A Novel, Drug Resistance-Independent, Fluorescence-Based Approach To Measure Mutation Rates in Microbial Pathogens

Erika Shor,a Jessica Schuyler,b David S. Perlin*a

aHMH Center for Discovery and Innovation, Nutley, New Jersey, USA
bPublic Health Research Institute, Rutgers University-New Jersey Medical School, Newark, New Jersey, USA

ABSTRACT All evolutionary processes are underpinned by a cellular capacity to mutate DNA. To identify factors affecting mutagenesis, it is necessary to compare mutation rates between different strains and conditions. Drug resistance-based mutation reporters are used extensively to measure mutation rates, but they are suitable only when the compared strains have identical drug tolerance levels—a condition that is not satisfied under many “real-world” circumstances, e.g., when comparing mutation rates among a series of environmental or clinical isolates. Candida glabrata is a fungal pathogen that shows a high degree of genetic diversity and fast emergence of antifungal drug resistance. To enable meaningful comparisons of mutation rates among C. glabrata clinical isolates, we developed a novel fluorescence-activated cell sorting-based approach to measure the mutation rate of a chromosomally integrated GFP gene. We found that in Saccharomyces cerevisiae this approach recapitulated the reported mutation rate of a wild-type strain and the mutator phenotype of a shuΔ mutant. In C. glabrata, the GFP reporter captured the mutation rate increases caused either by a genotoxic agent or by deletion of DNA mismatch repair gene MSH2, as well as the specific mutational signature associated with msh2Δ. Finally, the reporter was used to measure the mutation rates of C. glabrata clinical isolates carrying different alleles of MSH2. Together, these results show that fluorescence-based mutation reporters can be used to measure mutation rates in microbes under conditions of unequal drug susceptibility to reveal new insights about drivers of mutagenesis.

IMPORTANCE Measurements of mutation rates—i.e., how often proliferating cells acquire mutations in their DNA—are essential for understanding cellular processes that maintain genome stability. Many traditional mutation rate measurement assays are based on detecting mutations that cause resistance to a particular drug. Such assays typically work well for laboratory strains but have significant limitations when comparing clinical or environmental isolates that have various intrinsic levels of drug tolerance, which confounds the interpretation of results. Here we report the development and validation of a novel method of measuring mutation rates, which detects mutations that cause loss of fluorescence rather than acquisition of drug resistance. Using this method, we measured the mutation rates of clinical isolates of fungal pathogen Candida glabrata. This assay can be adapted to other organisms and used to compare mutation rates in contexts where unequal drug sensitivity is anticipated.

KEYWORDS Candida glabrata, GFP, MSH2, mutation rate

Acquisition of mutations underlies evolution in all contexts, and in pathogenic microbes this process can result in the emergence of dangerous drug-resistant strains. An organism’s capacity to acquire mutations is typically measured as the rate (per cell, per generation) at which a specific reporter gene accumulates mutations that
result in a detectable phenotypic change. Because most genomes are extremely stable and mutation rates are typically very low, the most convenient and widely used mutation measurement assays are set up as selections, where mutation of a reporter gene confers resistance to a particular drug (1). In such an assay, mutations in the reporter gene arise at some low rate in cells proliferating in culture and are then selected by plating the cultures on drug-containing medium, which kills wild-type cells. The CAN1 mutation assay in *Saccharomyces cerevisiae* is based on this principle and has been used extensively to gain insights into mechanisms controlling genome stability in yeast (2–5). Although drug resistance-based mutation assays have a number of advantages, most particularly in their relative ease and rapidity, their major limitation is that they allow direct comparisons of mutation rates only between strains that have the same level of drug sensitivity. This condition is largely satisfied when one compares the mutation rates of isogenic strains, e.g., a laboratory strain and a DNA repair mutant derived from it. However, this condition does not hold when one wishes to compare mutation rates among nonisogenic strains, e.g., a panel of clinical or environmental isolates, which may have various levels of drug tolerance. In this case, more tolerant strains are expected to survive for a longer period on selection medium, all the while continuing to produce resistance mutations, leading to an artificially high mutation rate estimate, and vice versa. Similar considerations preclude direct comparisons of mutation rates between different species, e.g., one that is highly drug susceptible versus one that is more drug tolerant, or between different growth conditions (e.g., growth in the presence of a stressor that may affect overall stress/drug tolerance). Thus, in order to rigorously measure and compare mutation rates in a way that is not restricted to a small number of laboratory strains and their derivatives, it is necessary to develop a drug resistance-independent method to measure mutation rates.

*Candida glabrata* is a yeast that is closely related to *S. cerevisiae* (6) and is also associated with the human microbiome (7). In contrast to *S. cerevisiae*, however, *C. glabrata* is an opportunistic pathogen that can cause life-threatening infections in immunocompromised individuals (8, 9). The prevalence of *C. glabrata* in infections has been increasing, and it is now the second most prevalent cause of invasive candidiasis in the United States and Europe (10, 11). One reason for this increase is that *C. glabrata* either is intrinsically resistant or acquires resistance relatively quickly to the limited number of antifungal drugs currently in clinical use (9, 12). In *C. glabrata*, drug resistance is predominantly caused by point mutations, either in transcription factors regulating drug efflux (13–15) or in genes encoding drug targets (16–18). Several studies have also documented the emergence of multidrug-resistant (MDR) *C. glabrata* infections, for which there are no treatment options and which are associated with extremely high mortality (19, 20).

Comparisons of DNA sequences (both of specific genes and of whole genomes) from different *C. glabrata* clinical isolates have revealed an exceedingly high level of genetic diversity, in terms of both single nucleotide polymorphisms (SNPs) and chromosomal arrangements (21–26). A multilocus sequencing typing (MLST) scheme based on SNPs at six different loci has identified over 100 distinct sequence types (STs) of *C. glabrata*, which cluster into seven clades (22, 25; https://pubmlst.org/cglabrata/). However, even strains within the same clade exhibit high genetic diversity (22), which, together with rapid emergence of mutations that cause drug resistance, has led to the hypothesis that *C. glabrata* may have a highly plastic, or mutable, genome. However, mutation rates in *C. glabrata* have not been measured or compared to other organisms.

In a previous study, we began to examine the role of DNA mismatch repair (MMR) in maintaining genome stability and emergence of drug resistance in *C. glabrata* (24). In particular, we found that different STs of *C. glabrata* are associated with specific SNPs in MMR gene *MSH2*, some of which result in amino acid changes and, when introduced into an *msh2Δ* reference strain on a plasmid, do not fully rescue that strain’s hypermutator phenotype. This result suggested that some *C. glabrata* isolates, e.g., those carrying certain variants of *MSH2*, may exhibit higher mutation rates and may therefore acquire drug resistance more rapidly. Indeed, in *Cryptococcus*, naturally
occurring mutations in *MSH2* have been shown to contribute to microevolution and population diversity (59, 60). Yet, recent clinical studies have not found an association between specific *MSH2* alleles and drug resistance (27–29, 61, 62), raising the question of whether clinical isolates carrying these alleles are true mutators. To answer this question, it is necessary to measure and directly compare mutation rates between clinical isolates of *C. glabrata*. However, as described above, comparisons of different clinical isolates are complicated by the variation in their drug resistance profiles, some of which is due to varying activity of drug efflux pumps (14, 15, 18), which is likely to render any drug resistance-based mutation assay inapplicable.

In this study, we developed and validated a GFP-based mutation reporter that allowed us to measure mutation rates in a drug resistance-independent way. The reporter was shown to recapitulate the mutation rate and spectrum of a DNA mismatch repair mutant and detect DNA damage-induced mutagenesis in *C. glabrata*, recapitulate the mutation rates of wild-type and mutator strains of *S. cerevisiae*, and compare spontaneous mutation rates in *C. glabrata* and *S. cerevisiae*. Finally, we used this reporter to measure the mutation rates of a number of clinical isolates of *C. glabrata*, including those carrying a specific *MSH2* variant previously suggested to increase mutagenesis.

### RESULTS

**Developing the GFP-based mutation rate reporter.** To measure mutation rates in *C. glabrata*, at first we attempted to use traditional drug resistance-based reporters, such as *CAN1*, which has been used extensively to measure mutation rates in *S. cerevisiae* (2–5). In that fungus, *CAN1* cells are sensitive to the drug canavanine, whereas mutations in the *can1* gene cause canavanine resistance and can be selected on canavanine-containing plates. However, although the *C. glabrata* genome contains several potential *CAN1* orthologs (CAGL0J08162g and CAGL0J08184g), commonly used reference strain ATCC 2001 (also known as CBS138) was completely resistant to canavanine up to concentrations of 1 mg/ml (see Fig. S1A in the supplemental material; also data not shown), whereas the typical selection concentration in *S. cerevisiae* is 60 μg/ml. We also tried using 5-fluoroanthranilic acid (5-FAA), which selects for mutations in the tryptophan biosynthetic pathway (30). Although ATCC 2001 and many clinical isolates of *C. glabrata* were sensitive to 5-FAA, we discovered that this sensitivity widely varied among different strains (Fig. S1B). Although this variation was not entirely surprising, as different clinical isolates are well known to show different levels of antifungal drug resistance, which is at least in part due to the activity of drug efflux pumps, it also eliminated the possibility of using 5-FAA—or likely any other drug resistance-based approach—to measuring mutation rates in *C. glabrata* clinical isolates.

To enable measurements of mutation rates in a way that was independent of drug resistance, we chose a fluorescence-based approach. We created a cassette where the gene encoding yeast enhanced green fluorescent protein (*yEGFP*) was driven by the strong constitutive promoter *pTEF1* of *S. cerevisiae* (31), which was also previously shown to strongly induce gene expression in *C. glabrata* (32) (Fig. 1A). In order to facilitate the chromosomal insertion and subsequent tracking of this construct, the cassette also contained the gene conferring nourseothricin resistance (*NAT*) driven by its own promoter (Fig. 1A). This cassette was inserted into the right arm of *C. glabrata* chromosome K between two uncharacterized ORFs (Fig. 1A) and validated by sequencing. The resulting strain was constitutively and strongly fluorescent (Fig. 1B) and was used to measure mutation rates of *yEGFP* using fluorescence-based cell sorting (FACS) in fluctuation experiments as described below (Fig. 1C; Fig. S2).

Briefly, in a typical experiment, a starter YPD culture was diluted into multiple (e.g., 8 to 12) parallel YPD cultures in a 96-well plate to a starting density of a few cells per well and incubated at 37°C overnight. The following morning, each culture, in its entirety, was diluted severalfold in YPD to ensure that cells collected for FACS analysis several hours later were in log phase, which was found to be necessary to achieve maximum expression of GFP and the optimal resolution between GFP-positive and
GFP-negative populations. The cells were then collected by filtration, resuspended in PBS, and analyzed by FACS. Prior to FACS analysis, propidium iodide (PI) was added to each sample to gate out the inviable, PI-positive cell subset (Fig. S2). Although GFP levels of fluorescent cultures varied slightly between experiments (e.g., between the same strain analyzed on different days and between different strains), the overall fluorescence levels of GFP\textsuperscript{H11001} cells were always significantly higher than those of GFP\textsuperscript{H11002} cells (Fig. S3), allowing for efficient sorting of GFP\textsuperscript{H11002} cells from GFP\textsuperscript{H11001} populations.

The number of cells per culture (\(n\)) was optimized by varying the final volume of the cultures. For strains with lower mutation rates, such as the reference \(C.\) glabrata strain ATCC 2001, the optimal \(n\) was found to be \(\geq 3 \times 10^6\) cells/well, whereas for cells with elevated mutation rates (such as \(msh2\Delta\)), \(1 \times 10^6\) cells/well was sufficient to obtain multiple cultures with mutants. Each culture, in its entirety, was analyzed by FACS, and GFP-negative cells were collected, immediately plated onto YPD plates, and allowed to form colonies. These colonies were then validated for (i) the presence of the NAT cassette (by replica plating onto nourseothricin medium) and (ii) reduction of GFP fluorescence (by flow cytometry). Next, \(y\text{EGFP}\) was sequenced in NAT\textsuperscript{H11001} GFP\textsuperscript{H11002} colonies to identify the mutations responsible for reduced fluorescence. For every strain, 200 GFP\textsuperscript{H11001} cells were also collected by FACS and plated on YPD to calculate plating efficiency. Finally, mutation rates and 95% confidence intervals were calculated using the MSS maximum likelihood method (1, 33) based on \(n\) (the number of processed cells \(\times\) PI-negative fraction \(\times\) plating efficiency) and the number of NAT\textsuperscript{+} GFP\textsuperscript{−} mutants in every culture (\(r\)). Importantly, we found that every NAT\textsuperscript{+} GFP\textsuperscript{−} colony in which \(y\text{EGFP}\) was sequenced contained a single mutation in the \(y\text{EGFP}\) ORF or, in a few cases, the \(p\text{TEF1}\) promoter (see below), indicating that loss of fluorescence is virtually always caused by mutations in \(y\text{EGFP}\) and that therefore this mutation assay is highly specific to a single locus.

Mutations in GFP do not affect cellular fitness. A key condition that has to be satisfied by any mutation rate measurement assay is that mutations in the reporter gene should not affect the fitness of the strain, either positively or negatively, as this would result in overestimating or underestimating the mutation rate, respectively (1). To check that this condition is fulfilled in the GFP-based mutation reporter, we isolated two different loss-of-function mutations in \(y\text{EGFP}\) using FACS and measured their fitness compared to the parent strain over 24 h of growth in YPD, which is the duration

**FIG 1** Description of the fluorescence-based mutation reporter. (A) A cassette containing pTEF1-driven yEGFP linked to the gene encoding nourseothricin resistance (NAT) under its own promoter was integrated into the indicated locus on chromosome K in \(C.\) glabrata laboratory strain ATCC 2001. (B) The resulting strain became constitutively fluorescent. (C) Schematic describing the fluctuation assay used to measure the mutation rate of yEGFP.
of a typical mutation rate experiment. Each strain was mixed 1:1 with the parent strain, producing a coculture where approximately half the population was fluorescent and half was not (Fig. 2, Time 0). Both cocultures were diluted into multiple wells to several hundred cells per well and grown for 24 h at 37°C, mimicking a typical fluctuation experiment. After 24 h, fluorescence measurements showed that the proportions of fluorescent and nonfluorescent cells in the cultures had not significantly changed (Fig. 2, 24 h), indicating that mutations in yEGFP did not affect fitness relative to the parental GFP-positive strain.

**GFP-based mutation reporter recapitulates mutation rates in** *Saccharomyces cerevisiae*. In addition to inserting the GFP cassette into the *C. glabrata* genome, we also inserted it at the CAN1 locus in *S. cerevisiae* (Fig. 3A) in order to test whether this reporter would recapitulate the previously determined mutation rates of two *S. cerevisiae* strains: a wild-type strain of the W303 background and an isogenic mutant carrying a deletion of the SHU1 gene. SHU1 functions in promoting error-free DNA repair by homologous recombination, and its loss was shown to increase CAN1 mutation rate by approximately 8.5-fold in one study (34) and by 4-fold in another study (35). The FACS-based assay described above (Fig. 1C) was used to measure the mutation rates of yEGFP in wild-type and *shu1Δ* strains. The assay recapitulated the increase in mutation rate in the *shu1Δ* mutant relative to the wild-type strain (4-fold; Fig. 3B). Furthermore, the absolute mutation rate obtained for the wild-type strain was ~2.7 per 10⁷
cells per generation, which agrees well with typical mutation rates obtained in *S. cerevisiae* (2, 5, 34).

Finally, using the fluorescence-based mutation reporter, we were able to directly compare spontaneous mutation rates at the *C. glabrata* locus carrying the cassette (Fig. 1A) and the *CAN1* locus of *S. cerevisiae*. Interestingly, we found that the mutation rate was 9-fold higher at the *S. cerevisiae* *CAN1* locus than at the analyzed locus in *C. glabrata* (Fig. 3B), suggesting that, at least during unperturbed growth in YPD, the examined strain of *C. glabrata* (ATCC 2001) does not behave as a spontaneous hypermutator.

**GFP-based mutation reporter captures the msh2Δ mutator phenotype and mutation spectrum.** To further validate the GFP-based mutation reporter in *C. glabrata*, we used CRISPR to insert the cassette into the same chromosomal location (Fig. 1A) in a *C. glabrata* strain derived from ATCC 2001 but carrying a deletion of DNA mismatch repair (MMR) gene *MSH2* (24). *MSH2* is the *C. glabrata* homolog of MutSβ, whose loss has been shown to result in a strong mutator phenotype in all organisms where it has been examined, including *C. glabrata* (24, 36). Both *MSH2* and *msh2Δ* strains were used in fluctuation experiments to measure their mutation rates as described above (Fig. 1C and Fig. S2). We found that, as expected, *msh2Δ* resulted in a strong mutator phenotype, increasing the rate of mutation of yEGFP by 40-fold (Fig. 4A), which is very similar to its effect on mutation rates in *S. cerevisiae*, where the *msh2Δ* effect on *CAN1* mutation rate ranges from a 16- to a 40-fold increase, depending on the study (37–40). Furthermore, sequencing GFP− colonies revealed that the *msh2Δ* strain produced a very different spectrum of mutations in yEGFP from the *MSH2* strain (Fig. 4B and C). Mutations in the wild-type (*MSH2*) strain were largely comprised of base

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**FIG 3** Fluorescence-based mutation reporter recapitulates the mutation rate increase of an *S. cerevisiae* mutant. (A) The yEGFP-NAT cassette was integrated at the *CAN1* locus of *S. cerevisiae* using homology-based targeting. (B) The assay detected the mutator phenotype of the *shu1Δ* mutant, previously shown to have an elevated *CAN1* mutation rate (34). The assay also showed that spontaneous forward mutation in *C. glabrata* during standard laboratory growth in YPD is not higher than that in *S. cerevisiae*. Sc, *S. cerevisiae*; Cg, *C. glabrata*. Error bars, 95% confidence intervals.
pair substitutions (bps), whereas the majority of mutations in the *msh2*Δ mutant were due to single nucleotide deletions or insertions in mononucleotide repeats (e.g., AAAA or TTTT), with the strongest mutation “hot spot” at a run of seven T’s (Fig. 4B). This mutation spectrum recapitulates that of the *msh2*Δ strain in *S. cerevisiae* and is thought to be due to DNA polymerase “slippage” errors at mononucleotide repeats, which are normally repaired by MMR (37, 38, 41). Thus, the GFP-based mutation reporter was able to accurately capture both the increase in mutation rate and the change in mutation spectrum of the *msh2*Δ mutant.

**GFP-based mutation reporter detects DNA-damage induced mutagenesis.** To investigate whether the GFP-based mutation reporter would capture DNA damage-induced mutagenesis, we performed the fluctuation assay on the ATCC 2001 strain grown in the presence of 0.01% methyl methanesulfonate (MMS), an alkylating agent. Indeed, we detected a 48-fold increase in *yEGFP* mutation rate in cells cultured in the presence of MMS (Fig. 5A). Sequencing mutations in GFP*/H11002* colonies recovered after growth in MMS showed that the spectrum of mutations did not change significantly from that in cells grown in the absence of MMS (Fig. 5B and C). Thus, MMS caused an overall increase in mutagenesis of *yEGFP*, as expected, but apparently did not significantly change the cellular pathways by which these mutations were generated. This was consistent with a previous report, where the MMS-induced mutation spectrum in yeast was similar to the spontaneous mutation spectrum (42).

**Mutation rates of C. glabrata clinical isolates.** We used the GFP-based mutation assay to measure mutation rates of six clinical *C. glabrata* isolates: three that, like ATCC 2001, belonged to sequence type (ST) 15 and carried the corresponding *MSH2* sequence and three belonging to ST16 and carrying the variant *MSH2E231G/L269F* (23, 24). For every clinical isolate, the reporter cassette was integrated into the same chromosomal locus using CRISPR and validated by flow cytometry and DNA sequencing, and the mutation rate was measured using FACS as described above. Interestingly, we found that none of the clinical isolates, including those carrying *MSH2E231G/L269F*, had elevated mutation rates relative to the reference strain ATCC 2001 (Fig. 6). Thus, even though our previous examination indicated that *MSH2E231G/L269F* did not fully rescue the mutator phenotype of the *msh2*Δ mutant in ATCC 2001 (24), a direct, rigorous assessment of the mutation rate of the clinical isolates showed that under standard laboratory conditions, the clinical isolates did not exhibit increased mutation rates compared to the reference strain.
conditions this variant, in its native genomic context, is not associated with an elevated spontaneous forward mutation rate.

**DISCUSSION**

The goal of this study was to develop, validate, and use a new method for measuring mutation rates in a way that did not rely on drug resistance-based reporters. To this end, we designed a FACS-based scheme to capture and quantify loss-of-function mutations...
in the gene encoding GFP and used it in fluctuation experiments to measure mutation rates in *C. glabrata* and *S. cerevisiae*. We found that this fluorescence-based mutation reporter recapitulated the previously reported mutator phenotype of the *S. cerevisiae* *shu1*Δ mutant and captured the expected increase in mutation rates due to loss of *MSH2* or treatment with a genotoxic agent in *C. glabrata*. This reporter also accurately captured the mutational spectrum (a predominance of single nucleotide insertions or deletions in homopolymeric runs) associated with loss of *MSH2*. Finally, the reporter was used to measure the mutation rates of several clinical *C. glabrata* isolates, including those carrying the *MSH2E231G/L269F* variant previously suggested to contribute to increased mutagenesis (24), and showed that all clinical isolates examined had very similar spontaneous mutation rates.

Our mutation assay showed that under standard laboratory growth conditions (YPD, 37°C), *C. glabrata* clinical isolates carrying the *MSH2E231G/L269F* variant do not show elevated spontaneous mutation rates relative to ATCC 2001 or to clinical isolates carrying the “wild-type” version of *MSH2* (i.e., one identical to that in ATCC 2001). This is consistent with several recent clinical studies that did not see an association between *MSH2* genotype and prevalence of drug resistance in *C. glabrata* (27–29, 61, 62). However, in our previous study, we found that several *MSH2* variants, including *MSH2E231G/L269F*, when introduced on a plasmid into an ATCC 2001-derived *msh2*Δ mutant, did not fully rescue that strain’s elevated mutation rate (24). There are several non-mutually exclusive possibilities that can reconcile previous data with this present study. First, it is possible that, although the plasmids were maintained by selection, a subpopulation of the culture had lost the plasmid and therefore lacked any copy of *MSH2*. Second, it is possible that the *MSH2* variants carried on a plasmid were present in more than one copy, which, depending on the nature of the mutation, might either help restore function (for a partial loss-of-function mutation) or further exacerbate the associated defects (for a dominant negative mutation). Finally, it is important to consider that in the clinical isolates studied here, the *MSH2E231G/L269F* variant is present in its normal genomic context, which is that of ST16 (23). ST16 is separated from ATCC 2001 (ST15) by hundreds to thousands of SNPs throughout the genome, including SNPs in genes that encode protein partners of Msh2, such as Msh3 and Msh6 (36). Thus, it is possible that each *MSH2* variant has evolved in concert with its partner genes in a way that maintains efficient MMR and low mutation rates. In this scenario, moving a particular variant to a different genomic context would force it to form suboptimal partnerships with noncognate interacting proteins, resulting in less efficient MMR and an increased spontaneous mutation rate. Consistent with this hypothesis, our analyses of *MSH3* and *MSH6* sequences have revealed that each of these genes has multiple SNPs between ST15 and ST16 strains, including five SNPs in each gene that result in amino acid changes (the ST15 and ST16 whole-genome sequences have been generated by us for a different project [unpublished data]).

The fluorescence-based mutation assay developed in our study has specific advantages and specific limitations that need to be considered when deciding whether to use it or another assay to measure mutation rates in a given experimental system. First, similar to other mutation assays based on a loss of function of a reporter, only mutations that affect GFP function, i.e., its ability to fluoresce in the detectable range, can be identified. Thus, if it is desirable to calculate true mutation rates independently of whether a mutation is expressed, it is more appropriate to use the recently developed whole-genome sequencing (WGS) approaches (43–45). Although this is a very powerful technique, it requires WGS of multiple isolates per strain and is therefore still considerably more expensive and computationally heavy than our method, which requires no computational expertise. One unique hurdle of our assay not shared by other methods is that it requires access to a FACS instrument that can sort millions of cells in minutes; otherwise, the time frame of a single experiment becomes unfeasibly long. Once the sorting is completed, however, the rest of the steps require a standard flow cytometer (or fluorescence microscope) and standard laboratory techniques. As discussed above, this assay is going to be more informative than drug resistance-based
assays when the strains under comparison have different drug tolerance profiles, as can be expected for environmental/clinical isolates or different species or when one wishes to measure the mutation rates of strains exposed to different types of environmental stress (e.g., antimicrobial drugs).

We validated our mutation assay by analyzing strains with elevated mutation rates (e.g., *shu1Δ* in *S. cerevisiae* and msh2Δ in *C. glabrata*). It should also be possible to use this assay to identify antimutators—i.e., genes whose loss reduces mutagenesis—which can be extremely informative for identifying cellular pathways that promote mutagenesis and genetic instability (46–48). However, this would require sorting cultures with significantly larger numbers of cells and would therefore take longer, with the sorting time required negatively correlating with the mutation rate of the strain. One potential improvement over the current methodology that would reduce the required time and labor is developing a fluorescence-based assay that uses flow cytometry to count the nonfluorescent cells but skips the sorting and plating steps. In our current setup, this was not possible because the number of GFP-negative cells recorded by the FACS instrument was typically much greater than, and did not correlate with, the number of colonies that grew from the sorted cells. In other words, despite our use of propidium iodide (PI) to gate out membrane-permeable cells, a large subset of PI-negative GFP-negative cells were inviable/nonculturable. Perhaps, with further optimization—e.g., using other fluorescent markers that can be used in conjunction with live/dead dyes—it will be possible to accurately record the number of nonfluorescent live cells using flow cytometry only, without the need for sorting and plating.

We have used the fluorescence-based assay to measure forward mutation rates; however, because the reporter cassette contains two genes (yEGFP and NAT), it can be adapted to measure large deletions by looking for simultaneous loss of both yEGFP and NAT, similar to the CAN1-URA3 loss assay developed in *S. cerevisiae* (49). Such an assay would be extremely useful in *C. glabrata* and other fungal pathogens characterized by frequent genomic rearrangements (26, 50, 51). In the current genomic location of the cassette, we did not identify any simultaneous deletions of yEGFP and NAT among the >170 analyzed *C. glabrata* cultures, including those containing genotoxic agent MMS, indicating that the spontaneous rate of deletions at this locus is extremely low. In future studies, the yEGFP-NAT cassette will be integrated at genomic loci more likely to undergo rearrangements, such as subtelomeric loci containing multiple and variable numbers of genes from the adhesin family (52, 53) and ribosomal DNA (rDNA) (54).

The high degree of genetic diversity in *C. glabrata* populations and the fast emergence of drug resistance both indicate that at least under some conditions, *C. glabrata* is able to rapidly mutate and diversify its genome. Our present study indicates that these conditions do not include unperturbed growth in YPD, suggesting that *C. glabrata* may be subject to stress-induced mutagenesis. Indeed, previously, stress-induced mutator phenotypes have been identified in a majority of natural isolates of *Escherichia coli*, whereas only 5% were shown to act as constitutive mutators (55). Future studies will examine whether mutagenesis is affected by stress conditions, including those encountered by *C. glabrata* in the host, such as oxidative stress and exposure to antifungal drugs. The fluorescence-based mutation assay is particularly well suited to address such questions because its outcome is independent of whether the strain’s sensitivity to stress is altered by an exogenous treatment (e.g., by an antifungal drug). This assay can also be adapted and used to address questions regarding mechanisms driving mutagenesis in other clinically relevant microbes, including bacterial pathogens and haploid pathogenic fungi, such as *Candida auris*, *Cryptococcus neoformans*, and *Candida lusitaniae*, where emergence of drug resistance poses a serious public health threat.

**MATERIALS AND METHODS**

Construction of the GFP reporter cassette and creating fluorescent strains for fluctuation analyses. The *S. cerevisiae* TEF1 promoter was amplified using primers CACACCAGAGCTCCAAAATGTTTCTACTCC and CCATTGGATCCAAAAACTTAGATTAGATTGC and subcloned into the BamHI-Sacl sites of pYC54 (56), placing it directly upstream of the YFP ORF. Next, the YFP ORF was replaced by that of yEGFP
as follows. yEGFP was amplified from pSR823 (57) using primers ACTAGTGGATCCCGGGGCTCGAGGAATTCATGGGCGGTATTTATCGTGTTCG and subcloned into the plasmid using BamHI-NheI sites. The final plasmid contained the pTEF1-yEGFP-NAT cassette (Fig. 1A). This cassette was amplified from the plasmid using primers CACCCGAGGACGAAGTTCC and TGTAATACGACTCAGCAGG and transformed into C. glabrata strain ATCC 2001 using nourseothricin resistance as selection.

Because there were no targeting homology sequences on the cassette, it integrated randomly into the C. glabrata genome. Several independent, constitutively fluorescent transformants were chosen and submitted for whole-genome sequencing at the New Jersey Medical School Molecular Resource Facility using the NextSeq (Illima, San Diego, CA) platform. Libraries were prepared with the Nextera XT kit (Illima, San Diego, CA) to produce paired-end reads of 150 bp for an approximate minimum coverage of 100×. Data analysis was performed using CLC Genomics Workbench (Qiagen, Hilden, Germany). Each transformant was found to contain a single integration of the cassette. The transformant carrying the cassette integrated between uncharacterized ORFs CAGL0K11132g and CAGL0K11198g on Chr K (strain EScG36), as shown in Fig. 1A, was chosen for further analysis.

In all other C. glabrata strains reported in this study, the cassette was integrated at the same genomic locus as in EScG36 using CRISPR-mediated targeted integration. The cassette was amplified from EScG36 using primers GCAGTCTTTCTGACACCATAC and CACAGAATTGGTAGGACGGG, which produced approximately 500-nt 5′ and 3′ homology each to the desired integration locus. CRISPR was performed as in reference 58, except that cells were made competent for electroporation using the Frozen-EZ yeast transformation kit (Zymo Research) according to the manufacturer’s instructions.

To replace the CAN1 ORF of S. cerevisiae with the yEGFP-NAT cassette, the cassette was amplified using primers AAAAAACCGGATACATGACAAATCAAAAGAAAGACCAGGGCTACATAGGAGCACGATGATTAGTAAATCGACCTC and AGGTATAAAAGCTCATATCTGACTAACAACATCCAAATTGTGTCTGTGACGGGCCCCCTCGAG, which introduced 48-nt regions of homology directly 5′ upstream and 3′ downstream of the CAN1 ORF. This PCR product was transformed into S. cerevisiae strains W4069-4C (wild-type W303 MATα) and W4220-15A (shu1Δ::HIS3 MATα) using the Frozen-EZ yeast transformation kit (Zymo Research) according to the manufacturer’s instructions. Transformants were selected on nourseothricin plates as described above and validated by sequencing of the CAN1 locus and by flow cytometry to verify the acquisition of green fluorescence.

All primers were ordered from Integrated DNA Technologies, and all Sanger sequencing of the above-described constructs was done by Genewiz.

**Measuring yEGFP mutation rates using fluctuation analysis and FACs.** A starter YPD culture of the strain whose mutation rate was being measured was diluted into multiple (e.g., 8 to 12) parallel YPD cultures in a 96-well plate to a starting density of several hundred cells per well and incubated overnight at 37°C (C. glabrata) or 30°C (S. cerevisiae). The following morning, each culture was diluted severalfold in YPD and grown at the same temperature to ensure that when the cells were collected for FACs several hours later, they were in log phase, which was found to be necessary to achieve maximum expression of GFP and the best resolution between GFP-positive and GFP-negative populations. The cells were then collected by filtration using 0.45-μm mixed cellulose ester membrane filters (Millipore), resuspended in PBS, and sorted using the BD FACSAria II (BD Biosciences) at the New Jersey Medical School Flow Cytometry and Immunology Core Laboratory. Ten to 15 min before sorting, each sample was stained with 10 μg/ml propidium iodide (PI; ThermoFisher) to identify and gate out inviable cells. FSC and SSC parameters were set on log scale. Cells were gated for singlets (SSC-W versus SSC-H) followed by live gating on PI negative. Finally, cells were sorted based on GFP expression. GFP-negative cells from each entire culture were collected into microcentrifuge tubes containing 250 μl YPD and then plated immediately onto YPD agar plates. Two hundred GFP-positive cells were also sorted and plated to calculate plating efficiency. The plates were incubated at 37°C (C. glabrata) or 30°C (S. cerevisiae) to allow the sorted cells to form colonies.

The resulting colonies were checked for the presence of the NAT marker by replica plating or patching onto plates containing 100 μg/ml nourseothricin (Jena Bioscience) and then checked for the level of green fluorescence using a BD Accuri C6 flow cytometer (BD Biosciences). Mutation rates and 95% confidence intervals were calculated using the MSS maximum likelihood method (1, 33) based on the number of NAT+ GFP− mutants in every culture (n) and the average number of viable cells per culture (N). To identify the mutations responsible for loss of GFP fluorescence, the yEGFP ORF and promoter were sequenced using primers CTTATTGGATATGAGTCCAATT and TGTAAATACGACTCAGCACG, respectively.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.00120-19.

**FIG S1**, PDF file, 0.9 MB.

**FIG S2**, PDF file, 1.2 MB.

**FIG S3**, PDF file, 1.7 MB.

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REFERENCES

1. Rosche WA, Foster PL. 2000. Determining mutation rates in bacterial populations. Methods 20:4–7. https://doi.org/10.1089/1063452007500901.

2. Lang GI, Murray AW. 2008. Estimating the per-base-pair mutation rate in the yeast Saccharomyces cerevisiae. Genetics 178:67–82. https://doi.org/10.1534/genetics.107.071506.

3. Lippert MJ, Kim N, Cho JE, Larson RP, Schoeny NE, O’Shea SH, Jinks-Robertson S. 2011. Role for topoisomerase 1 in transcription-associated atagenesis in yeast. Proc Natl Acad Sci U S A 108:698–703. https://doi.org/10.1073/pnas.1012363108.

4. Harrington JM, Kolodner RD. 2007. Saccharomyces cerevisiae Msh2-Msh3 acts in repair of base-base mispairs. Mol BiolCell 27:6546–6554. https://doi.org/10.1128/MCB.00855-07.

5. Buckland RJ, Watt DL, Chittoor B, Nilsson AK, Kunkel TA, Chabes A. 2014. Increased and imbalanced dNTP pools symmetrically promote both leading- and lagging-strand replicative helicities. PLoS Genet 10: e1004846. https://doi.org/10.1371/journal.pgen.1004846.

6. Roetzer A, Gabaldon T, Schuller C. 2011. From mutational innovation to a cell’s molecular blueprint. Mol Genet Genomics 285:1111–1124. https://doi.org/10.1007/s00438-011-3126-0.

7. Nguyen MH, Peacock JE, Jr, Morris AJ, Tanner DC, Nguyen ML, Snydman DR, Wagener MM, Rinaldi MG, Yu VL. 1996. The changing face of candidemia: emergence of non-Candida albicans species and antifungal resistance. Am J Med 100:517–623. https://doi.org/10.1016/S0002-9343(95)00010-0.

8. Antinori S, Milazzo L, Ischer F, Leibundgut-Landmann S, Sanglard D. 2013. Gain-of-function mutations in PDR1, a regulator of antifungal drug resistance in Candida glabrata, control adherence to host cells. Infect Immun 81:1709–1720. https://doi.org/10.1128/IAI.00074-13.

9. Glockner A, Cornely OA. 2015. Candida glabrata—unique features and properties. Front Microbiol 6:1995. https://doi.org/10.3389/fmicb.2016.01995.

10. Robertson S. 2011. Role for topoisomerase 1 in transcription-associated azole resistance in clinical isolates and petite mutants. PLoS One 6:e20111. https://doi.org/10.1371/journal.pone.0020112.

11. Arendrup MC, Patterson TF. 2017. Multidrug-resistant Candida: epidemiology, molecular mechanisms, and treatment. J Infect Dis 216:545–5451. https://doi.org/10.1093/infdis/jix131.

12. Alexander BD, Johnson MD, Pfeiffer CD, Jimenez-Ortigosa C, Catania J, Booker R, Castanheira M, Messer SA, Perlín DS, Pfaller MA. 2013. Increasing echinocandin resistance in Candida glabrata: clinical failure correlates with presence of FKS mutations and elevated minimum inhibitory concentrations. Clin Infect Dis 56:1724–1732. https://doi.org/10.1093/cid/cct136.

13. Vale-Silva L, Ischer F, Leibundgut-Landmann S, Sanglard D. 2013. Gain-of-function mutations in FKS1 and FKS2 mutations on echinocandin sensitivity in Candida glabrata isolates in India despite backmutation. PLoS One 8:e72218. https://doi.org/10.1371/journal.pone.0052218.

14. Lott TJ, Frade JP, Lockhart SR. 2010. Multilocus sequence type analysis reveals both clonality and recombination in populations of Candida glabrata bloodstream isolates from U.S. surveillance studies. Eukaryot Cell 9:619–625. https://doi.org/10.1128/EC.00002-10.

15. Polakova S, Blume C, Zarate JA, Mentel M, Jorck-Ramberg D, Stenderup J, Piskur J. 2009. Formation of new chromosomes as a virulence mechanism in yeast. Nature 462:599–606. https://doi.org/10.1038/nature08730.

16. Singh A, Healey KR, Yadav P, Upadhyaya G, Sachdeva N, Sarma S, Kumar A, Tarai B, Perlín DS, Chowdhary A. 2018. Absence of azole or echinocandin resistance in Candida glabrata isolates in India despite background prevalence of strains with defects in the DNA mismatch repair pathway. Antimicrob Agents Chemother 62:e00195-18. https://doi.org/10.1128/AAC.00195-18.

17. Deliere S, Healey K, Gits-Muselli M, Carrara B, Barbaro A, Guigue N, Leclef C, Touratier S, Desnos-Ollivier M, Perlín DS, Bretagne S, Alanoia A. 2016. Fluconazole and echinocandin resistance of Candida glabrata correlates better with antifungal drug exposure rather than with MS2H2 mutator genotype in a French cohort of patients harboring low rates of resistance. Front Microbiol 7:2038. https://doi.org/10.3389/fmicb.2016.02038.

18. Hall BM, Ma CX, Liang P, Singh KK. 2009. Fluclotin analysis of C. glabrata for determining the mutation rate using Luria-Delbruck
Long H, Weinstein J, Rothstein R. 2005. A genetic screen for top3 suppressors in *Saccharomyces cerevisiae* identifies SHU1, SHU2, PSY3 and CSM2: four genes involved in error-free DNA repair. Genetics 169: 1275–1289. https://doi.org/10.1534/genetics.104.036764.

Huang ME, Rio AG, Galibert MD, Galibert F. 2002. Pol32, a subunit of DNA polymerase delta, suppresses genomic deletions and is involved in the mutagenic bypass pathway. Genetics 160:1409–1422.

Ni TT, Marsischky GT, Kolodner RD. 1999. MSH2 and MSH6 are required for removal of adenine misincorporated opposite 8-oxo-guanine in *S. cerevisiae*. Mol Cell 4:343–444. https://doi.org/10.1016/S1097-2765(00)80346-9.

Gammie AE, Erdeniz N, Beaver J, Devlin B, Nanji A, Rose MD. 2007. Functional characterization of pathogenic human MSH2 missense mutations in *Saccharomyces cerevisiae*. Genetics 177:707–721. https://doi.org/10.1534/genetics.107.071084.

Marsischky GT, Filosi N, Kane MF, Kolodner R. 1996. Redundancy of *Saccharomyces cerevisiae* MSH3 and MSH6 in MSH2-dependent mismatch repair. Genes Dev 10:407–420. https://doi.org/10.1101/gad.10.4.407.

Tran HT, Keen JD, Kricker M, Resnick MA, Gordenin DA. 1997. Hypermutability of homonucleotide runs in mismatch repair and DNA polymerase proofreading yeast mutants. Mol Cell Biol 17:2859–2865. https://doi.org/10.1128/MCB.17.5.2859.

Lee GS, Blonsky KS, Van On DL, Savage EA, Morgan AR, von Borstel RC. 1992. Base alterations in yeast induced by alkylating agents with differing Swain-Scott substrate constants. J Biol Chem 223:617–626. https://doi.org/10.1016/0022-2836(92)90199-8.

Lee H, Popodi E, Tang H, Foster PL. 2012. Rate and molecular spectrum of spontaneous mutations in the bacterium *Escherichia coli* as determined by whole-genome sequencing. Proc Natl Acad Sci U S A 109: E2774–E2783. https://doi.org/10.1073/pnas.1003409109.

Long H, Behringer MG, Williams E, Te R, Lynch M. 2016. Similar mutation rates but highly diverse mutation spectra in *Saccharomyces cerevisiae* and their effect on recombination and mutagenesis. *Mutat Res* 807:135–145. https://doi.org/10.1016/j.mutres.2016.07.009.

Ragheb MN, Thomason MK, Hsu C, Nugent P, Gage J, Samadpour AN, Karissa A, Menikh CN, Miller SJ, Sherman DR, Menikh H. 2019. Inhibiting the evolution of antibiotic resistance. *Mol Cell* 73:157–165.e5. https://doi.org/10.1016/j.molcel.2018.10.015.

Ragheb MN, Putnam CD, Kolodner RD. 2018. Analyzing genome rearrangements in *Saccharomyces cerevisiae*. Methods Mol Biol 1672:43–61. https://doi.org/10.1007/978-1-4939-7306-4_5.

Sun S, Xu J. 2009. Chromosomal rearrangements between serotype A and D strains in *Cryptococcus neoformans*. PLoS One 4:e5524. https://doi.org/10.1371/journal.pone.0005524.

Forche A, Magee PT, Selmecki A, Berman J, May G. 2009. Evolution in *Candida albicans* populations during a single passage through a mouse host. Genetics 182:799–811. https://doi.org/10.1534/g3.113.006908.