TABLE 1

Bacterial strains used in this study

Name	Relevant genotype	Reference or source
FC29	$\Delta(lac-proB)$ XIII ara thi [F' $\Delta(lacI-lacZ)$]	CAIRNS and FOSTER (1991)
FC40	$\Delta(lac\text{-}proB)$ XIII ara thi Rif ^R [F' lac133 Ω lacZ $proAB^+$]	Cairns and Foster (1991)
FC231	FC40 lexA3(Ind ⁻)	CAIRNS and FOSTER (1991)
SMR868	$FC40 lexA3(Ind^-)$	McKenzie et al. (2000)
SMR4562	Identical to FC40, independent construction	McKenzie et al. (2000)
SMR5400	SMR4562 $sulA211 lexA51(Def) \Delta psiB::cat$	McKenzie et al. (2000)
SMR9436	SMR4562 ∆ruvC::FRTKanFRT	Magner <i>et al.</i> (2007)
SMR5889	SMR4562 $\Delta dinB50$::FRT [F' $\Delta dinB50$::FRT]	McKenzie et al. (2001)
SMR10292	SMR4562 [F' lafU2::FRTcalFRT]	This study
SMR10299	FC36 $\Delta(lafU-dinB)$ 2097(::FRTKanFRT)	This study
SMR10303	SMR4562 $\Delta(lafU-dinB)$ 2097(::FRTKanFRT) [F' $\Delta(lafU-dinB)$ 2096(::FRT)]	This study
SMR10304	SMR4562 Δ (lafU-dinB) 2097(::FRTKanFRT) [F' lafU2::FRTcatFRT dinBo-21(o ^c)]	This study
SMR10306	SMR4562 Δ (lafU-dinB) 2097(::FRTKanFRT) [F' lafU2::FRTcatFRT dinBo-92(os)]	This study
SMR10308	SMR4562 [F' $lafU2$::FRT cat FRT $dinBo-21(o^c)$]	SMR4562 \times P1(SMR10304)
SMR10309	SMR4562 [F' $lafU2$::FRT cat FRT $dinBo$ -22(o^c)]	$SMR4562 \times P1(SMR10306)$
SMR10310	SMR868 [F' lafU2::FRTcatFRT dinBo-21(o ^c)]	$SMR868 \times P1(SMR10304)$
SMR10311	SMR868 [F' lafU2::FRTcatFRT dinBo-22(o°)]	$SMR868 \times P1(SMR10306)$
SMR10314	SMR868 [F' lafU2::FRTcatFRT]	$SMR868 \times P1(SMR10292)$
SMR10760	FC231 [F' lafU2::FRTcatFRT]	$FC231 \times P1(SMR10292)$
SMR10761	FC231 [F' lafU2::FRTcatFRT dinBo-21(o ^c)]	$FC231 \times P1 (SMR10292)$ $FC231 \times P1 (SMR10304)$
SMR10762	FC231 [F' lafU2::FRTcatFRT dinBo-22(o ^c)]	$FC231 \times P1(SMR10304)$ FC231 × P1(SMR10306)
SMR10766	SMR4562 ΔruvC::FRTKanFRT [F' lafU2::FRTcatFRT]	$SMR10292 \times P1(SMR9436)$
SMR10767	FC231 ΔruvC::FRTKanFRT [F' lafU2::FRTcalFRT dinBo-21(o ^c)]	SMR10761× P1(SMR9436)
SMR10768	FC231 ΔruvC::FRTKanFRT [F' lafU2::FRTcatFRT]	$SMR10701 \times F1(SMR9430)$ $SMR10760 \times R1(SMR9430)$
SMR10838	SMR4562 [pPdinB]	$SMR10760 \times P1(SMR9436)$
SMR10839	SMR4562 [pPdinBOC1]	This study
MR10840	SMR4562 [pPdinBOC2]	This study This study
MR10841	SMR5400 [pPdinB]	
MR10842	SMR5400 [pPdinBOC1]	This study
MR10843	SMR5400 [pPdinBOC2]	This study
MR11023	SMR4562 [F' lafU2::FRTcatFRT ΔyafNOP::FRTKanFRT]	This study
MR11024	SMR4562 [F' lafU2::FRTcatFRT ΔyafNOP::FRTKanFRT dinBo-21(o ^c)]	This study
MR11026	FC231 [F' lafU2::FRTcatFRT ΔyafNOP::FRTKanFRT]	This study
MR11027	FC231 [F' lafU2::FRTcatFRT ΔyafNOP::FRTKanFRT dinBo-21(o ^c)]	$FC231 \times P1(SMR11023)$ $FC231 \times P1(SMR11024)$

ensure that the desired mutation was introduced and that no other mutation in dinB was generated inadvertently by PCR. One recombinant containing the $dinBo-21(o^c)$ mutation (SMR10304) and one containing the $dinBo-22(o^c)$ mutation (SMR10306) were chosen. Those strains were used as P1 donors of $\Delta lafU2::FRTcatFRT$ $dinBo-21(o^c)$ and $\Delta lafU2::FRTcatFRT$ $dinBo-21(o^c)$ and $\Delta lafU2::FRTcatFRT$ $dinBo-21(o^c)$ and $\Delta lafU2::FRTcatFRT$ $dinBo-21(o^c)$ alleles to all the genetic backgrounds of interest, including SMR4562 and strains FC231 and SMR868 carrying $lexA3(Ind^-)$.

Deletion of the yafNOP genes in the dinB operon was performed using short-homology recombination (Datsenko and Wanner 2000) as follows. Strains SMR10292 [SMR4562 (F' $\Delta lafU2$::FRTcatFRT)] and SMR10308 [SMR4562 (F' $\Delta lafU2$::FRTcatFRT $dinBo-21(o^c)$)] were used as recipients for deletion by transformation with a DNA fragment amplified from pKD13 with primers yafNwL and yafPwR. Homologous incorporation of this DNA fragment, which contains the FRTKanFRT marker, results in a deletion of the yafNOP genes. Kan^R recombinants were selected, and location of the marker

TABLE 2
Plasmids used in this study

Name	Description and source
pFZY pPdinB pPdinBOC1 pPdinBOC2	Low-copy plasmid with multicloning site abutting a promoterless <i>lacZ</i> (Koop <i>et al.</i> 1987) Bases -432 to -2 of <i>dinB</i> from strain SMR4562 cloned into pFZY, producing a P _{dinB} dacZ fusion Bases -432 to -2 of <i>dinB</i> from strain SMR10308 cloned into pFZY, producing a P _{dinBo-21(or)} lacZ fusion Bases -432 to -2 of <i>dinB</i> from strain SMR10309 cloned into pFZY, producing a P _{dinBo-22(or)} lacZ fusion

in the F' episome was confirmed both by ability to transfer the resistance during mating and by cotransduction of Kan^R and Cam^R (present in the linked *lafU2*::FRT*cat*FRT in both strains). The strains resulting from deletion of *yafNOP* from the episomes of SMR10292 and SMR10308 were named SMR11023 and SMR11024, respectively. Both strains were used respectively as P1 donors to transfer the *lafU2*::FRT*cat*FRT $\Delta yafNOP$::FRTKanFRT linkage and the *lafU2*::FRT*cat*FRT *dinBo-21*(o°) $\Delta yafNOP$:FRT:KanFRT linkage into the FC231 background, creating strains SMR11026 and SMR11027.

β-Galactosidase assays: β-Galactosidase assays were performed to determine the relative expression of lacZ under the control of the different versions of the dinB promoter cloned into the low-copy plasmid pFZY (Koop $et\ al.\ 1987$). Cells were grown in LBH medium until mid-log phase, and the levels of β-galactosidase were determined in samples of the cultures as described (MILLER 1992).

DinB Western blots: For DinB detection on Western blots, stationary-phase cultures grown from single colonies in 5ml of M9 B1 glycerol medium for 48 hr were harvested, and cells were suspended in sample loading/lysis buffer (62.5 mm Tris, pH 6.8, 25% glycerol, 2% SDS, 0.01% bromophenol blue, 0.5% β -mercaptoethanol), correcting for the OD_{600} of the terminal culture. For 1 ml of a culture at OD_{600} of 2 (measured at $OD_{600} \le 1$ with diluted samples), 100 µl of sample loading buffer was used. Twenty microliters of each sample was separated by electrophoresis on a SDS polyacrylamide gel (12.5%). Proteins were transferred to a Hybond-LFP PVDF membrane (Amersham Biosciences), and the membrane was probed with a polyclonal DinB rabbit antiserum (KIM et al. 2001). A goat anti-rabbit secondary antibody conjugated to the Cy5 fluorescent dye (Amersham Biosciences) was used for detection of DinB, using the Typhoon scanner (Amersham Biosciences).

Stress-induced mutagenesis assays: Stress-induced *lac* reversion assays were performed as described (HARRIS *et al.* 1996) with four independent cultures of each strain. The proportion of Lac⁺ point mutants and *lac*-amplified colonies was determined by plating cells from 20 colonies of each culture for each day in which Lac⁺ colonies were counted (days 2–5) on LBH rifampicin X-gal plates. This allows the distinction between Lac⁺ point mutants (solid-blue colonies) and *lac*-amplified cells, given the *lac*-unstable sectoring-colony phenotype diagnostic of *lac* amplification (HASTINGS *et al.* 2000).

Determination of the mutation sequences in the *lac* gene: Lac⁺ point mutants from experiment day 5 were identified as described above and purified on LBH plates containing rifampicin and X-Gal. A 300-nucleotide region spanning the *lac* +1 allele was amplified by PCR using primers lacIL2 (5'-AGCTATTCTGGTGGCCGGA-3' and lacD2 (5'-GCCTCTTCGCTATTACGCCAGCT-3'). DNA sequencing was performed by Seqwright (Houston) using primer lacU (5'-ATATCCCGCCGTTAACCACC-3').

RESULTS

Construction and characterization of the $dinB(o^c)$ alleles: To test the hypothesis that dinB might be the sole SOS gene required at induced levels for stress-induced point mutagenesis, we constructed dinB mutants in which the transcriptional repression by LexA, the repressor controlling the expression of the SOS regulon, is alleviated. This was achieved by site-directed muta-

wild-type dinB promoter

TGAAATCACTGTATACTTTACCAGTGTTGAGAGGTGAGCA ATGCGTA

dinBo-21(oc)

TGAAATCA $oldsymbol{A}oldsymbol{G}$ GTATACTTTACCAGTGTTGAGAGGTGAGCA $oldsymbol{A}$ TGCGTA

dinBo-22(oc)

TGAAATCACTGTATACTTTACC CT TGTTGAGAGGTGAGCA ATGCGTA

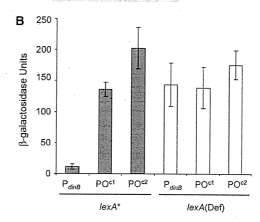


FIGURE 1.—Construction and characterization of two dinB(oc) alleles. (A) Location of the operator-constitutive mutations in the dinB promoter. The SOS operator (from Fernandez De HENESTROSA et al. 2000) is shaded, and the mutations introduced in each of the alleles are in boldface and italic type. The beginning of the dinB ORF is shown in boldface type. (B) Activity of the dinB promoter in transcriptional fusions with lacZ, measured in both wild-type (SMR4562) and its LexAdefective (null), lexA51(Def), derivative strain SMR5400, in which SOS is constitutively highly induced. The strains from left to right are SMR10838, SMR10839, SMR10840, SMR10841, SMR10842, and SMR10843. P_{dinB} indicates the wild-type dinBpromoter present in plasmid pPdinB, POcl indicates the dinBo-21(o^c) promoter contained in plasmid pPdinBOC1, and PO^{c2} indicates the dinBo-22(oc) promoter contained in plasmid pPdinBOC2. Mean ± 1 standard error of the mean (SEM) for three independent determinations.

genesis of the dinB promoter, altering the binding site of the LexA repressor. These are used (see below) to express dinB at SOS-induced levels in strains in which the rest of the SOS genes are repressed. The sequences of the operator-constitutive $dinB(o^c)$ mutations that were constructed are shown in Figure 1A.

To test whether these mutations behave as bona fide operator-constitutive alleles, we fused the dinB promoter regions from the two $dinB(o^c)$ alleles to lacZ and measured the levels of β -galactosidase expression from these $P_{dinB}lacZ$ fusions carried in a low-copy plasmid (Figure 1B). Introduction of these plasmids into wild-type cells resulted in ~ 10 -fold higher lacZ expression from both $P_{dinB(o^c)}lacZ$ fusions when compared with wild-type P_{dinB} . This is in agreement with previous estimates of transcriptional induction of dinB during the SOS response

(Courcelle et al. 2001; Kim et al. 2001). lexA51(Def) cells have no functional LexA repressor and show constitutive SOS expression (Mount 1977). We find that lacZ expression is increased in a lexA51(Def) strain when driven by the wild-type dinB promoter, but see no significant increase with the $dinB(o^c)$ promoters, showing that levels of dinB transcription similar to that achieved by true SOS derepression are achieved by the $dinB(o^c)$ mutations. The lexA51(Def) strain SMR5400 also carries a mutation in the sulA gene, which allows survival under constitutive SOS induction (Mount 1977), and a mutation in the F-encoded psiB gene, which has been shown to exert a negative effect on stressinduced mutagenesis (McKenzie et al. 2000) probably by affecting SOS induction (reviewed by Cox 2007).

In the Lac-assay strains such as FC40 and SMR4562, dinB is present both in the chromosome and in the F'128, at which locus it is more highly expressed (Kim et al. 2001). Introduction of both dinB(oc) alleles into the episomal dinB locus results in about five- to six-fold increased DinB-protein levels in stationary-phase cells compared with an otherwise isogenic SMR4562 derivative in both wild-type and lexA3(Ind⁻) backgrounds (Figure 2). This indicates that both $dinB(o^c)$ alleles are functional in vivo, conferring an increased basal dinB expression. Furthermore, both alleles confer levels of expression similar to those observed in lexA51(Def) cells (Figure 2), at least in the growth conditions used by us in the stress-induced mutagenesis experiments (cells grown for 48 hr in M9 B1 glycerol minimal medium). It was noted before (KIM et al. 2001) that expression of dinB in the F'128 plasmid is higher than that from the chromosomal dinB. Our finding that both dinB(o^c) alleles, when present only in the episome, increase DinB to levels similar to that observed in the lexA51(Def) strain (in which both the episomal and the chromosomal copy are constitutively highly expressed), also implies that the episomal expression is more pronounced than the chromosomal expression. To facilitate further strain construction and genetic analysis, we carried out the subsequent experiments in cells bearing a single $dinB(o^c)$ allele in the F'128 plasmid.

dinB(o°) mutations restore stress-induced point mutagenesis in SOS-off strains: Because DinB is a key player in stress-induced mutagenesis, we wanted to examine whether dinB is the only gene required at SOS-induced levels for stress-induced point mutagenesis in the Lac assay. The SOS response is induced when DNA damage is sensed in the form of single-strand DNA (reviewed by Friedberg et al. 2005). RecA binds the single-strand DNA and becomes active as a co-protease that facilitates cleavage of the LexA repressor, resulting in upregulation of the SOS genes, including dinB. To determine whether dinB upregulation constitutes the sole role of the SOS response in stress-induced point mutagenesis, we tested the effect of the dinB(o°) alleles on lac reversion in both wild-type and lexA3(Ind⁻)

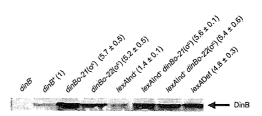


FIGURE 2.—DinB Western blots. Stationary-phase cells grown in M9 B1 glycerol medium were harvested and analyzed using a rabbit polyclonal DinB antiserum as described (MATERIALS AND METHODS). Values shown represent the average DinB protein levels relative to wild type determined in three independent experiments \pm SEM. Similar results were obtained with Western blots performed with a DinB monoclonal antibody. Strains are the following: dinB, SMR5889; dinB-22(o°), SMR10292; dinB-21(o°), SMR10308; dinB-22(o°), SMR10309; lexA3(Ind¬), SMR10760; lexA3(Ind¬) dinB-21(o°), SMR10761; lexA3(Ind¬) dinB-22(o°), SMR10761; lexA3(Ind¬) dinB-22(o°), SMR10761; lexA3(Ind¬) dinB-22(o°), SMR10761; lexA3(Ind¬) dinB-22(o°), SMR10762; and lexA(Def), SMR5400.

backgrounds. The lexA3(Ind-) mutation creates an uncleavable LexA/SOS repressor such that derepression of the SOS response genes during an SOS response is prevented (Mount et al. 1972). Previously, this allele was shown to cause reduced stress-induced point mutagenesis in the Lac assay (CAIRNS and FOSTER 1991; McKenzie et al. 2000, 2001), indicating that one or more SOS-controlled genes are needed at induced levels for efficient stress-induced mutagenesis. Representative results from single experiments with each of the two dinB(oc) alleles constructed are shown in Figure 3, A and B, and quantification of the stress-induced point mutagenesis rates from multiple experiments is shown in Figure 3C. Strikingly, either allele provides a complete suppression of the phenotype of the lexA3(Ind-) strain. These results show that the reduced stressinduced mutagenesis in a lexA3(Ind-) strain is caused specifically by the failure to upregulate dinB, and not any other gene in the LexA/SOS regulon. This finding places DinB as the central SOS-regulated protein in stress-induced mutagenesis and indicates that upregulation of other SOS genes such as recA, ruvA, and ruvB beyond their constitutive levels of expression is irrelevant.

SOS-induced levels of DinB are not sufficient to increase stress-induced point mutagenesis: We note that providing SOS-induced levels of DinB to all cells, with the $dinB(o^c)$ mutations, did not stimulate stress-induced mutagenesis above wild-type levels in the $lexA3(Ind^-)$ strain (Figure 3 and legend), even though normally SOS is expected to be induced spontaneously in only $\sim 1\%$ of cells (Pennington and Rosenberg 2007). Neither did DinB overproduction increase mutagenesis in the wild-type genetic background (Figure 3 and legend). These results indicate that DinB upregulation by the SOS response, although required, is not sufficient to differentiate the mutating subpopulation; $dinB(o^c)$ appears not to make all cells in the population

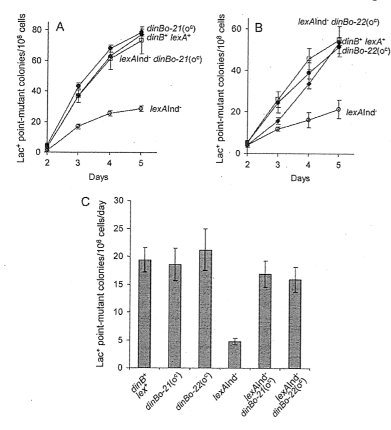


FIGURE 3.—Two dinB operator-constitutive alleles restore stress-induced Lac point mutagenesis to SOS-off lexA(Ind-) cells. (A) Effect of the dinBo-21(o^c) allele in stress-induced mutagenesis: a representative experiment. (B) Effect of the dinBo-22(oc) allele in stress-induced mutagenesis: a representative experiment. Note that, for both alleles, the stress-induced point mutagenesisdefective phenotype of lexA3(Ind-) cells is fully suppressed; however, overproduction of DinB with these alleles does not stimulate mutagenesis above wild-type levels. Data represent means ± SEM for four cultures. Strains are the following: dinB+ lex+, SMR10292 (solid diamonds in A and B); lexA(Ind-), SMR10760 (open circles in A and B); dinBo-21(oc), SMR10308 (solid circles in A); dinBo-22(o^c), SMR10309 (solid circles in B); lexA (Ind-) dinBo-21(oc), SMR10761 (open squares in A); and lexA(Ind-) dinBo-22(oc), SMR10672 (open squares in B). (C) Quantification of stress-induced point-mutation rates from six independent experiments, each with all genotypes done in parallel. Strains were as above except that three experiments were performed in the lexA3(Ind-) strains listed above, whereas an additional three experiments were performed, with similar results, in an independently constructed, identical set of lexA3(Ind-) strains: lexA(Ind-), SMR10314; lexA(Ind-) dinBo-21(oc), SMR10310; and lexA(Ind-) dinBo-22(oc), SMR10311. Rates represent the increase of Lac+ point mutant colonies per day observed between days 3 and 5 of each experiment. Means ± 1 SEM are shown. P-values were obtained for pairwise comparisons

by the nonparametric Mann–Whitney rank-sum test using the SYSTAT 11 statistics software by SYSTAT software and are as follows. The mutation rate of $dinB^+$ is not different from those of $dinBo-21(o^c)$ (P=0.699), $dinBo-22(o^c)$ (P=0.699), $lexA(Ind^-)$ $dinBo-21(o^c)$ (P=0.818), or $lexA(Ind^-)$ $dinBo-22(o^c)$ (P=0.18), but is significantly different from the rate of $lexA(Ind^-)$ (P=0.002), and the $lexA(Ind^-)$ rate differs from those of $lexA(Ind^-)$ $dinBo-21(o^c)$ (P=0.002) and $lexA(Ind^-)$ $dinBo-22(o^c)$ (P=0.002).

mutable. This might reflect either of two possible realities. First, in principle, it could be possible that during stress-induced mutagenesis conditions all cells are SOS-induced such that providing an operatorconstitutive dinB does not provide any more DinB protein than the population of cells already has, and so does not increase mutagenesis further. This is unlikely (discussed below). Second, and more likely, it could be that during stress-induced mutagenesis only a small fraction of cells is SOS induced, as is the case for growing cells in which ${\sim}10^{-2}$ are (Pennington and Rosenberg 2007), but that in this cell subpopulation some other condition must be met to allow mutagenesis. For example, it is likely that possession of a DNA double-strand break at which the mutagenic repair occurs (PONDER et al. 2005) is also required such that DinB upregulation alone is not sufficient.

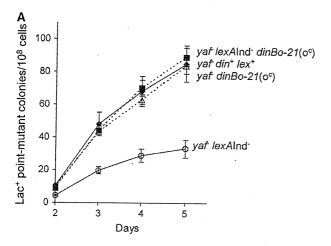
SOS induction of other genes in the dinB operon is irrelevant for stress-induced mutagenesis: dinB is part of a four-gene operon including dinB, yafN, yafO, and yafP (McKenzie et al. 2003). The functions of the three yaf genes are unknown. The whole operon, including the three genes downstream of dinB, is induced as part

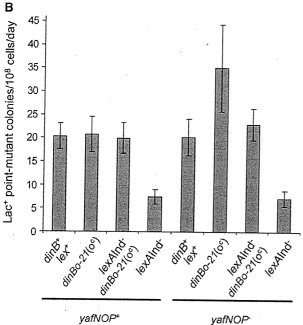
of the SOS response (Courcelle et al. 2001). Thus, in the experiments described above, all three yaf genes were also upregulated by the operator-constitutive mutations in the dinB promoter. We show that the restoration of mutability to SOS-off lexA(Ind-) cells conferred by the dinB(o^c) mutations was not conferred by increased yafNOP expression, only by increased dinB expression, because it also occurred in strains carrying a deletion of the yafNOP genes in cis with (downstream of) the dinBo-21(o°) mutation in F'128 (Figure 4). Although intact yaf genes are present in the chromosome of this strain, they will be repressed by the *lexA*(Ind⁻)-encoded uncleavable LexA/SOS repressor, such that only DinB is produced at SOS-induced levels in this strain. Therefore, dinB is indeed the only gene of the SOS regulon that is required at SOS-induced levels for stress-induced mutagenesis in the Lac assay. These experiments do not rule out a role for the yaf genes (expressed at uninduced levels) in mutagenesis, a topic that will be addressed in a future publication (L. Singletary, J. Gibson, E. Tanner, G. J. McKenzie, P. L. LEE and S. M. ROSENBERG, unpublished data).

The mechanism of stress-induced *lac* reversion in $dinB(o^c)$ cells is similar to that in wild-type cells: The

results obtained show that the $dinB(o^c)$ alleles are able to rescue completely the mutagenesis-defective phenotype of $lexA3(Ind^-)$ (SOS-off) cells in stress-induced mutagenesis (Figure 3). This could result from restoration of the same stress-induced mutagenesis pathway and mechanism that operates in wild-type cells. Alternatively, it was possible that constitutive expression of dinB might activate a different mutagenesis mechanism that coincidentally yielded similar mutant frequencies in the course of several days. We provide two lines of support for the first possibility that the normal pathway and mechanism of stress-induced point mutagenesis was restored to $lexA3(Ind^-)$ (SOS-off) cells by the $dinB(o^c)$ mutations.

First, we find that the Lac-reversion-mutation sequences in lexA3(Ind⁻) dinB(o^c) cells are indistinguishable from the characteristic point-mutation sequences seen normally in stress-induced point mutagenesis (in $lexA^+$ $dinB^+$ cells) (Figure 5A). The mutations are dominated





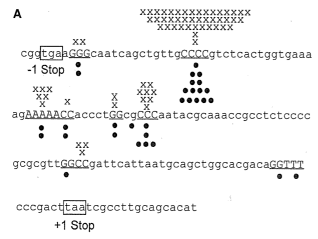
by -1 deletions at mononucleotide repeats that occur preferentially in the same preferred hotspot sequences as observed in $lexA^+$ $dinB^+$ cells. This characteristic mutation sequence spectrum is highly specific and different from, for example, spontaneous reversions of this lac allele during growth, which are more heterogeneous, including -1 deletions not at mononucleotide repeats and larger frameshift-reverting additions and deletions in about half the mutations (Foster and Trimarchi 1994; Rosenberg et al. 1994). These results imply that mutations occur via a similar or the same mechanism in both genetic backgrounds, supporting the idea that the rescue of the lexA3(Ind $^-$) phenotype by the dinB(o c) alleles restored the same mutagenesis mechanism that normally operates in $lexA^+$ $dinB^+$ cells.

Second, a hallmark of stress-induced mutagenesis in the Lac assay is its requirement for homologous-recombination, double-strand-break-repair functions, including recA, recB, and ruvAB, and ruvC (Harris $et\ al.\ 1994$, 1996; Foster $et\ al.\ 1996$; He $et\ al.\ 2006$), because the mutagenesis results from error-prone double-strand-break-repair events (Ponder $et\ al.\ 2005$). Similarly, we find that deletion of ruvC reduces stress-induced mutagenesis in $lexA3(Ind^-)$ (SOS-off) cells carrying a $dinB(o^c)$ allele (Figure 5B). Thus the mutagenesis restored to $lexA3(Ind^-)$ cells by the $dinB(o^c)$ mutation requires ruvC. This supports the conclusion that a similar or the same recombination-dependent mutagenesis pathway is operating in $lexA3(Ind^-)\ dinB(o^c)$ cells as is normal in cells wild-type for lexA and dinB.

DISCUSSION

How stress responses confer temporal regulation of mutagenesis: The coupling of mutagenesis programs to

FIGURE 4.—Stress-induced mutagenesis proficiency in SOSoff lexA3(Ind-) cells with a dinB(o) mutation does not require SOS induction of the yafNOP genes. (A) Representative experiment showing that deletion of the yafNOP genes in cis with dinBo-21(0°) does not affect the ability of this promoter mutation to rescue the phenotype of SOS-off lexA3(Ind⁻) cells. (B) Quantification of stress-induced point-mutation rates (calculated as in Figure 3) from four independent experiments. Means ± SEM are shown. P-values (calculated as in Figure 3) are as follows. For the yaf+ background, the dinB+ rate is not different from the rates observed with $dinBo-21(o^c)$ (P = 0.886) or $lexA(Ind^-)$ dinBo-21(o°) (P = 0.686) but differs from that of $lexA(Ind^-)$ (P = 0.029), and the rate of $lexA(Ind^-)$ differs from that of $lexA(Ind^-)$ dinBo-21(o°) (P = 0.029). Similarly, for the yaf background, the dinB+ rate is not significantly different from the rates of $dinBo-21(o^c)$ (P = 0.114) or $lexA(Ind^-)$ $dinBo-21(o^c)$ (P = 0.886) but differs from the $lexA(Ind^-)$ rate (P = 0.029), and the $lexA(Ind^-)$ rate differs from that of lex- $A(Ind^{-})$ dinBo-21(o^c) (P = 0.029). Strains are the following: SMR10292; lexA(Ind-), SMR10760; dinBo-21(0°), SMR10308; lexA(Ind⁻) dinBo-21(o^c), SMR10761; yaf⁻ dinB⁺ lexA⁺, SMR11023 (solid diamonds); yaf-dinBo-21(oc), SMR11024 (open triangles); yaf lexA(Ind), SMR11026 (open circles); yaf lexA (Ind⁻) dinBo-21(o^c), SMR11027 (solid squares).



mutations generated in the lexAlnd- dinBOo strains
 X mutations generated in the dinB+ strain

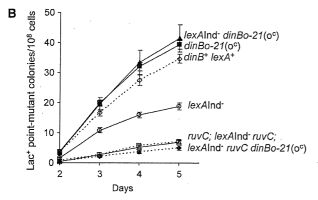


FIGURE 5.—Stress-induced mutagenesis in SOS-off lexA3 (Ind-) cells with dinB(oc) alleles occurs via a mechanism similar to normal stress-induced mutagenesis in the Lac assay. (A) Sequences of Lac⁺ mutations are the same in lexA3(Ind⁻) dinB(oc) strains and in lexA+ dinB+ cells. Both sets of reversion mutations are nearly all -1 deletions in mononucleotide repeats. The positions of the -1 deletions observed in the lexA3(Ind⁻) dinBo-21(o^c) and lexA3(Ind⁻) dinBo-22(o^c) strains (SMR10761 and SMR10762) are shown as circles, and the position of the -1 deletions observed in $lexA^+$ $dinB^+$ cells (data from Foster and Trimarchi 1994 and Rosenberg et al. 1994) are marked as X's. The region shown is part of the lacIZ fusion gene present in these strains. Compensatory frameshift mutations in a 130-nt region between the two out-of-frame stop codons (boxed) can restore gene function. (B) Stressinduced mutagenesis promoted by the dinBo-21(oc) allele in the lexA3(Ind-) background requires RuvC. A representative experiment performed with four independent cultures of each strain is shown. Means ± SEM. This result was repeated twice. Strains are the following: dinB+ lex+, SMR10292 (open diamonds); dinBo-21(oc), SMR10308 (solid squares); lexA(Ind-) dinBo-21(o^c), SMR10761 (solid triangles); lexA(Ind⁻), SMR10760 (open circles); ruvC, SMR10766 (open triangles); lexA(Ind-) ruvC, SMR10768 (open squares); and lexA(Ind-) ruvC dinBo-21(o^c), SMR10767 (closed diamonds).

cellular stress responses observed in bacterial and eukaryotic cells (reviewed in the Introduction and by GALHARDO et al. 2007) provides a temporal regulation of mutagenesis, limiting mutagenesis to times of stress. This may potentially accelerate genetic change, and thus the ability to evolve, specifically when cells and organisms are maladapted to their environments, i.e., are stressed. Here we demonstrate that, in the case of the E. coli Lac assay, the requirement for the SOS stress response can be deconvoluted to the need for induction of one specific gene, dinB. A number of other stress responses have been shown to upregulate mutagenesis, such as the RpoS response in E. coli, Salmonella, and Pseudomonas; the stringent response in E. coli and B. subtilis; the competence response of B. subtilis; and two human responses to hypoxic stress (see Introduction). All of these modulate the expression of tens to hundreds of different genes. It is not yet known whether any other of these stress responses can be narrowed down to relevant effects on one or a few

Roles of SOS in other stress-induced mutagenesis mechanisms: The SOS response is a major upregulator of mutagenesis during stress conditions but may not function identically in each case. For example, the SOS response is required for phage λ untargeted mutagenesis (ICHIKAWA-RYO and KONDO 1975), stress-induced point mutagenesis in E. coli in the Lac assay (McKenzie et al. 2000), ciprofloxacin (antibiotic)-resistance mutagenesis induced by exposure to ciprofloxacin (CIRZ et al. 2005), mutagenesis in aging colonies in a laboratory E. coli strain (TADDEI et al. 1995), and bile-resistance mutagenesis in Salmonella (PRIETO et al. 2006). Although DinB is required for λ untargeted mutagenesis (Kim et al. 1997) and for Salmonella bile-resistance mutagenesis (Prieto et al. 2006), it is not yet known whether the SOS requirement in either case is based on DinB upregulation. In the ciprofloxacin-induced mutagenesis, the SOS-controlled DNA polymerases DinB, Pol II, and Pol V are all required, as are the doublestrand-break-repair genes, including SOS-regulated recA, ruvA, and ruvB (CIRZ et al. 2005). Part of this mutation pathway's requirement for SOS is likely to be for production of DNA Pol V (CIRZ et al. 2005) because this polymerase is required for mutagenesis and is produced virtually not at all without an SOS response (reviewed by Nонмі 2006). It is not yet known whether upregulation of any, all, or none of the other two SOScontrolled DNA polymerases additionally account for the requirement for an SOS response for ciprofloxacininduced mutagenesis. Conversely, in the Lac assay, we measure frameshift reversion, which DinB promotes but Pols II and V do not, whereas the ciprofloxacin-induced mutations are base substitutions, which all three SOS polymerases promote (CIRZ and ROMESBERG 2007). CIRZ and ROMESBERG (2007) have pointed out that the mutagenesis pathway in the Lac system might be

identical to that in ciprofloxacin-resistance mutagenesis and might also require Pols II and V if base substitution mutations were assayed. In a different assay system, mutagenesis in aging colonies required an SOS response, did not require DinB, and required Pol I instead (TADDEI et al. 1997). Therefore, in the mutagenesis mechanism operating during that stress, the SOS requirement must be for some other function. Thus, although the SOS response is required for multiple examples of stress-inducible mutagenesis, its means of promoting mutagenesis in at least some of these different stress circumstances is different.

SOS induction of DinB is necessary but not sufficient for stress-induced mutability and for differentiation of a hypermutable cell subpopulation: Several lines of evidence indicate that although SOS induction of DinB is necessary, it is not sufficient for creating the transient mutable state in which most Lac reversions occur (Gonzales et al. 2008). Rather, the evidence also supports a model in which at least three events must occur: (1) a double-strand break and its repair, (2) SOS induction, and (3) induction of the RpoS stationary-phase and general stress response. The concerted induction of these stress responses in cells bearing double-strand DNA ends (DSEs) is proposed to differentiate the hypermutable subpopulation, as depicted in Figure 6A.

The first evidence for this model comes from experiments in which DNA double-strand breaks were delivered to the DNA by expression of the I-SceI double-strand endonuclease in vivo (PONDER et al. 2005). DinB-dependent, stress-induced mutagenesis was stimulated >1000-fold near the DSBs, but only weakly (3-fold) in a another DNA molecule (with no DSB) in the same cell. SOS is induced robustly by I-Scelinduced DSBs (Pennington and Rosenberg 2007). Therefore, these results indicate that the SOS-mediated DinB upregulation caused by I-SceI-mediated DNA cleavage was not sufficient for mutagenesis. A DSB was also required locally. The mutations appear to occur in acts of error-prone DSB repair (PONDER et al. 2005). Figure 6B shows a model for such mutagenic doublestrand-break repair occurring in stressed cells.

Second, RpoS is required for stress-induced Lac reversion (Layton and Foster 2003; Lombardo et al. 2004). Moreover, in the study using I-Scel-induced DSBs (Ponder et al. 2005), the DinB-dependent DSB-associated mutagenesis was provoked only in cells that either were in stationary phase or were expressing the RpoS stationary-phase and general stress-response transcriptional activator inappropriately during log phase. Again, because the SOS response is induced efficiently by I-Scel-mediated double-strand breakage (Pennington and Rosenberg 2007), this implies that repair of a DSB under the influence of the SOS response is also not sufficient for DinB-dependent mutagenesis; RpoS must also be induced. Additionally, although some kinds of

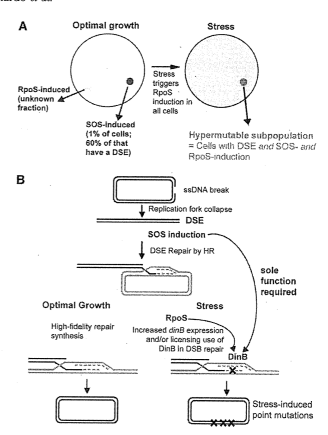


FIGURE 6.—Models for the role of SOS induction and DinB in differentiation of a hypermutable cell subpopulation and double-strand-break-repair-associated mutagenesis during stress. (A) Venn-diagram model of cell subpopulations that overlap to produce a transiently hypermutable cell subpopulation. In this model (modified from GALHARDO et al. 2007), differentiation of a HMS (green) is proposed to occur when three conditions are met: (1) induction of the SOS response in cells bearing (2) a double-strand break or a DSE (blue) and (3) induction of the RpoS regulon (yellow) by suboptimal growth conditions. During stress, all cells induce RpoS and, we suggest, the fraction of cells bearing both a DSE and SOS induction [\sim 6 \times 10⁻⁸ of growing cells (Pennington and Rosenberg 2007)] might remain roughly constant. In this model, the dinB(oc) mutations, which upregulate DinB in all cells, would not increase HMS size because the DSEs are not available in all cells. (B) Mutagenic double-strand-break repair during stress. Ponder et al. (2005) showed that repair of a double-strand break is a high-fidelity, nonmutagenic process in unstressed cells (left) but switches to a mutagenic mode during stress under the control of the RpoS general stress response (right). This process requires a DSE and its repair; induction of the SOS response (McKenzie et al. 2000), which we show here, is solely to provide DinB upregulation, and induction of the RpoS regulon (Ponder et al. 2005). Some as yet-unknown function regulated by the RpoS response licenses the use of DinB in those conditions. This function could be the documented increase in DinB expression (LAYTON and FOSTER 2003) or the induction of another regulatory factor, or a combination of both. ssDNA, single-strand DNA; HR, homologous recombination; X's, DNA polymerase errors/mutations; parallel lines, double-strand DNA; dashed lines, newly synthesized DNA strands.

DNA damage can induce RpoS (Merrikh *et al.* 2009), if it occurs in these experimental conditions, damage induction of RpoS appears not to be sufficient for mutagenesis; another RpoS-inducing input must contribute.

Finally, in growing cells, only $\sim 1\%$ of cells are SOS induced spontaneously (Pennington and Rosenberg 2007), and in this study we observed that making every cell in the population experience SOS-induced levels of DinB production, using the $dinB(o^c)$ alleles, did not increase stress-induced mutagenesis in otherwise wild-type cells (Figure 3). This could mean either of two things:

- 1. Unlike growing cells, in stationary phase, all of the cells are already SOS induced, and so the $dinB(o^c)$ alleles do not change the number of cells expressing DinB or the DinB levels in most cells. This possibility is unlikely, given the large increase in DinB levels that we observed with the $dinB(o^c)$ alleles measured in stationary-phase cells (Figure 2).
- 2. More plausibly, the data imply that, even though the $dinB(o^c)$ alleles confer SOS-induced levels of DinB to all cells, this is not sufficient for mutagenesis. These data support the model in which a DNA double-strand break and its repair, an RpoS response, and an SOS response (shown to act solely via DinB production) are all required for stress-induced point mutagenesis (Figure 6, A and B). We note that if all cells had become hypermutable when DinB was overproduced in all cells, mutation rate would have been higher than normal because the hypermutable cells would no longer be a small subpopulation. This was not observed (Figures 3 and 4).

Role of SOS/DinB in a hypermutable cell subpopulation: We previously suggested a model for the origin of the hypermutable cell subpopulation that appears to underlie most double-strand-break-repairassociated stress-induced point mutagenesis (Gonzales et al. 2008) on the basis of three requirements for stressinduced mutability discussed above: a genomic DSB/ DSE (and its repair), an SOS response, and an RpoS response (Galhardo et al. 2007). The simultaneous occurrence of these three events is proposed to differentiate the hypermutating cells. It is unknown what fraction of the cells in a stationary population experience an SOS response, but $\sim 1\%$ of the cells in log-phase cultures display spontaneous SOS induction, ~60% of those ($\sim 6 \times 10^{-3}$) due to a spontaneous DSB/DSE (Pennington and Rosenberg 2007). When these cells enter the stationary phase, RpoS is likely to be induced in all of them (HENGGE-ARONIS 2002). Thus, if the numbers for growing cells hold, then the ${\sim}6 \times 10^{-3}$ of cells with a DSB and an SOS response would become the HMS when RpoS induction occurred in the whole population in stationary phase (Figure 6A). We can now refine this model to note that the sole component of the SOS response required would be DinB upregulation.

The additional requirement for RpoS—to license the use of DinB in error-prone DSB/DSE repair (shown by PONDER et al. 2005)—could be based either solely on RpoS upregulation of DinB or on RpoS-controlled expression of other factors that permit DinB use (Figure 6B). The SOS and RpoS responses increase DinB expression ~10-fold and 2- to 3-fold, respectively (KIM et al. 2001; LAYTON and FOSTER 2003). The identities of potential DinB-licensing factors in the RpoS regulon are not yet known. This control would provide a restriction of the mutagenesis to periods of stress, and only to those few cells with a DSB/DSE. The restriction of mutagenesis to a cell subpopulation may allow clonal populations to hedge their bets during adaptation to changing environments, both conserving the original genome sequence, which is well adapted to the previous environment and useful if resources become available again suddenly, and simultaneously exploring the new adaptive landscape in the subpopulation.

Regulation of DinB mutator activity: In many other assay systems in which DinB-dependent mutagenesis has been observed, stress responses other than, or in addition to, SOS are required. In Salmonella bileinduced resistance mutagenesis, which is DinB dependent, the SOS and RpoS responses are required (J. Casadesus, personal communication, and Prieto et al. 2006). In B. subtilis starvation-associated mutagenesis, the ComK competence stress response is required for the DinB-dependent mutagenesis (Sung and YASBIN 2002). In E. coli, β -lactam antibiotics induce dinBtranscription independently of SOS (PEREZ-CAPILLA et al. 2005). P. putida DinB-dependent, stress-induced mutagenesis requires RpoS (SAUMAA et al. 2002). It is not known whether any stress response other than SOS is required for DinB-dependent, ciprofloxacin-induced resistance mutagenesis (CIRZ et al. 2005). Thus, it is plausible that DinB-dependent mutagenesis might usually require more than one stress-response input to occur. Although effects of DinB in SOS mutagenesis of E. coli (Kuban et al. 2006) have been observed, it is not known whether the DinB-dependent mutations may have arisen in cells also induced for RpoS or another stress response simultaneously.

What factors modulate DinB mutator activity? As a translesion DNA polymerase that inserts bases opposite several otherwise replication-blocking DNA adducts, DinB performs this role mostly in a high-fidelity fashion (Jarosz et al. 2006; Bjedov et al. 2007; Godoy et al. 2007). This, and the existence of dinB mutations that separate translesion from mutagenic functions (Godoy et al. 2007), imply that mutator activity occurs during synthesis that is not part of translesion synthesis (although it does occur during DSB repair; Ponder et al. 2005). Alternatively, DinB mutator activity could be taking place at sites of yet-unidentified DNA lesions.

A recent study implicated the SOS-induced UmuD protein as a candidate to inhibit DinB mutagenic

potential during SOS induction (GODOY et al. 2007). UmuD is produced virtually only during an SOS response (COURCELLE et al. 2001). Consequently, UmuD is not expected to be present in lexA(Ind⁻) dinB(o^c) cells, but stress-induced mutation rates were similar to those in wild-type cells (Figures 3 and 4). Therefore, UmuD appears not to inhibit DinB in its role in stress-induced point mutagenesis. Other levels of control are likely to exist.

One case of strongly DinB-dependent mutagenesis thought to occur with only one stress-response input in physiological conditions is SOS-mediated untargeted mutagenesis of phage λ (Kim et al. 1997). Under those conditions, mutagenesis of the phage DNA is heavily dependent on DinB, presumably relying on the physiological levels of dinB expression achieved in vivo during SOS induction. It is interesting to note that extensive double-strand-end-initiated recombination between the multiple copies of the phage DNA is expected to occur during a lytic cycle (Thaler and Stahl 1988). Those might be the sites of mutagenic action of DinB. Nevertheless, it is not known whether other factors, λ or host encoded, might also play a role in that process. Another example is the recent finding of increased DinB-dependent mutagenesis in cells lacking the ClpXP protease (AL MAMUN and HUMAYUN 2009). RpoS does not seem to be involved in the observed mutagenesis. However, the lack of a major protease is likely to have many pleiotropic effects, and there remains the possibility that other responses, which foster DinB activity, are triggered in those cells.

Evolution of stress-induced mutagenesis pathways: The occurrence of many different molecular mechanisms of stress-inducible mutagenesis (reviewed in the Introduction and by GALHARDO et al. 2007), and even of mechanisms by which a single stress response such as SOS promotes mutagenesis under different stress circumstances, suggests that these mutagenesis programs have evolved many times at least somewhat independently. This is also suggested by a survey of starvationstress-inducible mutability in 787 E. coli natural isolates (BJEDOV et al. 2003). Those authors found that whereas >80% of the natural isolates displayed stressinducible mutator activity, the ability to do so correlated well with ecological niche and poorly with strain phylogeny, suggesting multiple recent evolutions of the stress-inducible mutagenesis pathways. We have suggested that stress-inducible mutagenesis mechanisms are both somewhat varied and recently acquired because they confer a benefit to cells that is under periodic (alternating positive and negative) second-order selection (Galhardo et al. 2007). That is, these pathways are useful and selected in changing environments in which adaptation is promoted by mutability and responsiveness and superfluous and perhaps costly in static environments. Despite the variability and potential multiple origins, the basic themes of the regulation of

mutability temporally by stress responses, and potentially spatially in genomes (Galhardo *et al.* 2007), are widespread and appear to be potentially important evolutionary strategies.

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GENETICS

Supporting Information

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DinB Upregulation Is the Sole Role of the SOS Response in Stress-Induced Mutagenesis in *Escherichia coli*

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FILE S1

Primers used in this study

yafNwL:

5'TGTATATTCTGGTGTGCATTATTATGAGGGTATCACTGTATGCATCGAATTATTCCGGGGGATCCGTCGACC3'

yafPwR

5'A TACCAGGCGGCGTTATTTTCATTGCAAGCTGGATTTAATGTTGCGGTTTTGTAGGCTGGAGCTGCTTC3'

CatupdinB-F:

5'GAAGCGAATCTGGAGATGGAGATTGTTCCCCAGGGATTACGCGTGGTGTAGGCTGGAGCTGCTTC3'

CatupdinB-R:

5°CGCGCTGCCGCGTTCAAACATATTGCGGTTCTGGTCGTCTTTAATCATATGAATATCCTCCTTAG3°

dinBcatnock-R: 5' GATACCCTCATAATAATGC 3'

kandinBchrom-F:

5'CGCCACCGAGCTTGGTGAGCTGCAACCAGTATCAACACCATTGCGTGTAGGCTGGAGCTGCTTC3'

DinBRCAT:

5'GTGATACCCTCATAATAATGCACACCAGAATATACATAATAGTATCATATGAATATCCTCCTTA 3'

dinBOC1F: 5' CCCTGAAATCAAGGTATACTTTAC 3'

dinBOC1R: 5' GTAAAGTATACCTTGATTTCAGGG 3'

dinBOC2F: 5' GTATACTTTACCCTTGTTGAGAGGT 3'

dinBOC2R: 5' ACCTCTCAACAAGGGTAAAGTATAC 3'

For site directed mutagenesis, each of these two complementary primer pairs (last four primers listed) were used to amplify dinB fragments with the desired mutations in the borders. PCR amplification with external primers and both fragments as templates was used to generate full length products.



The Steric Gate Amino Acid Tyrosine 112 Is Required for Efficient Mismatched-Primer Extension by Human DNA Polymerase κ^{\dagger}

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ABSTRACT: Human DNA is continuously damaged by exogenous and endogenous genotoxic insults. To counteract DNA damage and ensure the completion of DNA replication, cells possess specialized DNA polymerases (Pols) that bypass a variety of DNA lesions. Human DNA polymerase κ (hPol κ) is a member of the Y-family of DNA Pols and a direct counterpart of DinB in Escherichia coli. hPolk is characterized by its ability to bypass several DNA adducts [e.g., benzo[a]pyrene diolepoxide-N2-deoxyguanine (BPDE-N2-dG) and thymine glycol] and efficiently extend primers with mismatches at the termini. hPolk is structurally distinct from E. coli DinB in that it possesses an ~100-amino acid extension at the N-terminus. Here, we report that tyrosine 112 (Y112), the steric gate amino acid of hPolk, which distinguishes dNTPs from rNTPs by sensing the 2'-hydroxy group of incoming nucleotides, plays a crucial role in extension reactions with mismatched primer termini. When Y112 was replaced with alanine, the amino acid change severely reduced the catalytic constant, i.e., k_{cat} , of the extending mismatched primers and lowered the efficiency, i.e., $k_{\text{cat}}/K_{\text{m}}$, of this process by ~400-fold compared with that of the wild-type enzyme. In contrast, the amino acid replacement did not reduce the insertion efficiency of dCMP opposite BPDE- N^2 -dG in template DNA, nor did it affect the ability of hPol κ to bind strongly to template-primer DNA with BPDE- N^2 -dG/dCMP. We conclude that the steric gate of hPolk is a major fidelity factor that regulates extension reactions from mismatched primer termini.

The human genome is continuously exposed to a variety of genotoxic agents such as polycyclic aromatic hydrocarbons, ultraviolet light, and reactive oxygen species (1). To counteract genotoxic insults, cells possess a number of defense strategies that enable them to complete chromosome replication and maintain the integrity of the genome. One of these strategies is translesion DNA synthesis (TLS)¹ accomplished by specialized DNA

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Abbreviations: Pol, DNA polymerase; hPolκ, human DNA polymerase κ; BPDE-N²-dG, benzo[a]pyrene diolepoxide-N²-deoxyguanine; Y112, tyrosine 112; TLS, translesion DNA synthesis; rNTP, ribonucleotide triphosphates; F12, phenylalanine 12; PCR, polymerase chain reaction; 8-OH-dG, 8-hydroxyguanine; O⁴-MedG, O⁴-methylthymine; MOE, molecular operating environment; Kd, equilibrium dissociation constant; HNE-dG, trans-4-hydroxy-2-nonenal-dG.

polymerases (Pols) (2, 3). These Pols can bypass a variety of DNA lesions, which would otherwise block DNA replication, to ensure the continuity of chromosome replication. Human cells possess more than 14 Pols, and approximately half of them are involved in TLS across DNA lesions, repair of DNA damage, or both (4, 5). In particular, Y-family Pols play major roles in the damage tolerance process by carrying out error-free TLS, although in some cases they mediate error-prone TLS, which can result in mutagenesis, carcinogenesis, and genetic diversity (6–8).

Of the four human Y-family Pols, i.e., Pol κ , - η , and - ι and REV1, Pol κ is unique in that its orthologues are present not only in Eukarya but also in bacteria and Archaea (9–11). The Escherichia coli orthologue is DinB or Pol IV, which bypasses several DNA lesions, such as N^2 -guanine adducts of benzo[a] pyrene diolepoxide (BPDE- N^2 -dG), in an error-free manner (12, 13). It is also active in spontaneous mutagenesis in λ phage and in stationary-phase E. coli cells (14–17). Like DinB, hPol κ bypasses BPDE- N^2 -dG in an error-free manner by incorporating dCMP opposite the lesion (18–23) and correctly mediates bypass DNA synthesis across other lesions, e.g., thymine glycol, at least in vitro (24). hPol κ is also distinct from other human Y-family Pols or even from E. coli DinB in that it efficiently extends

primers with terminal mismatches (25). $hPol\kappa$ extends mismatched primers with a frequency of $10^{-1}-10^{-2}$, which is more than 10 times greater than that seen with $hPol\eta$ or $E.\ coli\ DinB.$ Thus, $hPol\kappa$ may play a role in TLS in vivo as an extender of mismatched primers generated by other Pols (2). Structural analysis of the catalytic core of $hPol\kappa$ suggests that the N-terminus, approximately 75 amino acids that are absent in bacterial and archaeal counterparts (Figure 1 of the Supporting Information), forms an "N-clasp" domain that enhances the binding of $hPol\kappa$ to DNA (26). Despite distinct biochemical and structural characteristics, the amino acids in the vicinity of the terminal base in primers are similar to those of other family members. Therefore, the exact mechanism by which $hPol\kappa$ mediates efficient mismatch extension reactions remains unresolved.

In this study, we examined the role of tyrosine 112 (Y112), the steric gate amino acid of hPolk, in TLS and extension reactions from mismatched primer termini. The steric gate amino acid is the residue that distinguishes dNTPs from rNTPs by sensing the 2'-hydroxy group of rNTPs (27, 28). We have previously demonstrated that the steric gate of the Archaea orthologue of $hPol\kappa$, i.e., phenylalanine 12 (F12) of Sulfolobus acidocaldarium (Sac) Pol Y1 (also known as DBH), plays a role in fidelity during DNA synthesis by regulating the efficiency with which oxidized dNTPs are incorporated into DNA (29). Both F13, the steric gate amino acid of E. coli DinB, and F12 of DBH were also shown recently to play important roles in TLS across N^2 -deoxyguanine-DNA adducts (12). Here, we report that the steric gate amino acid of hPol κ is critical for mismatch extension reactions. When Y112 is replaced with alanine (A), the catalytic efficiency of extension from mismatched DNA substrates decreases more than 400-fold. Unlike in the bacterial and archaeal orthologues, the steric gate is unimportant in regulating correct counterbase insertion opposite N^2 -deoxyguanine—DNA adducts. We suggest that the steric gate amino acid might have evolved into a major fidelity factor regulating mismatch extension reactions when the ancestral E. coli DinB- and archaeal DBH-type proteins developed the extended N-terminal domain. We propose a possible mechanism by which hPolk extends mismatched primer termini depending on the steric gate.

EXPERIMENTAL PROCEDURES

More detailed experimental protocols are available in the Supporting Information.

Construction of the hPolk Overexpression Vector. The hPolk gene (NCBI GenBank POLK entry accession number XM_003930.2) was amplified by polymerase chain reaction (PCR) from the Clontech human testis large insert cDNA library. The amplified fragments were digested with NcoI and BamHI, and the resulting fragment was ligated into similarly digested vector pYG8582, which is the same as pET-16b (Novagen) but has the translational DB (downstream box) enhancer (30). The resulting plasmid pYG8583 carries the coding sequence for N-terminal 10-His-tagged full-length hPolκ. To construct a C-terminally truncated hPolk expression vector, a synthetic linker was ligated between the XbaI and BamHI sites of pYG8582. This construct was then digested with XbaI and AvrII. and after dephosphorylation of both ends, the digested plasmid was ligated to the XbaI fragment of pYG8583, which carried the N-terminal portion of the hPol κ coding sequence. The resulting construct, overexpressed C-terminally truncated 10-His-tagged

 $hPol\kappa_{1-559}$, was named pYG8591. We refer to it as $hPol\kappa$ throughout this study.

Construction of Mutant hPolκ Overexpression Vectors. The Y112A mutant of hPolκ was made from pYG8591 by site-directed mutagenesis using PCR. An amino acid change of Y112 to valine (V) was introduced by the same method. Overexpression vectors for Y112A and Y112V were named pYG331 and pYG332, respectively.

Overexpression and Purification. To express the wild-type and mutant proteins, plasmids pYG8591, pYG331, and pYG332 were transformed into Rosetta competent cells (Novagen), and expression was induced by adding IPTG. The harvested cells were resuspended in BugBuster lysis buffer (Novagen), and soluble proteins were collected by centrifugation. hPolk and the mutant proteins were purified by binding to BD TALON Superflow resin (BD Biosciences) and eluted in accordance with the manual provided by BD Biosciences. The eluted proteins were further purified by gel filtration, followed by ion exchange chromatography (HiTrap Heparin HP, GE Healthcare) using an FPLC system (AKTAexplorer 10S, GE Healthcare). The purified proteins were stored at -80 °C.

Primer Extension Assay. Standard polymerase reactions (10 μ L) were performed in 40 mM Tris-HCl (pH 8.0), 5 mM MgCl₂, 10 mM DTT, 0.1 mg/mL BSA, 60 mM KCl, 2.5% glycerol, and 250 μ M dNTPs. To this reaction mixture were added annealed oligonucleotides (100 nM), consisting of primers (14mers, 18mers, or 19mers) and template 5'-GC GCGCTTCTGGCCAATXGCAGAATTCCTAGGGAAG-3' (36mer), where X represents lesions, i.e., BPDE- N^2 -dG, 8-hydroxyguanine (8-OH-dG), O^6 -methylguanine (O^6 -MedG). O^4 -methylthymine (O^4 -MedT), or thymine glycol, at position 19. If no lesions were present, X represents dG. The 36mer templates containing the lesions were synthesized by Japan BioServices Co. (Saitama, Japan), except for BPDE-N²-dG. The template DNA bearing BPDE-N²-dG, i.e., (-)-trans-anti-benzo[a]pyrene diol epoxide adduct at N^2 -dG, at position 19 of the 36mer DNA was synthesized as reported previously (31). All other oligonucleotides were synthesized by BEX Corp. (Tokyo, Japan) and double purified by high-performance liquid chromatography. The primers were labeled with Cy3 at the 5'-terminus; the exact sequences are shown in Table 1 of the Supporting Information and in the top part of Figures 1, 2, and 4. Wild-type hPolk, Y112A, or Y112V was added at a concentration of 40, 10, or 5 nM, respectively, in the experiments shown in Figures 1, 2, and 4, followed by incubation for 15 min (Figures 1 and 2) or 20 min (Figure 4) at 37 °C. When ribonucleotide incorporation was assayed, rNTP instead of dNTP was included in the reaction mixtures at concentrations of 0, 50, 100, 250, or 500 μ M. After the reactions were terminated, the products were resolved by electrophoresis on a 15% polyacrylamide gel and visualized with the Molecular Imager FX (Bio-Rad). The band intensities were quantified with Quantity One (Bio-Rad).

Steady-State Kinetic Analyses. The constituents of the reaction mixtures were the same as those in the primer extension assay except that only one dNTP was included and the protein concentrations were 5 nM. The reactions were initiated by adding dNTP. Both the concentration of dNTP and the incubation time were varied depending on the DNA substrate (matched or mismatched primer termini with or without lesions in the template). The products were resolved by electrophoresis on a 15% polyacrylamide gel, and the band intensities were quantified as described in the primer extension assay. Less than 20% of the

primers were extended in the steady-state kinetic analyses, ensuring single-hit kinetics. For each DNA substrate, the rate of incorporation was plotted as a function of dNTP concentration, and the $V_{\rm max}$ and $K_{\rm m}$ values were determined by Enzyme Kinetics Module 1.1 of SigmaPlot 2001 software (SPSS Inc., Chicago, IL). $k_{\rm cat}$ was calculated by dividing $V_{\rm max}$ by the enzyme concentration. All values are means \pm standard errors from three experiments.

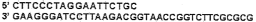
DNA Binding BIAcore Assay. The DNA binding assay was performed using the BIAcore3000 instrument (Biacore). The oligonucleotides that were used were basically the same as those used in the primer extension assay, except that primers were biotinylated at their 5'-termini to enable immobilization. The equilibrium dissociation constants (K_d) were calculated from the kinetic traces using BIAevaluation version 4.0 (Biacore) and employing local fitting according to the "1:1 binding with drifting baseline" predefined model.

Molecular Modeling. To visualize the relative position between Y112 or A112 and mismatched primer termini, active sites of wild-type hPolk and its derivative Y112A complexed with a G:T mismatch and dATP were modeled on the basis of the reported crystallographic structure [Protein Data Bank entry 20H2 (16)] using the 2007.09 version of the Molecular Operating Environment (MOE) (Chemical Computing Group Inc., Montreal, QC). Sequences of the template and primer were replaced with those which were applied to the experiments presented here, i.e., 3'-TCCTTAAGACGGTAA-5' and 5'-AG-GAATTCTGC \underline{T} -3', where \underline{G} and \underline{T} represent the mismatched template and primer bases, respectively. The incoming dTTP in the original structure was replaced with dATP, and Y112 was replaced with alanine in the model of Y112A. After all replacements, the total potential energy of the complex was minimized under the conditions where all coordinate sets of hPolk were fixed.

RESULTS

Y112 Is the Steric Gate Amino Acid of hPolκ. By aligning the amino acid sequences of various Y-family Pols, we postulated that Y112 of hPolκ was the steric gate amino acid. To explore this possibility, Y112 was substituted with A or V; the resulting proteins and the wild-type enzyme (Figure 1 of the Supporting Information) were subjected to primer extension assays with rNTPs and dNTPs. The truncated form of hPolκ consisting of the 559 N-terminal residues is far more stable than the full-length protein and was therefore used throughout the study. In the presence of four rNTPs, the Y112A and Y112V proteins efficiently incorporated rNTPs and extended the primer, while the wild-type enzyme did not (Figure 1). In the presence of dNTPs, all three proteins incorporated dNTPs and extended the primers. These results clearly indicate that Y112 functions as the steric gate of hPolκ.

Translesion Activity of hPol κ Y112A and Y112V. To examine whether the steric gate residue of hPol κ is required for TLS across lesions, we performed in vitro primer extension assays using Y112A, Y112V, and the wild-type enzyme (Figure 2). The DNA lesions we analyzed were BPDE- N^2 -dG, 8-OH-dG, O^6 -MedG, O^4 -MedT, and thymine glycol. hPol κ more efficiently bypasses the (-)-trans-anti-BPDE- N^2 -dG adduct than the (+)-trans-anti-BPDE- N^2 -dG adduct (32). To examine the effects of the amino acid substitution of Y112 on the bypass activities carefully, we employed the (-)-trans-anti-BPDE- N^2 -dG



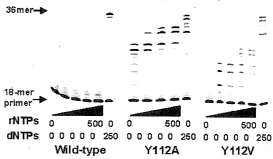


FIGURE 1: Incorporation of rNTPs by wild-type and mutant hPol κ proteins was analyzed by a primer extension assay using Cy3-labeled 18mer primers annealed to 36mer templates (i.e., 18/36G in Table 1 of the Supporting Information). The primer-template DNA (0.1 μ M) was incubated with 40 nM wild type, Y112A, or Y112V in the presence of rNTPs for 15 min at 37 °C. Concentrations of rNTP were increased up to 500 μ M. dNTP at a concentration of 250 μ M was added as the control. The samples were resolved by 15% denaturing polyacrylamide gel electrophoresis and analyzed as described in Experimental Procedures.

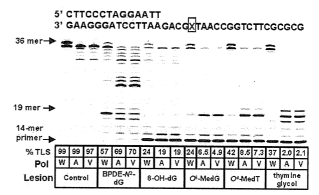


FIGURE 2: Effect of substituting A or V for Y112 on TLS across several lesions in DNA. The Cy3-labeled 14mer primer was annealed to each of six different 36mer templates containing an unmodified dG, dT, or lesions at position 19 (indicated as a boxed X). Reactions were allowed to proceed for 15 min at 37 °C with 100 nM primer-template and 10 nM hPolk protein. W, A, and V represent wild-type hPolk, Y112A, and Y112V, respectively. % TLS indicates the percentage of the amount of primers opposite and beyond the lesion relative to the total amount of the primer.

adduct in this study, although (+)-trans-anti-BPDE- N^2 -dG is the major adduct of benzo[a]pyrene (32). Both Y112A and Y112V exhibited primer extension activities across BPDE-N²-dG in DNA comparable to those of the wild-type enzyme. Rather. the mutant proteins displayed substantially reduced activities in inserting a base opposite other types of lesions except for 8-OHdG compared to the wild-type enzyme. In particular, Y112A and Y112V exhibited a severely reduced ability to deal with thymine glycol in DNA. To gain insight into the roles of the steric gate amino acid in TLS in a quantitative manner, we conducted steady-state kinetic analyses of insertion and extension steps across BPDE- N^2 -dG and thymine glycol in DNA (Table 1). The Y112A incorporation efficiency, i.e., k_{cat}/K_{m} , for dCMP opposite template BPDE- N^2 -dG was 8-fold greater than that of the wildtype enzyme. In contrast, the amino acid change decreased the efficiency of incorporation of dCMP opposite undamaged dG by 90%. In the extension step, replacement of Y112 with

Table 1: Steady-State Kinetic Parameters for Incorporation and Extension across BPDE-N²-dG and Thymine Glycol in DNA by the Wild Type and Y112A^a

template		$k_{\rm cat} ({\rm min}^{-1})$		Incorporation $K_{\rm m} (\mu { m M})$		$k_{\mathrm{cat}}/K_{\mathrm{m}}~(\mu\mathrm{M}^{-1}~\mathrm{min}^{-1})$		
	dNTP	WT ^b	Y112A	WT	Y112A	WT	Y112A	Y112A/WT
G BPG ^c	dCTP	7.9 ± 2.0 0.5 ± 0.088	8.2 ± 1.5 1.6 ± 0.30	6.7 ± 2.6 1900 ± 560	60 ± 22 750 ± 340	1.2 2.7 × 10 ⁻⁴	1.4×10^{-1} 2.1×10^{-3}	1/8.6 7.8/1
T ThG1 ^d	dATP	13 ± 2.2 5.6 ± 0.97	$13 \pm 2.0 \\ 0.20 \pm 0.060$	3.4 ± 1.4 190 ± 74	46 ± 15 2300 ± 1100	3.8 2.9×10^{-2}	2.9×10^{-1} 8.4×10^{-5}	1/13 1/350

	k_{cat} ((min ⁻¹)	Extension $K_{\rm m}$ ([μ M)	$k_{\rm cat}/K_{\rm m}~(\mu{\rm M}^{-1}~{\rm min}^{-1})$		
base pair at th 3'-primer termini (primer-template)	WT ^b	Y112A	WT	Ÿ112A	WT	Y112A	Y112A/WT
G/C BPG/C	18 ± 1.1 11 ± 0.69	15 ± 0.73 9.3 ± 1.7	3.5 ± 0.40	65 ± 6.1	5.2	2.3×10^{-1} 1.0×10^{-2}	1/23
T/A ThGI/A	15 ± 0.69 15 ± 0.72 4.4 ± 0.18	9.3 ± 1.7 18 ± 1.4 0.2 ± 0.026	44 ± 7.7 1.4 ± 0.16 240 ± 21	910 ± 380 17 ± 3.1 810 ± 240	0.25 11 1.8×10^{-2}	1.0×10^{-2} 1.1 2.5×10^{-4}	1/24 1/11 1/75

"Incorporation reactions were analyzed by determining the steady-state kinetic parameters for incorporation of dCMP opposite template guanine (G) or BPDE-N²-dG (BPG) and for incorporation of dAMP opposite template thymine (T) or thymine glycol (ThGl). Exact sequences of primer-template DNA, i.e., 18/36G or 18/36T, are described in Table 1 of the Supporting Information. The primer-template (100 nM) and the proteins (5 nM) were incubated at 37 °C. Extension reactions were analyzed by determining the steady-state kinetic parameters for incorporation of dAMP opposite template thymine (T) adjacent to G/C, BPG/C, T/A, or TG/A at the primer-template termini. Exact sequences of primer-template DNA, i.e., 19C/36G or 19A/36T, are described in Table 1 of the Supporting Information. The primer-template (100 nM) and the proteins (5 nM) were incubated at 37 °C. "WT, wild-type hPolk. "BPG, BPDE-N²-dG. "ThGl, thymine glycol.

A decreased the $k_{\rm cat}/K_{\rm m}$ of dAMP incorporation opposite template dT adjacent to either BPDE- N^2 -dG or dG by 95%. Therefore, the overall TLS across BPDE- N^2 -dG was only moderately compromised by the amino acid change. For TLS across thymine glycol, however, the amino acid change severely reduced both the insertion and extension efficiency. The insertion and extension efficiencies of Y112A were only ~0.3 and ~1%, respectively, compared to that of the wild-type enzyme. From these results, we concluded that the steric gate amino acid of hPolk is nonessential for TLS at least across (—)-transanti-BPDE- N^2 -dG in DNA. Instead, Y112 may play roles in incorporation steps opposite lesions other than N^2 -dG adducts, such as thymine glycol, and in extension steps from primer termini.

Strong Binding of hPolk to Primer-Template DNA with $BPDE-N^2-dG/dCMP$. Although the amino acid replacement of Y112 with A or V did not severely reduce TLS efficiency across BPDE- N^2 -dG in DNA, it might still modulate binding to primertemplate DNA. To test this possibility, we examined the physical interactions between hPolk and primer-template DNA with or without lesions using surface plasmon resonance and calculated the equilibrium dissociation constants (K_d) (Figure 3 and Table 2 of the Supporting Information). Wild-type hPolk bound to primer-template DNA with BPDE- N^2 -dG \sim 3-fold more strongly than it bound to DNA without the lesion. Strong binding depended on the length of the primer and on the dNMP opposite the lesion. Specifically, strong binding was observed when the length of the primer was the same as the length of the template between the 3'-end and the lesion, i.e., a 19mer primer. Alternatively, the length of the primer could be one or two base pairs longer than the length of template, i.e., a 20mer or 21mer primer. No strong binding was observed when the primer was shorter than 19 bp or longer than 21 bp. Additionally, for strong binding, the dNMP opposite the lesion had to be correct (dCMP); strong binding was not observed when the 19mer primers had dGMP,

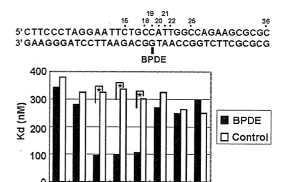


FIGURE 3: Affinity of hPolk for primer-template DNA with a BPDE- N^2 -dG adduct. Primers of different lengths, i.e., 15mers, 18mers, 19mers, 20mers, 21mers, 25mers, or 36mers, were annealed with template DNA (36mer) with BPDE- N^2 -dG at position 19. As a control, template DNA with an undamaged dG at position 19 was annealed to the various primers. The affinity between protein and DNA was measured with a BIAcore 3000, and the K_d was calculated with BIAevaluation version 4.0. The heights of bars represent the average K_d values of three experiments. An asterisk indicates P < 0.001 (t-test).

20 21

Length of primer (mer)

22 25

15

18 19

dTMP, or dAMP at the 3'-terminus. No strong binding was observed in primer-template DNA with other lesions, such as thymine glycol, or in primer-template DNA with mismatched termini (data not shown). Importantly, replacing the steric gate amino acid with A or V did not enhance or weaken binding to primer-template with BPDE- N^2 -dG lesions (Figure 2 and Table 3 of the Supporting Information), nor did they modulate the binding affinity for control DNA. These results suggest that the steric gate amino acid may play roles in TLS other than modulating the ability of hPol κ to bind to damaged or undamaged DNA.

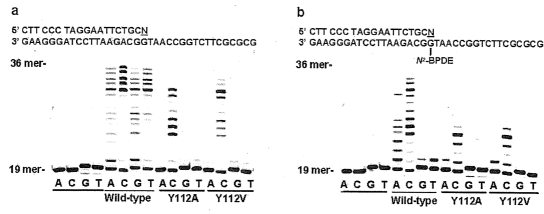


FIGURE 4: Extension from mismatched and matched DNA substrates by wild-type hPok, Y112A, and Y112V. Each of four 5'-Cy3-labeled primers (19mers) with either A, C, G, or T at the terminus (\underline{N}) was annealed to a 36mer undamaged template (a) or a template with BPDE- N^2 -dG at position 19 (b). Reactions were conducted for 20 min at 37 °C. The products were resolved by 15% polyacrylamide gel electrophoresis and visualized using a Molecular Imager FX (Bio-Rad) equipped with Quantity One software.

	k _{cat} (min ⁻¹)		$K_{\rm m} (\mu { m M})$		$k_{\rm cat}/K_{\rm m}~(\mu{ m M}^{-1}~{ m min}^{-1})$		
base pair at 3'-primer termini (primer-template)	WT ^b	Y112A	WT	Y112A	WT	Y112A	Y112A/WT
C/G	18 ± 1.1	15 ± 0.73	3.5 ± 0.40	65 ± 6.1	5.2	2.3×10^{-1}	1/23
A/G	3.1 ± 0.26	0.11 ± 0.020	89 ± 16	1400 ± 450	3.5×10^{-2}	8.2×10^{-5}	1/420
G/G	4.3 ± 0.40	0.24 ± 0.024	73 ± 16	1700 ± 300	5.9×10^{-2}	1.4×10^{-4}	1/430
T/G	3.6 ± 0.24	0.34 ± 0.039	44 ± 8.3	$1500 \pm 320^{\circ}$	8.2×10^{-2}	2.2×10^{-4}	1/360

^a Steady-state kinetic parameters were determined for incorporation of dAMP opposite template thymine (T) adjacent to matched (C/G) or mismatched (A/G, G/G, or T/G) primer termini. Exact sequences of matched and mismatched primer-template DNA, i.e., 19C/36G, 19A/36G, 19G/36G, and 19T/36G, are described in Table 1 of the Supporting Information. Primer-template DNA (100 nM) and the proteins (5 nM) were incubated at 37 °C. ^b WT, wild-type hPolκ.

Crucial Role of Y112 in Extension Reactions from Mismatched Termini. Because hPolκ efficiently extends from primer-terminal mismatches, we next examined the possibility that the steric gate amino acid might play a role in mismatch extension reactions. Remarkably, the ability of both Y112A and Y112V to extend primers from mismatched termini was severely reduced compared with extension from matched termini (Figure 4). Reduced extension from mismatched termini was observed when the template DNA had either BPDE-N²-dG or undamaged dG. To further analyze the effects of the amino acid changes on the mismatched extension reactions, we carried out steady-state kinetic analyses with an undamaged template (36mer) annealed to a 19mer primer with mismatched (A/G, G/G, and T/G) or matched (C/G) termini (Table 2). Y112A exhibited a severely reduced ability to extend primers from mismatched termini; the extension efficiency (k_{cat}/K_m) was 360-430 times lower than that of the wild-type enzyme. In contrast, the extension efficiency from matched termini was only 23-fold lower than that of the wild-type enzyme. Therefore, the amino acid change has an effect that is ~20 times greater on the ability of hPolk to extend primers from mismatched termini, compared to the extension of primers with matched termini. The reduction in extension efficiency from mismatched termini was due to a reduction in the catalytic constant, i.e., $k_{\rm cat}$. The $k_{\rm cat}$ values for mismatch extension were reduced 10-30-fold by the Y112A amino acid change, whereas the value for matchedprimer extension was not affected by the amino acid change. In contrast, the $K_{\rm m}$ values were increased 20–30-fold by the Y112 to A change in both mismatched and matched primer termini.

From these results, we conclude that the steric gate amino acid is crucial to the chemistry of extension reactions catalyzed by $hPol\kappa$ from mismatched primer termini.

DISCUSSION

In this study, we report for the first time, to the best of our knowledge, that the steric gate residue of hPolk plays a critical role in extension from mismatched DNA substrates. Mismatch extension reactions catalyzed by hPolk were substantially compromised by the replacement of the Y112 amino acid with A or V (Figure 4). The steric gate residue appears to be critical to the chemistry of mismatch extension, i.e., the nucleophilic attack of the misaligned primer 3'-OH group upon the α-phosphate of dNTP, because the replacement of Y112 with A specifically reduced the k_{cat} values for mismatch extension reactions (Table 2). No reduction was observed in the k_{cat} values for matched-primer extension (18 \pm 1.1 and 15 \pm 0.73 min⁻¹ in the wild-type and Y112A enzymes, respectively). Interestingly, amino acid replacement had similar distinct effects on the k_{cat} values for extension from the primer-template with thymine glycol/ dAMP or BPDE- N^2 -dG/dCMP at the termini (Table 1). The k_{cat} value for extension from primer-template DNA with thymine glycol/dAMP was reduced more than 20-fold by the amino acid replacement $(4.4 \pm 0.18 \text{ and } 0.20 \pm 0.026 \text{ min}^{-1} \text{ in the wild-type})$ and Y112A enzymes, respectively), while virtually no reduction was seen for extension from primer-template DNA with BPDE- N^2 -dG/dCMP (11 \pm 0.69 and 9.3 \pm 1.7 min⁻¹ in the wild-type and Y112A enzymes, respectively). Therefore, we propose that. at least partially, the termini of primer-template DNA with

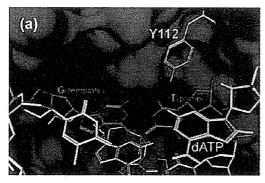
thymine glycol/dAMP or BPDE- N^2 -dG/dCMP lesions structurally resemble "mismatched" and "matched" termini, respectively, in the active site of hPol κ .

The first 18 N-terminal residues of hPol κ also play an important role in mismatch extension (26). hPolk missing the first 18 amino acids has a severely reduced ability to extend mismatched primer termini, although it retains primer extension activity from matched termini. Because the first 18 amino acids contribute to the proficiency of DNA binding by hPolk, it is surmised that the 18 amino acids and the subsequent N-clasp domain may contribute to the encirclement of DNA and may increase the time during which the mismatched primer 3'-OH group can acquire proper alignments for nucleophilic attack (26). Although the catalytic core of $hPol\kappa_{19-526}$ in a ternary complex with DNA and an incoming nucleotide has been crystallized and the structure has been determined, it lacks the first 18 amino acids (26). Therefore, the exact position adopted by these N-terminal amino acids in the complete structure is unclear. We suggest that Y112 may play a critical role in mismatch extension reactions in a manner distinct from that of the action of the 18 N-terminal amino acids. This is because replacing Y112 with A did not reduce the affinity of hPol κ for primer-template DNA (Figure 2 and Table 3 of the Supporting Information), while removing the 18 N-terminal amino acids strongly affects the enzyme's ability to bind to DNA. In addition, the steric gate amino acid is close to primer termini and incoming dNTPs, whereas the 18 N-terminal amino acids may not be. Given the location of Y112, we speculate that Y112 may directly interact with a mismatched terminal base in the primer, thereby preventing the terminal base from moving from the position of the cognate Watson-Crick pairing terminal bases (Figure 5a). This may be a prerequisite for extension from mismatched primer termini, because the mismatched terminal bases, which are otherwise moved from the normal base pairing positions, should be in the proximal normal positions in the active site to acquire proper alignment. The presence of the extended N-terminal domain of hPolk may enable the steric gate amino acid to interact with the terminal base in the primer, which thereby plays a decisive role in mismatch primer extension. Carlson et al. (33) reported that hPolk forms productive complexes with mismatched primer termini but not with matched primer termini. Both the 18 N-terminal amino acids and Y112 may play roles in the formation of the productive complexes with the mismatched termini. In addition, the steric gate may guide incoming dNTPs to

a particular position, where the chemistry between the primer 3'-OH group and α -phosphate of the dNTP can take place. Y112A may have a less constrained active site, which decreases the probability of achieving proper alignment for the phosphoryl transfer reactions, resulting in a decrease in $k_{\rm cat}$ values for mismatch primer extension (Figure 5b).

Immediately adjacent to the steric gate in hPol κ is the highly conserved amino acid F111. Replacing the corresponding amino acid with certain other amino acids in several B-family Pols such as yeast Pol results in a decrease in the fidelity of DNA synthesis and an increase in the efficiency of mismatch extension (34, 35). Although the exact mechanisms by which this conserved amino acid plays a role in the extension of mismatched termini are not known, amino acid substitutions may alter the geometry of the nascent base pair binding pocket and/or the chemistry of the reaction (35). To examine whether hPolk F111 affects the efficiency of extension reactions from mismatched termini, we changed F111 to A and purified the protein. However, F111A displayed significantly reduced DNA synthesis activity (data not shown). Replacing both F111 and Y112 with A resulted in greatly reduced activity. Thus, we could not determine the efficiency of extension reactions from mismatched termini with F111A or F111A/Y112A. Interestingly, F34L of yeast Poln, which is located next to the steric gate amino acid F35, exhibits substantially reduced DNA synthesis activity (34). Therefore, we suggest that unlike B-family Pols, the conserved amino acids adjacent to the steric gate in Y-family Pols, i.e., F111 in hPolκ and F34 in yeast $Pol\eta$, may primarily play roles in DNA synthesis activity.

Although Y112 was nonessential for TLS across BPDE- N^2 dG in DNA, replacing Y112 with A weakened the ability of hPolk to bypass several other lesions such as thymine glycol (Figure 2 and Table 1). The efficiency (k_{cat}/K_m) of incorporation of dAMP opposite thymine glycol was ~350 times lower than that of the wild-type enzyme, while the efficiency of incorporation of dAMP opposite undamaged thymine was only 13 times lower than that of the wild-type enzyme. In addition, the k_{cat}/K_{m} for incorrect incorporation of dNMP opposite an undamaged base was 40-60 times lower in Y112A than in the wildtype enzyme (Table 4 of the Supporting Information), while the $k_{\text{cat}}/K_{\text{m}}$ for correct incorporation of dCMP opposite template guanine was \sim 9 times lower than that of the wild-type enzyme. We propose therefore that replacing Y112 with A may alter the geometry of the nascent base pairing binding pocket, which in turn increases selectivity against both incorporation of dNTP



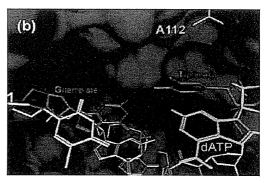


FIGURE 5: Images of the active site of wild-type hPolk (a) and Y112A (b). The steric gate Y112 may interact with primer T, thereby locking T in a position close to template G (a). A112 has no interactions with primer T, and thus, primer T is positioned far from template G (b). The distance between A112 and primer T is more than 5 Å greater than that between Y112 and primer T. Y112 (a) and A112 (b) are highlighted in orange. A mismatched base pairing of template G with primer T (red) was modeled in the active site with incoming dATP (light blue). The remaining template and primer strands are colored light brown. The active site of hPolk is displayed as a Connolly surface (blue).

opposite thymine glycol and incorrect incorporation of dNTP opposite an undamaged template base. In this respect, BPDE- N^2 -dG is exceptional, because changing Y112 to A or V enhanced the efficiency of incorporation of dCMP opposite the lesion (Figure 2 and Table 1). Template guanine base modified with BPDE might better fit the less constrained active site in Y112A or Y112V than in the native one.

hPol κ strongly binds to primer-template DNA with BPDE- N^2 dG when the primer possesses dCMP opposite the lesion (Figure 3 and Table 2 of the Supporting Information). No such strong binding was observed with primer-template DNA with any of the other lesions that were examined in this study. Structural analysis of the hPolk catalytic core suggests a model in which there is a cleft in the active site, through which the long chain of trans-4hydroxy-2-nonenal-dG (HNE-dG) in the template extends into the solvent (26). By analogy with the position of the HNE-dG adduct, we postulate that the BPDE adduct also fits into the cleft, thereby reducing the level of obstruction of DNA synthesis mediated by hPolk. Similar strong binding to primer-template DNA with a thymine dimer/dAMP base pair is observed with hPoln (36). Intriguingly, the strong binding by hPoln requires the presence of the correct nucleotide, i.e., dAMP, opposite the lesion. The strong binding disappears after Pol η has inserted two nucleotides beyond the lesion, which is remarkably similar to the mode of binding of hPol κ to DNA with BPDE- N^2 -dG. Because hPol κ and hPol η bypass BPDE- N^2 -dG and thymine dimer, respectively, in an error-free manner, it is tempting to speculate that they are cognate lesions for these two Pols. Alternatively, structurally similar but endogenous DNA lesions such as steroid hormone DNA adducts (37) could be the cognate lesions for hPolk. It has been proposed that the strong binding of hPoly may have implications for Pol switching and the restriction of error-prone Pols to damaged sites (36). Likewise, the strong binding of hPolk may contribute to the mechanism of transient access to primer-template DNA with BPDE-N²-dG by this intrinsically error-prone Pol.

In summary, we found that the steric gate amino acid Y112 was crucial to mismatch extension reactions catalyzed by hPolk. Y112 appears to play an important role in the chemistry of mismatch extension. It may directly interact with the mismatched terminal base in the primer strand and prevent movement from the normal matched base pairing position in the active site (Figure 5). Unlike bacterial and archaeal orthologues, in which the steric gates are essential for TLS across N^2 -dG adducts in DNA (12), hPolκ Y112 may be unnecessary in bypass reactions across BPDE-N²-dG in DNA. E. coli DinB, which lacks the Nterminal clasp domain (Figure 1 of the Supporting Information), does not display high efficiency in mismatch extension, although it has a steric gate (38). We speculate, therefore, that the steric gate amino acid may have evolved into a major fidelity factor that regulates mismatch extension in hPolk when the ancestral bacterial DinB and archaeal orthologue gained the extra Nterminal domain.

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SUPPORTING INFORMATION AVAILABLE

Detailed experimental protocols, sequences of primer-template DNA, affinity of hPol κ and the mutants for primer-template DNA with or without BPDE- N^2 -dG, kinetic parameters for incorporation of dNTP opposite template G by hPol κ and Y112A, and purification of hPol κ and the mutants. This material is available free of charge via the Internet at http://pubs.acs.org.

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