuclease or *in trans* by alternate cellular proofreading activities. Together, these studies reveal that natural selection can readily restore replication error rates to sustainable levels following an adaptive mutator phenotype.

Keywords: DNA replication fidelity, mutator, genetic adaptation, protein structure/function, genetic instability, cancer

Introduction

Genetic information must be accurately duplicated during cell division to ensure that daughter cells retain essential genes and the beneficial traits accrued through evolution. For this reason, organisms invest in an array of conserved mechanisms that promote genetic continuity (Friedberg *et al.*, 2006). DNA polymerase errors during cell division represent the largest potential source of genetic instability (Preston *et al.*, 2010). To minimize spontaneous mutation, these errors are corrected by exonucleolytic proofreading and post-replicative mismatch repair (MMR) (Iyer *et al.*, 2006; McCulloch and Kunkel, 2008; Reha-Krantz, 2010), thereby lowering the contribution of replication errors to the background mutation rate derived from other sources (Quah *et al.*, 1980; Morrison *et al.*, 1993; Roche *et al.*, 1994; Datta *et al.*, 2000). Defects in polymerase proofreading or MMR, as well as other DNA repair enzymes, elevate spontaneous mutation rate, producing 'mutator' phenotypes (Friedberg *et al.*, 2006).

While deleterious to long-term fitness, short-term genetic instability provides variation upon which natural selection can act (Sturtevant, 1937; Drake *et al.*, 1998; Sniegowski *et al.*, 2000; Giraud *et al.*, 2001b; de Visser, 2002; Elena and Lenski, 2003; Denamur and Matic, 2006). Mutator phenotypes due to defects in MMR or polymerase proofreading promote somatic evolution of cancer cells in mice and humans (Loeb *et al.*, 1974; Fishel *et al.*, 1993; Parsons *et al.*, 1993; de Wind *et al.*, 1995; Reitmaier *et al.*, 1996; Goldsby *et al.*, 2001; Goldsby *et al.*, 2002; Albertson and Preston, 2006; Loeb *et al.*, 2008; Daee *et al.*, 2010). Similarly, microbial mutators arise spontaneously when populations are subjected to repeated rounds of selection (Chao and Cox, 1983; Mao *et al.*, 1997; Sniegowski *et al.*, 1997; Giraud *et al.*, 2001a; Nilsson *et al.*, 2004). Interestingly, following adaptation, bacterial clones with lower mutation rates evolve from mutator lineages, suggesting that there is selection pressure for genetic suppressors that restore DNA replication

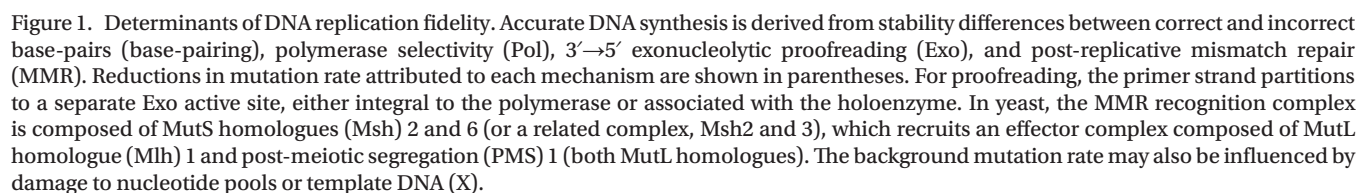
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This review focuses on DNA polymerase variants that impart an antimutator phenotype due to amino acid substitutions that increase DNA replication fidelity. To provide a framework for understanding the mechanisms of antimutagenesis, we begin our review with an exploration of the determinants of DNA replication fidelity. Several excellent in-depth reviews of DNA replication fidelity offer additional background information that readers may find useful (Pavlov *et al.*, 2006b; Kunkel, 2009; Johnson, 2010; Reha-Krantz, 2010). We continue by describing the origins of the four main antimutator polymerase collections, focusing on details that are pertinent for understanding the influence of antimutator substitutions on DNA polymerase fidelity. We discuss possible antimutator mechanisms in light of X-ray crystallographic structures of relevant DNA polymerase replication complexes. Finally, we consider the relevance of these findings for DNA polymerase evolution and plasticity in the natural world.

DNA replication is the sequential joining of nucleotides to the 3' end of a DNA primer sequence annealed to a complementary DNA template. Cells accomplish this task with astonishing precision (10^{-9} – 10^{-10} mutations per

Amino acid sequence alignments and polymerase crystal structures indicate that A, B, D, Y and RT families share a common core structure resembling a cupped



right hand (Beese *et al.*, 1993; Eom *et al.*, 1996; Wang *et al.*, 1997; Ding *et al.*, 1998; Huang *et al.*, 1998a; Li *et al.*, 1998; Franklin *et al.*, 2001; Ling *et al.*, 2001). The palm contains the catalytic residues, the thumb positions the double-stranded DNA in the active site, and the highly mobile fingers domain closes over deoxyribonucleoside-triphosphates (dNTPs) during nucleotide selection. C- and X-family polymerases are 'left-handed' based on the direction of DNA synthesis relative to the thumb and fingers domains, which also bind DNA and dNTPs, respectively, despite being structurally different than their right-handed counterparts (Sawaya *et al.*, 1997; Wing *et al.*, 2008).

With each elongation step, highly accurate DNA polymerases distinguish correct from incorrect dNTPs with an average error rate of 10^{-4} – 10^{-5} misinsertions per base-pair synthesized (Kunkel, 2009). Hydrogen bonds (H-bonds) between cognate base pairs were initially assumed to make the largest contribution to nucleotide selection (Watson and Crick, 1953a; Watson and Crick, 1953b). However, free energy differences of correct and incorrect base pairs in solution only account for a fraction (10^{-2}) of the observed specificity (Mildvan, 1974), because favorable gains in enthalpy due to correct base-pair formation are offset by unfavorable decreases in entropy of the solvent (Petruska and Goodman, 1995). Crystal structures of replication complexes reveal that high-fidelity DNA polymerases form tight nucleotide binding pockets that limit solvent access and select incipient base pairs that specifically conform to Watson-Crick geometry (Beese *et al.*, 1993; Eom *et al.*, 1996; Sawaya *et al.*, 1997; Doublié *et al.*, 1998; Kiefer *et al.*, 1998; Li *et al.*, 1998; Franklin *et al.*, 2001; Wing *et al.*, 2008; Swan *et al.*, 2009).

Watson-Crick base pairs share pseudo-dyad symmetry, similar distances between the C_{1'} atoms of the sugars, and equivalent glycosyl bond angles (Saenger, 1984). In addition, Watson-Crick base pairs have minor groove H-bond acceptors (O² of pyrimidines and N-3 of purines, Figure 1) in essentially identical positions (Saenger, 1984). While some mispairs form H-bond interactions of similar strength to Watson-Crick base pairs, their overall geometry differs dramatically. Thus, in theory, mispairs could be excluded on the basis of shape or absence of appropriately positioned minor groove contacts (Echols and Goodman, 1991; Beard and Wilson, 2003). Kool and colleagues found that close structural mimics of thymidine and adenine that lack H-bond donors or acceptors are more easily incorporated by DNA polymerases than noncognate nucleotides, suggesting that the shape of the incipient base pair plays a role in selection (Moran *et al.*, 1997; Morales and Kool, 1998; Morales and Kool, 2000; Potapova *et al.*, 2006; Lee *et al.*, 2008a). However, both A- and B-family polymerases also incorporate nucleotide analogues that do not conform to Watson-Crick geometry (Chiaramonte *et al.*, 2003; Kincaid *et al.*, 2005; Matsuda *et al.*, 2006; Leconte *et al.*, 2008). B-family polymerases, for instance, readily incorporate benzimidazole opposite all four natural nucleotides (Chiaramonte *et al.*, 2003;

Kincaid *et al.*, 2005). Benzimidazole resembles adenine and guanine except that it lacks all heterocyclic amines (N-1, N-3, N-7 and N-9, Figure 1) and is bereft of extracyclic chemical groups (N², N⁶ and O⁶). While incorporation of benzimidazole opposite T or C would satisfy Watson-Crick geometry, incorporation opposite A or G would clearly not. Systematic addition of substituents found in purines to benzimidazole reveals that they increase discrimination. N-1, N-3, and N⁶ help B-family polymerases exclude incorrect nucleotides, while groups involved in H-bonding (N-1, N², and N⁶) contribute positively to the selection of correct nucleotides (Beckman *et al.*, 2007; Cavanaugh *et al.*, 2009). A-family polymerases, which do not readily incorporate benzimidazole, utilize N-1, N-3, N², and N⁶ for both positive and negative selection (Chiaramonte *et al.*, 2003; Kincaid *et al.*, 2005; Trostler *et al.*, 2009). Thus, minor groove interactions appear to position nucleotides for optimal shape discrimination, while Watson-Crick H-bonding groups may contribute to negative discrimination through formation of non-standard base pairs or through solvent interactions that increase the effective size of the nucleotide (Kool, 2002).

Recent observations with the A-family polymerase T7 Pol suggests that, for some enzymes, negative discrimination of nucleotides is more than just passive steric exclusion (Tsai and Johnson, 2006; Johnson, 2010). The fingers domain resides in an open conformation in the absence of dNTP (Eom *et al.*, 1996; Wang *et al.*, 1997; Bailey *et al.*, 2006). Upon dNTP binding the fingers close, cradling the new base pair in a binding pocket and coordinating the triphosphate through conserved basic residues (Sawaya *et al.*, 1997; Doublié *et al.*, 1998; Kiefer *et al.*, 1998; Franklin *et al.*, 2001; Wing *et al.*, 2008). Initial kinetic experiments revealed only a small reduction in reaction rate with α -phosphothioate dNTP analogues, suggesting that a slow conformational change prior to phosphoryl transfer amplifies binding differences between correct and incorrect nucleotides [reviewed in (Echols and Goodman, 1991; Joyce and Benkovic, 2004)]. Crystallographic evidence for movement of the fingers domain presented an ideal mechanism for this proposed isomerization (Joyce and Benkovic, 2004). However, fluorescence resonance energy transfer (FRET) measurements with *Thermus aquaticus* (Taq) Pol I indicate that closing of the fingers domain is much faster than catalysis (Rothwell *et al.*, 2005). Further evidence for rapid closing was obtained by Tsai and Johnson who introduced a fluorescent label into the fingers domain of T7 Pol near an arginine involved in coordination with the triphosphate (Tsai and Johnson, 2006). Fluorescence corresponding to the open conformation rapidly decreased upon addition of correct dNTPs and unexpectedly *increased* with incorrect dNTPs. Although the structural reasons underlying the changes in fluorescence are unclear, they allowed Tsai and Johnson to monitor how rapidly the polymerase reverts to the open conformation following correct or incorrect dNTP binding. Reversal of the conformational change and dNTP release was 260-times slower for

correct nucleotides than for incorrect nucleotides. Thus, T7 Pol appears to harness the binding energy of incorrect dNTPs to adopt a conformation that actively promotes dissociation (Tsai and Johnson, 2006).

The studies of T7 Pol also show that the rate of phosphoryl transfer differs between correct and incorrect nucleotides (Tsai and Johnson, 2006). All DNA polymerases utilize two divalent metal ions that are coordinated by highly conserved carboxylate residues to catalyze nucleotidyl transfer (Steitz *et al.*, 1994; Beard and Wilson, 2003). With correct geometry, the first metal ion (Metal A) activates the 3'-OH of the primer terminus for in-line nucleophilic attack on the α -phosphate of the incoming dNTP. Metal A and a second metal (Metal B) stabilize the developing negative charge on the pyrophosphate leaving group (Steitz *et al.*, 1994). Crystal structures of some DNA polymerases, such as RB69 Pol and yeast Pol δ , reveal a third metal ion (Metal C) that may further stabilize the leaving group (Franklin *et al.*, 2001; Swan *et al.*, 2009). Proper alignment of the dNTP and stabilization of the leaving group are required for efficient chemistry (Beard and Wilson, 2003). In the T7 Pol studies, phosphoryl transfer was 100-times faster for correct compared to incorrect base pairs, suggesting that the incorrect nucleotides are misaligned in the active site, slowing phosphoryl transfer (Tsai and Johnson, 2006). Thus, T7 Pol increases base-pair selectivity some 10^4 -fold above that attributed to free energy differences in solution (10^{-2}) by two mechanisms: preferential stabilization and faster phosphoryl transfer of correct dNTPs (Tsai and Johnson, 2006; Johnson, 2010).

Proofreading and mismatch repair

Despite the above mechanisms to ensure accurate nucleotide selection, DNA polymerases make hundreds to thousands of mistakes every cell cycle, depending on the size of the genome. DNA sequence context influences the distribution of errors, creating 'hotspots' in the mutation spectra. Intra- or inter-strand base-base stacking interactions in some contexts may stabilize mispairs (Echols and Goodman, 1991). Other mutational hotspots result from sequences where the primer and template easily misalign, causing frameshifts, insertions, deletions or, if transient, base-base mismatches (Bebenek and Kunkel, 2000). To correct replication errors, many DNA polymerases have an exonucleolytic proofreading activity [reviewed in (Reha-Krantz, 2010)]. Kinetic studies indicate that mispaired primer termini reduce polymerization of added nucleotides [reviewed in (Echols and Goodman, 1991; Goodman *et al.*, 1993; Bebenek and Kunkel, 1995)]. Structural studies of an A-family polymerase with mispairs in any one of the first four positions of the upstream primer-template duplex, reveal that mispairs distort the active site by multiple mechanisms that slow catalysis (Johnson and Beese, 2004). Polymerase pausing permits the melting of double-stranded DNA and partitioning of primer termini to exonucleases for editing. These 'proof-reading' exonucleases can be integral to the polymerase

(as in many A- and B-family polymerases) or encoded as separate polypeptides that tightly associate with the polymerase (as in C-family polymerases) (Johnson and O'Donnell, 2005). Amino acid substitutions that inactivate exonuclease activity confer a mutator phenotype characterized predominantly by increased base-substitution errors and frameshift mutations (Simon *et al.*, 1991; Morrison *et al.*, 1993; Reha-Krantz and Nonay, 1993; Schaaper, 1993; Jin *et al.*, 2001; Fortune *et al.*, 2005). Thus, in the absence of proofreading, polymerases can slowly extend many mispaired primer termini, creating base-base mismatches or small bulges in duplex DNA.

Distortions in the DNA due to replication errors are substrates for post-replicative mismatch repair (MMR) [reviewed in (Kunkel and Erie, 2005; Iyer *et al.*, 2006)]. MMR complexes bind mismatched DNA and nick the nascent strand, providing entry points for enzymes that remove and resynthesize the error-containing DNA strand (Kadyrov *et al.*, 2006; Kadyrov *et al.*, 2009). If left unrepaired, mismatches lead to permanent mutations in the next round of replication when the mutated strand serves as a template. Defects in proofreading or MMR increase mutation rates between 10- and 1000-fold in organisms as diverse as phage, bacteria, yeast, mice, and humans (Muzyczka *et al.*, 1972; Simon *et al.*, 1991; Fishel *et al.*, 1993; Morrison *et al.*, 1993; Parsons *et al.*, 1993; Schaaper, 1993; Reitmaier *et al.*, 1997; Goldsby *et al.*, 2002; Albertson *et al.*, 2009). Simultaneous inactivation of both pathways increases mutation rates by more than 10,000-fold (Morrison *et al.*, 1993; Schaaper, 1993). Thus, polymerase nucleotide selection, proofreading, and MMR act sequentially to achieve an accuracy of better than 10^{-9} mutations/base pair/cell division (Drake *et al.*, 1998).

Discovery of antimutators

Given the low cellular mutation rate, the isolation of mutants with improved replication fidelity poses distinct challenges. Whereas mutators emerge when cells are subjected to adaptive pressure [reviewed in (Sturtevant, 1937; Drake *et al.*, 1998; Giraud *et al.*, 2001b; de Visser, 2002; Elena and Lenski, 2003; Denamur and Matic, 2006)], few selection schemes exist for antimutators. Furthermore, many processes contribute to the residual background mutation rate observed in wild-type cells so that an antimutator allele in any one pathway may not significantly improve overall fidelity (Drake, 1993). Cells contend with various forms of DNA damage through translesion DNA synthesis, base-excision repair, nucleotide excision repair, and recombination (Friedberg *et al.*, 2006). The actual contributions of individual pathways to background mutation rate vary with cell type, genetic background, and environmental conditions. In some cases, one pathway may predominate. For instance, in yeast, DNA polymerase zeta (Pol ζ) generates between 50 and 75% of all background mutations (Quah *et al.*, 1980; Roche *et al.*, 1994; Datta *et al.*, 2000), complicating the isolation of antimutators in other pathways. For the

above reasons, most antimutators have been identified by laborious screening procedures or as suppressors of a mutator phenotype or other deleterious DNA replication-associated defects.

T4 Pol

Drake uncovered the very first antimutator DNA polymerase variants while screening a collection of bacteriophage T4 gene 43 mutations for mutator phenotypes (Drake and Allen, 1968; Drake *et al.*, 1969). Gene 43 encodes the viral replicase, a B-family DNA polymerase with homology to eukaryotic replicative polymerases (Pol α , Pol δ , and Pol ϵ). Gene 43 mutant phage strains were monitored for reversion of *rII* alleles containing premature stop codons that prevent T4 from replicating in *Escherichia coli* that are lysogenic for phage lambda. The *rII* mutant growth defect provided an exquisitely sensitive assay for mutation frequency, allowing detection

of a single mutant in 10^9 – 10^{10} phage. Two temperature-sensitive mutants exhibited lower rates of reversion than wild-type T4 (Drake and Allen, 1968; Drake *et al.*, 1969), providing evidence that DNA polymerase fidelity could be improved by single amino acid substitutions (Figure 2). This revelation was startling at the time: given the deleterious nature of most mutations, why hadn't evolution favored these higher fidelity variants?

Early biochemical analyses in the Bessman and Nossal labs established a paradigm for how changes to the peptide sequence of a DNA polymerase could increase fidelity. T4 Pol contains a functional proofreading domain (Muzyczka *et al.*, 1972). The first antimutator variants, later shown to carry alanine to valine substitutions at positions 737 or 777 in the thumb domain (Figure 3A, 3H), hydrolyzed more dNTPs per base-pair synthesized than wild-type polymerase (Muzyczka *et al.*, 1972). Further analysis of the A737V variant revealed the polymerase to

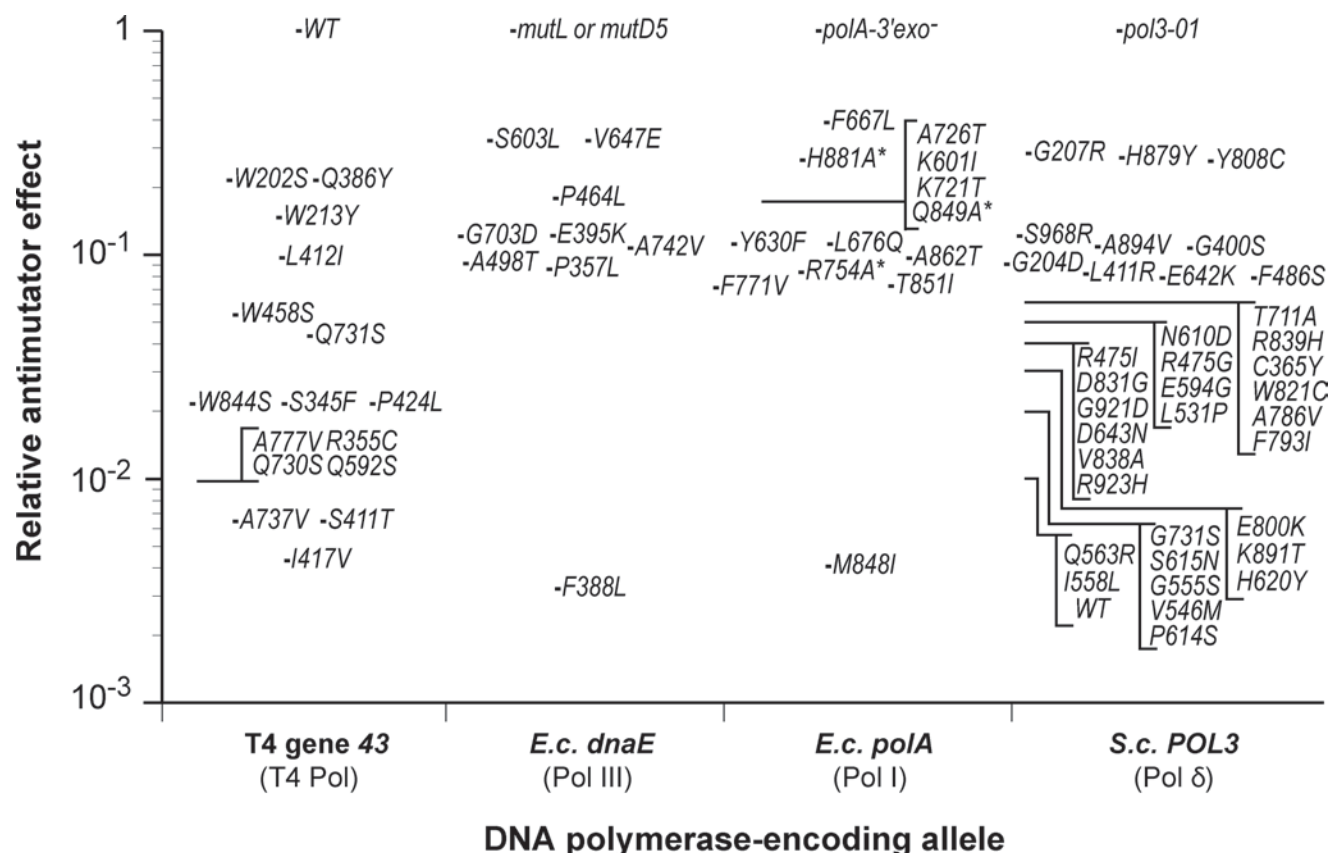


Figure 2. Antimutator effects of DNA polymerase variants. The relative mutation frequency or rate conferred by T4 Pol, *E. coli* Pol III and Pol I, and yeast Pol δ antimutator variants are shown. Each variant is indicated by the single amino acid substitution that confers antimutator activity. The T4 Pol antimutator effects are based on reversion frequency of *rII* UV199oc relative to wild-type T4 Pol (Reha-Krantz, 1988; Reha-Krantz and Nonay, 1993; Reha-Krantz and Nonay, 1994; Reha-Krantz and Wong, 1996). The value for the S411T single mutant is inferred from the reversion frequency of the S411T, L412M double mutant (Reha-Krantz and Nonay, 1993). Pol III antimutator effects are based on the average degree of *mutL* and/or *mutD5* mutator suppression, as measured by the frequency of Rifampicin-resistance, Nalidixic acid resistance, GalK2 (*ochre*) reversion and/or *lacZ*118 (*ochre*) reversion (Fijalkowska *et al.*, 1993; Fijalkowska and Schaaper, 1993; Fijalkowska and Schaaper, 1995; Fijalkowska and Schaaper, 1996). The average background reversion frequency for WT cells is reported to be 3×10^{-4} relative to *mutD5* (Schaaper, 1988). All Pol I antimutators were engineered into an *E. coli* Pol I variant lacking proofreading capability (D424A), with the exception of the *Taq* Pol I antimutator F667L (Suzuki *et al.*, 2000). The relative antimutator activities of Pol I variants marked with an asterisk are based on *in vitro* gap-filling mutation rate determinations (Minnick *et al.*, 1999). All other Pol I values are based on reversion frequencies of a premature nonsense codon in a plasmid-borne β -lactamase gene (Suzuki *et al.*, 2000; Shinkai and Loeb, 2001; Shinkai *et al.*, 2001; Loh *et al.*, 2007). The effect of Pol δ antimutators are based on suppression of the proofreading-defective *pol3-01* mutator allele, as measured by the rate of FOA-resistance mutations (Herr *et al.*, 2011).

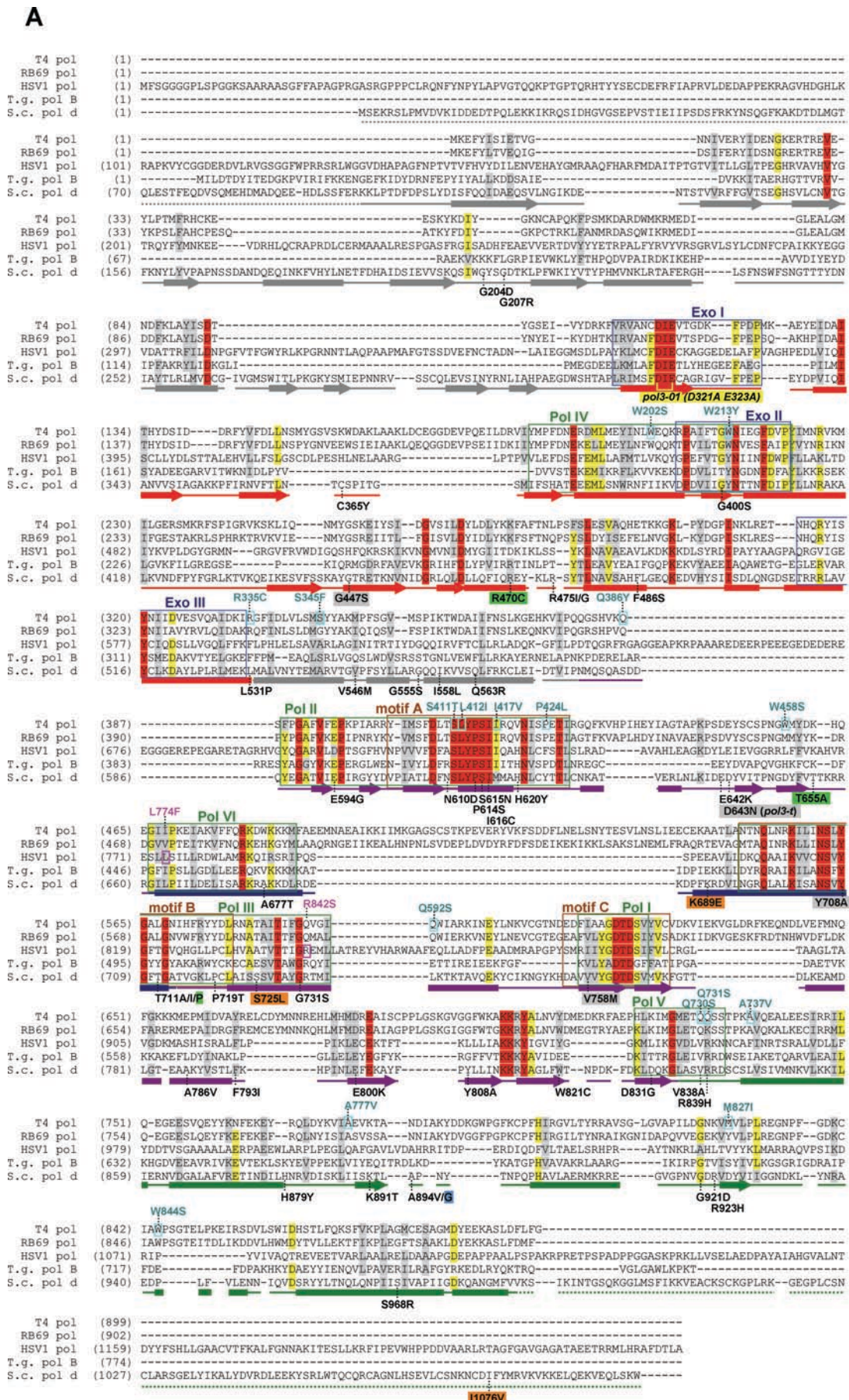


Figure 3. (Continued)

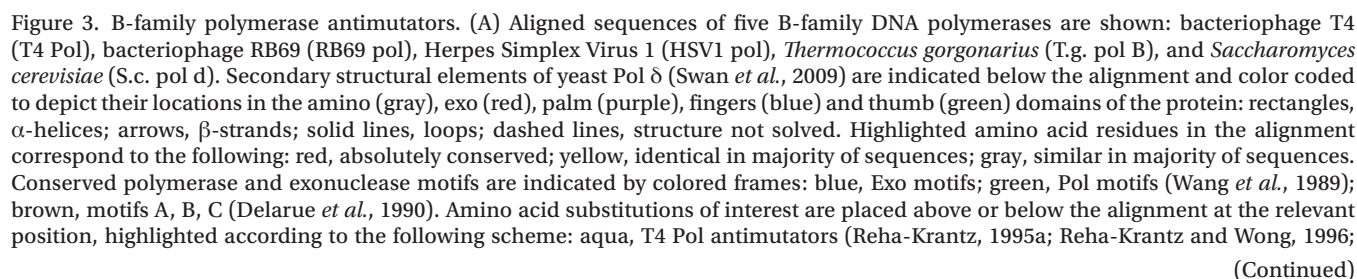


Figure 3. (Continued) Reha-Krantz, 2010); magenta, HSV1 Pol antimutators (Hall *et al.*, 1984; Gibbs *et al.*, 1988; Hwang *et al.*, 2004; Tian *et al.*, 2009); yellow, yeast *pol3-01*; no highlight, antimutators isolated as suppressors of the *pol3-01* mutator phenotype; green, antimutators isolated as suppressors of the *pol3-D407A* mutator phenotype; orange, antimutators isolated as suppressors of the *pol3-F406A* mutator phenotype (three substitutions in the same mutant); blue, antimutator isolated as a suppressor of the *pol3-D463A* mutator phenotype; gray, previously reported antimutators in Pol δ (*pol3-t* (D643N), G447S, Y708A, and V758M) (Tran *et al.*, 1999a; Hadjimarcou *et al.*, 2001; Pavlov *et al.*, 2004; Li *et al.*, 2005). The antimutators affecting *pol3-01*, *pol3-D407A*, *pol3-F406A*, and *pol3-D463A* are from Herr *et al.* (2011). (B–I) Locations of antimutator amino acid substitutions in T4 Pol [B, D, F, and H; mapped onto RB69Pol (Franklin *et al.*, 2001), Protein Data Bank accession code 1IG9] and Pol δ [C, E, G, and I; mapped onto Pol δ (Swan *et al.*, 2009), PDB accession code 3IAY]. The proteins are shown as schematic diagrams of the α -carbon backbone with α -carbons or amino acids of interest depicted as space-filling spheres. Wild type amino acid residues corresponding to antimutator mutations are shown as teal spheres and labeled to indicate the antimutator substitutions. Structural domains are color coded as in the alignment in panel A. The incoming dNTPs are denoted by green CPK sticks and the template nucleotides by orange CPK sticks. Active site carboxylate side chains are gray CPK sticks coming out of the purple (palm) or red (exo) ribbons. Metal ions are small black spheres. Important non-mutated residues proximal to the antimutator substitutions are also shown as space-filling spheres and color coded according to domain. (B and C) Distribution of antimutator substitutions. The α -carbons of wild type residues corresponding to antimutators are shown as teal spheres. (D and E) Antimutator substitutions near the polymerase active site. The fingers and thumb domains have been removed for clarity. T4 Pol S345 and Pol δ residues V546, G555, I558, and Q563 from the amino domain are in three closely associated α -helices that bind the template and buttress the fingers domain. The antimutator substitution (V758M) that conferred PAA-resistance to a Pol δ -L612M strain is shown in yellow. (F and G) Antimutator substitutions in the exo domain. All T4 Pol antimutator variants occurred in the context of a functional exonuclease, unlike Pol δ antimutator polymerases, which carried D321A and D323A substitutions that inactivate the exonuclease. The dotted red line in (G) corresponds to a missing loop in the Pol δ structure (amino acids 491–496). (H and I) Antimutator substitutions in the DNA binding track. The primer (gold) and template (brown) are held by a series of interactions along the DNA minor groove. Minor groove ‘sensing’ residues in the KKRYA motif of the palm domain (T4 Pol K702, K703, R704; Pol δ K813, K814 and R815) are shown as space-filling spheres labeled K and R. Unpaired 5′ nucleotides of the template along with the amino, exo, and fingers domains have been removed for clarity.

be less processive and efficient at strand-displacement synthesis (Gillin and Nossal, 1976a; Gillin and Nossal, 1976b). In contrast, mutator polymerase variants hydrolyzed less dNTPs per base-pair synthesized than wild type, strongly suggesting that replication fidelity was determined by the balance between proofreading and primer extension (Muzyczka *et al.*, 1972). The high rate of dNTP hydrolysis by the antimutator polymerases indicates that proofreading frequently removes even correctly paired 3′ primer termini. As a sign of this hyper-editing, T4 antimutator phage replicate poorly in an *optA1 E. coli* strain, which has lower dNTP pools due to high levels of a cellular dGTPase (Gauss *et al.*, 1983; Reha-Krantz and Nonay, 1994). The demonstration that increased proofreading efficiency comes with a pronounced cost in fitness provides one explanation as to why evolution has not favored the higher fidelity variants. Studies of the antimutator alleles using other mutation reporter sites revealed another answer. The antimutator polymerases lowered mutation rates for AT→GC transitions, but not for GC→AT transitions or transversions (Drake *et al.*, 1969; Ripley, 1975). Indeed, the rate of transversions was slightly higher at some sites. Thus, gains in fidelity conferred by some antimutators may be specific to certain sequence contexts. Moreover, these same polymerase variants may cause decreased fidelity in other contexts (Drake, 1993). For this reason, the relative magnitude of antimutator activity based on reversion assays reported in Figure 2 should be interpreted with caution. Forward mutation rate assays based on gene inactivation, which have numerous potential mutation sites, give a more balanced assessment of overall replication fidelity.

After this early work, Linda Reha-Krantz isolated a large collection of new antimutator T4 Pol variants [reviewed in (Reha-Krantz, 1995b)]. The first variants were identified by growing T4 gene 43 amber mutants in bacterial strains expressing different suppressor tRNAs

(Reha-Krantz, 1988). By varying the suppressor tRNA, the influence of different amino acids at the stop codon position could be assessed. Eight stop codon/suppressor tRNA combinations produced an antimutator phenotype (Figure 2). Those with the strongest effect (Q730S, Q731S, and W844S) clustered together in the thumb domain with A737V and A777V (Figure 3A, 3H), suggesting that they shared a common mechanism.

A strong antimutator substitution (I417V) in the palm domain of T4 Pol was discovered in a *mutator* phage encoding eight other amino acid substitutions. Analysis of each substitution's effect on polymerase fidelity revealed the presence of both mutator and antimutator changes in the same T4 Pol variant, suggesting that the I417V-encoding mutation may have arisen to mollify the mutator (Reha-Krantz, 1988). I417V affects motif A, which is highly conserved in DNA polymerases (Figure 3A) and helps to form the dNTP binding site (Figure 3D). In theory, the mutation could elevate replication fidelity by increasing nucleotide selectivity and/or reducing polymerase processivity, thereby allowing more time for proofreading. The I417V mutant phage grows poorly on the *optA1 E. coli* strain, and the purified polymerase displays reduced processivity (Reha-Krantz and Nonay, 1994). These observations are consistent with increased fidelity due to hyper-editing, although a concomitant effect on nucleotide selectivity has not been excluded.

Additional antimutagenic changes emerged with the development of the first genetic selection for antimutators. The selection utilized a T4 Pol variant allele encoding an L412M substitution, which was originally identified as a suppressor mutation of the *optA1* sensitivity conferred by I417V (Reha-Krantz and Nonay, 1994; Stocki *et al.*, 1995). As a single amino acid substitution in T4 Pol, L412M elevates mutation rate moderately and confers sensitivity to the antiviral drug and pyrophosphate analogue, phosphonoacetic acid (PAA) (Reha-Krantz *et al.*, 1993;

Reha-Krantz and Nonay, 1994). A PAA-resistant Herpes Simplex Virus I (HSV I) mutant polymerase encoding an R842S amino acid substitution (Figure 3A) was previously shown to be an antimutator (Hall *et al.*, 1984; Gibbs *et al.*, 1988). On the hunch that PAA-resistance was linked to antimutagenesis, Reha-Krantz selected for PAA-resistant variants of the L412M T4 mutant phage as well as a second PAA-sensitive mutant involving duplication of D836 (Reha-Krantz and Nonay, 1994; Reha-Krantz and Wong, 1996). They identified substitutions in motif A (S411T and L412I) and elsewhere in the polymerase (S335C and S345F) that bestowed antimutator activity even in the absence of the PAA-sensitizing mutation (Reha-Krantz and Nonay, 1994; Reha-Krantz and Wong, 1996). As expected, S335C also confers *optA1* sensitivity. Selection for genetic suppressors revealed a P424L substitution that restored the ability of the S335C phage to propagate on the *optA1* strain. Remarkably, the P424L single-mutant phage is an antimutator and unable to grow on *optA1 E. coli* (Li *et al.*, 2010a). The S335C,P424L double-mutant phage retains substantial antimutator properties, although the mutation rate is modestly higher than either of the single mutants alone (Li *et al.*, 2010a). The ability of the double mutant to reduce the antimutator and *optA1* sensitivity phenotypes of the single mutants suggests that the amino acid substitutions affect a common mechanism that influences fidelity. Genetic suppressor analysis of the *optA1*-sensitive phenotype of P424L mutant phage suggests that this mechanism may relate to closure of the fingers domain (Li *et al.*, 2010a).

Together, these elegant genetic and biochemical experiments with T4 Pol serve as a reference for all other antimutator studies, and indicate that a wide variety of single amino acid substitutions can increase replication fidelity (Figures 2, 3). Moreover, the recurrent phenotype of *optA1* sensitivity and PAA-resistance argues that most if not all T4 antimutator polymerases increase replication fidelity by shifting the balance from polymerization to proofreading.

E. coli Pol III

Studies with *E. coli* Pol III in the 1990s by Roel Schaaper and colleagues were the first to indicate that amino acid substitutions in DNA polymerases can elevate replication fidelity in the absence of proofreading activity. Because Pol III performs the bulk of both leading- and lagging-strand DNA synthesis in *E. coli* (Johnson and O'Donnell, 2005), mutator variants are especially detrimental. The polymerase activity resides in the α subunit of Pol III, encoded by the *dnaE* gene, while the proofreading activity is found in the tightly associated ϵ subunit, encoded by the adjacent gene on the chromosome, *dnaQ*. The first Pol III antimutator variant was found as a suppressor of the strong *dnaQ* mutator allele, *mutD5* (Schaaper and Cornacchio, 1992). *mutD5* exhibits a conditional mutator phenotype, producing moderate mutation rates in minimal media and high rates in rich conditions (Maruyama *et al.*, 1983; Scheuermann *et al.*, 1983), where polymerase

errors partially saturate MMR (Schaaper, 1988; Schaaper, 1989; Schaaper and Radman, 1989). Mutation accumulation in *mutD5* cells slows bacterial growth, and Schaaper isolated a spontaneous, faster growing clone of cells. Phage P1 transduction of *mutD5* from this clone resulted in both large and small colonies, suggesting that the isolate contained a suppressor mutation linked to *dnaQ* (Schaaper and Cornacchio, 1992). The mutation (later shown to encode a V648E substitution) mapped to *dnaE* and suppressed not only the growth defect, but also the *mutD5* mutator phenotype (Figure 2), as determined by stop codon reversion and 'forward' mutation frequency assays, which score mutations at multiple sites.

With the discovery that changes to the polymerase subunit of Pol III can offset proofreading defects, Schaaper and co-workers addressed whether Pol III antimutators suppress defects in MMR. In bacterial MMR, a MutS homodimeric protein recognizes the mismatched DNA and recruits a MutL homodimer, which in turn recruits the MutH endonuclease that nicks the nascent DNA and initiates repair (Iyer *et al.*, 2006). Mutagenized DNA from the *dnaE-dnaQ* chromosomal region was transduced into a *mutL* mutator strain, and mutants with an attenuated mutator phenotype were identified by scoring reversion frequency with a papillation assay (Fijalkowska *et al.*, 1993). Colonies with lower numbers of papillae also exhibited reduced mutation frequencies by forward mutation assays, confirming their antimutator character. The mutants carried mutations in *dnaE* that encoded single amino acid substitutions in the polymerase subunit (Figures 2, 4) (Fijalkowska and Schaaper, 1993). The ability to isolate *dnaE*-encoded suppressors of *mutL* indicates that most mutations in this strain derive from Pol III errors that escape proofreading. Thus, the antimutators reduce the frequency of the most common Pol III-derived MMR substrates.

The mechanism of the *mutL* Pol III antimutators could be interpreted within the paradigm established by the T4 studies in which antimutators lower mutation rates by increasing proofreading efficiency. In order to determine whether proofreading activity of the ϵ subunit was required for their antimutator phenotype, Schaaper and colleagues moved the new *dnaE* antimutators into the *mutD5* strain (Fijalkowska and Schaaper, 1995). The new antimutators effectively suppressed the mutator phenotype of the proofreading mutant (Figure 2), suggesting that these antimutators did not require the ϵ subunit for their gains in replication fidelity. However, the underlying mutation in the *mutD5* allele encodes a T to I change (**DTET** to **DTEI**, italics) adjacent to two highly conserved acidic residues required for catalysis (in bold). Because these essential catalytic residues were still intact, it remained possible that the antimutators simply increased the efficacy of the residual proofreading activity of the mutant ϵ subunit. Complete elimination of proofreading activity by substitution of the essential aspartate and glutamate residues to alanine (ATAT) produced a very strong dominant mutator phenotype

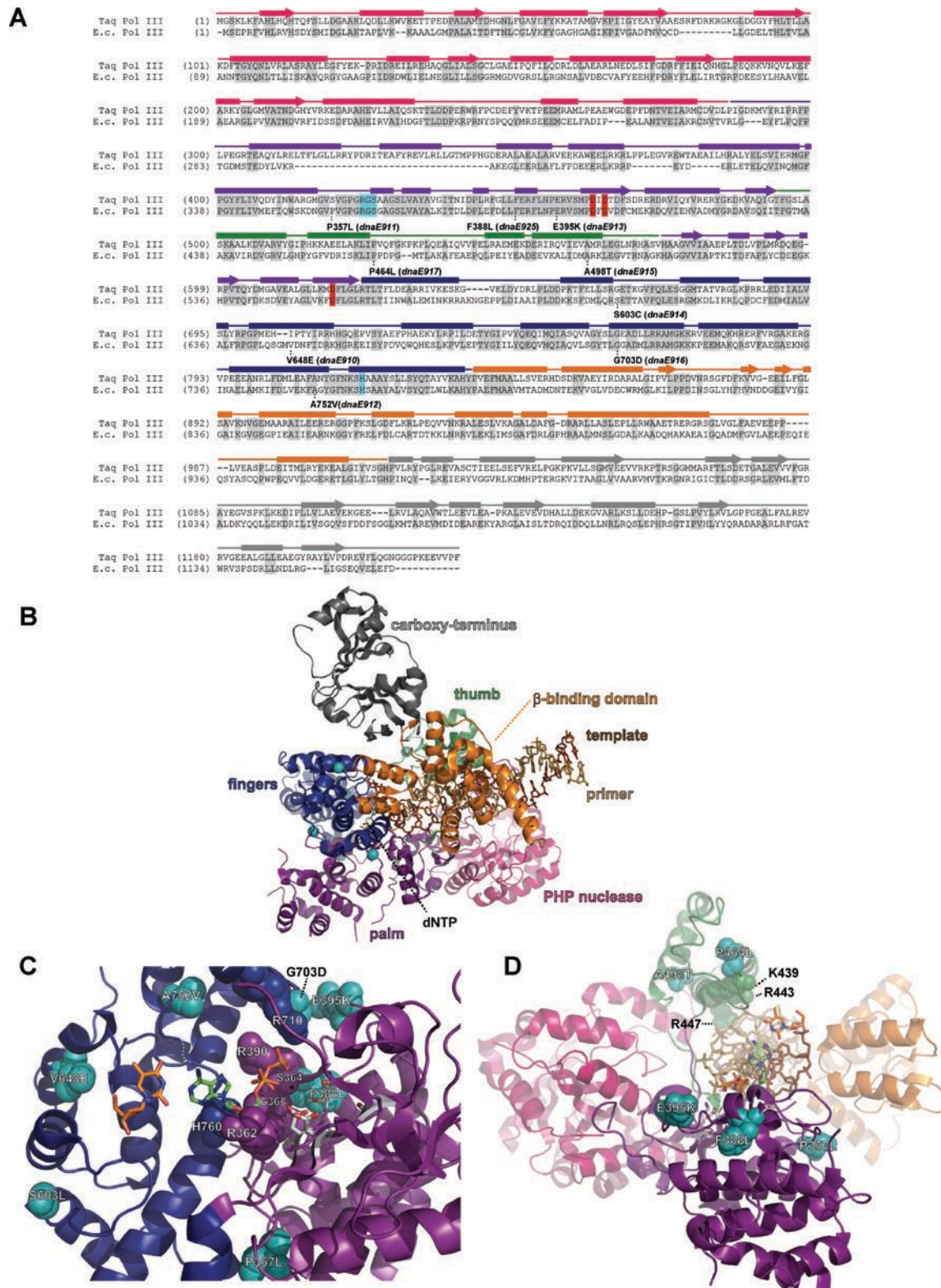


Figure 4. Pol III antimutators. (A) Alignment of Pol III peptide sequences from *Thermus aquaticus* (Taq Pol III) and *Escherichia coli* (E.c. Pol III). Secondary structural elements based on the Taq Pol III structure (Wing *et al.*, 2008) (PDB accession code 3E0D) are indicated below the alignment and color coded to depict their location (PHP, magenta; palm, purple; fingers, blue; thumb, green; β-binding, orange; carboxy-terminus, charcoal): rectangles, α-helices; arrows, β-strands; solid lines, loops. Amino acids shaded in gray are conserved between *Taq* and *E. coli*. Aspartates that coordinate metal ions for catalysis are shaded in red. Residues that stack with ribose of incoming dNTP are shaded in cyan. *E. coli* *dnaE* amino acid changes and allele numbers are depicted below the alignment. (B) Distribution of antimutator substitutions. The α-carbons of antimutator positions are shown as teal spheres. (C) Antimutators near the Pol III active site. The RGS motif (362–364) and H760 cradle the incoming dNTP, while R390 and R710 interact with the triphosphate. (D) Antimutators in the Pol III Thumb. A498T and P464L affect the positioning of an α-helix that includes three basic residues that interact with the primer-template backbone (K439, R443, R447).

that was lethal when the mutant protein was the sole source of the ϵ subunit (Fijalkowska and Schaaper, 1996). The antimutator substitutions in the α subunit rescued this lethality, confirming decisively that the amino acid alterations in Pol III lower mutation rates independently of the proofreading activity of the ϵ subunit (Fijalkowska and Schaaper, 1996).

Some Pol III antimutators may suppress polymerase errors by increasing nucleotide selectivity. Evidence of altered base selectivity comes from the observation that certain Pol III antimutators suppress the mutator phenotype of *mutT*, which encodes an 8-oxo-dGTPase. By 'cleansing' nucleotide pools, the *mutT* protein minimizes transversions caused by A•8-oxo-dGTP base pairs (Schaaper, 1996). Schaaper identified two new Pol III antimutator variants that suppressed the *mutT* mutator phenotype 10-fold, and these were also effective in suppressing the *mutL* mutator phenotype. One antimutator encoded by the *dnaE940* allele (mutation unknown) was three times more effective at suppressing *mutT* than other Pol III antimutators that were equally effective in suppressing *mutL* (Schaaper, 1996). The disproportionate effect of this allele on *mutT* suggests that the corresponding polymerase contains changes in the active site that excludes A•8-oxo-dGTP base pairs. Whether improved base-selectivity accounts for all Pol III antimutator effects is uncertain. As recognized by Schaaper, the antimutators improve replication fidelity in the context of the cellular milieu and may promote polymerase dissociation from the DNA, allowing an alternate proofreading activity to edit the error (Fijalkowska and Schaaper, 1995).

Recent reports indicate that the PHP domain of the *Thermus thermophilus* Pol III α subunit exhibits a Zn²⁺-dependent 3'→5' exonuclease activity capable of editing mispaired primer termini (Stano *et al.*, 2006). Substitution of two aspartates in the presumptive PHP active site from *Thermus aquaticus* Pol III inactivated nuclease activity, facilitating co-crystallization of the enzyme with primer-template DNA (Wing *et al.*, 2008). The resulting polymerase structure revealed the PHP active site is a full turn of the DNA helix away from the polymerase active site (Figure 4B), suggesting that dissociation of the duplex or significant unwinding of the primer strand would be required for PHP editing of the 3' end (Wing *et al.*, 2008). Is the *E. coli* Pol III PHP domain an active nuclease and could it contribute to antimutagenesis in Schaaper's mutants? Arginine residues replace two histidines in the *E. coli* PHP active site (R10 and R203; Figure 4A) (Aravind and Koonin, 1998). Moreover, purified recombinant *E. coli* Pol III α subunit did not degrade a primer-template with a terminal mismatch (Mo and Schaaper, 1996). These observations question whether the *E. coli* PHP is indeed a nuclease. However, it remains possible that the *E. coli* PHP degrades single-stranded DNA or requires the context of the holoenzyme to be active. A key test for this or any other candidate proofreader is to determine whether substitution of the active site aspartates (see *T. aquaticus*, above) counteracts the effects of the antimutators.

E. coli Pol I

The T4 Pol and *E. coli* Pol III polymerases play essential roles in the replication of their respective genomes, thereby facilitating the use of genetic approaches to identify active antimutator variants. In contrast, *E. coli* Pol I is not essential for survival of wild type cells (Kornberg and Baker, 1992). Thus, Pol I antimutator variants were initially identified by site-directed mutagenesis of an exonuclease-deficient Pol I allele (*pol3A-3'exo-*) followed by screening for biochemical activity and alterations in mutation rate (Minnick *et al.*, 1999). Using this approach, three amino acid substitutions were found (R754A, Q849A, and H881A) that individually confer significant antimutator activity (Figure 2) (Minnick *et al.*, 1999).

To facilitate the study of Pol I fidelity, Loeb and colleagues cleverly combined random mutagenesis of Pol I (Suzuki *et al.*, 2000; Patel *et al.*, 2001; Shinkai and Loeb, 2001; Loh *et al.*, 2007) with a genetic selection strategy first used to identify functional Pol β variants (Sweasy and Loeb, 1993). Pol I participates in DNA repair as well as in the initiation of ColE1 plasmid replication (Kornberg and Baker, 1992). Functional derivatives of Pol I were identified by their ability to complement a temperature-sensitive Pol I allele (*polA12*) in a synthetically lethal interaction with *recA718* in *E. coli*. The *polA12 recA718* double-mutant strain exhibits persistent single-stranded gaps in genomic DNA at the non-permissive temperature, suggesting that the complementing mutant polymerases perform gap repair (Witkin and Roegner-Maniscalco, 1992). Once functional Pol I variants were identified, they were screened for altered fidelity *in vitro* by a gap-filling forward mutation assay or *in vivo* by scoring reversion of a nonsense codon in the β -lactamase gene inserted close to the origin of replication on a ColE1 plasmid.

The first screen, targeting motif B of the fingers domain (O-helix) of *Thermus aquaticus* (*Taq*) Pol I, which is naturally proofreading-deficient, identified an F667L substitution that lowered the frequency of transversions *in vitro* (Suzuki *et al.*, 2000). The second mutagenesis screen, targeting motif A of proofreading-proficient *E. coli* Pol I, identified an E710D substitution that lowered β -lactamase reversion three-fold (Shinkai and Loeb, 2001; Shinkai *et al.*, 2001). A related screen targeting motif A of *Taq* Pol I identified a mutant with multiple changes (L605R, L606M, V607K, A608S, L609I, S612R) that encoded a higher fidelity enzyme (Patel *et al.*, 2001). Finally, Loeb and colleagues mutagenized the entire polymerase domain of exonuclease-deficient *E. coli* Pol I (Loh *et al.*, 2007). A surprising 12% of mutants suppressed β -lactamase reversion at least 10-fold. Some antimutator polymerases sustained single amino acid changes (Y630F, L676Q, K721T, F771V, M848I, T851I, and A862T). But most enzymes carried multiple substitutions, which complicates the interpretation. Analysis of one of these complex variants found that both amino acid substitutions in the clone (K601I and A726T) individually confer an antimutator phenotype (Loh *et al.*, 2007). Moreover, the substitutions acted in an additive fashion

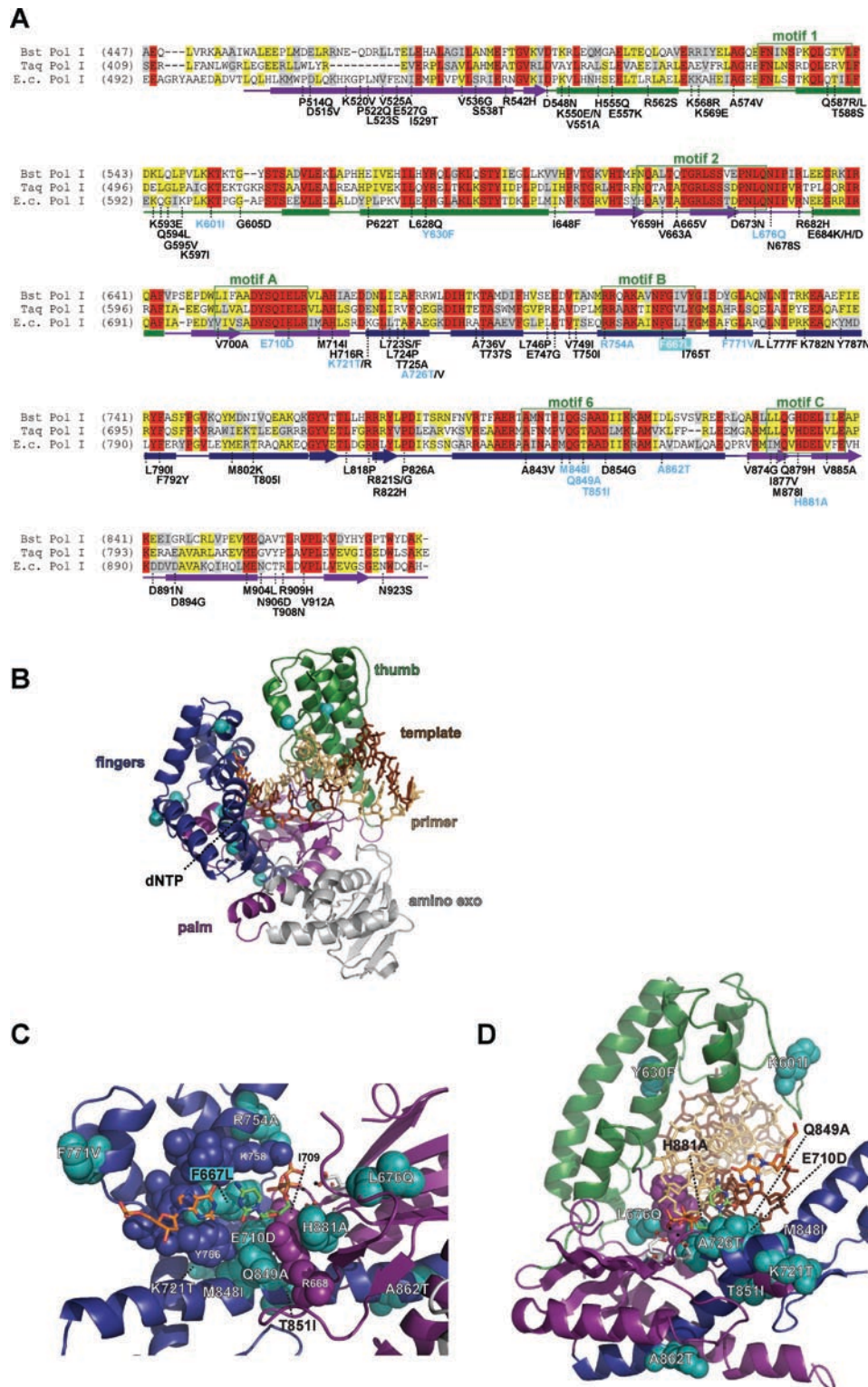


Figure 5. Pol I antimutators. (A) Alignment of Pol I peptide sequences from *Bacillus stearothermophilus* (Bst Pol I), *Thermus aquaticus* (Taq Pol I), and *Escherichia coli* (E.c. Pol I). Secondary structural elements based on the Taq Pol I structure (Li *et al.*, 1998) (PDB accession code 3KTQ) are indicated below the alignment and color coded to depict their locations in the amino (the proofreading exonuclease in *E. coli* Pol I; gray); palm (purple), thumb (green) and fingers (blue) domains of the protein: rectangles, α -helices; arrows, β -strands; solid lines, loops. Highlighted amino acid residues in the alignment correspond to the following: red, absolutely conserved; yellow, identical in majority of sequences; gray, similar in majority of sequences. Conserved Pol I motifs are indicated by green frames. Amino acid substitutions of interest are placed below the alignment at the relevant position, color coded according to the following scheme: teal, single amino acid antimutator substitutions in *E. coli* Pol I (Minnick *et al.*, 1999; Loh *et al.*, 2007); black, substitutions found in complex mutant alleles from Loh *et al.* (Loh *et al.*, 2007); white lettering with teal shaded box, Taq Pol I antimutator, (Suzuki *et al.*, 2000). (B) Distribution of antimutator substitutions. The α -carbons of antimutator positions are shown as teal spheres. (C) Pol I active site antimutator substitutions. (D) Pol I antimutators that may influence DNA binding. Purple spheres next to L676Q correspond to adjacent amino acids from Motif2 that bind the -3 base pair of the primer-template (PNL676QN).

in the double-mutant enzyme, suggesting that they lower mutation rates by independent means. Thus, other complex Pol I variants may also carry multiple antimutator amino acid substitutions.

The single amino acid changes sufficient for Pol I antimutagenesis cluster with substitutions encoded in the complex mutants (Figure 5A). Mutations in several other regions were repeatedly found in the complex antimutator clones, highlighting candidate regions for further study. These include sequences upstream of motif 1 (514–574), upstream of motif 6 (818–826), and downstream of motif C (904–912) (Figure 5A). The wide distribution of antimutator substitutions in Pol I mirrors the diverse locations of antimutator substitutions in T4 Pol and Pol III. As with the Pol III antimutators, increases in replication fidelity occurs independently of the associated proofreading exonuclease, which is inactivated in these screens. Thus, Pol I antimutator amino acid substitutions may either increase nucleotide selectivity or reduce extension of mispaired primer termini.

Kinetic analyses suggest that mutant Pol I enzymes with R754A and Q849A substitutions are less prone to extending mismatched primer termini than wild type Pol I (Minnick *et al.*, 1999). Likewise, the complex motif A antimutator variant of *Taq* Pol I (L605R, L606M, V607K, A608S, L609I, S612R) also exhibited reduced mispair extension (Patel *et al.*, 2001). In failing to extend mispairs efficiently, these mutant polymerases may lower apparent error rates by preventing misinsertions from being extended into duplex DNA; rare unextended mispairs are not detected in the gap-filling assay. A failure to extend mispairs could also account for reductions in mutation rates in the β -lactamase reversion assay *in vivo*. Pol III normally replaces Pol I following initiation of ColE1 plasmid replication (Kornberg and Baker, 1992). This usually occurs at a primosome assembly site (Kornberg and Baker, 1992), but can occur in its absence (Camps *et al.*, 2003). The reversion site used in these studies is positioned 230 nucleotides downstream of the origin. Thus, any defect that promotes Pol I switching to Pol III, at or before the reversion site, would produce an antimutator phenotype.

Yeast Pol δ

In current models of eukaryotic DNA replication, Pol δ performs the bulk of lagging-strand DNA synthesis, while Pol ϵ synthesizes the leading strand [reviewed in (Kunkel and Burgers, 2008)]. The first report of a eukaryotic antimutator polymerase variant was in yeast and resulted from combining the Pol δ frameshift mutator allele *pol3-t* (Tran *et al.*, 1995) with the proofreading-defective allele *pol3-01* (Morrison *et al.*, 1993). Rather than producing a compound mutator phenotype, the double mutant exhibited a three-fold reduction in mutation rate relative to *pol3-01* (Tran *et al.*, 1999a). Pavlov *et al.* demonstrated that another weak mutator allele (*pol3-Y708A*) suppressed *pol3-01* frameshift mutations, although not base-substitution errors (Pavlov *et al.*, 2004). Curiously,

the analogous substitution (Y831A) in proofreading-defective Pol ϵ (encoded by *pol2-4*) suppressed both types of errors (Pavlov *et al.*, 2004). The mutator phenotypes of *pol3-t* and *pol3-Y708A* partially depend on monoubiquitination of PCNA at K164 and the activity of the TLS polymerase, Pol ζ (Northam *et al.*, 2006), suggesting that the amino acid substitutions in these *pol3* mutants promote remodeling of the replication fork. The observation that *pol3-t* and *pol3-Y708A* suppress the *pol3-01* mutator phenotype suggests that error-free pathways may also gain access to primer termini in the mutant strains. Thus, the contrasting effects of antimutator substitutions on proofreading-defective Pol δ and Pol ϵ may reflect intrinsic differences between the polymerases or the ability of alternate replication enzymes to correct or extend errors on the leading and lagging strands.

Additional antimutator alleles of Pol δ were identified by Reha-Krantz and colleagues, who extended their T4 studies into yeast. Introduction of a G447S substitution into a β -hairpin structure in the exonuclease domain of Pol δ produced a frameshift-suppression phenotype (Hadjimarcou *et al.*, 2001). The analogous change in T4 Pol (G255S) had been isolated as a suppressor of the *optA1* sensitivity conferred by an antimutator (Stocki *et al.*, 1995). In T4 Pol, the substitution decreases fidelity by inhibiting hyper-partitioning of the primer strand to the exonuclease domain (Stocki *et al.*, 1995). This suggests that yeast Pol δ G447S suppresses misalignment errors that cause frameshifts by minimizing primer DNA dissociation. In a separate study, Reha-Krantz demonstrated that Pol δ could be made PAA-sensitive by the introduction of an L612M substitution (analogous to T4 Pol L412M; see above) (Li *et al.*, 2005). Selection for PAA-resistance in the *pol3-L612M* strain identified a V758M substitution that suppressed the L612M mutator phenotype in an MMR-deficient (*msh2 Δ*) mutant background (Li *et al.*, 2005). Intriguingly, substitution of the L612-equivalent in human Pol δ with lysine (L606K) increases polymerase fidelity *in vitro* in the presence or absence of proofreading activity (Schmitt *et al.*, 2010). Taken together, these observations indicate that Pol δ and T4 Pol have common antimutagenic pathways. This conclusion is strongly supported by the large collection of antimutator changes we recently isolated in Pol δ that suppress the proofreading defect encoded by *pol3-01* [(Herr *et al.*, 2011); Figures 2 and 3A].

We exploited the observation that in yeast, as in *E. coli*, excessive mutation accumulation undermines colony formation. Evidence for this ‘replication error-induced extinction’ came from early studies by Morrison and Sugino who demonstrated that proofreading and MMR acted sequentially to correct DNA polymerase errors (Morrison *et al.*, 1993; Morrison and Sugino, 1994; Tran *et al.*, 1999b). Inactivation of the integral proofreading activity of Pol δ (e.g., *pol3-01*) increased mutation rates by nearly two orders of magnitude, as did inactivation of MMR (Simon *et al.*, 1991; Morrison *et al.*, 1993; Prolla *et al.*, 1994). Diploids defective in both Pol δ proofreading

and MMR (*pol3-01/pol3-01 pms1/pms1*) exhibited a 10^4 -fold increase (Morrison and Sugino, 1993; Tran *et al.*, 1999b). Double-mutant haploid spores derived from *POL3/pol3-01 PMS1/pms1* diploids germinated, but formed microcolonies of around 100 cells arrested at all stages of the cell cycle (Morrison and Sugino, 1993; Tran *et al.*, 1999b). A pure replication defect would have arrested cells immediately in S-phase or perhaps at the G2/M transition, but more than 40% of *pol3-01 pms1* cells exhibited a G1-arrest phenotype, while cells arrested in S-phase comprised only 5% (Morrison and Sugino, 1993). The relative abundances of each morphology in the *pol3-01 pms1* microcolonies correlate with the distribution of cell cycle arrest phenotypes observed in a survey of null phenotypes of all essential yeast genes (Yu *et al.*, 2006), suggesting that random mutations in essential genes drive extinction of the double-mutant mutator yeast.

To define the maximal mutation rate in haploids, we explored the synthetic relationship between Pol δ proofreading and MMR using a plasmid shuffling scheme (Boeke *et al.*, 1984; Simon *et al.*, 1991) that created a conditional mutator phenotype. The yeast strain for these studies carried a deletion of the chromosomal gene for MSH6 (*msh6 Δ*), an essential MMR component for repair of most base–base mismatches (Marsischky *et al.*, 1996). We shuffled in a collection of mutant *pol3* plasmids carrying hypomorphic proofreading-defective alleles of *POL3*, as well as null alleles, such as *pol3-01*. Some *pol3 msh6 Δ* combinations were viable and displayed soaring mutation rates (1000-fold increase). However, the strongest *pol3* mutators were lethal in combination with *msh6 Δ* , consistent with previous observations of synthetic lethality between *pol3-01* and *msh6 Δ* (Sokolsky and Alani, 2000). By measuring mutation rates of the viable combinations and calculating the theoretical mutation rates of the lethal combinations, we determined that the threshold for error-induced extinction is around 1×10^{-3} *can1* mutants/cell division. Assuming *CAN1* (1500 base pairs) represents an average gene, this mutation rate predicts a mutational target size of 1000 essential genes, which is very close to the number of essential genes in yeast determined by systematic gene inactivation (Winzeler *et al.*, 1999). Thus, accumulation of mutations throughout the genome, in many different contexts, likely drives replication error-induced extinction in yeast.

While defining the upper limits of mutagenesis, we isolated numerous mutants that survived the replication error-induced extinction induced by combining *msh6 Δ* with *pol3-01* (*D321A E323A*), *pol3-F406A*, or *pol3-D407A* (Figure 3A). Mutants with increased growth capacity also arose in an *msh6 Δ* strain expressing a high, but sub-lethal mutator phenotype induced by *pol3-D463A*. A large-scale screen for *pol3-01 msh6 Δ* suppressors revealed that one third of these *error-induced extinction* (*eex*) mutants carried mutations that encode single amino acid changes in Pol δ . Importantly, the rest mapped to the nuclear genome, implicating other genes in antimutagenesis.

Some of these chromosomal mutations may alter checkpoint signaling genes (e.g., *DUN1*) responsible for a heightened *pol3-01* mutator state (Datta *et al.*, 2000). Others may influence the fidelity of the polymerase directly. The amino acid substitutions in Pol δ conferred a range of antimutator activities, suppressing *pol3-01* between 3- and 50-fold in forward mutation rate assays at *CAN1* and *URA3* (Figure 2). *DUN1* deficiency reduces the *pol3-01* mutator phenotype 5- to 10-fold (Datta *et al.*, 2000). Thus, the strongest antimutators from our study likely influence the fidelity of the polymerase directly, in addition to any effects on checkpoint signaling.

Many antimutator substitutions affected polymerase features previously implicated in antimutagenesis in the T4 studies (Figure 3A). Five mutants encoded changes to motif A of the palm domain (N610D, P614S, S615N, I616C, and H620Y). Additional suppressors affected regions of the palm domain with less conservation (A786V, F793I, E800K, Y808A, and W821C). Three substitutions affected motif Pol V at the boundary between the palm and the thumb domains, including two (V838A and R839H) that corresponded precisely with the T4 Pol Q730S and Q731S antimutators. As observed in the T4 studies, numerous antimutator substitutions affected other parts of the thumb domain (H879Y, K891T, A894V/G, G921D, R923H and S968R). And surprisingly, even though the Pol δ exonuclease was inactivated in the *pol3-01 msh6 Δ* strain, numerous suppressors altered the exonuclease domain (C365Y, G400S, R475I/G, F486S, and L531P), in a manner reminiscent of the T4 Pol antimutators W202S, W213Y, and R335C. Intriguingly, Pol δ L531P aligns with T4 Pol R335C at the boundary between the exonuclease domain and a sequence just before the palm domain that folds with the amino-terminal domain. Our screen revealed four additional substitutions in this linker sequence (V546M, G555S, I558L, and Q563R). These were among the strongest suppressors (Figure 2), suggesting that this region of the polymerase has an important role in replication fidelity.

MMR-proficient strains expressing the mutant alleles as the sole source of Pol δ grew normally, indicating that the mutant polymerases retain sufficient activity to fulfill the essential role of Pol δ in lagging-strand DNA synthesis. Because error-induced extinction is due to random mutations in multiple genes, the antimutators likely reduce Pol δ errors in many different sequence contexts. Moreover, like the Pol III and Pol I antimutators, the gains in replication fidelity must occur without the aid of the proofreading activity of the polymerase, which was inactivated in these screens.

Structural implications of antimutators

The antimutagenic amino acid substitutions described above are distributed throughout the structures of their respective DNA polymerases. But far from random, the underlying changes affect functionally important features that evoke distinct, yet potentially overlapping

antimutator mechanisms. Antimutators affect polymerase active sites, exonuclease domains, and structural elements that bind double-stranded DNA.

Antimutators in the active site

The active sites of all DNA polymerases are found at the junction between the fingers and palm domains. In A- and B-family polymerases, the catalytic carboxylates reside in β -strands and loops from motifs A and C that contribute to the palm β -sheet surface (Figures 3A, 5A). The ribose of the incoming dNTP stacks at the edge of the β -sheet surface on motif A residues proximal to the catalytic carboxylates (T4 Pol: L412, Y413; Pol δ : L612, Y613; *E. coli* Pol I: I709, E710; Figures 3D, 3E and 5C). These dNTP stacking residues are at one end of the motif A α -helix, which angles down from the β -sheet surface, wedged between two conserved α -helices. One interacting helix is from motif B of the fingers domain (Figures 3A, 5A) that rises above the plane of the β -sheet, defining the binding pocket for the dNTP base, together with the template nucleotide and the last primer-template base-pair. In a large domain movement, the fingers domain close over the incoming dNTP and the binding pocket from this motif B helix interrogates the incoming base. Basic residues from the fingers domain (T4 Pol R479, K557; Pol δ R674, K701; *E. coli* Pol I R754, K758) sandwich the triphosphate against metal ions coordinated by the catalytic carboxylates. This stabilizes fingers domain closure and helps to create the catalytic center for phosphoryl transfer. The second interacting helix immediately follows motif B in B-family polymerases (Figure 3A) and corresponds to motif 6 in the A-family in Pol I (Figure 5). Note that these two α -helices are structurally homologous, although classified as part of the fingers domain in the A-family polymerases and the palm domain in the B-family. Each helix runs at right angles to the motif A helix, supporting the dNTP sugar binding surface. Thus, three conserved, interacting α -helices create a dynamic dNTP binding site adjacent to the essential catalytic residues.

Alterations to the structure of the catalytic surface promote antimutagenesis in both A- and B-family DNA polymerases. Antimutators in Pol δ (N610D, A786V, and E800K) affect amino acids that influence formation of the β -sheet involved in metal ion coordination (Figure 3E). E800 is adjacent to E802, which interacts with Metal C (Swan *et al.*, 2009). A double-mutant enzyme bearing E800A and E802A substitutions exhibited reduced catalytic efficiency, which led to the conclusion that E802 coordination of Metal C contributes to pyrophosphate formation (Swan *et al.*, 2009). However, E800 plays a key structural role in knitting the β -sheet together, and the double-mutant enzyme may suffer defects to β -sheet stability as well as Metal C coordination. E800, supported by A786, interacts with N610 from the adjacent β -strand, which lies between the motif A helix and the Metal B-binding aspartate, D608 (Figure 3A, 3E, 3I) (Swan *et al.*, 2009). Both E800K and N610D antimutator substitutions

replace the H-bonding interaction that normally occurs between these residues with charge repulsion, disrupting an interaction also observed in the analogous positions in RB69 Pol (D684 with T413 via H₂O) and *Thermococcus gorgonarius* Pol (E578 with R406) (Wang *et al.*, 1997; Hopfner *et al.*, 1999; Franklin *et al.*, 2001). Alteration of the catalytic surface of the palm domain also induces an antimutagenic effect in Pol I. H881 (mutated to A) is adjacent to the Metal A-binding aspartate, D882 (Figure 5A, 5C) and stacks with the ribose of the primer terminus, helping to orient the 3' OH for chemistry. Thus, disruptions to the architecture of the β -sheet active site surface can be antimutagenic.

A related avenue of antimutagenesis involves modification of the dNTP binding site. Most striking perhaps are the numerous antimutator substitutions found in motif A in T4 Pol (S411T, L412I, I417V, P424L; Figure 3A, 3D, 3H), Pol δ (P614S, S615N, I616C, H620Y; Figure 3A, 3E, 3I), and Pol I (E710D; Figure 5). The antimutators T4 Pol L412I and Pol I E710D alter the surface of the binding pocket that contacts the sugar of the incoming nucleotide. Pol δ Y708A of motif B affects the same surface, contacting the minor-groove side of the base. The analogous mutant in RB69 Pol (Y567A) is a strong mutator, but this phenotype is suppressed by a second substitution in the dNTP binding site, S565G (Trzemecka *et al.*, 2010; Xia *et al.*, 2011). Structural and biochemical analyses show that G565 in the S565G,Y567A double mutant limits the flexibility of the template base position (Xia *et al.*, 2011) and that, under certain conditions *in vitro*, the double-mutant polymerase dissociates more rapidly from duplexes with mispaired primer termini (Trzemecka *et al.*, 2010). Thus, changes to the binding pocket may influence not only nucleotide selectivity, but also stability of the replication complex. Other antimutators influencing the dNTP binding pocket include Pol δ P614S, which positions N705 of motif B next to the base of the incoming dNTP, and Taq Pol I F667L, which aligns structurally with Pol δ N705 (compare Figures 3E and 5C). Some antimutators alter residues that support the minor groove binding surface or mediate inter-helical contacts that help create the functional architecture. For instance, Pol δ I616 of motif A, in addition to supporting amino acids that contact the dNTP sugar (L612, Y613), abuts the antimutator position G731 (mutated to S) from the helix immediately following motif B (Figure 3A, 3E). Similarly, Pol I substitutions in motif 6 (M848I, Q849A, and T851I; Figure 5A, 5C) affect residues within 4 Å of E710 at the interface between motifs A and 6. Q849 also plays a role in shaping the dNTP binding pocket by interacting with the last primer-template base-pair (Kiefer *et al.*, 1998). Two other Pol I antimutators (K721T and A726T; Figure 5) affect helix M, which is an A-family-specific structure in the fingers domain that supports the motif A helix from the opposite side of motif 6 (Loh *et al.*, 2007). Thus, structural perturbations throughout the dNTP binding site promote antimutagenesis in both A- and B-family polymerases.

A cluster of T4 Pol and Pol δ antimutator substitutions affect amino acids in a linker sequence between the exonuclease and palm domains. This sequence, which folds with the amino domain, binds the template and buttresses the fingers domain (Figure 3A, 3D, 3E). Pol δ Q563R alters a residue that contacts the fingers in the closed conformation (Figure 3E), suggesting that antimutator substitutions in this region may destabilize closure of the fingers domain. The importance of fingers closure for fidelity is also indicated by the A677T antimutator, which lies at the interface between the two α -helices of the fingers domain, one turn of the helix from the dNTP-interacting residue, R674 (Figure 3E). Other substitutions in the exonuclease-palm linker (Pol δ V546M and I558L) interact in the hydrophobic core of the fold (Figure 3E). Intriguingly, T4 Pol S345F occupies almost the same position in the structure as Pol δ V546M (compare Figures 3D and 3E). G555S in Pol δ adds a side chain to this core, immediately adjacent to two amino acids that interact with the template nucleotide: Q556 contacts the template phosphate, while Q557 helps define the shape of the binding pocket for the template sugar (Figure 3E). The location of these B-family antimutators is especially intriguing, because this region aligns structurally with the pre-insertion site for the template nucleotide observed in crystal structures of the A-family *B. stearothermophilus* Pol I (Johnson *et al.*, 2003). Closure of the Pol I fingers flips the template nucleotide from this location into the active site. The existence of an analogous pre-insertion site remains to be demonstrated for B-family polymerases.

Seven out of nine *E. coli* Pol III antimutator substitutions are found in the palm or fingers domains despite occurring in a polymerase with very different architecture (Figure 4). During Pol III synthesis the ribose of the incoming dNTP stacks on R362 (*E. coli* numbering) of the RGS motif in the palm and H760 of the fingers (Figure 4C) (Wing *et al.*, 2008). The triphosphate interacts with G363, S364, R390, R710, and the metals bound by the catalytic aspartates, D403 and D405 (Figure 4C). The strongest *dnaE* antimutator alters F388 (mutated to L) (Figure 2), which brings the helices containing the RGS motif and R390 into close proximity through a hydrophobic interaction (Figure 4C). Two other *dnaE* antimutators (E395K and G703D) alter amino acids that pin the palm and the fingers domains together, positioning R710 to interact with the dNTP (Figure 4C). Thus, in all three polymerase families, modifications to the network of interactions that position the incoming dNTP serve to increase replication fidelity.

Antimutagenic changes to the active sites of DNA polymerases could increase fidelity in several ways. Variants may increase negative selectivity in the binding pocket against non-Watson-Crick base-pairs. However, such variants may be rare if the benefits of improved selectivity against one non-Watson-Crick base-pair are blunted by less stringent selection against a different incorrect base-pair. Amino acid substitutions may also increase accuracy without modifying the dNTP binding

pocket. Antimutators affecting the structure of the catalytic surface may promote delays in phosphoryl transfer, providing more time for incorrect nucleotides to dissociate. Other changes may enhance the ability of the polymerase to adopt a conformation that promotes release of incorrect dNTPs. Similarly, destabilization of fingers closure may allow incorrect nucleotides to dissociate more rapidly than normal.

In addition to any increases in dNTP selectivity, antimutator substitutions in the active site likely promote partitioning of the primer strand to the exonuclease domain. All T4 Pol antimutators exhibit a hyper-editing phenotype that exhausts dNTP pools in the *optA1* strain (Reha-Krantz, 2010). This can be explained by frequent partitioning of correctly paired primer termini to the exonuclease domain and degradation. Primer partitioning in a polymerase poised for misincorporation would be especially efficient due to inherently slow catalysis, leading to dissociation of the incorrect dNTP. In enzymes with defective proofreading, the same mechanism could occur without costly degradation of the 3' end, allowing error-free synthesis to resume upon reannealing back into the polymerase active site. In the presence of a terminal mismatch, more complete dissociation of the duplex DNA from the polymerase may allow alternate editing enzymes to correct the error *in trans* (Perrino and Loeb, 1990; Morrison and Sugino, 1994; Fijalkowska and Schaaper, 1995; Pavlov *et al.*, 2004; Albertson and Preston, 2006; Pavlov *et al.*, 2006a).

The vast majority of antimutators affecting the polymerase active site were identified using genetic screens in replicating organisms. Detailed biochemical, structural and kinetic analyses of purified variant polymerases are needed to determine their mechanisms of increased fidelity.

B-family antimutators that change the exonuclease domain

Several T4 Pol and Pol δ antimutator substitutions involve amino acids with structural roles in the exonuclease domain. W202S and W213Y in T4 Pol and C365Y, G400S, and L411R in Pol δ change amino acids that contribute to the hydrophobic core (Figure 3F, 3G). Pol δ F486S and L531P alter a hydrophobic interaction between two α -helices of the exonuclease domain and F688 from the fingers domain (Figure 3G). The relevance of this interaction for antimutagenesis is reinforced by the T4 Pol antimutator R335C, which also lies at the interface between the fingers and exonuclease domains (Figure 3F). The positions of these antimutators, together with the Pol δ Q563R substitution described above, support the idea that noncovalent interactions between the fingers domain and other parts of the polymerase may stabilize the closed conformation and influence fidelity. It should be emphasized that the Pol δ antimutator substitutions occur in the context of a catalytically dead exonuclease domain. The substitutions are unlikely to restore editing activity; indeed, two of the Pol δ substitutions (G400S

and F486S) cause weak mutator phenotypes when the exonuclease catalytic residues are restored (Herr *et al.*, 2011), consistent with reduced proofreading efficiency. The structural defects to the core of the exonuclease domain may also alter closure of the fingers or positioning of the β -hairpin loop that interacts with the template strand (Hogg *et al.*, 2007; Swan *et al.*, 2009). Or during attempts to proofread an error, these structural changes may reduce the affinity of the exonuclease domain for the primer and promote polymerase dissociation.

Antimutator substitutions along the DNA binding track

Dissociation-based models best explain the antimutagenic properties of substitutions that change amino acids along the DNA binding track. Pol δ V838 (mutated to A) sits next to the primer backbone at the -3 base-pair¹. R839 (mutated to H) interacts with the minor groove at base pairs -4 and -5 (Figure 3I) (Swan *et al.*, 2009). These positions correspond precisely with the T4 Pol Q730S and Q731S antimutator substitutions (Figure 3A, 3H). Other B-family antimutator substitutions affect amino acids that support adjacent DNA-interacting residues. In Pol δ , D831 (mutated to G) forms a salt bridge with K833, positioning the carbon chain of the lysine against the deoxyribose of the -2 primer position (Figure 3I). R475 (mutated to I or G) from the exonuclease domain overlays R840, which interacts with the -3 phosphate² of the primer strand. W821C and Y808A affect β -strands on either side of the KKRYA loop, which interrogates the minor groove of base-pairs -2 (via K814) and -5 (via R815) (Swan *et al.*, 2009). Finally, E594 (mutated to G) helps to position K813 of the KKRYA motif to interact with the template phosphate backbone at the -4 position. The Pol I L676Q antimutator may also diminish DNA binding. L676 packs against residues from motif 2 (N675 and N678) that interact with the minor groove of the -3 base pair (Figure 5D). Motif 2 forms a β -hairpin turn that contours the template backbone (positions -2 through -5). The extensive interactions of DNA polymerases with the minor groove of the first several base pairs of DNA duplex has been hypothesized to facilitate proofreading of misinsertions even after they have been extended (Franklin *et al.*, 2001; Johnson and Beese, 2004; Swan *et al.*, 2009). Thus, by reducing overall binding affinity, these antimutator polymerases may be especially prone to releasing mispaired primer-templates.

In addition to mutations affecting minor groove binding close to the active site, antimutator substitutions affect the flexible tip of the thumb domain, which

clamps down on the DNA in the elongation mode and moves with the primer strand during partitioning to the exonuclease active site (Franklin *et al.*, 2001). Two Pol δ antimutator substitutions in this subdomain affect residues that interact directly with the DNA (Figure 3I). R923 (mutated to H) interacts with the -4 phosphate of the primer strand, and K891 (mutated to T) extends out across the minor groove towards the template strand. Although not in position to interact directly with the DNA, H879Y alters an amino acid near the hinge of the thumb tip. The T4 Pol A737V, A777V, and W844S antimutators (Figure 3H) change positions that appear to be involved in the folding of the thumb tip or stabilization of the 'clamped down' conformation, which is important for elongation (Franklin *et al.*, 2001).

Antimutator substitutions in Pol I and Pol III also alter residues in the thumb. Pol I K601 (mutated to I) resides in a loop of the thumb tip that folds back on itself and interacts with DNA (Figure 5D). K601 may stabilize the fold of this loop by forming a hydrogen bond to a carbonyl of the peptide backbone on the opposing strand of the loop. Pol I Y630F resembles the classic A737V T4 Pol antimutator. Located in the main body of the thumb, Y630 interacts with residues from motif 1 in the thumb tip, which may stabilize the closed conformation. The Pol III thumb domain (Figure 4B) is markedly different from the thumb domains of A- and B-family polymerases, yet performs a similar function in interacting with the primer-template duplex along the minor groove. The two antimutator substitutions in the Pol III thumb domain (P464L and A498T) affect amino acids that may stabilize the position of an α -helix containing three basic residues (K439, R443, and R447) that contact the duplex DNA (Figure 4D).

Summary and perspective

Although isolated from different organisms and different DNA polymerases, the antimutator substitutions described here exhibit commonalities that point to general mechanisms of DNA polymerase fidelity and antimutagenesis (Figure 6). Antimutator substitutions may improve nucleotide selectivity by making the dNTP binding pocket of the polymerase more restrictive, by slowing the rate of catalysis or destabilizing the catalytic complex so that incorrect dNTPs dissociate before misincorporation, or by enhancing polymerase dissociation from mispaired primer termini so that errors can be edited more efficiently either by an intrinsic exonuclease or by a cellular proofreading activity *in trans*.

The myriad of antimutagenic pathways revealed by these genetic studies is consistent with incremental optimization of DNA polymerase activity and fidelity during evolution. Coordination between polymerases and proofreading exonucleases represents perhaps the ultimate antimutagenic innovation, but not the only one. Archaea have evolved B-family DNA polymerases with a novel strategy to avoid DNA damage-induced mutations. These polymerases scan the incoming template DNA for uracil

¹Here we refer to nucleotide positions relative to the direction of DNA synthesis, where zero (0) corresponds to the template nucleotide•dNTP (T_0 •dNTP) and -1, -2, -3, etc. correspond to base pairs in the duplex DNA that are 1, 2, 3, etc. nucleotides 'upstream' of T_0 •dNTP. Phosphates in the 5' position are designated by the same numbering (e.g. the phosphate between the -3 and -4 bases on the primer strand is designated the -3 phosphate).

²See note 1.

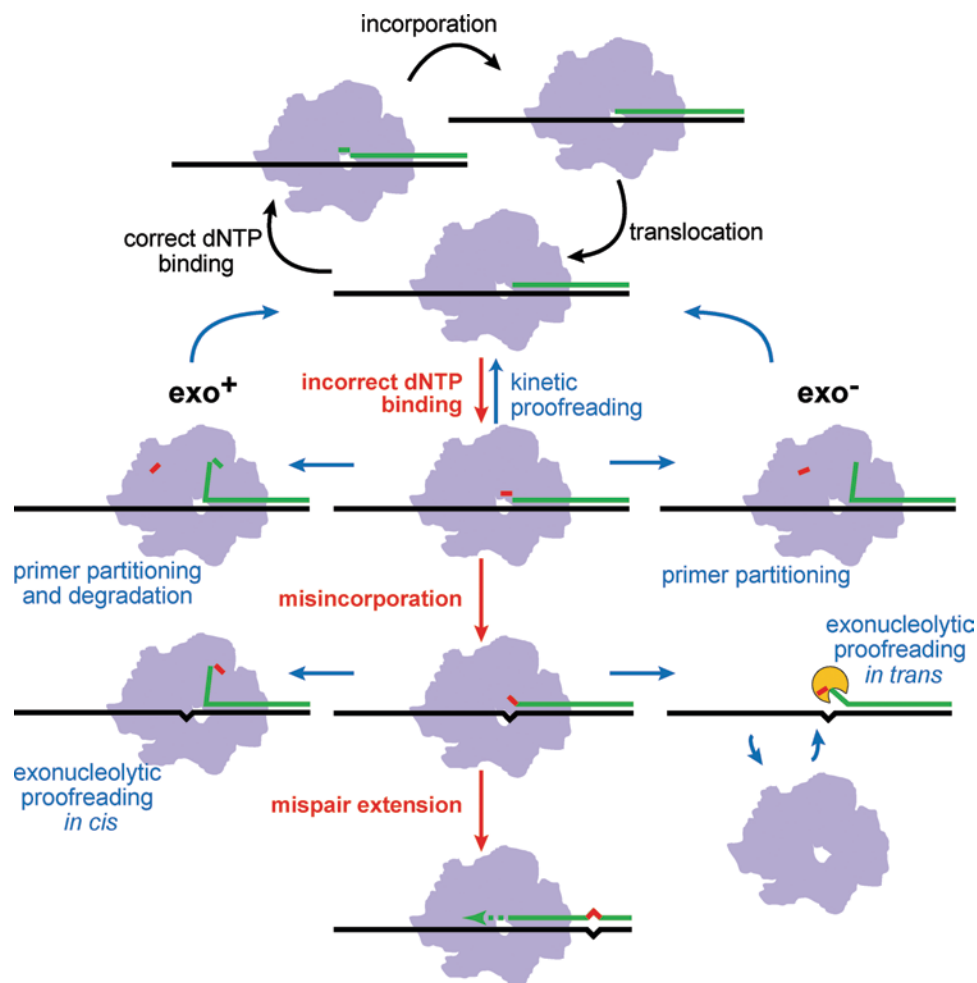


Figure 6. Mechanisms of DNA polymerase fidelity and antimutagenesis. A reiterative cycle of correct dNTP binding, nucleotide incorporation, and translocation comprises accurate DNA replication (top, black arrows). Antimutator mechanisms (blue arrows) enhance existing processes that oppose incorrect dNTP binding and misincorporation (red arrows) in DNA polymerases with functional (Exo^+) or defective intrinsic proofreading (Exo^-). Antimutator substitutions may increase the effectiveness of kinetic proofreading, by reducing incorrect dNTP binding stability or by enhancing partitioning of correctly paired primer strands to the exonuclease active site in the presence of bound, incorrect dNTPs. In proofreading-proficient polymerases, partitioning to the exonuclease domain may lead to degradation of correct primer ends, as exemplified by T4 Pol *optA1* sensitivity, as well as improved editing of misincorporations. In the absence of intrinsic proofreading, antimutators may enhance polymerase dissociation from the primer-template DNA, leading to editing by other cellular 3'→5' exonucleases (proofreading *in trans*).

and hypoxanthine residues, which result from deamination of cytidine and adenine and can lead to transition mutations (Greagg *et al.*, 1999). Uracil and hypoxanthine flip into a tight binding pocket in the amino domain and stall replication, likely leading to dissociation of the polymerase and initiation of DNA repair (Fogg *et al.*, 2002; Connolly *et al.*, 2003; Shuttleworth *et al.*, 2004; Gill *et al.*, 2007; Firbank *et al.*, 2008; Killelea *et al.*, 2010). Although this scanning functionality does not appear to extend to B-family polymerases from other Kingdoms (Wardle *et al.*, 2008), it illustrates that antimutagenic changes in DNA polymerases may optimize their response to environmental damage in addition to influencing inherent error rates.

In this review, we focused on antimutator variants of DNA polymerases. However, mutator phenotypes are also shaped by cellular responses to DNA replication errors.

For example, in yeast the mutator phenotype conferred by loss of Pol δ proofreading (*pol3-01* allele) depends in part on *DUN1*, a protein kinase gene that affects downstream events after S-phase checkpoint activation (Datta *et al.*, 2000). This *DUN1*-dependency suggests that pausing at mispaired primer termini by proofreading-defective Pol δ activates the S-phase checkpoint through Mec1, which stabilizes the stalled replication fork (Friedel *et al.*, 2009) and activates Dun1 through Rad53 (Chen *et al.*, 2010). Once activated, Dun1 phosphorylates a number of downstream targets (Chen *et al.*, 2010), including negative regulators of ribonucleotide reductase (Zhou and Elledge, 1992; Huang *et al.*, 1998b; Zhao and Rothstein, 2002; Lee *et al.*, 2008b; Wu and Huang, 2008). This signaling cascade ultimately increases dNTP pool concentrations in the cell, which helps DNA polymerases extend mispaired primer termini and copy past blocking lesions

(Reichard, 1988; Goodman *et al.*, 1993; Chabes *et al.*, 2003; Mathews, 2006; Kumar *et al.*, 2011). The *pol3-01* mutator phenotype does not depend on error-prone synthesis by TLS polymerases Pol ζ or Pol η (Datta *et al.*, 2000). Thus, loss of *DUN1* may suppress the *pol3-01* mutator phenotype by maintaining low dNTP levels, providing time for Pol δ to dissociate from mispaired primer termini and alternate repair pathways to correct 3' errors. Defects in other S-phase checkpoint molecules may have a similar antimutator effect. Therefore, in addition to direct effects on polymerase fidelity, alleles in polymerase or checkpoint genes may suppress mutator phenotypes by limiting Dun1 activation.

The plasticity of DNA replication fidelity may be important during times of intense selection pressure, which select for mutators that more quickly adapt to changing environments (Sturtevant, 1937; Drake *et al.*, 1998; Giraud *et al.*, 2001b; de Visser, 2002; Elena and Lenski, 2003; Denamur and Matic, 2006). During tumorigenesis, for instance, cells acquire multiple genetic changes that release restrictions on proliferation (Hanahan and Weinberg, 2011). Mutator phenotypes were predicted more than 30 years ago to accelerate tumorigenesis (Loeb *et al.*, 1974), and recent experiments suggest that many human tumors exhibit a mutator phenotype (Bielas *et al.*, 2006). Consistent with this hypothesis, defects in MMR, Pol δ proofreading, or Pol ϵ proofreading accelerate discrete cancer phenotypes in mice (de Wind *et al.*, 1995; Reitmair *et al.*, 1996; Goldsby *et al.*, 2001; Goldsby *et al.*, 2002; Albertson *et al.*, 2009). Defects in MMR and Pol δ fidelity are also associated with human tumors (Fishel *et al.*, 1993; Parsons *et al.*, 1993; Daele *et al.*, 2010). Once the requisite mutations for tumorigenesis are obtained, continued expression of a strong mutator phenotype may be deleterious for cancer cells as they are in microbial populations. The discovery of robust pathways of antimutagenesis offers one mechanism by which mutator phenotypes can be suppressed to tolerable levels that allow cancer cells to maintain replicative fitness.

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