https://doi.org/10.1093/genetics/iyac092 Advance Access Publication Date: 10 June 2022

Investigation

Complex mutation profiles in mismatch repair and ribonucleotide reductase mutants reveal novel repair substrate specificity of MutS homolog (MSH) complexes

Natalie A. Lamb \bigcirc , 1 Jonathan E. Bard \bigcirc , ^{1,2} Raphael Loll-Krippleber \bigcirc ,³ Grant W. Brown \bigcirc ,³ Jennifer A. Surtees \bigcirc ^{1,4}*

¹Department of Biochemistry, Jacobs School of Medicine and Biomedical Sciences, State University of New York at Buffalo, Buffalo, NY 14203, USA,

2 University at Buffalo Genomics and Bioinformatics Core, State University of New York at Buffalo, Buffalo, NY 14203, USA,

³Department of Biochemistry and Donnelly Centre, University of Toronto, Toronto, ON M5S 3E1, Canada,

OXFORD GENETICS

⁴Genetics, Genomics and Bioinformatics Graduate Program, Jacobs School of Medicine and Biomedical Sciences, State University of New York at Buffalo, Buffalo, NY 14203, USA

*Corresponding author: Department of Biochemistry, Jacobs School of Medicine and Biomedical Sciences, University at Buffalo, State University of New York, Rm 4215, 955 Main Street, Buffalo, NY 14203, USA. Email: jsurtees@buffalo.edu

Abstract

Determining mutation signatures is standard for understanding the etiology of human tumors and informing cancer treatment. Multiple determinants of DNA replication fidelity prevent mutagenesis that leads to carcinogenesis, including the regulation of free deoxyribonucleoside triphosphate pools by ribonucleotide reductase and repair of replication errors by the mismatch repair system. We identified genetic interactions between rnr1 alleles that skew and/or elevate deoxyribonucleoside triphosphate levels and mismatch repair gene deletions. These defects indicate that the mr1 alleles lead to increased mutation loads that are normally acted upon by mismatch repair. We then utilized a targeted deep-sequencing approach to determine mutational profiles associated with mismatch repair pathway defects. By combining rnr1 and msh mutations to alter and/or increase deoxyribonucleoside triphosphate levels and alter the mutational load, we uncovered previously unreported specificities of Msh2–Msh3 and Msh2–Msh6. Msh2–Msh3 is uniquely able to direct the repair of G/C single-base deletions in GC runs, while Msh2–Msh6 specifically directs the repair of substitutions that occur at G/C dinucleotides. We also identified broader sequence contexts that influence variant profiles in different genetic backgrounds. Finally, we observed that the mutation profiles in double mutants were not necessarily an additive relationship of mutation profiles in single mutants. Our results have implications for interpreting mutation signatures from human tumors, particularly when mismatch repair is defective.

Keywords: mismatch repair; dNTP pools; ribonucleotide reductase; deep sequencing; mutation profiles; replication fidelity

Introduction

Cancer is a genetic disease caused by mutation accumulation; DNA replication is an important source of mutation. Replicative polymerases, Pole and Pold, minimize errors via a highly selective nucleotide-binding domain to prevent deoxynucleotide triphosphate (dNTP) misincorporation a[nd an exonuc](#page-17-0)l[ease domain to](#page-17-0) [proofread an](#page-17-0)[d to remove errors \(K](#page-18-0)unkel 2004; [McCulloch and](#page-17-0) Kunkel 2008; St Charles et al. 2015; Ganai and Johansson 2016). Together, these DNA polymerase functions provide a selectivity in which misincorp[oration errors occur a](#page-18-0)t [rates that are on the](#page-17-0) [order](#page-17-0) of 1×10^{-7} (St Charles *et al.* 2015; Ganai and Johansson 2016). The mismatch repair (MMR) pathway acts on replication errors in vivo to prod[uce significantly lower error rat](#page-17-0)e[s of approx](#page-17-0)[imately 2](#page-17-0) \times 10^{–9} (McCulloch and Kunkel 2008; Ganai and Johansson 2016). Appropriate levels and ratios of the 4 dNTPs are also essential for maintaining high fidelity polymerase function. This has been demonstrated in yeast using mutations in allosteric sites of RNR1 that alter dNTP pools in different ways. RNR1

encodes the large subunit of the enzyme that catalyzes the ratel[imiting step in dNTP synthe](#page-18-0)[sis, ribonucleotid](#page-17-0)e reductase (RNR) (Nordlund and Reichard 2006; Hofer et al. 2012). Even a modest 2 times increase above normal levels, as seen in rnr1D57N, increased nucleotide misi[ncorporation by DN](#page-16-0)[A polymerases](#page-18-0) and elevated mutation rates (Chabes et al. 2003; Xu et al. 2008). More pronounced and skewed elevations in dNTP pools are generated in rnr1Y285F and rnr1Y285A, which [increase dCTP and](#page-17-0) dTTP 3 times and 20-times, respectively (Kumar et al. 2010). These increa[ses in dNTP pool levels further compromise rep](#page-17-0)l[ication fi](#page-16-0)[delity \(Ku](#page-16-0)mar et al. [2010;](#page-18-0) Kumar, Abdulovic, et al. 2011; Buckland et al. 2014; Watt et al. 2016). Cancer cells have an increased proliferative nature and thus elevated dNTP [pools may be necessa](#page-16-0)r[y to](#page-18-0) [support a](#page-18-0)[ccelerated DNA](#page-17-0) [replication \(David](#page-16-0)son et al. 2012; Poli et al. 2012; Mathews 2015; Connor et al. 2017), which may, in turn, increase mutagenesis, promote molecular evolution, and provide a selective advantage to the tumor.

We previously developed a targeted deep-sequencing approach to characterize mutation profiles of these 3 rnr1 alleles

Received: February 14, 2022. Accepted: May 24, 2022

V^C The Author(s) 2022. Published by Oxford University Press on behalf of Genetics Society of America. All rights reserved.

For permissions, please email: journals.permissions@oup.com

(Lamb et al. 2021). We specifically selected for mutations within CAN1 and sequenced the mutant can1 genes at an average depth of 16,000 reads per nucleotide. The depth of sequencing allowed a more robust a[nd nuanced ana](#page-18-0)l[ysis of mutation spectra than in](#page-17-0) [previous work \(Xu](#page-16-0) et al. 2008; [Kuma](#page-18-0)r, Abdulovic, et al. 2011; Buckland et al. 2014; Watt et al. 2016). We revealed genotypespecific mutation profiles, including mutation spectra, sequence context, and nucleotide motifs, with even modest changes in dNTP pools. Notably, all 3 rnr1 alleles exhibited increased CG>TA mutations and a shift in the relative distribution of single-base deletions toward G/C de[letions, which are](#page-17-0) typically rare events in wild-type backgrounds (Lamb et al. 2021). The frequency of single-base G/C deletions was particularly elevated in rnr1Y285A. We suggested that the effects of altered dNTP pools on mutation profiles should be incor[porated in the an](#page-17-0)alysis of mutation signatures in human cancer (Lamb et al. 2021).

The variants we observed in the rnr1 backgrounds are typically substrates for MMR, which functions as a spell-chec[k, recogniz](#page-17-0)[ing and d](#page-17-0)irecting the repair of errors in replication (Kunkel and Erie 2015), thereby incre[asing the fidelity of replication by an ad](#page-17-0)[ditional 10–1,00](#page-17-0)0 times (McCulloch and Kunkel 2008; Ganai and Johansson 2016). We and others have demonstrated co[mbinato](#page-18-0)[rial e](#page-18-0)[ffects of](#page-17-0) rnr1 and mmr alleles o[n mutation rates \(Xu](#page-16-0) [et al.](#page-18-0) [2008;](#page-18-0) Kumar, Abdulovic, et al. 2011; Buckland et al. 2014; Watt et al. 2016), consistent with the prediction that mutations generated with altered rnr1 are substrates for MMR. Once recognized by an MutS homolog (Msh) complex, the misincorporation structures are targeted for excision. In most eukaryotes, 2 heterodimeric complexes bind errors at the replication fork: Msh2–Msh3 and Msh2–Msh6, which recognize a broad spectrum of mismatches and insertion/deletion loops (indels) with different but overlapping specificities. The current model of postreplicative MMR posits that Msh2–Msh3 recogni[zes, binds, and](#page-18-0) [directs repair](#page-17-0) [of in](#page-17-0)dels up to 17-nucleotide long (Sia et al. 1997; Jensen et al. 2005), while Msh2–Msh6 targets mismatches and indels of 1–2 nucleotides. Msh2–Msh3 also has affini[ty for some mismatches,](#page-17-0) [espec](#page-17-0)[ially C–C, A–A, and \(](#page-18-0)possibly) G–G (Harrington and Kolodner 2007; Srivatsan et al. 2014).

MMR is deficient in \sim 25% of [sporadic cancers caused by in](#page-17-0)creased rate of mutagenesis (Mastrocola and Heinen 2010). Deficiencies in MMR genes MSH2, MSH6, PMS2, and MLH1 cause hereditary Lynch syndrome, which leads to a strong predisposition to cancers of the gastrointestinal tract, endometrial cancer, [and lymphomas](#page-18-0), [and are def](#page-17-0)ined by microsatellite instability (Pino et al. 2009; Heinen 2010). M[MR mutations are also impli](#page-17-0)[cated](#page-17-0) in breast and ovarian cancer (Davies et al. 2017; Fusco et al. 2018). While Msh2–Msh3 has not been directly linked to Lynch syndrome, mutations in MSH3 lead to cance[r predisposition](#page-17-0) [withi](#page-17-0)[n and outside the ga](#page-18-0)[strointestinal trac](#page-16-0)[t \(Edelmann](#page-18-0) et al. 2000; [van Oers](#page-18-0) et al. 2014; [Adam](#page-18-0) et al. 2016; Morak et al. 2017; Santos et al. [2018;](#page-18-0) Valle et al. [2019\) as well as](#page-18-0) [chemoresistant](#page-18-0) [tumo](#page-18-0)rs (Takahashi et al. 2011; Park et al. 2013; Nogueira et al. 2018). Loss of MSH3 function leads to elevated alterations in selected tetranucleotide repeats, which is distinct from microsatellite instability and has been associated with a number of [different cancers, in](#page-16-0)cluding up to $~60\%$ of colorectal cancers (Carethers et al. 2015). Therefore, while MSH3 also plays a role in tumorigenesis, its role is distinct from that of Msh2–Msh6.

The multiplicative and synergistic effects of combining defects in both MMR and RNR1 [on m](#page-18-0)utation rates of $mr1D57N$ msh2 Δ , [rnr1D57N m](#page-16-0)sh6 Δ (Xu [et al.](#page-18-0) 2008), and rnr1Y285A msh2 Δ (Buckland et al. 2014; Watt et al. 2016) indicated a genetic interaction between these pathways. We predicted that additional pathways would interact genetically with rnr1 [alleles. We performed s](#page-16-0)ynthetic genetic array (SGA) screens (Baryshnikova et al. 2010a) to identify pathways that interact genetically with all 3 m r1 alleles, identifying a number of pathways involved in DNA metabolism. Most strikingly, in the rnr1Y285A SGA screen, we identified essentially the entire MMR pathway. Therefore, we focused on the characterization of rnr1–MMR genetic interactions at the [nucleo](#page-18-0)[tide l](#page-18-0)e[vel, using our targeted deep-se](#page-17-0)[quencing approach \(X](#page-16-0)u [et al.](#page-17-0) [2008;](#page-17-0) Kumar, Abdulovic et al. 2011; Buckland et al. 2014; Lamb et al. 2021) with an eye to developing a mechanistic understanding of mutation signatures that are observed in tumors. At the [same time, the al](#page-17-0)tered mutation profiles generated by mr1 alleles (Lamb et al. 2021) allowed us to evaluate the role of Msh2–Msh3 and Msh2–Msh6 in directing the repair of typically rare replication errors. We characterized single $msh\Delta$ mutants and evaluated combinatorial effects on the mutation profiles when combined with rnr1 alleles. We identified novel and specific DNA substrates for Msh2–Msh3- vs. Msh2–Msh6-mediated MMR and demonstrated that mutation profiles of $m r1$ msh Δ double mutants were not necessarily additive of the single mutant profiles, which has implications for the analysis of mutation signatures in human cancers.

Materials and methods Strains and plasmids

All strains used for sequencing and m[utator assays were derive](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)d from the W303 RAD5+ background (Supplementary Table 1). Strains [used for the synthetic leth](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)al screens were derived from S288C (Supplementary Table 1). Construction of rnr1D57N, rnr1Y285F and rnr1Y285A[, with and](#page-17-0) without pGAL-RNR1, was described previously (Lamb et al. 2021). MMR genes were deleted by amplifying msh2 Δ ::kanMX, msh6 Δ ::kanMX, and msh3 Δ ::kanMX chromosomal fragments from deletion c[ollection strains utilizing](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) primers A and D specific to each locus (Supplementary Table 2). PCR products from these strains were used for transformation to replace the endogenous MSH gene with the kanMX cassette, which confers resistance to the drug, G418. Transformants were selected on YPD plates containing G418 and deletions were confirmed by PCR. We did not generate msh3 Δ in the m1Y285F/ApGAL-RNR1 backgrounds.

Measuring mutation rates at CAN1

Mutation r[ates were mea](#page-18-0)[sured at the](#page-17-0) CAN1 locus as previously described (Xu et al. 2008; Lamb et al. 2021). Briefly, strains were grown on complete media (YPD) until colonies reach 2 mm in size. Colonies were then suspended in 100 μ l of 1 \times TE (10 mM Tris-HCl, pH 7.5; 1 mM EDTA) and diluted 1:10,000. Twenty microliters of the undiluted colony suspension was plated on SC-ARG + canavanine and 100 μ l of the 10⁻⁴ dilution was plated on synthetic complete plates lacking arginine. The plates were incubated at 30°C until colonies reached \sim 1 mm in size. Colonies were counted and mutation rates and 95% confidence intervals [were calculated thou](#page-18-0)gh FluCalc fluctuation analysis software (Radchenko et al. 2018). Assays were performed on multiple, independent isolates for each genotype on separate days. For $mr1Y285A$ msh Δ genotypes, we only tested isolates that retained the slow growth phenotype.

Synthetic genetic array analysis

[The genetic screens w](#page-16-0)ere performed using SGA technology (Baryshnikova et al. 2010b). Briefly, query strains carrying each of the rnr1 alleles were crossed to an ordered array of all the viable

yeast deletion strains and an array of temperature sensitive alleles of yeast essential genes. Diploid cells were transferred to a sporulation-inducing medium, after which the germinated spores were selected for the simultaneous presence of the gene deletion and the rnr1 allele. Colony size was quantified as a measure of fitne[ss, and SGA scores and](#page-16-0) P-values were calculated as described in Baryshnikova et al. (2010b). SGA scores from deletion mutants and ts mutants were merged by scaling the ts screen scores according to the SGA scores of the 186 del[etion mutants that](#page-16-0) are present in the ts allele array, as described (Costanzo et al. 2016). A z-score was calculated for all the genes in each screen, and a cutoff of $z = -2$ was applied to identify negative genetic interactions [\(Supplementary Table 3\).](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) The raw SGA data are presented in Supplementary Tables 4–9. Negative genetic interactions identified in each screen, and the overlaps between [and among the](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) [negative](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) genetic interactions, are presented in Supplementary Table 10.

Gene ontology enrichment analysis

Gene ontology ([GO\) term analysis was p](http://go.princeton.edu/)erformed using the GO term finder tool (http://go.princeton.edu/) using a P-value cutoff of 0.01 and applying Bonferroni correction, querying biological process enrichment for each gene set. G[O term enrichm](#page-18-0)ent results were further processed with REViGO (Supek et al. 2011) using the "Medium (0.7)" term similarity filter and simRel score as the semantic similarity measure. Terms with a frequency of greater than 15% in the REViGO output were eliminated as too general.

Spatial analysis of functional enrichment

Network annotations were made with the Python implementa[tion of Spatial Anal](#page-16-0)ys[is of Functional Enrichment \(SAFE\)](https://github.com/baryshnikova-lab/safepy) [\(Barysh](https://github.com/baryshnikova-lab/safepy)nikova 2016; https://github.com/baryshnikova-lab/ safepy). The yeast genetic interaction similarity network and i[ts functional dom](#page-16-0)ain annotations were obtained from (Costanzo et al. 2016).

Sample preparation and analysis pipeline

A detailed description of sample preparation a[nd the analytica](#page-17-0)l pipeline used for data analysis can be found in (Lamb et al. 2021). Briefly, we pooled \sim 2,000 colonies Can^R colonies for each biological replicate and extracted genomic DNA from the pool. Each genotype w[as represented by at lea](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)st 4 independent biological replicates (Supplementary Table 11). The CAN1 gene was amplified by PCR using KAPA HiFi (Roche) in 6 overlapping fragments that were purified using the Zymo ZR-96 DNA Clean-up Kit. Nextera barcode adaptors were added to the amplicons, followed by attachment of Illumina Nextera XT index primers set A (Illumina). Excess adapters were removed using Ampure XP beads (Beckman Coulter). Pooled samples were diluted to 4 nM, denatured using NaOH and loaded onto an Illumina MiSeq sequencing platform (PE300, V3) with 20% PhiX control in 2 separate runs to increase coverage and as a check for reproducibility. Paired end (2 \times 300) deep sequencing of CAN1 provided enough sequencing depth to determine mutation spectra for 150 unique samples, representing over 30 different genotypes, and including biological and technical replicates. CAN1 was sequenced at an average depth of approximately 16,000 reads per base in CAN1 per sample allowing for detailed characterization of mutation spectra.

Sequence reads were trimmed (CutAdapt version 1.14), specifying a quality score of Q30, and then processed using CLC Genomics Workbench Version 11. Paired-end reads were merged, primer locations were trimmed, and processed reads were aligned to the SacCer3 reference genome. CLC low frequency variant caller was used to call variants, with required significance of 0.01%. Variant files were exported from CLC as VCF files and downstream analysis was performed in RStudio (version 1.2.1335), paired with custom python [scripting. Total variant](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) frequency for each genotype is shown in [Supplementary Fig. 1.](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) All sequence variants are provided in Supplementary Table 24. Variants in wild-type, rnr1D57N, rnr1Y285F, rnr1Y285F pGAL-RNR1, rnr1[Y285A](#page-17-0) and rnr1Y285A pGAL-RNR1 were characterized previously (Lamb et al. 2021) and are used here to compare single and double mutants.

Permissive variant filter

To assess the validity of our pipeline we included permissive samples grown in the absence of canavanine selection. We determined total variant frequency, by taking the number of variants that occurred in a sample and dividing by the number of reads. The permissive samples had an approximately 10 times lower total variant frequency, 8.7% com[pared to the samples s](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)elected on Canavanine, on average 99.4% (Supplementary Fig. 1). We used the permissive samples to inform a filtering approach to remove false positives from our data set. We developed the permissive filter to remove any variants that occurred below the average permissive variant allele frequency of 0.109%, as well as position specific mutations that occurred at a frequency above [this cutoff.](#page-17-0) [A de](#page-17-0)tailed description of this filter can be found in (Lamb et al. 2021).

Cosine similarity analysis

[We performed cosine](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) similarity among all genotypes (Fig. 2a and Supplementary Fig. 2) to assess the relationship between genotype and mutation profile, using the lsa package in R. First, we [performed this analy](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)sis [across all samples in](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) this study (Supplementary Fig. 2 and Supplementary Table 13). We generated a data matrix that contained all the unique variants within a sample and then determined cosine similarity scores based on the presence or absence of the variant within a sample. We then used the Heatmap2 package in R to show cosine similarity scores. Biological replicates from strains with lower mutation rates (e.g. wild type, msh3 Δ , mr1D57N) had lower cosine scores than those from strains with higher mutation rates (e.g. $msh2\Delta$, $msh6\Delta$, $mr1Y285A$), indicating that mutation events occurred more systematically (less stochastically) in these genetic backgrounds.

We then combined biological replicates and performed cosine [similari](#page-6-0)ty [analysis on the variant](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) counts within a genotype (Fig. 2a and Supplementary Table 14). The number of times a variant occurred in all the biological replicates within a genotype was scored. The different classes of variants include 6 classes of single-nucleotide variants (SNVs), single-base A/T or G/C insertions and deletions, complex insertions and deletions, as well as mononucleotide variants (MNVs) and replacements (Replac.). MNVs are dinucleotide SNVs, where 2 neighboring nucleotides are both mutated, ex: CC> AT. Replacements are complex insertions or deletions, where the deleted or replaced base is a variant. Two examples include $AAC > G$ and $C > AT$. Both MNVs and replacements are extremely low frequency events and rarely occur in our data set; neither had a significant impact on clustering.

The data were condensed based on genotype by combining biological replicates and adding the total number of times a variant was seen in the total number of replicates. This analysis does not take frequency into account and instead totals how many

unique types of variants occur in a sample. If the same variant occurred in multiple biological replicates within a genotype it was counted as such. A data matrix with the number of times a variant occurred in biological replicates for a particular genotype was generated. We then performed cosine similarity analysis on this matrix and displayed these data using the Heatmap2 package in R.

Principal component analysis

Principal component analysis (PCA) was performed on the number of times a unique variant was observed within a genotype. PCA was plotted using the "factoextra" package in RStudio, with a singular value decomposition (SVD) approach using the prcomp() function.

Determining SNV in trinucleotide context

A 3-bp window surrounding each CAN1 SNV was identified. The average number of each SNV [within this conte](#page-17-0)xt, a total of 96 possible trinucleotide contexts (Lamb et al. 2021), was calculated for each genotype a[nd divided by the num](#page-16-0)ber of times that context occurs in CAN1 (Alexandrov et al. 2015). The number of trinucleotide sequence contexts in CAN1 was calculated using a sliding window approach utilizing python scripting. For each of the 96 different SNV changes in triplet context, the average number of SNVs in a genotype was divided by the number of times the triplet sequence context occurs in CAN1. This dataset was imported into R-studio and plotted via the barplot() function. We also performed cosine similarity analysis on the normalized fre[quencies of SNVs in tr](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)inucleotide context in each genotype (Supplementary Table 22).

Cluster analysis for genotype-specific correlations

To identify variants specific to a particular genotype and to eliminate frequency bias from variants that occurred early in the growth of a particular sample, we condensed unique variants based on the number of biological replicates sequenced for that genotype. If a particular variant occurred in 4 out of 4 biological replicates it was represented as 1, if it occurred in 3 out of 6 replicates it was represented as 0.5. This gives an unbiased approach to score for the probability that a particular variant was present in a particular genotype, without considering the absolute frequency at which that variant occurred. These data were clustered on rows (or unique variants), after applying a row sum cutoff of greater than 2. This cutoff eliminates low frequency variants which are less likely to be driving the differences in mutation spectra that we observe. By clustering the data only on unique variants, it allows us to see groups of different types of variants in specific sequence contexts that are potentially diagnostic for a particular genotype.

To infer variants that were enriched in a particular genotype we divided the probabilities into 4 different bins (0–0.25, 0.26– 0.50, 0.51–0.75, and 0.76–1.0). A variant was positively enriched in a genotype if it occurred at 0.76 or greater probability, and negatively enriched if it occurred below a 0.25 probability. Variants were grouped for motif enrichment (black boxes) based on the main branches in the dendrogram, paired with similar patches of enrichment on the heatmap. It is worth noting the majority of variants that were negatively enriched did not occur at all in a given genotype (light blue on heatmap). On average there was greater than a 2 times increase in probability for variants that were positively enriched.

CC dinucleotide cluster analysis

A python script using a sliding window approach was used to identify all reference positions containing CC dinucleotides within CAN1. Our dataset was then subset to include only variants that occurred in these dinucleotide CC positions. Of the 138 CC dinucleotide contexts across CAN1, 110 $(\sim80\%)$ were mutated, compared to 857/1,711 base pairs or \sim 50% of the base pairs in CAN1. Unique CC run variants were clustered based on the number of times that variant occurred in each genotype, while accounting for (normalizing by) the number of biological replicates sequenced for each genotype, as described above. Heatmaps were plotted using the pheatmap package in RStudio a[nd motif enrichm](#page-16-0)ent was performed using Berkeley web logos (Crooks et al. 2004).

COSMIC single-base substitution signature cluster analysis

We performed cosine similarity analysis to compare the mutation spectra from our study with human mutation signatures, through an unbiased approach. The single-base substitutions (SBS) COSMIC signatures from GRCh38 (v3.2, March 2021, https:// cancer.sanger.ac.uk/signatures/sbs/) were [combin](#page-9-0)e[d with the](#page-16-0) [normalize](#page-16-0)d SNVs in trinucleotide context (Fig. 4) (Alexandrov et al. 2020). Cosine similarity scores were calculated using the lsa packa[ge in R and data was visu](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)alized as a heatmap, as described above (Supplementary Table 23).

Results

Pathways in DNA metabolism, including MMR, interact gen[etically with](#page-18-0) rnr1 [alleles](#page-16-0)

[Previ](#page-18-0)ous work (Xu et al. 2008; Buckland et al. 2014; Watt et al. 2016) demonstrating the combined effect of rnr1 alleles and msh deletions on mutation rates suggested that together these genes contribute to increased replication fidelity, and therefore they might show synergistic effects on cell fitness. We predicted that other pathways also interact genetically with rnr1 alleles that altered dNTP pools. Therefore, we performed synthetic genetic [array](#page-4-0) (SGA) [analysis using each of 3 dif](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)ferent rnr1 alleles (Fig. 1 and Supplementary Tables 3–10). The rnr1Y285A query had the greatest number of genetic interactions, consistent with the Y285A mutation having the highest increase in dNTP levels. The 3 rnr1 alleles showe[d surpris](#page-4-0)ingly [little overlap in their](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) genetic interactions (Fig. 1a and Supplementary Table 10), supporting the idea that different dNTP levels and pool balances stress cells in different ways. Both the rnr1Y285A screen and the rnr1D57N [screen](#page-4-0) showed enrichment for the GO term "DNA replication" (Fig. 1b) in addition to displaying unique enrichments for "maintenance of DNA repeat elements" and "DNA repair" (rnr1Y285A) and "lagging strand replication" (rnr1D57N). The rnr1Y285F screen did not show any statistically supported GO term enrichment and interacted with only a single "DNA replication" gene, POL31. To further assess the functional propertie[s of each](#page-16-0) rnr1 [allele](#page-16-0)'s genetic interactions, we applied SAFE (Baryshnikova 2016) to determine if any regions of [the functional geneti](#page-16-0)c interaction similarity yeast cell map (Costanzo et al. 2016) are [overrep](#page-4-0)resented for the negative genetic interaction gene sets (Fig. 1c). We found a statistically supported overrepresentation of the negative interacting genes in the DNA replication and repair neighborhood of the genetic interaction cell map for all 3 rnr1 alleles, indicating that dNTP pool alterations impinge

Fig. 1. Genome-scale screens for synthetic fitness defects with mr1 alleles. a) The overlap of the mr1-negative genetic interactions for the 3 SGA screens is plotted as a Venn diagram. The number of genes identified in each screen is indicated, as is the mr1 allele for each screen. b) GO-term enrichments for the negative interacting genes from each mr1 screen are tabulated. The enrichment for each term is indicated. Note that $mr1Y285F$ did not display any statistically supported enrichment. [c\) SAFE. On the left,](#page-16-0) the yeast genetic interaction similarity network is annotated with GO biological process terms to identify major functional domains (Costanzo et al. 2016). Thirteen of the 17 domains are labeled and delineated by colored outlines. On the right, the network is annotated with negative genetic interactions from each rnr1 SGA screen. The overlays indicate the functional domains annotated on the left. Only nodes with statistically supported enrichments (SAFE score > 0.08, P < 0.05) are shown. d) Tetrad analysis of m 1 × msh6 Δ crosses. Ten tetrads were dissected for each cross, and colonies were imaged after 3 days. Each column of 4 colonies is the 4 spores from a single meiotic ascus. Genotypes are indicated by circles (msh6∆) and squares (mr1). e) Tetrad analysis of mr1Y285A × msh2∆ and mr1Y285A × msh3∆ crosses. Genotypes are indicated by circles (msh2 Δ or msh3 Δ) and squares (mr1Y285A).

most dramatically on the DNA replication and DNA repair capacity of the cell. As with GO term enrichment, differences among the 3 rnr1 alleles were also apparent in the SAFE analysis, with rnr1Y285A and rnr1D57N interactors being overrepresented in the chromatin organization neighborhood compared with rnr1Y285F, and rnr1Y285F and rnr1Y285A interactors

showing more overrepresentation in the mitosis neighborhood than did rnr1D57N interactors.

Most notably, we found that almost the entire MMR pathway (MSH2, MSH6, MLH1, PMS1, and EXO1) was identified specifically in the rnr1Y285A screen and the "MMR" GO term was very strongly and specifically enriched in the rnr1Y285A screen

(Fig. 1b). This indicates a strong requirement for the MMR pathway when dNTP pools are highly unbalanced. We validated the specificity of the genetic interaction between MMR and rnr1Y285A by performing t[he tetra](#page-4-0)d analysis of msh6 Δ crosses with each of the 3 rnr1 alleles (Fig. 1d). Fitness defects in double mutant colonies were only evident in the rnr1Y285A cross. In tetrad analysis, msh6 Δ , and particularly msh2 Δ , showed stronger [fitness](#page-4-0) defects when combined with $mr1Y285A$ than did msh3 Δ (Fig. 1e), consistent with the reduced viability observed in mutation rate experiments (Table 1) [and with](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) msh3 Δ being just below our cutoff in the rnr1Y285A SGA (Supplementary Table 3). Our genetic interaction data indicate that all 3 mr1 alleles interface with DNA replication and repair pathways and that rnr1Y285A might be expected to have a particularly dramatic effect in MMR-deficient cells.

Combining $rnr1$ and msh \triangle alleles significantly increased mutation rates

The striking MMR interaction with rnr1Y285A, along [with the](#page-18-0) [comb](#page-18-0)inatorial effects of mr1D57N with msh2 Δ or msh6 Δ (Xu et al. 2008), encouraged us to focus on the combined effects of $msh\Delta$ and rnr1 alleles on mutation profiles. To determine whether the rnr1 msh double mutants displayed the anticipated increases in mutagenesis, we measured the forward mutation rates at CAN1 (Table 1). It is worth noting that it has been demonstrated previously that $msh2\Delta$ does [not alter dNTP pools, alone or](#page-17-0) i[n combina](#page-18-0)[tion](#page-18-0) with rnr1Y285A (Kumar, Abdulovic, et al. 2011; Watt et al. 2016). The relative mutation rates for msh [deletions wer](#page-17-0)[e consis](#page-18-0)[tent](#page-18-0) [with previous observations \(Marsischky](#page-17-0) et al. 1996; Xu et al. 2008; Kumar, Abdulovic, et al. 2011; Lang et al. 2013), with msh2 Δ exhibiting the highest mutation rate, msh6 Δ exhibiting a lower but still increased mutation rate, and $msh3\Delta$ exhibiting a mild mutator effect in this assay (Table 1). The mr1 muta[tion rates in](#page-17-0)[creas](#page-17-0)ed with increasing dNTP levels as expected (Kumar et al. 2010), with rnr1Y285A showing a 20 times increase above the wild-type mutation rate (Table 1). When mr1D57N and mr1Y285F were combined with msh deletions, the mutation rates exhibited the same hierarchy ($msh2\Delta > msh6\Delta > msh3\Delta$), but with higher rates than either of the relevant single mutants (Table 1). In most cases, the mutation rate of the double mutant approximated the product of the single mutant mutation rates. The rnr1Y285F $msh6\Delta$ strain had a particularly large increase, 85 times above wild type, compared to the expected 19 times increase given the mutation rates of the respective single mutants (Table 1). Notably, mr1Y285A msh3 Δ exhibited a strong, synergistic mutator phenotype, indicating that the mutations generated in the presence of rnr1Y285A are substrates for Msh2–Msh3-mediated MMR. It is possible that the mutation rates for $mr1Y285A$ msh2 Δ and $mr1Y285A$ msh6 Δ are underestimates, due to [fitness](#page-4-0) defects in these strains, consistent wit[h our SGA resu](#page-18-0)lts (Fig. 1) and previously published fitness data (Watt et al. 2016).

MMR genotype-specific variant profiles with msh3 Δ genotypes separated from msh2 Δ , msh6 Δ genotypes

We applied targeted deep sequencing of CAN1 (Lamb et al. 2021) to define and to characterize mutation profiles in m r1 msh Δ backgrounds, to derive mutation profiles from first principles.

We first selected ${\sim}2{,}000$ canavanine resistant (Can^R) colonies (selected samples) for each different genetic backgrounds. By definition, each of these colonies should have sustained a mutation in CAN1 that inactivates its gene product and allows cells to grow in the presence of canavani[ne. Multiple biological rep](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)licates for each strain were sequenced (Supplementary Table 11). The Can^R colonies were pooled and genomic DNA was extracted. The CAN1 gene was then amplified in 6 overlapping fragments[, bar-coded,](#page-17-0) [and](#page-17-0) subjected to Illumina paired-end sequencing (Lamb et al. 2021). To assess the effectiveness of our selection method and gain insight into baseline mutations, we calculated total variant frequency for each genotype, by summing the variants in each replicate/genotype and dividing by total sequencing reads. When strains were selected in the presence of cana[vanine, we observed](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) [an](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) average total variant frequency of 99.4% (Supplementary Fig. 1a). In contrast, strains grown in the absence of selection (permissive samples) exhibited an average total variant frequency of 8.7%, likely a result of a combination of low-level CAN1 mutations and PCR/sequencing artifacts. To address the latter, we applied a permissive variant filter that removed any variants at a frequency of below 0.109%. This cutoff is conservative and is

Fig. 2. Distinct genotypes share [unique features. a\) Cosin](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)e similarity analysis was performed on the unique counts of the 14 different classes of variants that occurred in each genotype (Supplementary Table 14). The histogram shows the distribution of cosine similarity scores across samples. b) Principal component analysis performed on all unique variants from biological replicates within a genotype.

based on the average frequency of individual variants in permissive samples (grown in t[he absence of se](#page-17-0)l[ection\) and the position](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)s in which they occurred (Lamb et al. 2021) (Supplementary Fig. 1b). To minimize false positives, it is likely that this approach will have eliminated some legitimate low frequency variants.

This targeted deep-sequencing approach identified primarily base substitution and small insertion/deleti[on mutations; rear](#page-17-0)rangements will not be easily identified (Lamb et al. 2021). Furthermore, there is a bias toward mutations that inactivate the CAN1 gene product, arginine permease, although we did observe about 3% synonymous mutations (data not shown). Nonetheless, the depth of sequencing (~16,000 \times coverage at each nucleotide) [and larg](#page-10-0)e number of total variants observed in each genotype (Table 2) revealed novel insights into DNA substrate preferences of Msh2–Msh3- or Msh2–Msh6-mediated MMR.

[We ch](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)aracterized mutation eve[nts in 2 ways \(Su](#page-17-0)pplementary Table 12), as previously described (Lamb et al. 2021). First, we determined the number of a specific variant type, i.e. the number of C>A chan[ges, at different positions](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) along can1 ("unique counts") (counts in Supplementary Table 12). Second, we calculated the frequency at which each of these unique variants occurred, i.e. the combined frequency of all C>A ch[anges at any position alon](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)g can1 ("sum of frequencies") (freq. in Supplementary Table 12). These analyses allowed us to determine whether different types of mutations occurred at specific locations in a genotypedependent manner, independent of frequency, and whether variant frequencies [were altered in a signifi](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)cant way by genotype (counts/freq. in Supplementary Table 12). A decreased number for "unique counts" combined with unchanged or increased "sum of frequencies" would indicate that variant type is more localized, possibly indicating a mutational hotspot. For instance, $msh6\Delta$ exhibited the highest proportion of unique events [contributing to](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) [the muta](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)tion spectrum (counts/freq. $= 1.62$; Supplementary Table 12). In contrast, $mr1Y285F$ msh3 Δ and $mr1Y285A$ msh3 Δ exhibited the lowest proportion of unique variants; the mutation spectra were instead dominate[d by G/C single-base de](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)letions, which occur at high frequencies (Supplementary Table 12).

We performed cosine similarity analysis to determine the quantitative relationship between mutation profile and genotype (see Materials and methods). All unique variants for all genotypes were assessed in parallel, based on both the presence and frequency of unique variants, as described above. In general, biological replicates of the same genotype had high cosine similarity scores and cluster[ed because their mut](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)ationa[l profiles were](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) [highly co](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)rrelated (Supplementary Fig. 2 and Supplementary Table 13). Therefore, we combined variants from all biological replicates within a genotype for the remainder of our analyses. Cosine similarity analysis performed on unique variants between geno[types w](#page-6-0)[as consistent with ge](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)noty[pe-specific mutation pro](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)files (Fig. 2a, Supplementary Fig. 2 and S[uppleme](#page-6-0)ntary Table 14), as was PCA based on unique variants (Fig. 2b). Most notably, msh2 Δ and msh6 Δ backgrounds segregated away from msh3 Δ backgrounds. Combined, these results indicated that it is possible to distinguish among genotypes based on unique variant profiles observed from can1 deep sequencing.

Previous work characterized $mr1Y285F$ and $mr1Y285A$ (\pm MSH2) in backgrounds that carried a silenced [pGAL-RNR1](#page-17-0) gene [\(i.e. grown in the](#page-16-0) absence of galactose) (Kumar et al. 2010; Buckland et al. 2014). We previously demonstrated that the mutation profiles of these strains were comparable to [rnr1Y285F](#page-17-0) and rnr1Y285A without the pGAL-RNR1 construct (Lamb et al. 2021). They are included here, with and without MSH genes to provide a comparison with that previous work. In ge[neral,](#page-5-0) pGAL-RNR1 did [not impact the observe](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)d mutation rates (Table 1) or profiles (Supplementary Fig[. 2–4\), although there](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) were some differences in specific variants (Supplementary Fig. 4).

Targeted deep sequencing revealed novel substrate preferences for Msh2–Msh3- vs. Msh2– Msh6-mediated MMR

Previous studies of mutation [spectra in the a](#page-17-0)[bsence of MMR have](#page-16-0) [focused primarily](#page-18-0) on msh2 Δ [\(Lang](#page-18-0) et al. 2013; Buckland et al. 2014; Serero et al. 2014; Watt et al. 2016), which effectively eliminates all M[MR activity and retain](#page-17-0)s errors made by replicative polymerases (Kunkel and Erie 2005). In contrast, studies of mutation spectra in msh3 Δ and msh6 Δ [analyze](#page-18-0)[d fewer than 100 colonies via](#page-17-0) [Sanger sequenc](#page-18-0)ing (Sia et al. 1997; Harrington and Kolodner 2007; Xu et al. 2008), preventing statistically supported conclusions about variant types. We examined mutation profiles in the absence of MSH2 ($msh2\Delta$). We also removed only Msh2-Msh6 (msh6 Δ) or Msh2–Msh3 (msh3 Δ) to assess the substrate specificity and repair efficiency of each complex. In general, the $msh2\Delta$, msh3 Δ , and msh6 Δ mutation profiles in the current study were consistent with previous studies performed in yeast: $msh2\Delta$ and $msh3\Delta$ profiles were dominated by indels, particularly deletions in rep[etitive region](#page-18-0)s, while msh6 Δ [profiles were](#page-17-0) [dominated by](#page-18-0) [SNVs \(Sia](#page-17-0) et al. 1997; Harring[ton and Kolodne](#page-17-0)[r 2007;](#page-16-0) Xu et al. 2008; [Kumar, Piacente](#page-18-0), et al. 2011; Lang et al. 2013; Buckland et al. 2014; Serero et al. 2014). We did not, however, observe the same bias toward tr[ansitions vs. trans](#page-17-0)v[ersions previously](#page-18-0) o[bserved in WGS o](#page-18-0)f msh2 Δ (Lang et al. 2013; Serero et al. 2014; Watt et al. 2016). Nonetheless, the substantially increased sample number and sequencing depth revealed some novel specificities. Deleting MSH2 resulted in a relative decrease in SN[Vs and incr](#page-8-0)[eased deletions](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) [and in](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)sertions compared to wild type (Fig. 3a and Supplementary Fig. 4), with A/[T single-base deletions \(A/](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)[T-1\) dom](#page-8-0)inating the mu[tation profile \(Supple](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)mentary Table 12, Fig. 3c, green bar, and Supplementary Fig. 4). In contrast, the $msh6\Delta$ spectrum was dom[inated by SNVs, which](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) [represented](#page-8-0) >80% [of the mutations](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) [\(S](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)upplementary Table 12, Fig. 3, a and b, and Supplementary Fig. 4), although the types [of SNVs gener](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)[ated we](#page-8-0)re si[milar for](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) $msh2\Delta$ [and](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) msh6 Δ (Supplementary Table 12, Fig. 3b, and Supplementary Fig. 4). In msh3 Δ cells, deletion events prevailed, similar to msh2 Δ , but there was a marked increase in G/C-1 bp deletions and complex deletions and i[nsertion](#page-8-0)s $(>1$ bp) compared to wild type, msh2 Δ , or [msh6](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) Δ (Fig. 3c, Supplementary Table 12, and Supplementary Fig. 4), consiste[nt with the preference](#page-17-0) [of Msh2–](#page-18-0) [Msh3 for binding larger inde](#page-18-0)ls (Habraken et al. 1996; Sia et al. 1997; Surtees and Alani 2006). Approximately 30% of the mutations that accumulated in $msh3\Delta$ were SNVs, but again the distribution was distinct, with increased CG>GC and TA>[AT changes](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) [compa](#page-8-0)red with [msh2](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) Δ and msh6 Δ (Supplementary Table 12, Fig. 3a, and Supplementary Fig. 4), despite the fact t[hat these](#page-17-0) [errors sho](#page-17-0)[uld be efficiently](#page-16-0) repaired by Msh2–Msh6 (Genschel et al. 1998; Bowers et al. 1999). This is consistent w[ith Msh2–Msh3](#page-18-0) [in vit](#page-18-0)ro affinity of C–C, A–A, and G–G mispairs (Srivatsan et al. 2014). In contrast, there were fewer TA>[CG changes in](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) $msh3\Delta$ [compa](#page-8-0)red to msh2 Δ and msh6 Δ (Supplementary Table 12 and Fig. 3a). These observations are consistent with Msh2–Msh3 playing a r[ole in correcting a specific subset of misincorporation](#page-17-0) [even](#page-17-0)ts (Harrington and Kolodner 2007; Kumar, Piacente, et al. 2011).

In assessing the ability of Msh2–Msh3 and Msh2–Msh6 to direct the repair of replication errors using msh alleles, we are limited to evaluating repair of those mutations that arise because of replication error. Due to the inherent proofreading activity of Pold [and Pol](#page-17-0)e, this [rate is low and certain](#page-16-0) [mutations ar](#page-17-0)e very rare (Kunkel 2009; Arana and Kunkel 2010; Kunkel 2011). However, rnr1 alleles generate distinct mutation pr[ofiles, including variants](#page-17-0) [that](#page-17-0) [are rare in wild-type](#page-16-0) [backgrounds \(Ku](#page-18-0)[mar, Abdulovic,](#page-17-0) et al. 2011; Buckland et al. 2014; Watt et al. 2016; Lamb et al. 2021). To evaluate the specificity of MMR MSH complexes for a broader range of mutations and to assess the impact of combined genotypes on mutation profiles, we combined msh deletions with rnr1 [alleles. Notabl](#page-16-0)[y, dNTP pools](#page-17-0) are likely elevated in cancer cells (Aye et al. 2015; Mathews 2015), including those caused by defects

Fig. 3. Mutation spectra are distinct across genotypes. a) The relative distribution of: deletions (coral), insertions (olive), MNVs (green), replacements (blue) and single-nucleotide polymorphisms (SNVs; coral) by genotype. b) The SNV spectra normalized relative total variants. Our sequencing approach does not allow us to determine which strand was originally mutated and, therefore, we use notation that indicates the base pair that is changed. Using this notation, there are 6 possible SNVs: C/G to A/T variants (CG>AT; coral), C/G to G/C variants (CG>GC; copper), C/G to T/A variants (CG>TA; green), T/A to A/T variants (TA>AT; teal), T/A to C/G variants (TA>CG; blue) and T/A to G/C variants (TA>GC; magenta). c) The deletion spectra normalized relative to total variants. We observed 3 types of deletions: deletions of more than 1 base pair (>1bpdel; coral), deletion of 1 A/T base pair (A/T-1; green), and deletion of 1 G/C base pair (G/C-1; blue). d) The insertion spectra normalized relative to total variants. We observed 3 types of insertions: insertions of more than 1 base pair (>1bp-ins; coral), insertions of 1 A/T base pair (A/T + 1; green), and insertions of 1 G/C base pair (G/C + [1; blue\). To compare th](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)e individual frequencies between genotypes, we took the average frequency and calculated the standard error of the mean (SEM) (Supplementary Fig. 4). Error bars are not shown here for ease of viewing the data.

in MMR. Therefore, the effect of the combination of $msh\Delta$ and altered dNTPs on replication fidelity could result in unique mutation signatures observed in tumors.

When $msh2\Delta$ was combined with each mrl allele, distinct mutation s[pectra \(i.e. the proportion o](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)f each type of variant) were observed (Supplementary Table 12 and Fig. 3a). The rnr1D57N, rnr1Y285F, and rnr1Y285A profiles exhibited an increase in the proportion of G/C-1 deletions relative to A/T-1 deletions compared to wild type (Fig. 3c). The opposite trend was observed in $msh2\Delta$ and in the rnr1 msh2 Δ backgrounds (Fig. 3c). This indicated that, while elevated/skewed dNTPs increase the proportion of G/C-1 variants, A/T-1 deletions are preferentially repaired in the presence of MMR. There was also a significant increase in $A/T + 1$ insertions in $mr1D57N$ msh2 Δ relative t[o either single mutan](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)t (Fig. 3d, Supplementary Table 12, and Supplementary Fig. 4). Finally, while the rnr1Y285F and rnr1Y285A SNV profiles were biased toward CG>TA and CG>AT SNVs, $mr1Y285F$ msh2 Δ and $mr1Y285A$ msh2 Δ were almost completely dominated by these [variants, with proportions](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) that differed fr[om either single mutan](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)t (Supplementary Table 12, Fig. 3b, and Supplementary Fig. 4).

These SNV profiles indicated that: (1) DNA polymerases primarily generated these errors when dCTP and dTTP were modestly skewed and elevated (rnr1Y285F) and (2) these elevated frequencies began to saturate MMR activity and/or these errors are inefficiently repaired by MMR.

Deleting MSH6 in combination with rnr1 alleles, so that only Msh2–Msh3-mediated MMR was [present, resulted in un](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)ique shifts in mutagenesis across can1 (Supplementary Table 12 and Fig. 3a). The effect was most dramatic in $mr1Y285F$ msh6 Δ with a profile that was al[most completely domina](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)ted by CG>TA and CG>[AT SNV errors \(Su](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)pplementary Table 12, Fig. 3, b and c, and Supplementary Fig. 4), indicating that Msh2–Msh3 is not efficient in correcting these mismatches. The proportion of CG>AT transversions was even higher in $mr1Y285A$ msh6 Δ , although most of the variant classes were still observed. In both $mr1Y285F$ msh6 Δ and $mr1Y285A$ msh6 Δ , there was almost a complete loss of the G/C-1-bp deletions that were ob[served at an in](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)[creased f](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)requency in rnr1Y285F and [rnr1Y285A](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) (Supplementary Table 12, Fig. 3, c and d, and Supplementary Fig. 4), consistent with the efficient repair of G/C-1-bp slippage events by Msh2–

Fig. 4. The average number of each SNV as it occurs in unique triplet nucleotide context differs by genotype. Bars are colored according to the 6 different types of SNVs. a) The 16 different triplet contexts are lettered for display purposes. The variant change (C>A, bar) occurs at the middle nucleotide marked X in each triplet context. The same triplet context is repeated for each possible variant in the following panels. b) Wild-type [and single](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) $mr1$ [alleles. c](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)) Genotypes with MSH2 deleted. d) Genotypes with MSH3 deleted. e) Genotypes with MSH6 deleted. Cosine similarity scores (Supplementary Table 22) demonstrate differences among these genotypes. To evaluate specific variants, we averaged the variant frequency in a specific trinucleotide context and calculated the SEM (data not shown).

Msh3-mediated MMR. Strikingly, the opposite was observed in the absence of Msh2-Msh3 (msh3 Δ). In both rnr1Y285F msh3 Δ and $mr1Y285A$ msh3 Δ , there was a dramatic increase in G/C-1bp deletions compared to either single mutant, almost to the exclusion of any other variants, indi[cating](#page-8-0) [that Msh2–Msh6 was](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) [un](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)able [to correct this type of](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) error (Fig. 3c, Supplementary Table 12, and Supplementary Fig. 4).

Genotype-specific position susceptibility to mutation within CAN1

We observed genotype-specific positions within CAN1 that [were susceptible to mutati](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)on, i.e. mutational "hotspots" (Supplementary Tables 15–21). We previously identified novel [susceptible posi](#page-17-0)tions in rnr1D57N, rnr1Y285F, and rnr1Y285A (Lamb et al. 2021), many of which were associated with repetitive DNA sequences, particularly insertions and deletions. In this study, we defined susceptible positions as those in which the same variant was observed at the same position in >50% of bio-logical replicates. Of the [49 susceptible positions in](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) $msh2\Delta$, 9 were unique to this genotype (Supplementary Table 18). The remaining positions overlapped with those observed in msh3 Δ and/or $msh6\Delta$. Approximately 2/3 (61 of 96) of the susceptible mutations in $msh6\Delta$ were specific to this genotype and were predominantly SNVs. None of the 18 msh3 Δ susceptible sites were unique to msh3 Δ ; 4 were also observed in msh2 Δ while the remaining 9 were also observe in msh6 Δ . All variants were more likely to occur at or adjacent to repetitive sequences, although the specific repetitive sequences varied by genotype.

We observed more distinct profiles of susceptible positions in CAN1 when msh deletions were combined with rnr1 alleles, typically differing from the individual single mutants in the number of replicates affected and/or the frequency of mutation. The susceptible positions in $mr1Y285A$ msh2 Δ identified in ou[r study](#page-17-0) [largely overlapped wi](#page-17-0)[th previously identif](#page-16-0)ied "hotspots" (Kumar, Abdulovic, et al. 2011; Buckland et al. [2014\), but our analysis also](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) [reveale](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)d new susceptible positions (Supplementary Tables 15 and 22). One noteworthy position is 527 (chromosome V coordinate 32,940), which occurs in A/T-rich sequence (5'-TTTT<u>C</u>TTGG/ CCAAGAAAA) and is susceptible to C/G>A/T mutation in rnr1Y285A msh2 Δ and rnr1Y285A-pGAL msh2 Δ . The variant frequency at this position increased synergistically in rnr1Y285ApGAL msh2D and rnr1Y285A msh2D double mutants, occurring at a frequency at least 10 times greater than the single mutants. C/ $G>T/A$ SNVs also occurred in a variety of msh6 Δ genotypes at this position, indicating decreased replication fidelity in this context. Notably, the majority of susceptible positions in rnr1Y285A $msh2\Delta$ are also susceptible in $mr1Y285A$ but not $msh2\Delta$, indicating that highly skewed/elevated dCTP and dTTP levels promoted

Table 2. Total number of mutations at any position within CAN1 observed per genotype.

^a These variants were characterized in Lamb et al. (2021),

specific errors to a level where MMR was invoked, and approached saturation of the MMR system. In contrast, mr1Y285A msh6 Δ and mr1Y285A msh3 Δ tended to exhibit susceptible positions that were distinct from either single mutant, consistent with different specificities of Msh2–Msh3 and Msh2– Msh6-directed repair in the presence of a distinct mutational baseline observed in the presence of rnr1 alleles.

We also noted increased insertion and [deletion events in](#page-17-0) [re](#page-18-0)[petitive runs, sim](#page-18-0)ilar to previous work (Lujan et al. 2014; St Charles et al. 2015), although there are only short homopolymer runs in CAN1. Notably, some G/C deletions occurred adjacent to repetitive runs in a genotype-specific manner. For example, a $G\Delta$ at CAN1 position 449 (chromosome V: 33018) was specific to $msh3\Delta$, with dinucleotides n[earby upstream and down](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)stream (5 $^{\prime}$ -GTTACATCC/GGATGTAAC) (Supplementary Table 24).

Insertions and deletions involving more than 1 nucleotide were rare but occurred at increased frequency in $msh3\Delta$. The complex insertion of CT in the repetitive run stretching from CAN1 positions 255-261 (chromosome V: 33209-33215) (5'-GAGAGAGACTTAAG/CTTAAGCTCTCTC) is noteworthy. It was observed almost exclusively in $msh2\Delta$ genotypes, and more frequently when paired with rnr1 alleles. The increased CT insertion in mrl msh2 Δ genotypes indicates that positions 255–261 were particularly susceptible to mutation when dNTPs were elevated, even by a small amount as is the case in rnr1D57N. However, the CT insertion was very efficiently repaired by either Msh2–Msh3 or Msh2–Ms[h6-directed MMR as it wa](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)s not observed in either msh3 Δ or msh6∆ (Supplementary Table 24).

Unique mutation signatures revealed by variant sequence context analysis

Genotype-specific susceptible positions indicated that sequence context was affecting the mutation profile in different genetic backgrounds. Therefore, we took 2 related approaches: (1) trinucleotide context and (2) motif analysis, to gain information about sequence context that might influence either replication errors or MMR activity. First, we assessed the SNV profile as a function of

trinucleotide context, i.e. the ide[ntity o](#page-9-0)f the nucleotide immediately 5' and 3' to a given variant (Fig. 4). This is an accepted way to analyze and compare base [substitution profiles \(insertions and](#page-16-0) [delet](#page-16-0)i[ons were not inc](#page-18-0)[luded\) \(Blokzijl](#page-17-0) et al. 2018; Alexandrov et al. 2020; Tracy et al. 2020; Lamb et al. 2021). We determined the average number of times an SNV was observed in a particular triplet context per genotype, normaliz[ed by](#page-9-0) the number of times the triplet context occurs in CAN1 (Fig. 4). H[ere, we](#page-9-0) highlight a few examples. First, $C \rightarrow T$ changes (red bars, Fig. 4), particularly in GCC and GCG sequence contexts, dominated in all genotypes, but most dramatically in [rnr1Y285F](#page-17-0) and rnr1Y285A samples, as we previously observed (Lamb et al. 20[21\) and consiste](#page-18-0)nt with whole genome sequencing in rnr1Y285A (Watt et al. 2016), possibly as a result of increased extension after misincorp[oration of dTTP,](#page-17-0) [whic](#page-17-0)h is in excess, superseding proofreading (Kunkel and Soni 1988). Notably, no C>T SNVs occurred in the GCT context in wild type, msh3 Δ , rnr1D57N msh3 Δ , or rnr1D57N, but were observed in all rnr1Y285A and rnr1Y285F double mutant backgrounds, albeit at relatively low frequencies. This example shows an error that was specific to skewed increases in dCTP and dTTP and was efficiently repaired by Msh2–Msh6 (msh3D) but not Msh2–Msh3 $(msh6\Delta)$.

We also observed a decrease in unique SNVs when rnr1Y285F or $mr1Y285A$ alleles were paired with msh2 Δ , msh3 Δ , or msh6 Δ . For example, we observed a complete loss of C>G SNVs in all trinucleotide contexts in mr1Y285A msh3 Δ , mr1Y285[A msh6](#page-8-0) Δ , and mr1Y285F msh6∆ backgrounds, as noted above (Fig. 3b and Supplementary Table 12). C>G variants rarely occurred in $msh2\Delta$ backgrounds, indicating that the replicative polymerases rarely generate these errors when dNTP pools are unperturbed. However, rnr1 single and double mutant profiles revealed distinct changes in the C $>G$ variants, compared to single msh Δ mutants. For example, rnr1Y285A sustained several C>G mutations in multiple trinucleotide contexts, which were also observed in $mr1Y285A$ msh 2Δ . The C $>G$ variants were completely absent in $mr1Y285A$ msh3 Δ and $mr1Y285A$ msh6 Δ , indicating that either Msh2–Ms3 or Msh2–Msh6 can direct the repair of these variants. Thus, different rnr1 alleles paired with different MMR deletions result in distinct mutational fingerprints.

Next, we performed motif analysis to identify l[arger sequ](#page-11-0)ence contexts driving distinct mutational signatures (Figs. 5–7). We were particularly interested in sequence motifs that might define Msh2–Msh3- vs. Msh2–Msh6-mediated substrate specificity and MMR activity. Genotypes were clustered based on the types of variants that were differentially enriched as a function of genotype, using hierarchical cl[usterin](#page-11-0)g as a means of identifying genotype[specific](#page-12-0) dif[fe](#page-13-0)rences (Fig. 5), which we then investigated further (Figs. 6 and 7). It is w[orth no](#page-6-0)ting that overall, clusters were similar to those observed in Fig. 2, which suggests that unique variants were the main drivers of genotype-specific mutation profiles. We identified specific variants that were differentially enriched by genotype and performed motif analysis (12 base window) to determine whether broader sequence context influenced the occurrence of these variants. We identified several motifs that were positively or negatively enriched in different genetic backgrounds. MMR status appeared to be the primary driver for enrichment. G/C-1 variants within G/C homopolymeric runs were positively enriched in msh3 Δ genotypes. In contrast, A/T-1 variants within A/T runs were positively enriched in $msh2\Delta$ and $msh6\Delta$ genotypes, as were SNVs, especially C $>$ T variants, within [sever](#page-11-0)al different but related sequence motifs (see examples in Fig. 5).

Fig. 5. Variants that occur in unique sequence contexts cluster together. Hierarchical cluster analysis of all unique variants within our study by genotype. a) A heatmap displaying enrichment of a given variant between genotypes, with notable clusters boxed. b) Twelve base window motif enrichment on sequence contexts surrounding the notable clusters. c) The type of variant observed in the center of sequence context from B. d) Summary of genotypes showing negative or positive correlation in each cluster.

Some of the most pronounced differential enrichment between genotypes were SNVs, espe[cially](#page-12-0) C>A and C>T, that oc[curred in CC din](#page-18-0)[ucleotide context](#page-17-0) (Fig. 6). As observed previously (Watt et al. 2016; Lamb et al. 2021), CC/GG dinucleotides bordered by G on either the 5' side or the 3' side were frequently mutated in the presence of rnr1 alleles, particularly rnr1Y285F and $mr1D57N$. This was not observed with $msh\Delta$ alleles. MMR status, especially msh3 Δ or msh6 Δ , in combination with these mr1 alleles [appear](#page-4-0)[ed](#page-12-0) to modify this bias, resulting in negative enrichment (Fig. 6, I, shaded blue). The majority of SNVs that occurred in other CC dinucleotide contexts were positively enriched in $msh2\Delta$ $msh2\Delta$ and msh6 Δ samples but negatively enriched in msh3 Δ (Fig. 6, II, III, IV & V), indicating that Msh2–Msh6 was uniquely required to recognize and direct the repair of misincorporation events in the CC dinucleotide context. Notably, these motifs were A/T-rich, particularly on the 5' side of the CC dinucleotide. A subset of C>T and C>[A var](#page-12-0)iant motifs were uniquely enriched in rnr1Y285A msh2 Δ (Fig. 6, VI), indicating either Msh2-Msh3 or Msh2-Msh6 can direct repair.

Motif enrichment analysis also revealed important combinatorial effects on mutation profiles. In double mutants, the combined mutation profiles were not necessarily additive. We observed examples of additive, antagonistic, epistatic, and synergistic effects on mutation profiles. For example, motifs in Box IV are negatively enriched in msh3 Δ and positively enriched in mr1Y285F. These effects appeared to offset each other such that little enrichment was observed in rnr1Y285F msh3 Δ . In contrast, the positive enrichment of Box I in rnr1D57N and rnr1Y285F is completely neutralized in the presence of any $msh\Delta$ allele. In another example, the rnr1Y285A msh2 Δ background was highly enriched in Box VI but is essentially unaffected in either single mutant. We noted a similar effect with mr1Y285A msh6 Δ .

One striking example of nonadditive effects on variant profiles was the synergistic increases in G/C-1 variants within specific sequenc[e cont](#page-13-0)exts when mr1Y285F/A and msh3 Δ alleles were combined (Fig. 7). We previously found that mr1Y285F and mr1Y285A [single mutants s](#page-17-0)howed an increase in G/C single-base del[etions](#page-8-0) (Lamb et al. [2021\). Here, we f](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)ound that msh3 Δ does as well (Fig. 3 and Supplementary Table 12). Individual position[s that sustained](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) [G/C error](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)s along CAN1 were often msh3 Δ [specific \(S](#page-17-0)upplementary Table 12) or rnr1Y285F/A specific (Lamb et al. 2021). Notably, the frequency of G/C-1 variants within sequence contexts that contain G or C runs increa[sed sy](#page-13-0)nergistically in the rnr1Y285F/A $msh3\Delta$ double mutants (Fig. 7). In contrast, G/C-1 variant frequency was neutral or negatively enriched in all other genotypes, consistent with the apparent specificity of Ms[h2–Msh](#page-8-0)3 for direct[ing the repair of G/C sin](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)gle-base deletions (Figs. 3 and 5 and Supplementary Table 12). Loss of Msh2–Msh3 resulted in increased G/C single-base deletions in homopol[ymer runs](#page-13-0) bordered by G/C -rich sequence on the 5' side of the run (Fig. 7, a–c). This error in this context occurred rarely in rnr1Y285F/A alone. There was a significant increase in G/C-1 mutations in $rr1Y285F$ msh3 Δ

Fig. 6. SNVs occur in C–C dinucleotide sequence contexts with differential enrichment between genotypes. a) Hierarchical cluster analysis of all SNVs that occur at CC dinucleotides. Clusters of interest are boxed, labeled by roman numerals. b) Motif enrichment of a 12 base window surrounding the mutated nucleotide was performed using Berkley web logos. The mutated base is at the 7th nucleotide position in all logos, indicated by the arrow. c) The most predominant type(s) of SNV in the cluster are displayed. d) A summary of genotypes that shows negative or positive correlation in each cluster.

double mutants. In contrast, G/C runs bordered by A/T nucleotides were more prone [to mutage](#page-13-0)nesis in rnr1Y285F/A than in msh3 Δ single mutants (Fig. 7, d-f). The frequency of these variants directly bordered by A/T increased synergistically when MSH3 was deleted in the presence of rnr1Y295F/A, but not when MSH6 was deleted, indicating that Msh2–Msh3 has specificity in directing the repair of G/C-1 deletions in repetitive G/C context.

Discussion

[Utilizing a](#page-17-0) [CAN1](#page-17-0) selection-based deep-sequencing approach (Lamb et al. 2021), we characterized mutation spectra in $msh\Delta$ and $mr1$ msh Δ double mutant genotypes. While we likely missed non-inactivating mutations and more significant rearrangements that might occur, the sequencing depth afforded by our approach allowed us to expand our understanding of MMR substrate recognition as well as the combined effects of MMR defects and altered dNTP pools on replication fidelity. By using rnr1 backgrounds that alter the type and frequency of mutations sustained, we revealed previously unrecognized specificities for the MMR initiation complexes, Msh2–Msh3 and Msh2–Msh6. The combinatorial effects that we find highlight the importance of studying mutation signatures in different genetic contexts.

Different mechanisms of mutagenesis result from distinct elevations in dNTP levels

In $mr1D57N$ msh2 Δ , the mutation rate increased 74 times higher than wild type and 3 times above $msh2\Delta$, yet the mutation spectrum of $rm1D57N$ msh2 Δ is closely related to msh2 Δ , with the

[except](#page-8-0)ion of an increase in $A/T + 1$ insertions in the double mutant (Fig. 3). The same is true of $mr1D57N$ msh 3Δ and $mr1D57N$ msh 6Δ ; their mutation spectra are most closely related to those of msh3 Δ [and](#page-8-0) $msh6\Delta$ [, res](#page-5-0)pecti[vely, despite high increas](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)es in mutation rates (Fig. 3, Table 1, and Supplementary Table 12). Therefore, the elevated dNTP pools in rnr1D57N[, which resulted](#page-17-0) in a mutation spectrum similar to wild type (Lamb et al. 2021), with the notable exception of G/C-1 deletions, did not substantially drive the type of mutation generated. The low-frequency variants that accumulate in rnr1D57N were effectively repaired by MMR, in general, and even the abse[nce of M](#page-4-0)MR [did not result in an over](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)t fitness defect in rnr1D57N (Fig. 1d and Supplementary Table 7). We conclude that the balanced dNTP increases in rnr1D57N alter mutagenesis without a remarkable change in mutation spectrum.

The rnr1Y285F allele has a modest effect on mutation rate, yet yielded a distinct mutation spectrum especially when paired with MSH deletions. In fact, the $mr1Y285F$ msh Δ spectrum closely resembles that of rnr1Y285A mshA, despite rnr1Y285A [hav](#page-5-0)ing a higher skew in d[NTP pools, a hi](#page-4-0)gher mutation rate (Table 1), and a fitness defect (Fig. 1, d and e). We conclude that even modest skewed increases in dNTPs (rnr1Y285F) result in distinct error accumulation, likely due to both a decrease in selectivity of the replicative polymerases and an increase [in efficient mismatch](#page-17-0) [exten](#page-17-0)[sion at the expe](#page-18-0)[nse of p](#page-14-0)roofreading (Kumar, Abdulovic et al. 2011; Watt et al. 2016). Fig. 8a illustrates an example of 2 different positions in the $mr1Y285A$ msh6 Δ background that are predicted to be mutated at increased frequencies via this mechanism.

The increase in G/C deletions in both rnr1Y285F and rnr1Y285A genotypes can be explained by limiting the levels of

Fig. 7. G/C single-base deletions that show synergistic increases in variant frequency in $mr1Y285A$ msh3 Δ genotypes. a) Motif enrichment of a 12-base window surrounding 2 G deletions (starred nucleotide) that are specific to genotypes with msh3D. The asterisk indicates the deleted base. b) The average variant frequencies from biological replicates in wild-type, mr1Y285F/A, and msh3A genotypes for the single-base deletion that occurred at position 31979 is plotted. c) The average variant frequencies across replicates for the G deletion at 32272. In both cases (b and c), there are very few events in the single mutants, but significant frequencies in double mutant backgrounds. d) Motif enrichment for 2 C deletions that are specific to genotypes with rnr1Y285F and rnr1Y285A but increase synergistically in double mutants with msh3D in combination. The asterisk indicates the deleted base. e) The average variant frequencies across replicates for the G deletion at 32747. f) The average variant frequencies across replicates for the G deletion at 32658. In (e) and (f), this event occurs in mr1Y285A, but not in mr1Y285F or msh3 Λ . It occurs at increased frequencies in double mutant backgrounds. Error bars indicate the standard error of the mean for frequencies. The P-values were generated by t-tests comparing average frequencies. A double asterisk indicates that a t-test was not possible because one of the genotypes had an average frequency of 0. The percentage of G/ C-1 variants in each genotype is: WT—7.7%, msh3D—26.3%, rnr1Y285F—4.9%, rnr1Y285F msh3D—79.1%, rnr1Y285A—28.9%, rnr1Y285A msh3D—94.3%.

dGTP (Fig. 8[b\). dGTP levels are](#page-16-0) [limiting in both y](#page-18-0)[east and mam](#page-17-0)malian cells (Chabes et al. 2003; Wilson et al. 2011; Mathews 2015) and notably led to distinct patterns of mutagenesis when the concentration decreases further, relative to the increase in the other 3 nucleotides, in rnr1Y285F and rnr1Y285A and even in rnr1D57N. This effect is exacerbated by the loss of MMR, especially Msh2–Msh3. Notably, rnr1 alleles that cause increases in [dGTP](#page-18-0) are also found to be extremely mutagenic (Schmidt et al. 2019), highlighting the importance of maintaining the proper level and relative abundance of dGTP.

New insights into MMR specificity

Msh2–Msh3 and Msh2–Msh6 have separate but overlapping DNA substrate specificities, leading to an expanded repertoire of

Fig. 8. Mechanisms of mutagenesis for the incorporation of errors specific to either Msh2-Ms[h6 or M](#page-12-0)sh2-Msh3 repair. The mutated base of interest is represented in red. a) Two examples of sequence context surrounding CC dinucleotides from Fig. 6b, where mis-insertion is due to the nucleotide in excess in rnr1Y285F/A backgrounds. These errors are efficiently [repaired by Ms](#page-13-0)h2-Msh6. This specificity becomes apparent when msh6D is paired with rnr1Y285F/A alleles. b) Two examples of sequence context from Fig. 7, a and d, where misalignment events occur due to the severely limiting amount of dGTP in mr1Y285F/A genetic backgrounds. The run where the deletion occurred is underlined. These single-base G/C deletions are efficiently repaired by Msh2-Msh3, but not Msh2-Msh6, a previously unidentified specificity of the repair complex.

repair. Previous studies, using a variety of reporter assays, have demonstrated that both can re[cognize and d](#page-18-0)i[rect the repair of](#page-17-0) [small insertions and deletions \(Sia](#page-17-0) et al. 1997; [Flores-Rozas and](#page-17-0) [Kolod](#page-17-0)ner 1998; Harfe and Jinks-Robertson 1999; Kunkel and Erie 2005), but we still have an incomplete understanding of the mechanistic differences in Msh2–Msh3- vs. Msh2–Msh6-directed repair. The deep sequencing reported here highlights previously unreported specificities for these pathways and provides new information about sequence context effects on MMR. Previous studies used single-strand oligonucleotide transform[ation effi](#page-17-0)[cienc](#page-17-0)[y to define Msh2–Msh3 and M](#page-18-0)sh2–Msh6 activities (Kow et al. 2007; Romanova and Crouse 2013). This approach indicated that Msh2–Msh6 preferentially corrects insertions, while Msh2–Msh3 preferentially corrects deletions. We did not observe this bias in our data. Although insertions were relatively infrequent in our data sets, [ms](#page-8-0)h3 Δ and msh6 Δ exhibited similar levels of insertion events (Fig. 3), although msh3 Δ exhibited more >1-bp insertions.

By altering dNTP pools, we altered the frequency and types of replication errors generated by DNA polymerases, revealing new substrate specificities for Msh2–Msh3 and Msh2–Msh6. It was striking how the mutation profiles for $msh3\Delta$ and $msh6\Delta$ clustered

into such distinct groups, in terms of both the types of variants observed and the sequence contexts in which they occurred. While G/C SNVs are common in wild-type backgrounds, G/C-1 deletions are relatively rare, making it difficult to determine the relative efficiencies of MSH complexes in repairing this type of error. However, elevating the dNTP pools to any extent increased the proportion of G/C-1 deletions, allowing us to assess MMR efficacy in their repair. In particular, the meta-analysis of our dataset showed most single-base deletions in rnr1Y285F, rnr1Y285A, and $msh3\Delta$ [occurre](#page-13-0)d in G/C rich contexts, especially homopolymeric runs (Figs. 7 and 8b). The double mutants $mr1Y285F$ msh3 Δ and $mr1Y285A$ msh3 Δ exhibited mutation profiles that were completely dominated by G/C-1 deletions. Therefore, Msh2–Msh6 was not able to compensate for the loss of Msh2–Msh3 for the repair of G/C-1 deletions within G/C-rich sequence contexts, despite the fact that both complexes have b[een implicated in](#page-18-0) directing the repair of single-base deletions (Meier et al. 2018). This suggests a previously unexplored role of Msh2–Msh3 in promoting replication fidelity within G/C-rich genomic regions. It will be interesting to see how different MLH complexes contribute to this specificity. Previous work has indicated that 2 MLH

complexes, Mlh1–Pms[1 and Mlh1–Mlh3, are impor](#page-18-0)tant for repair of deletion mutations (Romanova and Crouse 2013).

Similarly, SNVs at GG/CC dinucleotides were more pre[valent](#page-12-0) in rn[r1D](#page-13-0)57N and rnr1Y285F backgrounds than in wild type (Figs. 6 and 7a). We demonstrated that mutations in these G/C-rich patches were enriched in msh2 Δ and msh6 Δ backgrounds but depleted in msh3Δ, indicating that Msh2-Msh6 directs more efficient repair of mutations in GG/CC dinucleotides than Msh2– Msh3. GG dinucle[otides are mutated as](#page-18-0) a signature mutation in colorectal cancer (Rubin and Green 2009), cancers which are defined by defects in MMR. Mutat[ions at CC dinucleotid](#page-17-0)e[s are also](#page-17-0) [found](#page-17-0) in human lung cancer (Greenman et al. 2007; Lee et al. 2010) and could be generated due to defects in MMR and elevations in dNTP levels in combination. Interestingly, the combination of overexpression of the RNR R2 subunit and deletion of MSH6 caused a synergistic increase in lung carcinogenesis in a mouse mod[el, although](#page-18-0) no link with altered dNTP pools was established (Xu et al. 2008).

Sequence context analysis indicated that repetitive homoplymeric sequences were the strongest predictor of both indels and SNVs. Previous work found specific sequence context e[ffects for](#page-17-0) [human M](#page-17-0)s[h2–Msh6 \(hMsh2–hM](#page-17-0)sh6) activation in vitro (Mazurek et al. 2009). Mazurek et al. (2009) found that mispairs surrounded by symmetric 3' purines were preferentially bound by hMsh2hMsh6. These substrates also enhanced activation of hMsh2– hMsh6 ATPase activity. Both effects were predictive of enhanced repair and were hypothesized to increase the flexibility of the DNA substrate to allow efficient MSH-DNA complex formation. Our deep sequencing of can1 did not reveal a strong Msh2–Msh6 bias for these sequence contexts. There was a subset of mispairs [surro](#page-9-0)unded by 3' purines (trinucleotide contexts e, g, m, and o, Fig. 4) that were enhanced in the absence of MSH6, but the effect was relatively small $(\sim 2$ times increase) and was not systematic. In all genotypes, the greatest predictor of an SNV appeared to be the presence of dinucleotide repeat within t[he trinucleotide con](#page-17-0)text (GG, CC, TT, or AA). As previously noted (Mazurek et al. 2009), we observed no distinct broader sequence co[ntext imp](#page-11-0)licated in directing MMR beyond homopolymeric runs (Figs. 5–7) although we did note the presence of surrounding A's in our motif analysis, which may be important for increasing DNA flexibility and bending by MSH complexes.

[There are few stra](#page-16-0)nd-specific effects from altering dNTP levels (Buckland et al. 2014), but numerous studies h[ave indicated that](#page-18-0) [MMR is more efficient on the la](#page-17-0)gging strand (Pavlov et al. 2003; Kow et al. 2007; Lujan et al. 2012). However, Msh2–Msh3 does not appear to have a lagging strand [bias and may,](#page-17-0) in fact, preferentially act on the leading strand (Kow et al. 2007). This leads us to hypothesize that Msh2–Msh3 may have greater specificity for lagging strand DNA repair. Our targeted sequencing approach could be applied to strains with CAN1 in the reverse orientation to explore the differential activity and/or specificity of Msh2–Msh3 and Msh2–Msh6 on leading vs. lagging strands. This may be due to distinct i[nteractions of](#page-17-0) [Msh2–Msh3 vs. Msh2–Msh6 with M](#page-17-0)LH compl[exes \(Kow](#page-17-0) et al. [2007;](#page-17-0) Iyer et al. 2010; Kadyrova et al. 2020) or PCNA (Lau et al. 2002; Iyer et al. 2010).

Pairing elevations in dNTP levels with MMR deletions led to increased mutation rates and distinct mutation spectra, similar to previous observations specifically with $msh2\Delta$. However, when measuring the rate of canavanine resistance in $mr1Y285A$ msh2 Δ and $mr1Y285A$ msh6 Δ backgrounds, we observed substantially lower colony numbers under permissive conditions, indicating reduced fitness even under conditions when all cells should be able to grow. Reduced fitness of mr1Y285A msh2A and mr1Y285A

 $msh6\Delta$ was also seen in SGA analysis and in tetrad dissections, consistent with the phenomenon of error extinction where the threshold of [mutation rate that a](#page-18-0)l[lows wild-type](#page-17-0) c[ellular fitness](#page-18-0) [is su](#page-18-0)rpassed (Williams et al. 2013; Herr et al. 2014; Schmidt et al. 2017). A simila[r growth defect i](#page-18-0)n $mr1Y285A$ msh2 Δ was noted in previous work (Watt et al. 2016) and other rnr1 [alleles combine](#page-18-0)d with msh2 Δ also exhibited growth defects (Schmidt et al. 2019). We expect that the fitness defects we observe in the absence of selection would also reduce the number of cells that are able to grow under the selective conditions of mutation rate experi[ments,](#page-5-0) resulting in mutation rates that are underestimates (Table 1). Our results support a model in which MMR protects cellular fitness in the presence of rnr1Y285A. This could be by reducing the l[evel of mutag](#page-17-0)enesis to avoid mutation-induced extinction (Herr et al. 2014). Alternatively, it is possible that the $mr1Y285A$ msh 2Δ co[mbination result](#page-16-0)s in a cell cycle defect, as observed with pol3-01 (Datta et al. 2000). In this context, it is intriguing that the SGA screens identified synthetic effects between rnr1Y285F/A and several genes involved in mitosis.

Application to mutation signatures in human cancer

The msh2 Δ , msh3 Δ , and msh6 Δ mutation [spectra all had feature](#page-16-0)s of MMR deficient human tumor samples (Alexandrov et al. 2013). Notably, the msh6 Δ spectrum i[n our study clo](#page-18-0)sely resembles that of Msh6^{-/-} in a HAP1 cell line (Zou et al. 2018). The percentage of substitutions and indels in the $msh6\Delta$ mutation spectrum is consistent with what is seen in a $Msh6^{-/-}$ cell line. While T>C variants did not dominate the yeast msh 6Δ mutation signature as it did in the $\mathit{Msh6}^{-/-}$ cell line, the overall frequency and proportion of T>C changes did increase significantly. These data indicate that mutation signatures developed through defined mutations and deletions in Saccharomyces cerevisiae will be broadly applicable to mammalian systems. Mutation signatures from Caenorhabditis elegans also resemble human ca[ncer s](#page-17-0)i[gnatu](#page-18-0)res, albeit with some minor discrepancies (Meier et al. 2014, 2018). We note that C. elegans lacks an MSH3 [homol](#page-17-0)og, which is present in both yeast and humans (Denver et al. 2005).

Mutation signatures observed in human cancers are routinely used to predict mechanisms of tumorigenesis. We determined cosine similarity scores for our SNV tr[inucleotide context profiles](#page-16-0) [and the COSMIC S](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)BS [dataset \(Alexandrov](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data) et al. 2020) (Supplementary Fig. 5 and Supplementary Table 23). Overall, the C>A and C>T changes appeared to drive the clustering. The majority of our samples formed a distinct cluster. The exceptions were rnr1Y285F msh3 Δ , rnr1Y285A msh3 Δ and rnr1Y285A pGAL-RNR1 msh3 Δ , which formed a smaller cluster with the COSMIC signatures SBS6 and SBS15. SBS6 and SBS15 also have high cosine similarity scores with essentially all genotypes except wild type, msh3 Δ , and mr1D57N msh3 Δ . SBS6 most commonly occurs in colorectal and uterine cancers and is associated with defective MMR. It is possible that elevations in dCTP and dTTP contribute to this mutation signature in human cancers leading to synergistic increases in these types of mutations in the absence of certain MMR genes. SBS15 is characterized by C>T SNVs within NpCpG sequences in lung and stomach cancers with unknown origin. The origin of these cancers could be in part, due to defects in MMR coupled with skewed increases in dCTP and dTTP, consistent with the increased frequency of these SNVs at dinucleotides observed in rnr1Y285F/A msh3 Δ backgrounds.

SBS20 showed high similarity with msh6 Δ , rnr1D57N msh6 Δ , $mr1Y285F$ msh6 Δ , and msh2 Δ . This signature is associated with combined MMR and POLD1 mutations; the latter is deficient in

polymerase δ proofreading activity. This indicates that reduced replication fidelity resulting from either increased dNTP pools or decreased proofreading combine with MMR defects to generate similar mutation signatures. We also noted similarity between SBS24, which is associated with aflatoxin exposure, and $msh6\Delta$ and $msh2\Delta$ individually or in combination with $mr1Y285F$ or $mr1Y285A$. Several genotypes were similar to SBS94 (msh2 Δ , msh6 Δ , rnr1D57N msh6 Δ , and rnr1Y285F pGAL msh6 Δ) and SBS23 ($msh2\Delta$, $msh6\Delta$, and $mr1D57N$ $msh6\Delta$), both of which are of unknown etiology.

In addition, we also compared mutation signatures from our study to the COSMIC insertion and deletion (ID) signatures (Alexandrov et al. 2020). A major caveat to comparing CAN1 ID mutations to human mutation signatures is the lack of homopolymeric sequences within CAN1. Nonetheless, within the sequence contexts available, we noted that increased G/C singlebase deletions in the G/C context were present in COSMIC ID signatures ID3, ID7, ID9, and ID15. The proposed etiology of ID Signature 3 is cigarette smoking, ID7 is MMR defects, and ID9 and ID15 have unknown etiology. Elevated dNTP levels have not been part of a clinical diagnosis but skewed increases in dNTPs likely also contribute to these signatures of unknown etiology.

Our results highlight the importance of considering altered dNTP pools and combinatorial effects of genetic backgrounds, when defining the source of tumor mutation signatures.

Data availability statement

All strains and plasmids are available upon request. All CAN1 sequences were uploaded [to SRA BioProject PRJNA785873. SAFE](https://github.com/baryshnikova-lab/safepy) [analys](https://github.com/baryshnikova-lab/safepy)is can be found at https://github.com/baryshnikova-lab/ safepy. All mutation profile and SGA data are also available in [the supplementary tables. Representat](https://github.com/nalamb/CAN1-Paper)ive code is available at: ht[tps://github.com/nalam](https://academic.oup.com/genetics/article-lookup/doi/10.1093/genetics/iyac092#supplementary-data)b/CAN1-Paper.

Supplemental material is available at GENETICS online.

Acknowledgments

We gratefully acknowledge Dr. Anastasia Baryshnikova for her help with the SAFE analysis. We thank the UB Bioinformatics and Genomics Core for their expertise performing the sequencing in this study.

Funding

NAL is a University at Buffalo Presidential Scholar. This work was supported by the American Cancer Society (RSG-14-2350-01) to JAS. JAS is an ACS Research Scholar. JAS is also grateful for support from the University at Buffalo's Genome, Environment and Microbiome Community of Excellence. This work was also supported by a grant from the Canadian Institutes of Health Research (FDN-159913 to GWB).

Conflicts of interest

None declared.

Literature cited

Adam R, Spier I, Zhao B, Kloth M, Marquez J, Hinrichsen I, Kirfel J, Tafazzoli A, Horpaopan S, Uhlhaas S, et al. Exome sequencing identifies biallelic MSH3 germline mutations as a recessive

subtype of colorectal adenomatous polyposis. Am J Hum Genet. 2016;99(2):337–351.

- Alexandrov LB, Jones PH, Wedge DC, Sale JE, Campbell PJ, Nik-Zainal S, Stratton MR. Clock-like mutational processes in human somatic cells. Nat Genet. 2015;47(12):1402–1407.
- Alexandrov LB, Kim J, Haradhvala NJ, Huang MN, Tian Ng AW, Wu Y, Boot A, Covington KR, Gordenin DA, Bergstrom EN, et al.; PCAWG Consortium. The repertoire of mutational signatures in human cancer. Nature. 2020;578(7793):94–101.
- Alexandrov LB, Nik-Zainal S, Wedge DC, Aparicio SAJR, Behjati S, Biankin AV, Bignell GR, Bolli N, Borg A, Børresen-Dale A-L, et al.; Australian Pancreatic Cancer Genome Initiative Signatures of mutational processes in human cancer. Nature. 2013;500(7463): 415–421.
- Arana ME, Kunkel TA. Mutator phenotypes due to DNA replication infidelity. Semin Cancer Biol. 2010;20(5):304–311.
- Aye Y, Li M, Long MJ, Weiss RS. Ribonucleotide reductase and cancer: biological mechanisms and targeted therapies. Oncogene. 2015; 34(16):2011–2021.
- Baryshnikova A. Systematic functional annotation and visualization of biological networks. Cell Syst. 2016;2(6):412–421.
- Baryshnikova A, Costanzo M, Dixon S, Vizeacoumar FJ, Myers CL, Andrews B, Boone C. Synthetic genetic array (SGA) analysis in Saccharomyces cerevisiae and Schizosaccharomyces pombe. Methods Enzymol. 2010a;470:145–179.
- Baryshnikova A, Costanzo M, Kim Y, Ding H, Koh J, Toufighi K, Youn J-Y, Ou J, San Luis B-J, Bandyopadhyay S, et al. Quantitative analysis of fitness and genetic interactions in yeast on a genome scale. Nat Methods. 2010;7(12):1017–1024.
- Blokzijl F, Janssen R, van Boxtel R, Cuppen E. MutationalPatterns: comprehensive genome-wide analysis of mutational processes. Genome Med. 2018;10(1):33–33.
- Bowers J, Sokolsky T, Quach T, Alani E. A mutation in the MSH6 subunit of the Saccharomyces cerevisiae MSH2-MSH6 complex disrupts mismatch recognition. J Biol Chem. 1999;274(23):16115–16125.
- Buckland RJ, Watt DL, Chittoor B, Nilsson AK, Kunkel TA, Chabes A. Increased and imbalanced dNTP pools symmetrically promote both leading and lagging strand replication infidelity. PLoS Genet. 2014;10(12):e1004846.
- Carethers JM, Koi M, Tseng-Rogenski SS. EMAST is a form of microsatellite instability that is initiated by inflammation and modulates colorectal cancer progression. Genes (Basel). 2015;6(2):185–205.
- Chabes A, Georgieva B, Domkin V, Zhao X, Rothstein R, Thelander L. Survival of DNA damage in yeast directly depends on increased dNTP levels allowed by relaxed feedback inhibition of ribonucleotide reductase. Cell. 2003;112(3):391–401.
- Connor AA, Denroche RE, Jang GH, Timms L, Kalimuthu SN, Selander I, McPherson T, Wilson GW, Chan-Seng-Yue MA, Borozan I, et al. Association of distinct mutational signatures with correlates of increased immune activity in pancreatic ductal adenocarcinoma. JAMA Oncol. 2017;3(6):774–783.
- Costanzo M, VanderSluis B, Koch EN, Baryshnikova A, Pons C, Tan G, Wang W, Usaj M, Hanchard J, Lee SD, et al. A global genetic interaction network maps a wiring diagram of cellular function. Science. 2016;353(6306):aaf1420.
- Crooks GE, Hon G, Chandonia JM, Brenner SE. WebLogo: a sequence logo generator. Genome Res. 2004;14(6):1188–1190.
- Datta A, Schmeits JL, Amin NS, Lau PJ, Myung K, Kolodner RD. Checkpoint-dependent activation of mutagenic repair in Saccharomyces cerevisiae pol3-01 mutants. Mol Cell. 2000; 6(3):593–603.
- Davidson MB, Katou Y, Keszthelyi A, Sing TL, Xia T, Ou J, Vaisica JA, Thevakumaran N, Marjavaara L, Myers CL, et al. Endogenous

DNA replication stress results in expansion of dNTP pools and a mutator phenotype. EMBO J. 2012;31(4):895–907.

- Davies H, Morganella S, Purdie CA, Jang SJ, Borgen E, Russnes H, Glodzik D, Zou X, Viari A, Richardson AL, et al. Whole-genome sequencing reveals breast cancers with mismatch repair deficiency. Cancer Res. 2017;77(18):4755–4762.
- Denver DR, Feinberg S, Estes S, Thomas WK, Lynch M. Mutation rates, spectra and hotspots in mismatch repair-deficient Caenorhabditis elegans. Genetics. 2005;170(1):107–113.
- Edelmann W, Umar A, Yang K, Heyer J, Kucherlapati M, Lia M, Kneitz B, Avdievich E, Fan K, Wong E, et al. The DNA mismatch repair genes Msh3 and Msh6 cooperate in intestinal tumor suppression. Cancer Res. 2000;60(4):803–807.
- Flores-Rozas H, Kolodner RD. The Saccharomyces cerevisiae MLH3 gene functions in MSH3-dependent suppression of frameshift mutations. Proc Natl Acad Sci U S A. 1998;95(21):12404–12409.
- Fusco N, Lopez G, Corti C, Pesenti C, Colapietro P, Ercoli G, Gaudioso G, Faversani A, Gambini D, Michelotti A, et al. Mismatch repair protein loss as a prognostic and predictive biomarker in breast cancers regardless of microsatellite instability. JNCI Cancer Spectr. 2018;2:pky056.
- Ganai RA, Johansson E. DNA replication—a matter of fidelity. Mol Cell. 2016;62(5):745–755.
- Genschel J, Littman SJ, Drummond JT, Modrich P. Isolation of MutSbeta from human cells and comparison of the mismatch repair specificities of MutSbeta and MutSalpha. J Biol Chem. 1998; 273(31):19895–19901.
- Greenman C, Stephens P, Smith R, Dalgliesh GL, Hunter C, Bignell G, Davies H, Teague J, Butler A, Stevens C, et al. Patterns of somatic mutation in human cancer genomes. Nature. 2007;446(7132):153–158.
- Habraken Y, Sung P, Prakash L, Prakash S. Binding of insertion/deletion DNA mismatches by the heterodimer of yeast mismatch repair proteins MSH2 and MSH3. Curr Biol. 1996;6(9):1185–1187.
- Harfe BD, Jinks-Robertson S. Removal of frameshift intermediates by mismatch repair proteins in Saccharomyces cerevisiae. Mol Cell Biol. 1999;19(7):4766–4773.
- Harrington JM, Kolodner RD. Saccharomyces cerevisiae Msh2-Msh3 acts in repair of base-base mispairs. Mol Cell Biol. 2007;27(18): 6546–6554.
- Heinen CD. Genotype to phenotype: analyzing the effects of inherited mutations in colorectal cancer families. Mutat Res. 2010; 693(1–2):32–45.
- Herr AJ, Kennedy SR, Knowels GM, Schultz EM, Preston BD. DNA replication error-induced extinction of diploid yeast. Genetics. 2014; 196(3):677–691.
- Hofer A, Crona M, Logan DT, Sjöberg BM. DNA building blocks: keeping control of manufacture. Crit Rev Biochem Mol Biol. 2012; 47(1):50–63.
- Iyer RR, Pluciennik A, Genschel J, Tsai M-S, Beese LS, Modrich P. MutLalpha and proliferating cell nuclear antigen share binding sites on MutSbeta. J Biol Chem. 2010;285(15):11730–11739.
- Jensen LE, Jauert PA, Kirkpatrick DT. The large loop repair and mismatch repair pathways of Saccharomyces cerevisiae act on distinct substrates during meiosis. Genetics. 2005;170(3):1033–1043.
- Kadyrova LY, Gujar V, Burdett V, Modrich PL, Kadyrov FA. Human MutL γ , the MLH1-MLH3 heterodimer, is an endonuclease that promotes DNA expansion. Proc Natl Acad Sci U S A. 2020;117(7): 3535–3542.
- Kow YW, Bao G, Reeves JW, Jinks-Robertson S, Crouse GF. Oligonucleotide transformation of yeast reveals mismatch repair complexes to be differentially active on DNA replication strands. Proc Natl Acad Sci U S A. 2007;104(27):11352–11357.
- Kumar C, Piacente SC, Sibert J, Bukata AR, O'Connor J, Alani E, Surtees JA. Multiple factors insulate Msh2-Msh6 mismatch repair activity from defects in Msh2 domain I. J Mol Biol. 2011;411(4): 765–780.
- Kumar D, Abdulovic AL, Viberg J, Nilsson AK, Kunkel TA, Chabes A. Mechanisms of mutagenesis in vivo due to imbalanced dNTP pools. Nucleic Acids Res. 2011;39(4):1360–1371.
- Kumar D, Viberg J, Nilsson AK, Chabes A. Highly mutagenic and severely imbalanced dNTP pools can escape detection by the Sphase checkpoint. Nucleic Acids Res. 2010;38(12):3975–3983.
- Kunkel TA. DNA replication fidelity. J Biol Chem. 2004;279(17): 16895–16898.
- Kunkel TA. Evolving views of DNA replication (in)fidelity. Cold Spring Harb Symp Quant Biol. 2009;74:91–101.
- Kunkel TA. Balancing eukaryotic replication asymmetry with replication fidelity. Curr Opin Chem Biol. 2011;15(5):620–626.
- Kunkel TA, Erie DA. DNA mismatch repair. Annu Rev Biochem. 2005; 74:681–710.
- Kunkel TA, Erie DA. Eukaryotic mismatch repair in relation to DNA replication. Annu Rev Genet. 2015;49:291–313.
- Kunkel TA, Soni A. Mutagenesis by transient misalignment. J Biol Chem. 1988;263(29):14784–14789.
- Lamb NA, Bard JE, Buck MJ, Surtees JA. A selection-based next generation sequencing approach to develop robust, genotype-specific mutation profiles in S[accharomyces cerevisiae.](https://doi.org/10.1093/g3journal/jkab099) G3 (Bethesda). 2021;11(6):jkab099. doi: 10.1093/g3journal/jkab099.
- Lang GI, Parsons L, Gammie AE. Mutation rates, spectra, and genome-wide distribution of spontaneous mutations in mismatch repair deficient yeast. G3 (Bethesda). 2013;3(9):1453–1465.
- Lau PJ, Flores-Rozas H, Kolodner RD. Isolation and characterization of new proliferating cell nuclear antigen (POL30) mutator mutants that are defective in DNA mismatch repair. Mol Cell Biol. 2002;22(19):6669–6680.
- Lee W, Jiang Z, Liu J, Haverty PM, Guan Y, Stinson J, Yue P, Zhang Y, Pant KP, Bhatt D, et al. The mutation spectrum revealed by paired genome sequences from a lung cancer patient. Nature. 2010; 465(7297):473–477.
- Lujan SA, Clausen AR, Clark AB, MacAlpine HK, MacAlpine DM, Malc EP, Mieczkowski PA, Burkholder AB, Fargo DC, Gordenin DA, et al. Heterogeneous polymerase fidelity and mismatch repair bias genome variation and composition. Genome Res. 2014;24(11): 1751–1764.
- Lujan SA, Williams JS, Pursell ZF, Abdulovic-Cui AA, Clark AB, Nick McElhinny SA, Kunkel TA. Mismatch repair balances leading and lagging strand DNA replication fidelity. PLoS Genet. 2012;8(10): e1003016.
- Marsischky GT, Filosi N, Kane MF, Kolodner R. Redundancy of Saccharomyces cerevisiae MSH3 and MSH6 in MSH2-dependent mismatch repair. Genes Dev. 1996;10(4):407–420.
- Mastrocola AS, Heinen CD. Lynch syndrome-associated mutations in MSH2 alter DNA repair and checkpoint response functions in vivo. Hum Mutat. 2010;31(10):E1699–1708.
- Mathews CK. Deoxyribonucleotide metabolism, mutagenesis and cancer. Nat Rev Cancer. 2015;15(9):528–539.
- Mazurek A, Johnson CN, Germann MW, Fishel R. Sequence context effect for hMSH2-hMSH6 mismatch-dependent activation. Proc Natl Acad Sci U S A. 2009;106(11):4177–4182.
- McCulloch SD, Kunkel TA. The fidelity of DNA synthesis by eukaryotic replicative and translesion synthesis polymerases. Cell Res. 2008;18(1):148–161.
- Meier B, Cooke SL, Weiss J, Bailly AP, Alexandrov LB, Marshall J, Raine K, Maddison M, Anderson E, Stratton MR, et al. C. elegans whole-genome sequencing reveals mutational signatures related

to carcinogens and DNA repair deficiency. Genome Res. 2014; 24(10):1624–1636.

- Meier B, Volkova NV, Hong Y, Schofield P, Campbell PJ, Gerstung M, Gartner A. Mutational signatures of DNA mismatch repair deficiency in C. elegans and human cancers. Genome Res. 2018;28(5): 666–675.
- Morak M, Käsbauer S, Kerscher M, Laner A, Nissen AM, Benet-Pagès A, Schackert HK, Keller G, Massdorf T, Holinski-Feder E, et al. Loss of MSH2 and MSH6 due to heterozygous germline defects in MSH3 and MSH6. Fam Cancer. 2017;16(4):491–500.
- Nogueira GAS, Costa EFD, Lopes-Aguiar L, Lima TRP, Visacri MB, Pincinato EC, Lourenço GJ, Calonga L, Mariano FV, Altemani A, et al. Polymorphisms in DNA mismatch repair pathway genes predict toxicity and response to cisplatin chemoradiation in head and neck squamous cell carcinoma patients. Oncotarget. 2018; 9(51):29538–29547.
- Nordlund P, Reichard P. Ribonucleotide reductases. Annu Rev Biochem. 2006;75:681–706.
- Park JM, Huang S, Tougeron D, Sinicrope FA. MSH3 mismatch repair protein regulates sensitivity to cytotoxic drugs and a histone deacetylase inhibitor in human colon carcinoma cells. PLoS One. 2013;8(5):e65369.
- Pavlov YI, Mian IM, Kunkel TA. Evidence for preferential mismatch repair of lagging strand DNA replication errors in yeast. Curr Biol. 2003;13(9):744–748.
- Pino MS, Mino-Kenudson M, Wildemore BM, Ganguly A, Batten J, Sperduti I, Iafrate AJ, Chung DC. Deficient DNA mismatch repair is common in Lynch syndrome-associated colorectal adenomas. J Mol Diagn. 2009;11(3):238–247.
- Poli J, Tsaponina O, Crabbé L, Keszthelyi A, Pantesco V, Chabes A, Lengronne A, Pasero P. dNTP pools determine fork progression and origin usage under replication stress. EMBO J. 2012;31(4): 883–894.
- Radchenko EA, McGinty RJ, Aksenova AY, Neil AJ, Mirkin SM. Quantitative analysis of the rates for repeat-mediated genome instability in a yeast experimental system. Methods Mol Biol. 2018;1672:421–438.
- Romanova NV, Crouse GF. Different roles of eukaryotic MutS and MutL complexes in repair of small insertion and deletion loops in yeast. PLoS Genet. 2013;9(10):e1003920.
- Rubin AF, Green P. Mutation patterns in cancer genomes. Proc Natl Acad Sci U S A. 2009;106(51):21766–21770.
- Santos LS, Silva SN, Gil OM, Ferreira TC, Limbert E, Rueff J. Mismatch repair single nucleotide polymorphisms and thyroid cancer susceptibility. Oncol Lett. 2018;15(5):6715–6726.
- Schmidt TT, Reyes G, Gries K, Ceylan CÜ, Sharma S, Meurer M, Knop M, Chabes A, Hombauer H. Alterations in cellular metabolism triggered by URA7 or GLN3 inactivation cause imbalanced dNTP pools and increased mutagenesis. Proc Natl Acad Sci U S A. 2017; 114(22):E4442–e4451.
- Schmidt TT, Sharma S, Reyes GX, Gries K, Gross M, Zhao B, Yuan J-H, Wade R, Chabes A, Hombauer H, et al. A genetic screen pinpoints ribonucleotide reductase residues that sustain dNTP homeostasis and specifies a highly mutagenic type of dNTP imbalance. Nucleic Acids Res. 2019;47(1):237–252.
- Serero A, Jubin C, Loeillet S, Legoix-Né P, Nicolas AG. Mutational landscape of yeast mutator strains. Proc Natl Acad Sci U S A. 2014;111(5):1897–1902.
- Sia EA, Kokoska RJ, Dominska M, Greenwell P, Petes TD. Microsatellite instability in yeast: dependence on repeat unit size and DNA mismatch repair genes. Mol Cell Biol. 1997;17(5):2851–2858.
- Srivatsan A, Bowen N, Kolodner RD. Mispair-specific recruitment of the Mlh1-Pms1 complex identifies repair substrates of the Saccharomyces cerevisiae Msh2-Msh3 complex. J Biol Chem. 2014;289(13):9352–9364.
- St Charles JA, Liberti SE, Williams JS, Lujan SA, Kunkel TA. Quantifying the contributions of base selectivity, proofreading and mismatch repair to nuclear DNA replication in Saccharomyces cerevisiae. DNA Repair (Amst). 2015;31:41–51.
- Supek F, Bošnjak M, Škunca N, Šmuc T. REVIGO summarizes and visualizes long lists of gene ontology terms. PLoS One. 2011;6(7):e21800.
- Surtees JA, Alani E. Mismatch repair factor MSH2-MSH3 binds and alters the conformation of branched DNA structures predicted to form during genetic recombination. J Mol Biol. 2006;360(3):523–536.
- Takahashi M, Koi M, Balaguer F, Boland CR, Goel A. MSH3 mediates sensitization of colorectal cancer cells to cisplatin, oxaliplatin, and a poly(ADP-ribose) polymerase inhibitor. J Biol Chem. 2011; 286(14):12157–12165.
- Tracy MA, Lee MB, Hearn BL, Dowsett IT, Thurber LC, Loo J, Loeb AM, Preston K, Tuncel MI, Ghodsian N, et al. Spontaneous polyploids and antimutators compete during the evolution of Saccharomyces cerevisiae mutator cells. Genetics. 2020;215(4):959–974.
- Valle L, de Voer RM, Goldberg Y, Sjursen W, Försti A, Ruiz-Ponte C, Caldés T, Garré P, Olsen MF, Nordling M, et al. Update on genetic predisposition to colorectal cancer and polyposis. Mol Aspects Med. 2019;69:10–26.
- van Oers JMM, Edwards Y, Chahwan R, Zhang W, Smith C, Pechuan X , Schaetzlein S, Jin B, Wang Y, Bergman A, et al. The MutS β complex is a modulator of p53-driven tumorigenesis through its functions in both DNA double-strand break repair and mismatch repair. Oncogene. 2014;33(30):3939–3946.
- Watt DL, Buckland RJ, Lujan SA, Kunkel TA, Chabes A. Genome-wide analysis of the specificity and mechanisms of replication infidelity driven by imbalanced dNTP pools. Nucleic Acids Res. 2016; 44(4):1669–1680.
- Williams LN, Herr AJ, Preston BD. Emergence of DNA polymerase e antimutators that escape error-induced extinction in yeast. Genetics. 2013;193(3):751–770.
- Wilson PM, LaBonte MJ, Russell J, Louie S, Ghobrial AA, Ladner RD. A novel fluorescence-based assay for the rapid detection and quantification of cellular deoxyribonucleoside triphosphates. Nucleic Acids Res. 2011;39(17):e112.
- Xu X, Page JL, Surtees JA, Liu H, Lagedrost S, Lu Y, Bronson R, Alani E, Nikitin AY, Weiss RS, et al. Broad overexpression of ribonucleotide reductase genes in mice specifically induces lung neoplasms. Cancer Res. 2008;68(8):2652–2660.
- Zou X, Owusu M, Harris R, Jackson SP, Loizou JI, Nik-Zainal S. Validating the concept of mutational signatures with isogenic cell models. Nat Commun. 2018;9(1):1744.

Communicating editor: J. A. Nickoloff