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## Estimating the Impact of Human Base Excision Repair Protein Variants on the

# **Repair of Oxidative DNA Base Damage**

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## ABSTRACT

Epidemiological studies have revealed a complex association between human genetic variance and cancer risk. Quantitative biological modeling based on experimental data can play a critical role in interpreting the impact of genetic variation on biochemical pathways relevant to cancer development and progression. Defects in human DNA base excision repair (BER) proteins can reduce cellular tolerance to oxidative DNA base damage caused by endogenous and exogenous sources, such as exposure to toxins and ionizing radiation. If not repaired, DNA base damage leads to cell dysfunction and mutagenesis, consequently leading to cancer, disease, and aging. Population screens have identified numerous single nucleotide polymorphism (SNP) variants in many BER proteins, and some have been purified and found to exhibit mild kinetic defects. Epidemiological studies have led to conflicting conclusions on the association between SNP variants in BER proteins and cancer risk. Using experimental data for cellular concentration and the kinetics of normal and variant BER proteins, we apply a previously developed and tested human BER pathway model to (i) estimate the impact of mild variants on BER of abasic sites and 8-oxoguanine, a prominent oxidative DNA base modification, (ii) identify ranges of variation associated with substantial BER capacity loss, and (iii) reveal non-intuitive consequences of multiple simultaneous variants. Our findings support previous work suggesting that mild BER variants have a minimal effect on pathway capacity, while more severe defects and simultaneous variation in several BER proteins can lead to inefficient repair and potentially deleterious consequences of cellular damage.

#### INTRODUCTION

DNA is continuously being damaged by endogenous sources, such as oxidative base modification resulting from reactive oxygen species (ROS) attack (reviewed in (1)). ROS are formed as metabolites during normal cellular respiration, with increased levels of ROS, as well as increased DNA base damage, associated with ischemia (2, 3), hyperoxia in human lung cells (4), human artherosclerotic plaques (5), and chronic inflammation in ulcerative colitis patients (6). ROS are also formed as secondary damaging agents as a result of exposure to toxins and ionizing radiation (e.g., sunlight, environmental IR). DNA base damage can lead to genotoxicity, replication stalling, apoptosis, and mutagenesis, with physiological consequences including disease, aging, and cancer, as reviewed in (7). Modified bases not only affect transcriptional integrity and replication, but also DNA-protein binding (8, 9). Consequently, oxidative DNA base damage is normally removed and repaired by the efficient base excision repair (BER) pathway system of prokaryotic and eukaryotic cells, reviewed in (1, 10, 11).

Population screens have identified extensive genetic variation, in particular single nucleotide polymorphisms (SNPs), altering the primary amino acid sequence of DNA base excision repair proteins (e.g., (12)). This genetic variation has been found in some instances to diminish the stability or efficiency of the encoded repair protein, reducing its kinetic effectiveness (13-15). Furthermore, sequence variants with potential kinetic impact have been predicted computationally (16). Splice variants of repair genes have also been identified in cancerous cells, some of which may lead to a dominant negative protein form with reduced repair efficiency (17, 18).

Non-functional DNA repair alleles, such as BRCA1/2 for breast cancer (19) and MSH2 and MLH1 for hereditary non-polyposis colorectal cancer (20), are "cancer genes" strongly linked to increased cancer risk. However, "cancer genes" account for only about 5% of known cancer cases (21, 22). Consequently, reduced function repair protein variants may be exposure-dependent susceptibility alleles responsible for the majority of observed increased cancer risk associated with family history (23) as well as sporadic cancer incidence. Epidemiological evidence for a correlation of BER gene variants with cancer risk is abundant, albeit conflicting, as reviewed in (24, 25). Nonetheless, extracts from cells of head and neck cancer patients (26) and smokers with lung cancer have been found to exhibit reduced DNA repair efficiency (27) associated with the presence of BER protein variants and defective BER protein activity respectively.

The BER pathway system includes proteins with multiple enzymatic activities, cooperativity, and compensatory sub-pathways. This complexity makes it difficult to intuitively estimate how variation in a single repair protein will change a cell's overall ability to tolerate DNA base damage. Moreover, there is a high likelihood that individuals will possess multiple variations, which in combination can further increase or reduce pathway capacity. Consequently, we apply a mathematical model of human BER that integrates biological knowledge of enzyme mechanisms and biochemical data on enzyme kinetics and protein concentration obtained from the literature. This model has been used to interpret published data from *in vitro* pathway reconstitution and cell extracts to evaluate mechanistic hypotheses of enzyme cooperativity and coordination and predict the relative significance of the BER sub-pathways (28). Notably, the model recently predicted that under normal conditions, background oxidative DNA base damage level is

at the low end of the widely varying published measurements, i.e., up to hundreds of lesions per cell, and that the background level is relatively stable to small changes in enzyme kinetics (29). In the context of the normally robust operation of BER, we applied the model based on published experimental data to: (i) estimate the potential impact if any of the previously characterized BER variant proteins, (ii) identify the magnitude of changes in individual protein kinetics and concentration required to observe a substantial difference in overall BER kinetics and capacity, and (iii) use the model to predict the effect of simultaneous variation in multiple repair proteins on total BER kinetics.

#### **MODEL & METHODS**

BER Pathway Model Structure. The comprehensive BER pathway model is based on current biological knowledge, illustrated in Fig. 1. The BER pathway in the nucleus is initiated by a lesion-specific DNA glycosylase, which recognizes and removes damaged bases (30). For example, 8-oxoguanine (8-oxoG;  $y_1$  in Fig. 1) is removed by 8-oxoG DNA glycosylase (Ogg1; e<sub>1</sub>). Some DNA glycosylases (including Ogg1) also exhibit an AP lyase activity, incising 3' to the abasic (AP) residue, leading to a subpathway (Pathway A) in which the phosphodiesterase activity of human AP endonuclease (Ape1;  $e_2$ ) removes the 3' block ( $y_{10}$ ) (31, 32). In most cases, "short-patch" BER (Pathway B) occurs (33), where Ape1 incises 5' to the lesion (34). The resultant 5'-deoxyribose phosphate (dRp;  $y_4$ ) group is removed by the dRp lyase activity of DNA polymerase  $\beta$ (Pol $\beta$ ; e<sub>3</sub>), which then executes single base replacement at the gapped site (y<sub>5</sub> or y<sub>11</sub>). In our model, the Polß-catalyzed activities may occur in either order (thus the fork in Pathway B). An alternative "long-patch" BER pathway (Pathway C) exists in which gapfilling of 2-7 nucleotides is performed by Pol $\delta$  (e<sub>4</sub>) or possibly Pol $\epsilon$  (excluded from our model due to a lack of kinetic data) in complex with PCNA followed by flap excision by Fen1 ( $e_5$ ) and subsequent ligation (35). The final step in all sub-pathways is performed by DNA Ligase 1 (Lig1;  $e_6$ ), or Lig3 ( $e_7$ ) stabilized in complex with Xrcc1 (36). Lig1 and Lig3/Xrcc1 activities appear largely interchangeable (37), though a few recent results indicate that Lig1 may be more significant for long-patch repair (38).

**Model Assumptions.** In general, reactions in the model are modeled using Michaelis-Menten enzyme kinetics, with  $k_{cat}$  and  $K_M$  parameters defining a reaction velocity ( $v_i$  in Fig. 1). This implicitly assumes homogeneity and deterministic, continuous reaction kinetics, which are justified by the high concentration of repair proteins as discussed in (28). If Michaelis-Menten assumptions hold for a biochemical system, then  $k_{cat}$  is interpreted as the catalytic turnover rate of the enzyme (its "activity"), and  $K_M$  is a function of the affinity of the enzyme for the substrate.

To account for experimental measurement of 5 to 10-fold reduced diffusion velocities in the crowded cell nucleus (39),  $K_M$  of all reactions are increased 10-fold (due to the reduction in collision probability). The model includes experimentally measured protein cooperativity by  $k_{cat}$  increases as in previous modeling (28, 29), as well as including coordination of consecutive single enzyme activities (such as Pol $\beta$  and the dual-function DNA glycosylases, e.g., Ogg1) by modeling the second step as a first-order reaction, as described in (28). We also assume no product inhibition for individuals under normal conditions in the pathway, as suggested by observations (32, 40). We do not however include the potential role of Xrcc1 as a mediator of a multiprotein coordination

complex (41), as the contribution of Xrcc1 to such a complex forming and any consequent enhancement of BER kinetics are unclear (42, 43). In addition, many of the potential coordinative effects are already in the model. Table 1 shows kinetic parameters used in the model, including rate enhancements due to cooperativity.

Model Protein Concentrations. Table 2 shows BER protein concentrations determined from published data calculated by assuming the number of proteins measured per cell are homogenously distributed in a nucleus with a diameter of 5  $\mu$ m, as described in (28, 29). Because of the limited data currently available, these protein levels were drawn from various cell lines. Where multiple measurements were available, a conservative assumption was used, since some are from tumor cell lines, which have been measured to have higher protein expression levels than normal cells (76). Furthermore, Ape1 (79-80) and DNA glycosylases, such as Ogg1 (81), may be distributed in both the nucleus and other cell compartments (i.e., cytosol and mitochondria). In our analysis, we neglect the contribution to overall cellular DNA damage of the oxidative damage in mitochondria, which is repaired equally or more efficiently than nuclear DNA and does not occur at levels measurable in whole cell assays (82-84). Consequently, protein concentrations in Table 2 do not include Ogg1 and Apel localized to mitochondria (74, 81, 85). We have found in our analysis of the sensitivity of our results to protein concentration that it closely follows that of sensitivity to k<sub>cat</sub>, which is published here (results not shown; similar results were also found in our previous work on estimating steady-state BER capacity (29)). This is expected from the mathematical form of the Michaelis-Menten equations used to model most of the BER pathway.

**Solution of Model Equations.** The mathematical structure of the model is identical to that in (29). Differential equations were solved using the 'ode15s' stiff solver of MATLAB R14 using parameters from Tables 1 and 2, adjusted by the analysis described below. Steady-state reactant levels were the equilibrium point found by simulating the differential equations on a time scale of 24 hours and then setting equations for repair intermediate concentrations equal to zero and using the nonlinear least squares MATLAB routine 'fsolve' to solve the resulting algebraic system (results not shown).

#### RESULTS

**Simulation of 8-oxoguanine (8-oxoG) levels as an estimate for oxidative DNA base repair capacity.** We focus on the repair of 8-oxoguanine (8-oxoG) initiated by Ogg1, which includes the full set of pathway steps in Fig. 1. This is motivated by the large amount of data from experimental sources for the kinetics of wild type and polymorphic forms of Ogg1 (28), as well as cellular and physiological levels of 8-oxoG (e.g., (86); also reviewed in (29)). 8-oxoG is frequently used as marker for oxidative DNA damage associated with disease consequences (87). As further data accumulate for the kinetics and cellular abundance of other DNA glycosylases (88), we can extend our analysis in the future using the approach described herein. In our studies, we assume that the contribution of NEIL1/2 enzymes recently found to have 8-oxoG DNA glycosylase activity (89) are quantitatively insignificant as compared to Ogg1; recent studies have

suggested that their biological role may be primarily for specialized cases, such as damage within bubble DNA structures (90, 91).

We account for the production of other base lesions in our modeling by including an additional damage load of abasic (apurinic/apyrimidinic; AP) sites, comprising the products of other DNA glycosylases, as well as spontaneous hydrolysis and base loss, the majority of which result in apurinic sites (92). Overall, we predict the combined numbers of original lesions and repair intermediates, which are a significant component of observed lesions (93) and may be highly genotoxic and mutagenic themselves (94, 95).

Estimated impact of mild protein variants found in the population on BER kinetics and capacity. If a protein variant is found to affect the kinetics of a BER reaction, then its impact on the overall pathway is estimated by modifying the appropriate  $k_{cat}$  and  $K_M$  parameters for that reaction in the comprehensive BER system model. Published data are available for the kinetics of some BER proteins coded by sequence variants identified in population screens. Ape1 variants were characterized with AP endonuclease efficiencies from 35-110% of wild type (13). These are calculated based on the amount of repair of an initial lesion concentration over a fixed period of time, and they may be interpreted as being equivalent to the catalytic rate  $(k_{cat})$  of Ape1. The commonly found Ogg1 polymorphism Ser326Cys has a reported k<sub>cat</sub> 63% of wild type (15), and other Ogg1 variants were found to have a k<sub>cat</sub> 44% (Arg154His) and 85% (Arg36Gln) of wild type (14). In the case of Pol $\beta$ , one quantitatively characterized allelic variant is a dominant negative inhibitor of wild type activity (17). Based on the measured equilibrium binding affinity ( $K_D$ ) of the variant with the Pol $\beta$  substrate and potential cell concentration, the effective  $K_M$  for Pol $\beta$  gap-filling reaction is increased 3 to 5-fold (calculated based on competitive inhibition in the Michaelis-Menten model). The estimated impact of protein variants in these ranges of variation are shown in Fig. 2, where we have simulated the effect of Ape1 with 10% of normal  $k_{cat}$  (the highest amount at which any difference could be observed), Ogg1 with 50% of normal k<sub>cat</sub>, and Pol $\beta$  with a 5-fold increase in K<sub>M</sub>.

Fig. 2 illustrates two possible kinds of impact on the BER pathway: the capacity of BER to tolerate a constant rate of base lesion formation (8-oxoG and AP site formation combined), and the kinetics of BER repairing an instantaneous pulse of oxidative DNA damage (also consisting of both 8-oxoG and AP site lesions). Such a pulse simulates acute exposure: for example, a dose of radiation therapy, increased metabolism during intense exercise, or acute inflammation during an infection. We consider repair of transient damage pulses in the context of continuing background base lesion formation. Base lesion formation rates (2,000 8-oxoG/cell/day and 20,000 AP sites/cell/day) are based on experimental data for normal background rates of formation (92, 96, 97). They are adjusted upwards to reflect (i) increased 8-oxoG corresponding to exercise and other variations in metabolism (98) and (ii) AP sites produced by the action of additional DNA glycosylases processing other base lesions. Damage "clearance" is defined as the time at which the background steady state level of lesions corresponding to the constant damage formation rate is restored. Notably, as Fig. 2 shows, 8-oxoG repair rate is more sensitive to small changes in protein kinetics than the steady state repair capacity.

Table 3 summarizes the impact on BER capacity (steady state number of lesions per cell upon exposure to the formation of 2,000 8-oxoG and 20,000 AP sites/cell/day) and BER kinetics (time to clear 10,000 8-oxoG and 100,000 AP sites) of each protein in

the pathway with 10% and 50% of their normal (Table 1)  $k_{cat}$ . Results are given for each of the activities of multiple-function proteins separately. Data are provided as a percentage increase in steady state damage and clearance time with respect to normal protein kinetics. Table entries of "0%" represent a change less than 0.01% (results not shown). Notably, because of the dominance of Pathway B for our model of 8-oxoG repair, assuming normal Pol $\beta$  function, Pathway C proteins, Pol $\delta$  and Fen1, will have negligible impact on kinetics, as was found in our previous modeling analysis (28).

Sensitivity analysis of BER pathway capacity to hypothetical protein variants. In Fig. 2, curves showing an increased lesion burden and clearance time as a result of increased damaged load show a quantitatively small difference for mild kinetic variants of BER proteins. This suggests that the BER pathway is apparently robust to mild kinetic variation. Accordingly, we explored a wider range of protein kinetic parameters to identify levels at which variants may have a qualitatively observable impact on BER pathway capacity, as shown in Fig. 3 for k<sub>cat</sub> and K<sub>M</sub> (modified relative to the normal values in Table 1; not shown are variants for BER reactions that were found to have no effect on steady state damaged base levels, e.g., Fen1  $k_{cat}$  and  $K_{M}$ ). While the analysis in Fig. 3 is specifically for steady state damage levels given physiologically relevant levels of 8-oxoG and AP site formation, it qualitatively reflects the general sensitivity of the BER pathway to its individual components. Regions on the plots in Fig. 3 that show a rapidly changing curve represent parameter values for which BER capacity is highly sensitive and less robust to increases in damage level. Regions for which no curve is plotted at all represent parameter ranges of BER pathway kinetics that preclude a steady state equilibrium point. For these hypothetical kinetic variants, the BER pathway breaks down, with lesions and repair intermediates accumulating uncontrollably. For normal BER reaction kinetics, this breaking point is approximately 125,000 8oxoG/cell/day (29), a level that seems physiologically unrealistic. By comparison, in cell culture, 8-oxoG formation was measured at  $0.01/10^6$  bases per Gy (99), and 8-oxoG levels on the order of  $100/10^6$  have been measured in naked DNA exposed to highly oxidative agents or >5 Gy radiation (100). But, for some modeled changes in the  $k_{cat}$  and  $K_M$  of Ogg1, Ape1, Pol $\beta$ , and Lig1, it can be on the order of  $10^4$  8-oxoG/cell/day (data not shown) or less - for example, where the curves are not shown on Fig. 3. Such variants will result in a BER pathway incapable of tolerating even normal damage formation rates (i.e.,  $10^3$  8-oxoG/cell/day and  $10^4$  AP sites/cell/day) without accumulating unrepaired lesions.

Fig. 3 shows the sensitivity of repair capacity to changes in only Lig1 kinetic efficiency. The concentration and kinetics of Lig3 were assumed to be equivalent to those of Lig1. Consequently, simulated predictions for Lig3 are identical to those for Lig1, demonstrating redundancy in the DNA ligation step of the repair pathway. This agrees partially with experimental results in (37), which do suggest that in some cases one DNA ligase may be preferred to another. Due to the redundant role of Lig1 and Lig3 in our model, simulated data predict that BER capacity is sensitive to reduction in Lig1 kinetics to a certain point, after which further reduction no longer affects the pathway, as shown in Fig. 3.

**Observation of unexpected synergies for hypothetical multiple protein variants in the BER pathway.** In some cases, the complexity of individual protein kinetics within the context of the whole BER pathway leads to unexpected synergistic variation. To illustrate this, we estimate the consequence of simultaneous variation of Ape1 5'-incision activity and Pol $\beta$  gap-filling and 5'dRp lyase catalytic activities ( $k_{cat}$ ) on overall BER dynamics for the repair of a transient pulse of DNA damage, assuming that the Pol $\beta$  rate enhancement by Ape1 is unaffected. In the steady state analysis (not shown), there is only a 3.75 to 4 lesion/cell difference (including repair intermediates) between Pol $\beta$  variants with 200-fold reduced  $k_{cat}$  and 10-fold reduced  $k_{cat}$  (assuming constant rates of formation of 2,000 8-oxoG/cell/day and 20,000 AP sites/cell/day), with the exception of the situation with a 1000-fold Ape1  $k_{cat}$  reduction, in which case the number of steady state lesions differ by 27 per cell. This contrasts with the findings shown in Fig. 4 for transient repair dynamics. The time to fully clear a pulse of 8-oxoG and abasic lesions and repair intermediates for the Pol $\beta$ -inefficient pathway is at its maximum when Ape1  $k_{cat}$  is 70% of normal. As Fig. 4B shows, this point of respective Ape1 and Polb activity modification is a peak (darkest region). Repair efficiency improves (more lightly shaded regions of Fig. 4B) as Ape1 kinetics either increase or decrease, and the overall system shows less sensitivity to reduced Pol $\beta$  kinetics.

(par. break) Based on a study of the estimated levels of BER intermediates during the simulation, it appears that at that particular critical level of Ape1, there is a shifting of repair to Pathways A and C, which are slower than Pathway B – and sites being repaired through Pathway A are particularly sensitive to Pol $\beta$  kinetics following the first steps. Consequently, there is a persistence of repair intermediates and prolongation of the time to reach the normal steady state background. As this model analysis demonstrates, the complexity of multiple sub-pathways in the BER pathway can be a mechanism for unexpected synergies between protein variants, which are not merely additive results of reduced kinetic rates.

#### DISCUSSION

There have been numerous investigations of potential correlations between BER gene variants (e.g., SNPs, truncations, frameshifts) with levels of biomarkers, including DNA damage and mitotic delay, prevalence in tumor cells, and increased cancer susceptibility. To date, many of these studies have been inconclusive or contradictory. For instance, the commonly found Ser326Cys variant of Ogg1 has been found to be correlated epidemiologically with increased cancer risk, increased measurements of cellular DNA base damage, and reduced base repair kinetics in several studies and assays (101-108), while null or statistically insignificant correlations have been found in others (102, 109-112); also a recent extensive review (113). Applying a previously developed and tested comprehensive BER pathway model integrating reaction mechanisms and parameters obtained from published experimental data, we estimate that Ogg1 base excision k<sub>cat</sub> reduced by 50%, comparable to that of the Ogg1 Ser326Cys polymorphism (15), results in a decrease of approximately 5-10% in BER capacity, as defined by the steady state number of lesions per cell for a wide range of constant damage formation rates (as shown in Fig. 2). The transient kinetics of repairing a pulse of damage are somewhat more sensitive to mild kinetic variants, but effects are still minimal.

Overall, our modeling supports a complex role for genetics in tolerating exposure to genotoxic stresses in determining DNA base damage tolerance and cancer risk. Under normal and increased damage loads and rates (simulated based on experimental measurements) results favor the null or marginal experimental findings for the impact of Ogg1 Ser326Cys on cellular tolerance of oxidative damage and cancer susceptibility. Furthermore, Ape1 population variants characterized in (13) are estimated by our modeling to have a negligible effect on BER kinetics and capacity. These results are supported by a null finding for cancer susceptibility with the Ape1 variant Asp146Glu in pooled studies (108).

Our analysis (Fig. 3) reveals that further increases in individual repair protein kinetics have a minor effect on overall BER pathway kinetics, as suggested by the long evolutionary history of BER, conserved to a relatively high degree from *E. coli* through human cells (7). Of all its steps, the BER pathway is most sensitive to decreases in Ogg1 base excision, Ape1 5'-incision of the abasic site, and Pol $\beta$  gap-filling and 5'-dRp lyase reaction kinetics, with limited compensated sensitivity to DNA ligase activity. The high sensitivity to Ogg1 kinetics is supported by evidence that allelic loss (which reduces Ogg1 concentration and reaction velocity) is prevalent in tumor cells with reduced 8-oxoG repair activity (102, 109, 114, 115). Furthermore, while there is little evidence for Pol $\beta$  variants in cells of normal tissues in the human population, several variants with diminished activity have been found in tumor cell lines and are associated with reduced overall BER pathway capacity and a Pol $\beta$  mutator phenotype, in accord with estimates presented here (17, 18, 116-121). This suggests a potentially significant role for inherited Pol $\beta$  variants in increasing cellular susceptibility to DNA base damage and cancer risk, suggesting targets for further epidemiological studies.

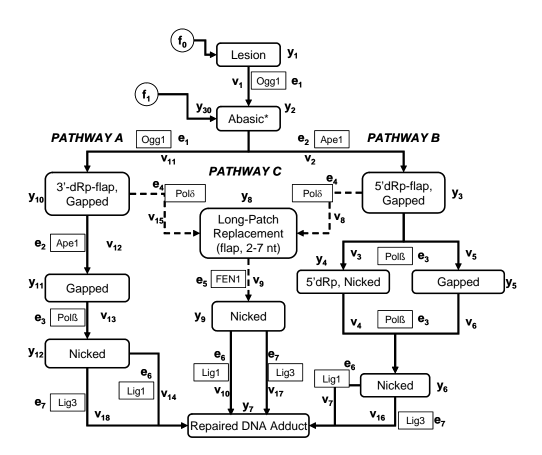
While the assumption of redundant Lig1 and Lig3 activities results in a modeling prediction of some compensation for the case of variants, there is still notable sensitivity to kinetics that would be exacerbated if activities are not completely overlapping. A Lig1 Arg771Trp variant has been characterized in an immunodeficient patient, with cell cultures demonstrating hypersensitivity to DNA damaging agents and impaired BER activity with 3-5% Lig1 kinetics (122), and a noncoding variant of Lig3 has been associated with increased esophageal cancer risk (123). Also, Lig3 activity requires Xrcc1, and there are reports associating Xrcc1 variants with increased cancer susceptibility (103, 123-127). However, other studies indicate null or statistically marginal results for Xrcc1 variants, highlighting the complexity of variant protein roles in the BER pathway (128-130).

While the association of most proteins involved in oxidative DNA base damage repair with cancer remains a controversial question, there is recent evidence of a direct link between protein sequence of MYH (the DNA glycosylase that removes a mismatched adenine across from the 8-oxoguanine lesion) and colorectal cancer (131-133), reviewed in (134). There is also some evidence that MYH variants associated with increased cancer risk and polyposis have reduced catalytic efficiency, but published studies have been on human MYH sequence variants generated by site-directed mutagenesis not found in the population (135) or mutants of the *E. coli* homolog corresponding to the population variants (136). As further enzymatic data are obtained for MYH activity, our comprehensive BER system model can be extended to include MYH along with Ogg1 and other DNA glycosylases, allowing for estimation of variant impact on the whole pathway.

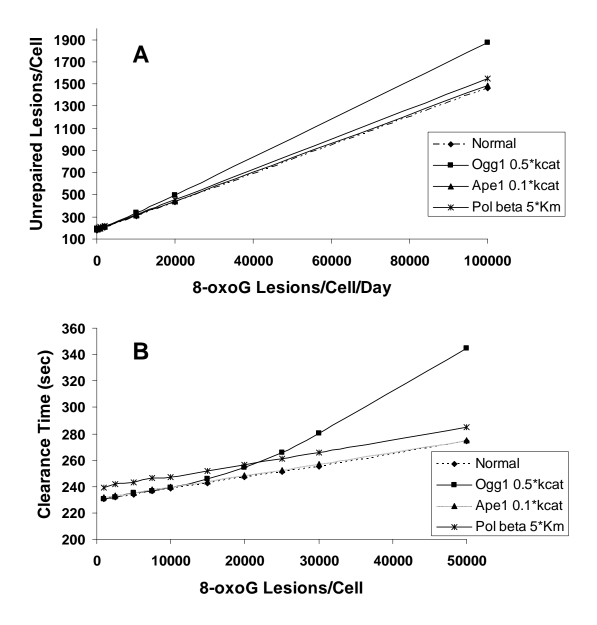
In the post-genome era, there has been tremendous investment in the search for genetic variants in the population related to increased cancer risk and chemoresistance,

particularly SNPs (137). The relative prevalence of SNPs in the population suggests that many are low-penetrance alleles, which will reduce the efficiency of biological pathways without eliminating their function entirely. As this study shows, biochemically modeling the impact of variants reveals complex behavior and pathway sensitivity, and can provide insight to interpret incomplete and contradictory results obtained from both epidemiology and laboratory experiments. Progress in understanding how complex genotypes can predict the response to endogenous background and exogenous environmental exposure to DNA damaging stresses will allow an individualized, predictive approach to interpreting the medical significance of DNA damage quantification (138), identify chemoprevention strategies for individuals with known susceptibility factors (139, 140), and unveil mechanisms of chemoresistance (141) to therapeutics. In general, the analysis presented here underscores the critical role of quantitative biological system modeling and experiments will play in the long-term goal of the post-genomic era, effectively interpreting complex genetic studies and translating their findings to benefit human health.

**Figure 1.** Schematic of the human BER pathway model, described in the main text, showing substrates and intermediates (y), enzymes (e), and reactions (v). (\*Abasic sites can be formed directly,  $y_{30}$ , or as intermediates of a bifunctional DNA glycosylase, e.g. Ogg1,  $y_{2.}$ )

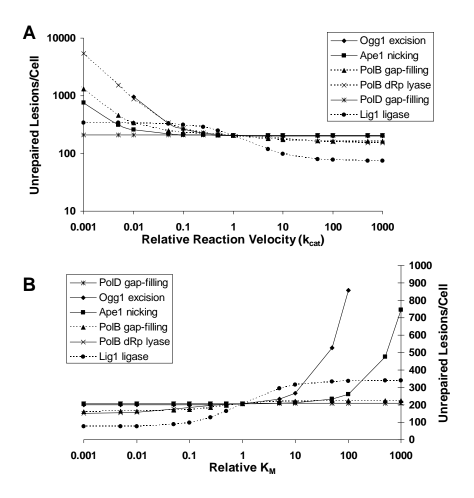


**Figure 2.** Sensitivity of 8-oxoG BER capacity and kinetics to mild kinetic defects in Ogg1 (50% of normal  $k_{cat}$ ), Ape1 (10% of  $k_{cat}$ ), and Pol $\beta$  (5-fold increased  $K_M$ ). (A) Total number of 8-oxoG, abasic, and repair intermediate bases per cell given increasing 8-oxoG rates of formation (assuming simultaneous production of an additional 20,000 AP sites/cell/day). (B) Time required to clear an instantaneous pulse of 100,000 AP sites/cell and an increasing number of 8-oxoG lesions/cell and restore damage to the background steady state level for continuous formation of 2,000 8-oxoG/cell/day and 20,000 AP sites/cell/day. Simulated data for the pathway with normal kinetics and with a 10%  $k_{cat}$  Ape1 variant are nearly overlapping.

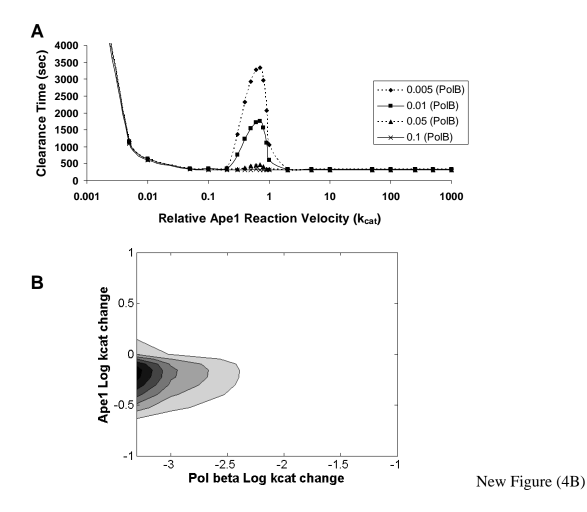


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**Figure 3.** Sensitivity over a broad range of  $k_{cat}$  (**A**) and  $K_M$  (**B**) of enzymatic activities in BER (Ogg1 excision of 8-oxoG, Ape1 nicking at the 5' end, Pol $\beta$  gap-filling and 5'-dRp lyase reactions, Pol $\delta$  gap-filling, and the ligase activity of Lig1. Changes in kinetic parameters are fractions and multiples of the values for those reactions in Table 1 (with all K<sub>M</sub> increased 10-fold as described in the text). The steady state total number of 8-oxoG, AP sites, and repair intermediates is calculated based on a constant total damage formation rate of 2,000 8-oxoG lesions/cell/day and 20,000 AP sites/cell/day. The plot is truncated at the point where a continued KM increase results in no physically possible steady state.



**Figure 4.** Simulation of the effect of varying Ape1 and Polb kcat simultaneously on the time required to clear 10,000 8-oxoG sites and 100,000 AP sites, with a constant background damage formation rate of 2000 8-oxoG/cell/day and 20,000 AP sites/cell/day. (A) Clearance time for varying Ape1  $k_{cat}$  (multiple of normal parameter value in Table 1) for fractions of normal Pol $\beta$   $k_{cat}$ . (B) Contour plot of clearance time for simultaneous changes (log ratio to normal  $k_{cat}$ ) in Ape1 and Pol $\beta$ ; darker regions correspond to higher clearance times.



|                      |          | K <sub>M</sub><br>(nM) ‡ | References          | Rate Enhancements   | References |  |
|----------------------|----------|--------------------------|---------------------|---|------------|--|
| 1                    | 0.0052   | 121.5                    | (14, 15, 32, 44-46) | 10x decr. K <sub>M</sub> (Ape1);<br>8x incr. k <sub>cat</sub> (Ape1;<br>posttranslational<br>modifications) | (32, 47)   |  |
| 2                    | 3.2028   | 34.7                     | (32, 48-55)         |   |            |  |
| 3, 6*                | 0.817    | 210                      | (56-59)             |   |            |  |
| 4*, 5                | 0.075    | 500                      | (60)                | 6x incr. k <sub>cat</sub> (Ape1)  | (61, 62)   |  |
| 7, 10, 14,<br>16-18† | 0.0213   | 56.7                     | (56, 63, 64)        | 4x incr. k <sub>cat</sub> (PCNA,<br>RpA)  | (64, 65)   |  |
| 8, 15                | 0.6      | 100                      | (66-68)             | (Rate in presence of PCNA)  |            |  |
| 9                    | 0.1335   | 39                       | (59, 69)            | 2x incr. k <sub>cat</sub> (Ape1)  | (70)       |  |
| 11                   | 0.000887 | 7.2                      | (32)                | 2.5x incr. k <sub>cat</sub> (Ape1;<br>posttranslational<br>modification)                                    | (47, 71)   |  |
| 12                   | 0.0608   | 169                      | (49, 72, 73)        |   |            |  |

# Table 1 Normal BER Protein Kinetics

\* Assuming pathway coordination, this reaction is first order with a rate of k<sub>cat</sub>.
† Lig3 kinetics assumed equal to those for Lig1 kinetics (37).
‡ Where multiple sources are used in the literature, the median is used to avoid excessive influence by outliers.

| Protein | Concentration<br>(nM) | References   |
|---------|-----------------------|--------------|
| Ogg1    | 406                   | (46, 74)     |
| Ape1    | 2000                  | (72, 74, 75) |
| Polβ    | 419                   | (76, 77)     |
| Polð    | 600                   | *            |
| Fen1    | 450                   | *            |
| Lig1    | 254                   | (78) †       |
| Lig3    | 254                   | *<br>*       |

 Table 2 BER Protein Concentrations

\* In absence of data, assumed based on concentration of other nuclear proteins.

† Also, A. E. Tomkinson, personal communication.

‡ Assumed equal to Lig1 (37).

| Protein | Activity         |               | 10% k <sub>cat</sub> |      | 50% k <sub>cat</sub> |      |
|---------|------------------|---------------|----------------------|------|----------------------|------|
|         | ( <b>vi</b> )    |               | Damage               | Time | Damage               | Time |
| Ogg1    | 1                | base excision | 29%                  | 341% | 3%                   | 9%   |
| Ape1    | 2                | 5'-incision   | 2%                   | 1%   | 0%                   | 0%   |
| Polβ    | 3, 6             | gap-filling   | 13%                  | 7%   | 4%                   | 2%   |
| Ροlβ    | 4,5              | 5'-dRp lyase  | 32%                  | 56%  | 4%                   | 1%   |
| Lig1†   | 7, 10, 14, 16-18 |               | 53%                  | 90%  | 21%                  | 34%  |
| Polð    | 8, 15            |               | 1%                   | 0%   | 0%                   | 0%   |
| Fen1    | 9                |               | 0%                   | 0%   | 0%                   | 0%   |
| Ogg1    | 11               | AP lyase      | 0%                   | 0%   | 0%                   | 0%   |
| Ape1    | 12               | 3'-dRp lyase  | 0%                   | 0%   | 0%                   | 0%   |

**Table 3** Predicted Impact of Protein Kinetic Variants on Background Damage Levels and

 Time to Clear Lesions (% of wild type)\*

\* Steady state number of lesions per cell is based on 2,000 8-oxoG and 20,000 AP sites/cell/day; time to clear is calculated for10,000 8-oxoG and 100,000 AP sites/cell. † Based on model assumptions, Lig3 will have the same results.

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